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# WILLIAM LLEWELLYN'S

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Anabolic Overview

Anabolic steroids are a class of medications that contain a synthetically manufactured form of the hormone testosterone, or a related compound that is derived from (or similar in structure and action to this hormone. In order to fully grasp how anabolic steroids work, it is, therefore, important to understand the basic functioning of testosterone.

Testosterone is the primary male sex hormone. It is manufactured by the Leydig's cells in the testes at avaing amounts throughout a person's life span. The effects of this hormone become most evident during the time of puberty, when an increased output of testosterone will elicit dramatic physiological changes in the male body. This includes the onsect of secondary male characteristics such as a deepened voice, body and facial har growth, increased oil output by the sebaceous glands, development of sexual organs, mutantant of sperim, and an increased flish. Indeed the male reproductive and the secondary of the

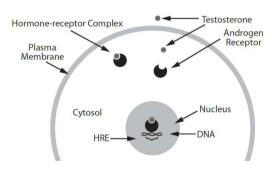
Increased testosterone production will also cause growth promoting or "anabolic" changes in the body, including an enhanced rate of protein synthesic (leading to muscle accumulation). Testosterone is the reason males carry more muscle mass than women, as the two sexes have vasily contrasting amounts of this hormone. More specifically, he adult male body will manufacture between 2.5 and 11 mg per day! while females only produce about 14/mg. The dominant sex hormone for women is estrogen, which has a significantly different effect on the body. Among other things, a lower androgen and higher estrogen level will cause women to store more body fall, accumulate less muscle tissue, have a shorter stature, and become more apt to bone weakening with ange (steaporosis).

The actual mechanism in which testosterone elicits these changes is somewhat complex. When free in the blood stream, the testosterone molecule is available to interact with various cells in the body. This includes skeletal muscle cells, as well as skin, scalp, kidney, bone, central nervous system, and prostate tissues. Testosterone binds with a cellular target in order to exert its activity, and will, therefore, effect only those body cells that posses the proper hormone receptor site (specifically the androgen receptor). This process can be likened to a lock and key system, with each receptor (lock) only being activated by a particular type of hormone (key). During this interaction, the testosterone molecule will become bound to the intracellular receptor site (located in the cytosol, not on the membrane surface), forming a new "receptor complex." This complex (hormone + receptor site) will then migrate to the cell's nucleus, where it will attach to a specific section of the cell's DNA, referred to as the hormone response element. This will activate the transcription of specific genes, which in the case of a skeletal muscle cell will ultimately cause (among other things) an increase in the synthesis of the two primary contractile proteins, actin and myosin (muscular growth). Carbohydrate storage in muscle tissue may be increased due to androgen action as well

Once this messaging process is completed, the complex will be released, and the receptor and farmone will disassociate. Both are then free to migrate back into the cytosol for further activity. The testosterone molecule is also free to diffuse back into circulation to interact with other cells. The entire receptor cycle, including hormone binding receptor-hormone complex migration, gene transcription and subsequent return to cytosol is a slow process, taking busts, not minutes, to complete. For example, in studies using

a single injection of nandrolone, it is measured to be 4 to 6 hours before free androgen receptors migrate back to the cytosol after activation. It is also suggested that this cycle includes the splitting and formation of new androgen receptors once returned to cytosol, a possible explanation for the many observations that androgens are integral in the formation of their own receptor sizes.<sup>2</sup>

In the kidneys, this same process works to allow androgens to augment erythropoissic for blood cell production.<sup>3</sup> It is this effect that leads to an increase in red blood cell concentrations, and possibly increased oxygen transport capacity, during anholic/androgenic steroid therapy. Many athletes ability, due to specific uses or mentions of this effect in fund literature. In fact, straulation of erythropoissis occurs with nearly all anabolic/androgenic steroids, as this effect is simply tied with activation of the androgen receptor in kidney cells. The only real exceptions might be compounds such as displottedostostorous and some of its derivatives, <sup>4</sup> which are ripidly brocked colors upon interaction with the 3alpha-dytoxysteroid dehydrogenuse enzymes (kidney tissue has a similar enzyme distribution to muscle tissue, see "anabolic/androgenic dissociation" section), and therefore display low activity is these tissues.



CELLULAR DIAGRAM: Testosterone freely diffuses through the plasma membrane and binds with an intracellular androgen receptor. The hormone-receptor complex then enters the cell nucleus to bind with a specific segment of DNA (the Hormone Response Element), activating the transcription of specific genes.

support the lipolytic (fat mobilizing) capacity of cells.5 This may be accomplished by an androgen-tied regulation of beta-adrenergic receptor concentrations or general cellular activity (through adenylate cyclase).6 We also note that the level of androgens in the body will closely correlate (inversely) with the level of stored body fat. As the level of androgenic hormones drops typically the deposition of body fat will increase 7 Likewise as we enhance the androgen level body fat may be depleted at a more active rate. The ratio of androgen to estrogen action is in fact most important, as estrogen plays a counter role by acting to increase the storage of body fat in many sites of action 8 Likewise if one wished to lose fat during steroid use estrogen levels should be kent low. This is clearly evidenced by the fact that non-aromatizing steroids have always been favored by bodybuilders looking to increase the look of definition and muscularity while aromatizing compounds are typically relegated to bulking phases of training due to their tendency to increase body fat storage. Aromatization is discussed in more detail in a following section (see: Estrogen Aromatization)

As mentioned, testosterone also elicits androgenic activity, which occurs by its activating recentors in what are considered to be androgen responsive tissues (often through prior conversion to dihydrotestosterone. See: DHT Conversion) This includes the sehaceous glands, which are responsible for the secretion of oils in the skin. As the androgen level rises, so does the release of oils. As oil output increases, so does the chance for nores becoming clossed (we can see why acne is such a common side effect of steroid use). The production of body and facial hair is also linked to androgen. recentor activation in skin and scaln tissues. This becomes most noticeable as hove mature into nuberty, a period when testosterone levels rise rapidly, and androgen activity begins to stimulate the growth of hair on the body and face. Some time later in life and with the contribution of a genetic predisposition androgen activity in the scalp may also help to initiate male-pattern hair loss. It is a misconcention that dihydrotestosterone is an isolated culprit in the promotion of hair loss, however, as in actuality it is the general activation of the androgen receptor that is to blame (see: DHT Conversion). The functioning of sex glands and libido are also tied to the activity of androgens as are numerous other regions of the central nervous/neuromuscular system 1. Role of androgens in growth and development of the fetus, child, and adolescent.

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# Direct and Indirect Anabolic Effects

experimented with for many decades now, there is still some debate today as to exactly how steroids affect muscle mass. At this point in time, the primary mode of anabolic action with all anabolic-indrogenic steroids is understood to be direct activation of the cellular androgen receptor and increases in protein synthesis. As follows, if we are able to increase our androgen level from an external source by supplementing testosterone or a similar anabolic steroid, we can greatly enhance the rate in which protein is retained by the muscles. This is clearly the primary cause for muscle growth with all anabolic/androgenic steroids. As our hommone levels increase, so does androgen receptor activation, and ultimately the rate of protein synthesis.

But other indirect mechanisms could possibly affect muscle growth outside of the normally understood androgen action on protein synthesis. An indirect mechanism is one that is not brought about by activation of the androgen receptor, but the affect androgens might have on other hormones, or even the release of locally acting hormones or growth promoters inside cells (perhaps mediated by other membrane bound receptors). We must remember also that muscle mass disposition involves not only protein synthesis, but also other factors such as tissue nutrient transport and protein breakdown. We need to look at androgenic interaction with these factors as well to get a complete picture. Concerning the first possibility, we note that studies with testosterone suggest that this hormone does not increase tissue amino acid transport. 9 This fact probably explains the profound synergy bodybuilders have noted in recent years with insulin, a hormone that strongly increases transport of nutrients into muscle cells. But regarding protein breakdown, we do see a second important pathway in which androgens might affect muscle growth

# Anti-Glucocorticoid Effect of Testosterone

Testosterone (and synthetic anabolic/androgenic steroids) may help to increase mass and strength by having an anti-catabolic effect on muscle cells. Considered one of the most important indirect mechanisms of androgen action, these hormones are shown to affect the actions of another type of steroid hormone in the body, glascoorticoids (controls is the primary representative of this group). Of Chaccoorticoid hormones actually have the exact opposite effect on the muscle cell than androgens, namely sending an order to release stored protein. This process is referred to as catabolism, and represents a breaking down of muscle tissue. Muscle growth is achieved when the anabolic effects of testosterone are more pronounced overall than the degenerative effects of cortisols (Whi intense training and a proper dict, the body will typically store more protein than it removes, but this underlying battic s always constant.

When administering anabolic steroids, however, a much higher androgen level can place glucocorticoids at an onable disadvantage. With their effect reduced, fewer cells will be given a message to release protein, and more will be accumulated in the long run. The primarily mechanism believed to bring this effect out is androgen displacement of glucocorticoids bound to the glucocorticoid receptor. In fact, in-vitro studies have supported this notion by demonstrating that testosterone has a very high affinity for this receptor, If and further suggesting that some of its anabolic activity is directly mediated through this action. <sup>12</sup> It is also suggested that androgens may indirectly interfere with DNA holiding to the glucocorticoid response chemnel. <sup>13</sup> Although the exact underlying mechanism is still in debate, what is clear is that steroid administration inhibits protein breakdown, even in the fasted state, which seems clearly indicative of an anti-catabolic effect.

#### Testosterone and Creating

In addition to protein synthesis, a rise in androgen levels should also enhance the synthesis of creatine in skeletal muscle tissues. 14 Creatine, as creatine nhosphate (CP) plays a crucial role in the manufacture of ATP (adenosine triphosphate) which is a main store of energy for the muscles. As the muscle cells are stimulated to contract. ATP molecules are broken down into ADP (adenosine dinhosphate), which releases energy. The cells will then undergo a process using creatine phosphate to rapidly restore ADP to its original structure, in order to replenish ATP concentrations. During periods of intense activity however this process will not be fast enough to compensate and ATP levels will become lowered. This will cause the muscles to become fatigued and less able to effort a strenuous contraction. With increased levels of CP available to the cells. ATP is replenished at an enhanced rate and the muscle is both stronger and more enduring. This effect will account for some nortion of the early strength increases seen during steroid therapy. Although nerhans not technically considered an anabolic effect as tissue hypertrophy is not a direct result androgen support of creatine synthesis is certainly still looked at as a positive and growth-supporting result in the mind of the bodybuilder

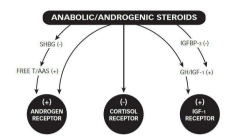
#### Testosterone and IGF-1

It has also been suggested that there is an indirect mechanism of testosterone action on muscle mass mediated by Insulla-Like Growth Tacort. To be more specific, studies note a clear link between androgens and tissue release of, 15 and responsiveness to, this anabolic hormone. For example, it has been demonstrated that increases in IGF-1 receptor concentrations in skeletal muscle are noted when eldedy men are given replacement dosses of testosterone. 16 In essence, the cells are becoming primed for the actions of IGF-1, by restosterone. Alternately we see marked decreases in IGF-1 receptor protein levels with androgen deficiency in young men. It also appears that androgens are necessary for the head production and function of IGF-1 in skeletal imuscle cells, adeependent of circulating growth hormone, and IGF-1 levels. <sup>15</sup> Since we do know for certain that IGF-1 is at least a minor anabolic hormone in muscle tissue, it seems reasonable to conclude with secroid themse.

#### Direct and Indirect Steroids?

In looking over the proposed indirect effects of testosterone, and pondering the effectiveness of the synthetic anabolicalmogenis stretols, we must resist the temptation to believe we can categorize steroids as those which directly, and those which indirectly, promote musel growth. The belief that there are two dichotomous groups or classes of steroids ignores the fact that all commercial steroids promote not only musele growth but also androgenistic productions of the state of the

way varous compounts count toter growtn morrenty, such trait anyuntages might even be found with certain synergistic drug combinations, the primary might even be found with certain synergistic drug combinations, the primary mode of action with all of these compounds is the androgen receptor. The notion that streotly X and Y must never be stacked together because they both compete for the same receptor when stimulating growth, while X and Z is also also also combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they work via different mechanisms, should be combined because they would be compared to the combined of the combined because they work with the combined of the combined because they would be combined to the combined t



MECHANISM OF ACTION DIAGRAM: The mechanism of anabolic action due to the administration of anabolic action due to the administration of anabolic accuses not only direct stimulation of the androgen receptor, but also usupports muscle growth by increasing the levels of free androgen receptor, the description of the analysis of the androgen receptor due to the analysis of the androgen receptor density, inhibiting corticosteroid action, increasing called the analysis of the ana

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#### Free vs. Round Testosterone

A very small amount of testosterone actually exists in a free state, where interaction with cellular recentors is possible. The majority will be bound to the proteins SHBG (sex hormone binding globulin, also referred to as sex steroid binding globulin and testosterone-estradiol binding globulin) and albumin which temporarily prevent the hormone from exerting activity Steroid hormones actually bind much more avidly to SHBG than albumin (with approximately 1 000 times greater affinity) however albumin is present in a level 1 000 times greater than SHBG. Therefore, the activity of both binding proteins in the body is relatively equal. The distribution of testosterone in men is typically 45% of testosterone bound to SHBG and about 53% bound to albumin. The remaining 2% of the average blood concentration exists in a free unbound state. In women, the percentage of free testosterone is lower measured to be approximately 1% A hinding protein called ABP (androgen binding protein) also helps to mediate androgen activity in the reproductive system, although since it is found exclusively in these tissues it is not relevant to muscle growth.

The level of free testosterone available in the blood is likewise an important factor mediating is activity, as only a small percentage is really active at any analoperating as really active at any analoperating analoperatin

For example, Proviron® (1-methyl dihydrotestosterone) binds with SHBG many times more avidly than testosterone. <sup>19</sup> while mibolerone (7,17 dimethyl-nandrolone) and bolasterone (7,17 dimethyl-testosterone) show virtually no affinity for this protein at all (clearly the reason these steroids are such notent androcens).

The level of SHBG present in the body is also variable, and can be altered by a number of factors. The most prominent seems to be the concentration of estrogen and thyroid hormones present in the blood. We generally see a reduction in the amount of this plasma binding protein as estrogen and thyroid content decreases, and a rise in SHBG as they increase. A heightened androgen level due to the administration of anabolic-landongenis estroids has also been shown to lower levels of this protein considerably. This is clearly supported by a 1989 German study, which noted a strong tendency for SHBG reduction with the oral anabolic steroid stanozolol (Winstrolle), 20 AHC on 3) 4 alony of administration and a faily dose of 2 cargle poly-weight (about 18mg for a 200th man), SHBG was lowered nearly 50% in normal subjects. Similar results have been obtained with the use of nijectable testosterone enanthate; however, milligram for milligram, the effect of stanozold was much present in comparison. The form of administration may

have been important in reaching this level of response. Although the injectable was not tried in the German study, we can refer to others comparing the effect of oral vs. transdermal estrogen. <sup>21</sup> These show a much greater response in SHBG levels when the drug is given orally. This is perhaps explained by the fact that SHBG is produced in the here. Therefore, we cannot assume that injectable Winstrol® (or injectable steroids in general) will disbut whe same level of potners or this regard.

Lowering the level of plasma binding proteins is also not the only mechanism that allows for an increased level of the estosaterone. Servicials that display a high affinity for these proteins may also increase the level of free testosterone by competing with a for binding. Obvolusely if estosterone finish a more difficult to locate available plasma proteins in the presence of the additional including disploydotestosterone. Proviscentification of Oral-Turinshol (chlorodelydromethylestosterone) display a strong tendency for this effect. If the level of free-testosterone can be alrered by the use of different anabolic androgenic steroids, the possibility also exists that one steroid can increase the potency of another through these same mechanisms. For example, Proviront 8 is a poor anabolic, but its externedy high affinity for SIIG might made in surfail by allowing the displacement of other stronds.

We must not let this discussion lead us into thinking that binding proteins serve no valuable function. In fact they play a vial role in the transport and functioning of endogenous androgens. Binding proteins act to protect the steroid against rapid metabolism, enusue a more stable blood hormone concentration, and facilitate an even distribution of hormone to various body organs. The recent discovery of a specific receptor for Sex Hormone Binding Globulin (SHBG-R) located on the membrane surface of steroid responsive body cells also aggests a much more complicated role for this protein than the stable of the stable st

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# Estrogen Aromatization

Testosterone is the primary substrate used in the male body for the synthesis of estrogan (estratable), the principal fermale sex hormone. Although the presence of estrogan may seem quite unusual in men. it is structurally very similar to testosterone. With a slight alteration by the enzyme aromatuse, estrogan is produced in the male body. Aromatuse activity occurs in various regions of the male body, including adipose, <sup>22</sup> lover, <sup>23</sup> gondal, <sup>24</sup> central nervous system; <sup>25</sup> and skeletal musicle<sup>26</sup> tissues. In the context of the average healthy male, the amount of estrogan produced is generally not very significant to one's body disposition, and may even be beneficial in terms of cholesterol values (See Siske Effects: Cantion accolar System). However, in larger amounts it does have potential to cause many unwanteed effects.

including water retention, female breast tissue development (gynecomastia), and and body flat accumulation. For these reasons, many focus on minimized to build-up or activity of estrogen in the body with aromatsase inhibitors such as the build-up or activity of estrogen in the body with aromatsase inhibitors such as the strength of the strength of

We must, however, not be led into thinking that estrogen serves no henefit. It is a tautilly a desirable hormone in many regards. Athletes have known for years that estrogenic steroids are the best mass builders, but it is only recently that we are frailsy coming to understand the underlying mechanisms why. It appears that reasons go beyond the simple size, weight, and strength that the strength of the size of mabelian. This is manifest through increases in glacose utilization, growth hormone secretion, and androgen receptor proliferation.

#### Glucose Utilization and Estrogen

Estrogan may play a very important role in the promotion of an anabolic state by affecting glacose utilization in muscle tissue. This occurs via an altering of the level of available glacose 6-phosphate dehydrogenase, an enzyme directly tict to the use of glacose for muscle tissue growth and recruenting 1.72 st. More specifically, G6PD is a vial part of the pentose phosphate pathway, which is integral in determining the rate nucles each and plasts are to be synthesized in ethe for tissue repair. During the period of regeneration after which is believed to represent amentations for the body to enhance recovery when needed. Surprisingly, we find that estrogen is directly tied to the level of G6PD that is to be made available to cells in this recovery window.

The link between estrogen and G6PD was established in a study demonstrating levels of this dehydrogenase enzyme to rise after administration of testosterone propionate. The investigation further showed that the aromatization of testosterone to estradiol was directly responsible for this increase, and not the androgenic action of this steroid 29 The nonaromatizable steroids dihydrotestosterone and fluoxymesterone were tested alongside testosterone propionate, but failed to duplicate the effect of testosterone. Furthermore, the positive effect of testosterone propionate was blocked when the aromatase inhibitor 4-hydroxyandrostenedione (formestane) was added, while 17-beta estradiol administration alone caused a similar increase in G6PD to tesosterone propionate. The inactive estrogen isomer alpha estradiol, which is unable to bind the estrogen receptor, failed to do anything. Further tests using testosterone propionate and the antiandrogen flutamide showed that this drug also did nothing to block the positive action of testosterone, establishing it as an effect independent of the androgen receptor.

#### Estrogen and GH/IGF-1

Estrogan may also play an important role in the production of growth hormone and IGF1-I IGF-I (install-like growth factor) is an anabolic hormone released in the liver and various peripheral tissues via the stimulus of growth hormone (See Dung Profiles: Growth Hormone). IGF-I is responsible for the anabolic activity of growth hormone such as increased mitrogen retention/protein synthesis and cell phyerplassi (profileration). Doe of the first studies to bring this issue to our attention looked at the effects of the anti-estrogan monositien on IGF-I levels, demonstrating it to have a in 1993, which looked at the effects of testosterone replacement therapy on GH and GFI-1 levels alone, and compared them to the effects of restosterone combined again with tamoxifen. 31 When tamoxifen was given. GH and IGF-1 levels were notably suppressed, while both values were elevated with the administration of restosterone cannibate alone. Another study has shown 300 anomal men. Here the 300 mg of testosterone eater caused an elevation of centraled levels, which would be expected at such a door. This was compared to the effect of the same dosage of nandrolene decanotate; however, this strend falled to produce the same increase. This result is quite interesting, especially when we note that estrogen levels were actually lowered<sup>32</sup> when this steroid hadd to produce the same increase. This result is quite interesting, especially when we note that estrogen levels were actually lowered<sup>32</sup> when this steroid was given. Yet another demonstrated that GH and IGFI-1 secreted with testosterone (non-aromatizable) seems to suppress GH and IGFI-1 secretom. 3

suppressive effect 50 A second, perhaps more noteworthy, study took place

#### Estrogen and the Androgen Receptor

It has also been demonstrated that estrogen can increase the concentration of androgen receptors in certain tissues. This was shown in studies with rats. which looked at the effects of estrogen on cellular androgen receptors in animals that underwent orchiectomy (removal of testes, often done to diminish endogenous androgen production). According to the study, administration of estrogen resulted in a striking 480% increase in methyltrienolone (a potent oral androgen often used to reference receptor binding in studies) binding in the levator ani muscle 34. The suggested explanation is that estrogen must either be directly stimulating androgen recentor production, or perhaps diminishing the rate of recentor breakdown. Although the growth of the levator ani muscle is commonly used as a reference for the anabolic activity of steroid compounds, it is admittedly a sex organ muscle, and different from skeletal muscle tissue in that it possesses a much higher concentration of androgen receptors. This study, however, did look at the effect of estrogen in fast-twitch skeletal muscle tissues (tibialis anterior and extensor digitorum longus) as well, but did not note the same increase as the levator ani. Although discouraging at first glance, the fact that estrogen can increase androgen receptor binding in any tissue remains an extremely significant finding, especially in light of the fact that we now know androgens to have some positive effects on muscle growth that are mediated outside of muscle tissue.

Extrogen and Fatfgue

"Steroid Fatigue" is a common catchphrase these days, and refers to another
important function of estrogen in both the male and female body, namely its
ability to promote wakefulness and a mentally alert state. Given the common
availability of potent third-generation aromatuse inhibitors, bodybuates
today are (at times) noteing more extreme estrogen suppression than they
conditions, the athlete, though on a productive cycle of rings, may not be
able to maximize his or her gains due to an inability to train at full vago. This
effect is sometimes also dubbed "servioid lethrage," The reason is that
extrogan plays an important supporting role in the activity of sectorian.
Servioin is one of the body's principle neutralsmattines, vital to mental
alertness and the skeptwake cycle, 35 36 Interference with this
neutralsmatter is also associated with crottoe fittings exponence. The service is that
meutralsmatter is also associated with chronic fittings exponence, 378 so we
can see how vital it is to fatigue specifically. Estrogen suppression in

menopause has also been associated with fatigue. 39 as has the clinical use of newer (more potent) aromatase inhibitors like anastrozole. 40 letrozole. 41

exemestane. 42 and fadrozole 43 in some patients. These things may be important to consider when planning your next cycle. Although not everyone notices this problem when estrogen is low for those that do a little testosterone or estrogen can go a long way in correcting this. It is also of note that the use of strictly non-aromatizable steroids sometimes causes this effect as well likely due to the suppression of natural testosterone production (cutting off the main substrate used by the male body to make estrogen).

#### Anti-Estrogens and the Athlete

So what does this all mean to the bodybuilder looking to gain ontimal size? Basically I think it calls for a cautious approach to the use of estrogen maintenance drugs if mass is the key objective (things change of course if we are talking about cutting). Obviously, anti-estrogens should be used if there is a clear need for them due to the onset of estrogenic side effects, or at the very least, the drugs being administered should be substituted for nonestrogenic compounds. Gynecomastia is certainly an unwanted problem for the steroid user as are noticeable fat mass gains. But if these problems have not presented themselves, the added estrogen due to a cycle of testosterone

or Dianabol, for example, might indeed be aiding in the buildup of muscle mass or keening you energetic. An individual confident they will notice or are not prone to getting estrogenic side effects, may therefore want to hold off using estrogen maintenance drugs so as to achieve the maximum possible gains in tissue mass

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## DHT Conversion

As we see from our discussion with estrogen, in considering the physiological effects of any steroid, we must look at all of its active metabolises, and not just the initial compound. This includes not only estrogenic products, but androgenic metabolises as well. With his initial, it simportant to note that the potency of testosterone is considerably increased in many androgen responsive tissues when it converts to dibydrotestsetone. More commonly referred to by the three-letter abbreviation DHT, this hommone is, in fact, measured to be approximately three for our times stronger than testosteron. It is the most potent steroid found naturally in the human body, and important to discuss if we are to understand the full activity of testosterone, as well as other anabolic/androgenic steroids that undergo a similar

Testosterone is converted to dihydrotestosterone upon interaction with the 5-alpha reductase exympe. More specifically, this exzympe removes the C4-5 double-hoad of testosterone by the addition of two hydrogen atoms to its structure (hence the name di-hydro testosterone). The removal of this bond is important, as in this case is creates a steroid that binds to the androgen receptor much more avoilty than its purent steroid. Scalpha reductase is present in high amounts in tissues of the prostate, skin, scalp, liver, and various regions of the certain leverous system, and as such represents a mechanism for the body to increase the potency of testosterone specifically testosterone that the statisterone will be attainly made its way to the receptor without being converted to dihydrotestosterone, making DHT by far the active form of androgen here.

#### DHT and Androgenic Side Effects

In some regards this local potentiation of testosterone's activity may be unwelcome, as higher androgenia earlivity in certain tissues may produce a number of undesirable side effects. Acne, for example, is often triggered by dideytoetostosterone activity in the schezous glands, and the local formation of dihydrotestosterone activity in the schezous glands, and the local formation of dihydrotestosterone in the scalp is typically blanned for triggering male pattern hair loss, two should know that it is a terrible misconception among bodybrailers that dhydrotestosterone is an isolated captive when it comes to manifold and androgenic, through the same cellular androgen receptor. Dhydrotestosterone is no different than any other steroid except that it is a more potent activator of this receptor than most, and can be formed locally in certain androgen-sensitive tissues. All steroids can cause androgenic side effects in direct relation to their affirmly for this receptor, and DHT has no

# Renefits of DHT

While a lot of attention is being paid to the negative side effects of the androgen dihydrotestosterone, you should know that there are some known benefits to the strong androgenic activity brought about by this hormone as well. For example, DHT plays an important role in the organization and functioning of the central nervous system. Many neural cells contain active androgen receptors, and it is thought that there may even be a specific importance of dihydrotestosterone in this area of the body. Studies have shown DHT to have a profoundly greater impact in these cells compared to testosterone. More specifically, animal models demonstrated that both testosterone and DHT would result in increased androgen receptor proliferation in neural cells three and seven hours after being administered. however only DHT was able to sustain this increase at the twenty-one hour mark. 44 Although some might contend that this difference is simply due to DHT forming a more stable and lasting complex with the androgen receptor. others suggest that DHT and testosterone might even be affecting neural cells differently, such that the dihydrotestosterone-receptor complex and testosterone-receptor complex might be activating the transcription of different target genes.

The strong interaction between the central nervous system and skeletal muscles, collectively referred to as the neuromuscular system, so of key importance to the athiet. There appears to be little doubt that the ability of the body to adapt to training, and to activate nerve endings in muscle itsue, is reliant on the interactions of the neuromuscular system. Inhibiting the formation of DHT during a testostence cycle may therefore inadverently interfere with strength and muscle mass gains. This would explain why bodybuilders commonly report at dops in steroid potency when they add the S-alpha reductase inhibitor finasteride to a testosterone cycle. Many complain strength and even muscle mass gains solve spill ending the strength and even the strength and even the size with sufficiently responsible for goods. Clearly more is activation in muscle tissue were solely responsible for goods. Clearly more is involved, and we cannot look at dihydrotestosterone simply as a side-effect hormone.

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#### Brief History of Anabolic/Androgenic Steroids

While is had been clear for many centuries that the testicles were crucial for the male body to properly develop, it was not until modern times that an understanding of testosterone began to form. The first solid scientific experiments in this rear, which eventually led to the discovery and replactant of testosterone (and related androgens), were undertaken in the 1800s. During this century a number of animal experiments were published, most of which involved the removal and/or implantation of testicular material from/in a sulpect. Although very rurde in design by today's standards, these studies certainly last the foundation for the modern field of endocrinology (the study of homonosa). By the turn of the century, scientists were alse hor produce the first experimental androgen injections. These were actualized either through the filtering of large quantities of urine (for article homonosa), or by extracting testosterone from animal testicles. Again, the methods were rough but the final results proved to be very enlightening. Chemists finally synthesized the structure of testosterone in the mid-1930's, sparking a new wave of interest in this homone. With the medical community paying a tremendous amount of attention to this achievement, the possible therapeutic uses for a readily available synthetic testosterone quickly became an extremely popular focus. Many believed the applications for this type of a medication would be extremely fare-reaching, with uses ranging from the maintenance of an androgan deficiency, to that of a good health and well-being treatment for the sixkly or delderly. During the infinery of such experimentation, many believed they had crossed paths with a true "fountain of youth."

Dihydrotestosterone and nandrolone, two other naturally occurring steroids. were also isolated and synthesized in the early years of steroid development. To make things even more interesting, scientists soon realized that the androgenic, estrogenic, and anabolic activity of steroid hormones could be adjusted by altering their molecular structure. The goal of many researchers thereafter became to manufacture a steroid with extremely strong anabolic activity, but will display little or no androgenic/estrogenic properties. This could be very beneficial, because side effects will often become pronounced when steroid hormones are administered in supraphysiological amounts. A "pure" anabolic would theoretically allow the patient to receive only the beneficial effects of androgens (lean muscle mass gain, increased energy and recuperation, etc.), regardless of the dosage. Some early success with the creation of new structures convinced many scientists that they were on the right track. Unfortunately none of this progress led researchers their ultimate goal. By the mid-1950's, well over one thousand testosterone, nandrolone, and dihydrotestosterone analogues had been produced, but none proved to be purely anabolic compounds.

The failure to reach this goal was primarily due to an initial flawed understanding of testosterone's action. Scientists had noticed high levels of DHT in certain tissues, and believed this indicated an unusual receptor affinity for this hormone. This led to the belief that the human body had two different androgen receptors. According to this theory, one receptor site would respond only to testosterone (eliciting the beneficial anabolic effects). while the other is activated specifically by the metabolite. dihydrotestosterone. With this understanding, eliminating the conversion of testosterone to DHT was thought capable of solving the problem of androgenic side effects, as these receptors would have little or none of this hormone available for binding. More recently, however, scientists have come to understand that only one type of androgen receptor exists in the human body. It is also accepted that no anabolic/androgenic steroid can possibly be synthesized that would participate only with receptors in tissues related to anabolism DHT which was once thought not to bind to the same recentor as testosterone, is now known to do so at approximately three to four times the affinity of its parent, and the unusual recovery of DHT from androgen responsive tissues is now attributed to the distribution characteristics of the 5a-reductase enzyme.

# Synthetic AAS Development

In order to develop products that would be effective therapeutically, chemists needed to solve a number of problems with using natural steroid hormones for treatment. For example, oral dosing was a problem, as our basis steroids testosterone, nandrolone, and dihydrotestosterone are ineffective when

administered this way. The fiver would efficiently break down their structure before reaching circulation, so some form of alteration was required in order for a tablet or capsale to be produced. Our natural seroid hormones also have very short tall-fives in the body, so when administered by njection, an extremely frequent and uncomfortable dosing schedule is required if as sealy blood level is to be achieved. Therefore, extending settered striptives are major goal for many chemists during the early years of synthetic AAS development. Scientists also focused on the nagging problems of possible excess extragmic buildup in the blood, particularly with testosterone, which can become your uncomfortable for negations and some products and the same of the produce of the produced produced to the produced pr

#### Methylated Compounds and Oral Dosing

Chemists realized that by replacing the hydrogen atom at the steroid's 17th alpha position with a carbon atom (a process referred to as alkylation), its structure would be notably resistant to breakdown by the liver. The carbon atom is typically added in the form of a methyl group (CH3), although we see oral steroids with an added ethyl

(C2HS) grouping as well. A steroid with this alteration is commonly described as a C-17 alpha algylated oral, although the terms of methylated or ethylated or alstrand are also used. The alkyl group cannot be removed metabolically, and therefore inhibits reduction of the steroid to is inactive 17-ketosteroid form by occupying one of the necessary carbon bonds. Before long, pharmaceutical companies had utilized this advance (and others) to manufacture an array of effective oral steroids including methylesisotatorus, Anavar. The principle drawbuck to these compounds is that they place a notable amount of stress on the liver, which in some instances can lead to actual damage to this organ.

Because the alkyl group cannot be removed, it mediates the action of the steroid in the body. Methylesotsortem, for example, is not simply an oral equivalent of testoterone, as the added alkylation changes the activity of this steroid considerably. One major change we see is an increased tendency for the steroid to produce estrogenic side effects, despite the fact that it actually lowers the ability of the hormone to interact with aromanses. <sup>45</sup> Appurently with 17-alkylation present on a steroid, aromatization (when possible) produces a more active form of extogen (typically 17-alpha-methyl or 17-alpha-echyl estradiol). These estrogens are more biologically active than estradiol due to their longer half-life and weaker tendency to bind with serim proteins. In some instances, 17-alpha-alphylation will also enhance the ability of the initial steroid compound to bind with and activate the estrogen or

progesierone receptor. Au emanicement or estrogenic properties is ano obvious when we look at methandrostenolone, which is an alkylated form of boldenone (Equipoise®), and Nilevar, which is an alkylated form of the mild anabolic nandrolone. Dianabol is clearly more estrogenic than Equipoise®, a drug not noted for producing strong side

Methyltestosterone (+ 17alpha methyl)

effects of this nature. The same holds true for the comparison of Nilevar to Deca-Durabolin, a compound that we also know to be extremely mild in this regard.

C17 alpha alkylation also typically lowers the affinity in which the steroid binds of the androgen receptor, as is noted with the weak relative binding affinity of such popular agents as Dianabol and Winstrol (stanozolo). However, since this alteration also greatly prolongs the half-life of a steroid, as well as increases the tendency for it to exist in an unbound state, it creates as more potent anabolic/androgenie agent in bort cases. This explains why Dianabol and stanozolol are notably effective in relatively lower weekly doses often the original control of the cont

In an attempt to solve the mentioned problems with liver toxicity we see with

#### Non-Alkylated Orals

c17-alpha alkylated compounds, a number of other orals with different chemical alterations (such as Primobolan®, Proviron®, AndriolE, and Anabolicum Vister) were created Primobolan® and Proviron® are alkylated at the one position (methyl), a trait which also slows ketosteroid reduction. Andriol® uses a 17beta carboxylic acid ester (used with injectable compounds, discussed below), however, here the oil-dissolved steroid is sealed in a capsule and is intended for oral administration. This is supposed to promote steroid absorption through intestinal lymphatic ducts, bypassing the first pass through the liver. In addition to 1 methylation, Primobolan® also utilizes a 17 beta ester (acetate) to further protect against reduction to inactive form (here there is no lymphatic system absorption). Anabolicum Vister uses 17beta enol ether linkage to protect the steroid, which is very similar to esterification as the ether breaks off to release a steroid base (boldenone in this case). While all of these types of compounds do not place the same stress on the liver, they are also much less resistant to breakdown than 17 alkylated orals, and are ultimately less active milligram for milligram.

#### Esters and Injectable Compounds

You may notice that many injectable steroids will list long clemical names like testosterone cylinotate and testosterone canditate, instead of just testosterone. In these cases, the cypionate and enanthate are esters (carboxpia cash) that have been attached to the 17-bet hydroxyl group of the testosterone molecule, which increase the active life span of the steroid preparation. Such alterators will reduce the steroid's level of water subshibly, and increase its oil solubility. Once an esterified compound has been injected, it will form a deposit in the muscle lessue (depty) from which a will solvely enter circulation. Generally the larger the ester chain, the more oil soluble the reference of the credit of the control of the control of the control of the control of the credit of the credit

There are a wide number of esters, which can provide varying release times, used in medicine today. To compare, an ester like decanonce can extent the release of active parent drug into the blood stream for three to four weeks, while it may only be extended for a few days with an actate or propionate ester. The use of an ester allows for a much less frequent injection schedule than if using a water-based (straight) testosterone, which is much more comfortable for the patient. We must remember when calculating dosages, that the ester is figured in the steroid's measured weight. 100 mg of testosterone enanthate, therefore, contains much less base hormone than 100 mg of a straight testosterone suspension (in this case it equals 72 mg of testosterone). In some instances, an ester may account for roughly 40% or more of the total steroid weight, tutt the typical measure is somewhere around 15% to 35%. Below are the free base equivalents for several popular steroid compounds.

100 mg of steroid as:	Approximate Free Equivalent:	
Trenbolone acetate	87 mg	
Testosterone propionate	83 mg	
Testosterone enanthate	72 mg	
Testosterone cypionate	70 mg	
Testosterone undecanoate	63 mg	
Nandrolone phenylpropionate	67 mg	
Nandrolone decanoate	64 mg	

It is also important to stress the fact that esters do not alter the activity of the parent steroid in any way. They work only to slow its release. It is quite common to hear people speak about the properties of different esters, almost as if they can magginal pater a steroid's effectiveness. This is really nonsense. Enanthate is not more powerful than cypionate (perhaps a few extra milligurans of testosterone released per injection, but nothing to note), nor is Sustanon some type of incredible testosterone blend. Personally, I have always considered Sustanon a very poor buy in the face of cheaper 250 mg enanthate ampules. Your muscle cells see only testosterone, ultimately there is no difference. Reports of varying levels of muscle gain, androgenic side effects, water retention, etc. are only issues of timing. Faster releasing testosterone esters will produce estroyen building faster trebassing testosterone esters will produce estroyen building faster trebassing testosterone tree in the blood from the start of the cycle. The same is true when we state that Durabolim's is a midler nandroboe for women

compared to Deca. It is simply easier to control the blood level with a faster acting drug. Were virilization symptoms to become apparent, hormone levels will drop much faster once we stop administration. This should not be confused with the notion that the anadrolone in Durabolin® acts differently in the body than that released from a shot of Deca-Durabolin®.

It is also worth noting that while the ester is typically hydrolyzed in general circulation, some will be hydrolyzed at the injection site where the steroid depot first contacts blood. This will cause a slightly higher concentration of both free steroid and ester in the muscle where the drug had been administered. On the plus side, this may equate to slightly better growth in this muscle, as more hormone is made available to nearby cells. Many bodybuilders have come to swear by the use of injection sites such as the deltoids, biceps, and triceps, truly believing better growth can be achieved if the steroid is injected directly into these muscles. The negative to this is that the ester itself may be irritating to the tissues at the site of injection once it is broken free. In some instances it can be so caustic that the muscle itself will become swollen and sore due to the presence of the ester, and the user may even suffer a lowgrade fever as the body fights off the irritant (the onset of such symptoms typically occurs 24-72 hours after injection). This effect is more common with small chain esters such as propionate and acetate, and can actually make a popular steroid such as Sustanon (which contains testosterone propionate) off-limits for some users who experience too much discomfort to justify using the drug. Longer chain esters such as decanoate and evojonate are typically much less irritating at the site of injection, and therefore are preferred by sensitive individuals.

# Anabolic/Androgenic Dissociation

Although never complete, scientists had some success in their quest to separate the androgenic and anabolic properties of testosterone. A number of synthetic anabolic steroids had been developed as a result, with many being notably weaker and stronger than our base androgen. In order to first assess the anabolic and androgenic potential of each newly developed steroid. scientists had generally used rats as a model. To judge androgenic potency the typical procedure involved the post-administration measure (% growth) of the seminal vesicles and ventral prostate. These two tissues will often respond unequally to a given steroid, however, so an average of the two figures is used. Anabolic activity was most commonly determined by measuring the growth of the levator ani, a sex organ (not skeletal) muscle. This tissue may not be the most ideal one to use though, as it contains more androgen receptor than most skeletal muscles (the AR is still less abundant here than in target tissues such as the ventral prostate). 47 48 In integrating both measures, the anabolic index is used, which relates the ratio of anabolic to androgenic response for a given steroid. An anabolic index greater than one indicates a higher tendency for anabolic effect, and therefore classifies the drug as an anabolic steroid. A measure lower than one in turn assesses the steroid as androgenic. There is some variance between experimental results and the actual real world experiences with humans, but (with a few exceptions) designations based on the anabolic index are generally accepted. Below are discussed a few factors that greatly affect anabolic/androgenic dissociation

#### Nandrolone and 19-norandrogens

The section of this book dealing with DHT conversion is important, because it helps us understand the anabolic steroid nandrolone and many of its derivatives. Nandrolone is identical to testosterone except it lacks a carbon

atom in the 19th position, hence its other given name 19-nortestosterone. Nandrolone is very interesting because it offers the greatest ratio of anabolic to androgenic effect of the three natural steroids (see: Sentheric: AAS Chemistra). This is because it is metholoized into a less potent structure (dihydronandrolone) in androgen target tissues with high concentrations of the 5-alpha reductions enzyme, which is the exact opposite of what happens with testosterone. Apparently the removal of the c4-5 double bond, which normally increases the androgen receptor briding capability of testosterone, causes an unusual lowering of this ability with nandrolone. Instead of becoming three to four times more potent, it becomes several times weaker. This is a very desirable trait if you want to target anabolic effects over androgenic. This characteristic also carries over to most synthetic stronds derived from nandrolone, making this an attractive base steroid to use in the synthesis of new, primarily anabole, scrends.

## 5-alpha Irreducible Steroids

When we look at the other mild anabolic steroids Primobolan®. Winstrol®. and Anavar, none of which are derived from nandrolone, we see another interesting commonality. These steroids are DHT derivatives that are unaffected by 5alpha-reductase, and therefore become neither weaker nor stronger in androgen responsive target tissues with high concentrations of this enzyme. In essence, they have a very balanced effect between muscle and androgen tissues, making them outwardly less androgenic than testosterone. This is why these steroids are technically classified as anabolics, and are undeniably less troublesome than many other steroids in terms of promoting androgenic side effects. However, if we wanted to look for the absolute least androgenic steroid, the title would still go to nandrolone (or perhaps one of its derivatives). Female bodybuilders should likewise take note that despite the recommendations of others, steroids like Anavar, Winstrol and Primo are not the least risky steroids to use. This is of great importance, as male sex hormones can produce many undesirable and permanent side effects when incorrectly taken by females (See: Side Effects, Virilization).

The 3-alpha hydroxysteroid dehydrogenase enzyme is also important, because it can work to reduce the anabolic potency of certain steroids considerably. As follows, not all potent binders of the androgen receptor are.

#### 3-alpha Hydroxysteroid Dehydrogenase

as a rule, great muscle-building drugs, and this enzyme is an important factor. Dihydrotestosterone is a clear example. Just as the body converts testosterone to DHT as a way to potentiate its action in certain tissues (skin, scalp, prostate, etc.), it also has ways of countering the strong activity of DHT, in other tissues where it is unneeded. This is accomplished by the rapid reduction of DHT to its inactive active metabolites, namely androstanediol, before it reaches the androsen recentor. This activity occurs via interaction with the 3-alpha hydroxysteroid dehydrogenase enzyme. This enzyme is present in high concentrations in certain tissues, including skeletal muscle, and DHT is much more open to alteration by it than other steroids that possess a c4-5 double-bond (like testosterone).49 This causes dihydrotestosterone to be an extremely poor anabolic, despite the fact that it actually exhibits a much higher affinity for the cellular androgen receptor than most other steroids. Were it able to reach the cellular androgen receptor without first being metabolized by 3a-HSD, it certainly would be a formidable muscle-building steroid. Unfortunately this is not the case. explaining why injectable dihydrotestosterone preparations (no longer commercially produced) were never favorite drugs among athletes looking to build mass. This trait is also shared by the currently popular oral androgen Drawings which is in account just an aral form of DUT (1 mathul PTOVIOUS, WHICH IS, III ESSENCE, JUST AN OTHER OF LOTH (1-INCHIP) dihydrotestosterone to be specific) and known to be an extremely poor tissue builder.

#### Anabolics and Potency

One must remember that being classified as an anabolic just means that the steroid is more inclined to produce muscle growth than androgenic side effects. Since both effects are mediated through the same receptor, and growth is not produced by androgen receptor activation in muscle tissue alone (other CNS tissues, for example, are integral to this process as well). we find that a reduction in the androgenic activity of a compound will often coincide with a similar lowering of its muscle-building effectiveness. If we are just looking at overall muscle growth, androgenic steroids (usually potent due to their displaying a high affinity to bind with the androgen receptor in all tissues) are typically much more productive muscle-builders than anabolics. which usually bind with lower affinity in many tissues. In fact, with all of the analogues produced throughout the years, the base androgen testosterone is still considered to be one of the most effective bulking agents. The user must simply endure more side effects when acquiring his or her new muscle with this type of drug. Individuals wishing to avoid the stronger steroids will. therefore, make a trade-off, accepting less overall muscle gain in order to run a more comfortable cycle.

# RBA Assay:

Another way of evaluating the potential ratio of anabolic to androgenic activity is the practice of comparing the relative binding affinity (RBA) of various steroids for the androgen receptor in rat skeletal muscle versus prostate. When we look at the detailed study published in 1984, we see some recognizable (and expected) trends. Aside from dihydrotestosterone and Proviron® (mesterolone), which undergo rapid enzymatic reduction in muscle tissue to inactive metabolites, the remaining anabolic/androgenic steroids seem to bind with near equal affinity to receptors in both tissues. They seem to be relatively "balanced" in effect. This study also discusses the unique activity of testosterone and nandrolone compounds, which are good substrates for the 5a-reductase enzyme found in androgen target tissues (such as the prostate), and seem to provide the most notable variance between anabolic and androgenic effect in humans due to this local metabolism. When it comes to real-world use in humans, anabolic steroids do not always behave in 100% uniformity with their anabolic and androgenic profiles as determined by animal models, so all such figures need to be taken with a small grain of calt

Compound	Human SHBG	Rabbit Muscle	Rat Muscle	Rat Prostate	Ratio M vs. P
methyltrienolone	<.01	1	1	1	1
dihydrotestosterone	1	.07	<.01	.46	.03
mesterolone	4.4	.21	.08	.25	.32
testosterone	.19	.07	.23	.15	1.53
nandrolone	.01	.20	.24	.60	.4
methyltestosterone	.05	.1	.11	.13	.85
methenolone	.03	.09	.24	.14	1.67
stanozolol	.01	.03	.02	.03	.6
methandrostenolone	.02	.02	.02	.03	.75
fluoxymesterone	<.01	.02	.01	.02	.77
oxymetholone	<.01	<.01	<.01	<.01	1.54
ethylestrenol	<.01	.01	<.01	<.01	2

RBA of various anabolic/androgenic steroids as competitors for human SIBG binding of pHT, and for receptor binding of methyltrienolog of methyltrienolog of methyltrienology of methyltrienology in Endocration (pt 14) and prostate. Source: Endocration(pt) 14(6):210-60. 1984 June, "Relative Binding Affilial", Anabolic-Androgenic Steroids...", Saartok T; Dahlberg E; Gustafsson JA, 2. Allo Section II. Hedits SAID 11/1/1/10 1987 PM Perc 1981.

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46. Unique steroid congeners for receptor studies. Ojasoo T, Raynaud J. Cancer Research 38 (1978) 4186-98

<sup>47</sup> Cytosolic androgen receptor in regenerating rat levator ani muscle. Max S.R. Mufti S, Carlson B.M. J Biochem 200 (1981) 77

<sup>48</sup>. In vitro binding and metabolism of androgens in various organs: a comparative study. Kreig M., Voigt K.D. J Steroid Biochem 7 (1976) 1005

<sup>49</sup> Androgen concentrations in sexual and non-sexual skin as well as striated muscle in man. Deslypere J.P., Sayed A., Verdonck L., Vermeulen A. J Steroid Biochem 13 (1980) 1455-

# Synthetic AAS Chemistry

Steran Nucleus (All natural and synthetic AAS hormones share this base structure)

Testosterone

5-alpha reductase.

Dihydrotestosterone

All analoxicaltoguics acroicks are preparations containing one of the above three stands about hormouse, or chemically alread desirables thereof in correga new synthetic compounds, one of the three minnel hormouse is selected as a starting post, spitsally due to the possession of particular trains that may be hearfulfial for the new corporant. For instance, of the three minnel steends above, disploits estimates in the only steened decoded of the possession of particular and 5-bella treation. It was likewise a very people close in probability of accommission and 5-bella reduction. It was likewise a very people close in analogue; to analogue and the particular analogue and the selection, due to its washening upon internation with the 5-bella reduction engine. Nandrobne was upstably used when over lower analogue; action is desired, due to its washening upon internation with the 5-bella reduction engine. Nandrobne show anomalies much more showly than toxication of the selection of

# Testosterone derivatives

#### Roldenone

Bolkimone is testosterone with an added double-bond between earbon atoms one and two. However, this bond changes the activity of the steed considerable, First, it dumnifically shows automationis, such that bolkemone converts to estandio at about half the rate of testosterone. Secondly, this bond causes the steed to be a very poor substante for the S-alpha reduced contractive engine. The more active S-alpha reduced metabolic Salpha-dephtedelerone spreaded only in very small marrants in harman. The harmone resident contractive and the steed of the net steed of the steed o

#### Methyltestosterone

This is the most basic derivative of testosterone, differing only by the added 17- alpha methylation that makes the steroid only active. Conversion to 17-alpha methylstradiol makes this steroid extremely estrogenic, despite the fact that this alteration actually reduces interaction with the aromatisse enzyme.

#### Methandrostenolone

In muny regards, methanthoustroubers is very similar to boldenone, as it too exhibits reduced settengenic and modernic actived for the e1-2 double-bond However, this steroid does have a reputation of being semewhat estrongenic, owing to the fact that it corrects to a highly active form of extension (Trapharmethylesthoid) See: <u>Medhaplated Correctors</u> and Orial Diseas). Methandroutershoke is also much more does the militigation relingance, as the 17-sharmethylest open and the second origination of the control of the second origination of the control origination or the control origination of the control origination of the control origination of the control origination or the control origination of the control origination or the control origin

#### Fluoxymesterone

Flaxoymesterone is a c-17alpha alcylated oral derivative of testosterone. The 11-beta group functions to iribib aromatication, so there is no estrogen conversion at all with this steroid. It also works to lower the afflisty of this steroid toward restrictive serum binding proteins, increasing its relative activity. Introduction of fluorine at the 9-position also potentiates the section of this extra of the steroid potential.

# Nandrolone derivatives

#### Nomthandrolone

Nordmarkolone is simply anathone with an added 17-alpha ethyl group. This alternion is merely used with an abded 17-alpha ethyl group. This alternion is merely used with anathocian-disequine steroids, and is much more commonly found with synthetic estrogens and progenitis. Although 17-redshiptin in although 18-bit 17-ketostouir fordaction just a used an 17-redshiption, and therefore allow this steroid to exhibit a similarly high loved programs of the steroid of the steroid steroid in the steroid steroid of the steroid steroid of the steroid steroid of the steroid steroid steroid of the steroid stero

#### Ethylestrenol

Ellykstroud is an oral derivative of randrolone, very similar in structure to lete group, which is visit to arringon receptor bringing. As such, edlykstroud is possibly the weakest stavoid milligarm for milligarm ere sold commercially. Any activity this steroid does exhibit is Body from its conversion to norediradnose, which does sent to occur with some similar (expresserly the 3 oxygan group is metabloically added to this compound visition until commercial to the control of the co

#### Trenbolone

Although a derivative of mancholme, the two additional double-bonds present on teroboles make any similarities to its parent frommer externed difficult to see. First, the 5-10 bond sithibits aromatization. Nandolone is very slowly aromatized, however, some entorgan is sail produced from the storick. Not not solve retrobolen. The 11-25 bond additionally increases produced from the storick. Not so solve retrobolen. The 11-25 bond additionally increases the storic produced from the storic. Not solve the storic produced from the storic. Not solve the storic produced from the storic. Not solve the storic produced from the storic produced from

## Dihydrotestosterone derivatives

#### Mesterolone

Mesterobne is a potent orally active derivative of dihydrotestosterone. Similar to methenolone, it possesses a non-toxic 1-methyl group, which increases its resistance to hepatic breakdown. This alteration does not increase the stability of the 3-keto group however, and as such this steroid is a noor anabolic like its natert.

#### Droctanolone

Drostanolone is simply dihydrotestosterone with an added 2-methyl group. This addition greatly increases the stability of the 3-keto group, vital to androgen binding. As such, the activity of this steroid in muscle tissue is greatly enhanced (see: <u>Anabolic/Anabogonic</u> Dissociation).

## Oxymetholone

Oxymetholese is an orally active derivative of dispotentsotstorme. The 17-methyl group is well understood at this poirt as we have discussed it with mury stroids, however, the 2-hydroxymethylene group is not seen on any other commercial steroid. We do know that this group greatly entireness analysis postery by traversing the ashibity of the 3-bette group, and that the configuration of this substituent also appears to allow this steroid to brind and actionte the estoneen recently.

#### Stanovolol

Stanozolol is a potent anabolic steroid, owing to the fact that the 3-2 pyrazol group creates a stable configuration off the A-ring that allows for androgen receptor binding (this steroid is one of the few that does not possess an actual 3-keto group). As such, it is highly active in muscle tissue, unlike dishufcetes/sterone.

#### Methenolone

Methenolone also is a potent anabolic steroid, due to the fact that the c1-2 double bond increases the stability of the 3-keto group. The 1-methyl group works to increase its oral biovasiability, making methenolone (as methenolone accetae) one of the few orally active non-17-aloylated orals. The c1-2 bond may also help increase hepatic resistance (slightly) to 17-ketoteroid deachisation as well.

#### Ovandrolone

Oxandrolone is an orally active derivative of dilydrotestosterone, due to its 17-methylation. It also differs from DHT by the substitution of its 2-carbon molecule with oxygen. This is the only commercial steroid to carry this group, and further, the only to have a modification to the base carbon structure of the Steran nucleus. The 2-oxo group increases resistance of the 3-keto group to metabolsm considerably, multiple oxandrolone a notest ranabolic.

#### Steroid Nomenclature

Perhaps not obvious at first glance, there is a marning convention in place that was used to create identities for the various anabolic/androgenic steroid hormones. This typically involves

forming a root word to convey the structural base of the steroid, and signifying other unique structural characteristics by including appropriate prefixes or suffixes. Below, we will look at the common roots, prefixes, and suffixes used in steroid nomenclature, and identify them as they are used in the various commercial compound names. As you will see, the adoption of names like nandrolone, methandrostenolone, and ethylestrenol were not as arbitrary as one might imagine. This section is also helpful if you wish to understand the deeper chemical designations for the various substances that one might find in the medical literature, which involve the exclusive use of this terminology (such as is the representation of methandrostenolone as 17b-hydroxy-17a-methylandrosta-1, 4-dien-3-one).

Common prefixes and suffixes used in steroid naming:

Structural Property Prefix Suffix Carbonyl (C=O) oxo-: keto--one Hydroxyl hydroxy--ol

Double Bnd (C=C) meth- methyl-Methyl eth-: ethyl-

Common mote used in stemid naming

Ethyl

Androstane Base carbon structure of dihydrotestosterone (no double-bond) Base carbon structure of or similar to testosterone (one double-bond)

Androstene Androstadiene Base carbon structure similar to methandrostenolone (two double-bonds: di-ene)

-ene:-en

Incorporated Into Name As

METH ANDROSTEN OF ONE

BOL DEN ONE

ETHYL ESTREN OL

ME STER OL ONE

METH ANDIEN ONE

METH EN OL ONE

OX ANDR OL ONE

STANO ZOL OL

TREN BOL ONE

OXY METH OLONE

NANDROL ONE NOR ETH ANDR OL ONE

FLU OXY ME STER ONE

Base structure of nandrolone (19-norandrostene) and estrogen Estren: Estra also: Norandrostene

Common Commercial Compound Names:

Taken From

Roldonono [17b-ol. androstadiene, 3-one]

Ethylestrenol [17a ethyl estren 17h-ol]

Fluoxymesterone [9-fluoro, 11b-hydroxyl, 17a-methyl, testosterone, 3-one]

Mesterolone [1-methyl, dihydrotestosterone, 17b-ol, 3-one] Methandienone [17a-methyl, androstadiene, 3-one]

Methandrostenolone [17a-methyl androstadiene 17b-ol 3-one] Methenolone [1-methyl, c1-2 double bond (en), 17b-ol. 3-one]

Nandrolone Inorandrostene, 17b-ol, 3-one1 Norethandrolone [19-nor, 17a-ethyl, (nor)androstene, 17b-ol, 3-one]

Oxandrolone [2-oxy, androstane, 17b-ol, 2-one]

Oxymetholone [2-hydroxymethylene, 17a-Methyl, 17b-ol, 3-one]

Trenbolone

Stanozolol [Stanolone (androstanolone, DHT), 2-pyrazol, 17b-ol] [tri-en, 17b-ol, 3-one]

Clinical Applications

Anabolic/androgenic steroids are approved for sale by prescription in virtually every pharmaceutical market around the world. Having been applied for many decades to treat a variety of diseased states, today these drugs have a number of well-established medical uses. They have been used to treat most patient populations, including men and women of almost all ages, ranging from children to the elderly. In many instances anabolic/androgenic steroids have proven to be life saving medications, which is a fact easily overlooked with all of the discussion about steroid abuse. This section details some of the most common and accepted medical applications for anabolic/androgenic steroids.

#### Androgen Replacement Therapy/Hypogonadism

The most widely used medical application for anabolic/androgenic steroids in the world is that of androgen replacement therapy. Also referred to as Hormone Replacement Therapy (HRT) or Testosterone Replacement Therapy (TRT), this therapy involves the currelementation of the reimary male hormone testactorane to alleviate currentence of loss supportant or the primary time neutron encounter to mercure projectors or troformers levels (clinical referred to as lapportation); Patricar my be adolescert miles suffering from childhood hypogendism or a speciel, disorder that causes andoparister the properties of the properties of the properties of adult men over the age of the properties of the properties of the properties of adult men over the age of the properties of the properties be the properties of the propertie

The mast common complaints associated with two testasteneous in subfar non-include checked bilds, centrels dyndrates, loss of energy, descented strength and or enthrance, reduced ability to plys sports, mord fluctuations, rothcod height flower lossly reduced work performance, memory loss and muck loss. (When associated withing these sportness are collectively placed under the label of "mediepasses" in a clinical esting this sticned: is referred to as it incorred propagational flowed besistence loss below 250 gold, are usually regarded as clinically significant, atthough some physicians will use a lovel as low as 250 gold; as the tradevold for remail. Hypogramidium s, justificatings, skil widely under-diagnosed. Most physicians will also not recommend treatment for low testosteneous labes a protein's complaining about symposium (symposium at antique, and contributions).

Androgan replacement therapy effectively alleviates most symptoms of low testostrone levels. To begin with, mising testostrone levels above SNOBHELL (the very low out of the recent image) will one measure normal measure function and life of its men with dysfurctions recent in the contraction of th

theory also increases red blood cell concentration (toyan earying capacity), improving energy and serse of web-leng Henryal so apports the restricts (the body mes, and improve mancel strength and endeannee. Likelike strend also becomes replacement florage) may have breafts with regard to endorscatche disease risk. For example, studies tend to show hormore replacement as howing a positive effect on ensuringlish. This relaxion are reduction (Legod) choicated levels, corbined with no significant change in HDL (good) choicated levels,  $^{5/2}$  52 Technications anapharentian also reduces melaction do-sist, and improves instain sensitivity and glocenic control. These are important factors in metabolic syndrome, which respectively the control of the co

is in port health. One study commind the safety of IRIF in men aged 65 and older with infinited melbilg and various health conditions such as obesty, begreission, diebetes, or hypotripolium.<sup>35</sup> Each subject took a fraundormal testistence agd (10g/100mg) or placebotory of the control search place callescent-orbidal adviser ceiters. The was compared to only 9 in the placebo group, Another study with middle-aged hypogenoidal men found the testistence replacement therapy (because currently active place) which will be a subject to the control of the control

There are some concerns with initiating testosterone replacement therapy when the individual

There are other areas of concern with cludy patients. To begin with, isototenerse administration may increase prostets observe and PAA values. <sup>250</sup> While this does not appear to be of clinical significance with normal leadily patients, being prostate lepenturely, and prostate careare in a best similated by selectories. More with prototace carear, laip PAA values, or breast carear are generally not prescribed testosterous. Androgon separementation has abote mixed to sheep apone, which can interfere with the most readily (REM) place of sleep, <sup>600</sup> The staties have produced confucing data, however, and the notestial relativeship terms that solved committee the solved means the solved or made the solved means the

therapy has demonstrated regardice, positive, and neutral effects on cognitive fractioning in cludy range. <sup>2006</sup> Studies do suggest that the donce an distance the based of responsibility, the most positive effects need when the antisigns head reaches the risks outpressings, from the strengthy-slopingful. <sup>2006</sup> Belley trainers with preceding deficial in cognitive function should have their cognitive performance and blood hormone beech mentioned closely during bottoms represent the range.

#### Common Treatment Protocols:

Transform? Transformal application is the most commonly prescribed method for supplimenting testionscen in the litied Seless and Camada, and is generally the fist consort of therapy situated with antiquam replacement therapy patients. This method of long delivery offices a naturel or all advantages to the patient when compared to in periods. Search to transformal application is painties, patient complainer and confirst is increased in order to a fine-transformation of the patients. The search of the patients and patients are particularly and the patients are common protocol among hommor explacement decroes is to prescribe estosterous extent. The most common protocol among hommor explacement decroes is only not provide testionare used and one of or admissive transformal patient that is replaced daily. Note that the to metabolishim in the daily transformation of testionation extents for increase search mellipsochestoweme (IAHT) levels more profoundly than testionare neighbors. This may exceed mellipsochestoweme (IAHT) levels more profoundly than testionare neighbors. This may exceed mellipsochestoweme and indepress described and patients and patients are considered in the patients.

Injection: Testosterone enauthate and testosterone explorate are the most widely prescribed injectable testosterone drugs in the United States and Canada. In many other markets the blended ester products Sustanon 100 and Sustanon 250 are also commonly prescribed Injection of one of these testosterone ester products will provide the patient supplemental androgen levels for approximately 2 to 3 weeks after each application. The most common protocol among hormone replacement doctors is to administer 200 mg of testosterone enanthate or explorate once every 2 to 3 weeks. It is important to remember that testosterone esters will deliver varying levels of testosterone to the body on a day-to-day basis throughout each application window. Levels will be highest the first several days after injection, and will slowly decline to baseline over the following weeks. Physicians are usually encouraged to monitor their patients closely to ensure androgen supplementation is sustaining hormone levels within the normal range (and alleviating symptoms of hypogenadism) throughout the entire therapeutic period. The longer acting injectable testosterone preparation Nebido (testosterone undecanoate) is undergoing review in the U.S. and has already been approved in other markets. This drug requires only 4 to 5 injections per year for most natients.

Only Testostence undecanout (Antirol) is the only prescription medication that delivers testostence via and capuals. His medication is not approved for sise in the Linda States, but is a prescription drug in Canada and many other markets around the world. Pleast complience and cornelit are high with the fism of therapy, as there are to special rotations or requirements saids from taking a few capuals each day with meals. Only which the contraction of the contractio

#### Angioedema, Hereditary

Anabolic steroids are commonly prescribed for the treatment of heroidiary angioedems, a run ean potentially life-fluenteing identified roll the immune system. Heroidiary angioedem is caused by genetic mutations of blood clotting factors, characterized by a decrease in the level of infutioning of the protein CI elestrase iribblor. This protein cortos CI, which is a "complement system" protein that plays an important role in the control of inflammation. Symptoms of heroidiary angioedem in achieve in neutrinotis to fraigh swelling of the hands, ams, Isps., Isps. eyes, Iongae, or fireot. Sweling may also be noteed in the digistive tract, residing in also final carming, masses, or vorining, in the most serious cases, the patient may notice a swelling of the threat and a blockage of the airway passages, resulting in applyacition and salden death. Many statels occur without a specific trigger, allough stress, traums, sargary, and derial work are commonly associated with angioederm attacks.

One 1-71 jable adjusted antholic fauthogenic streads have been shown to be a useful form of procurent features, subhings complement system protein levels and reducing the frequency and severity of machine frequency and severity of severity for severity to 2 days, the severity to 2 days, the severy to 2 days, the severity of severity of severity to 2 days, the severity of severity to 2 days, the severity to 2 days, the severity of severity to 2 days, the severity of severity of severity to 2 days, the severity of se

# Anemia

As a class of drugs, arabolic introlegaris storoids stratute the synthesis of crystopocietis in the kidneys, a hormone that supports the manufacture of new red blood cells. By design making them of morphic therapeutic value for mening certain forms of arounds (desired making them of menghe therapeutic value for mening certain forms of arounds (desired charactered by sendificient red blood cell production). Forms of arounds (a desired charactered by sendificient red blood cell production) forms of arounds (a desired characteris the production) arounds a sendificient red blood cell production. The sending the sen

In the United States, both conyenthelone (Anadrud 50) and mandrolene decanote (Decandrolin) are approached by the FDA for the tenturent of severe norms. The galdelines for using conyenthelone with both male and femule arean: partients (children and adults) recommend a closure of 1-2 mylager perf. This would equate to a daily dosage of 7-15 mylager perf. This would equate to a daily dosage of 7-15 mylager perf. This would expent to a daily dosage of 7-15 mg for an individual weighing about 160 hs. Doses as high as 5 mylagely are sometimes excessively to achieve the desired therappeter response. The agadiness for mandrolund decanoties recommend a dosage of 50-100 mg per work for women and 100-200 mg per work 5 man. Called the 20 to 13 years of ago were economical a dosage of 25-50 mg works.

Inscent years, the advent of recordinate explicacycles in a prescription drug law changed the face of ments in tenurited considerably. When attributed margines stands still offer theraperits value been, and are sell mutested and sold to rend awarine patients, they are presently regarded as subject of fallback medications for use only when therapy with an exploragionia above has failed to achieve a desired response. The hemateria increase from anticohordronogenic actions is gueenable just production between the considerable margines are conferenced in the desired assess to similar the entropy on reducions to deed for the tenure in the best measures in similar telephonogens, especially in some and challents from any change the telephonogens actually, given that there are revoer and directly margin the telephonogens actually, given that there are revoer and directly margin the telephonogens to the challenge and the challenge of th

#### Breast Cancer

Anabolizandroganic steroids are sometimes prescribed to treat beast cancer in postmerapusal owners or permenguated somens who have had their outsire serround. These drugs are of value when the cancer is homeone responsive, which means that its growth can be affected (postisely or regulation) by hormonal manipulation. Analogass and estingers have opposing actions on harmone-responsive turnors, with estimates supporting the growth of Protect accord times and analogase inhibiting of. I'm expolarations of an exposure and the production of the product

in women, however, so therapy is usually initized with goat causion. An oral arriogen such as flavorymenterone is usually preferred to a slower setting injectable, set offers the contraction of the co

In nevery years the development of never and trave trapted anti-estroparic drags such as sective strongen recoper modulations (SRAs) and aromatine shiring drugs have almost completely eliminated the use of arthodoclarathogueis storoids for breast camer treatment. Medicative tensionare for breast camer color, usually consists of a SEMI the Nobolacut tensionality, which may be used with a strong aromatine inhibitor such as Avriatics tensionality, which may be used with a strong aromatine inhibitor such as Avriatics tensionality, which may be used with a strong aromatine inhibitor such as Avriatics tensionally are almost a strong aromatine inhibitor such as Avriatics to the contract of the strong of the strong of the strong are almost and are sometimes all applied. They are very mater agended as adjust or influed; material aromatic and aromatine such when the aromatic aromati

# Decreased Fibrinolytic Activity

Anabois steroids may be preserbed to treat conditions associated with decreased flaring-like action, Firebrushs is the process in which a blood congation, with the two oystems may be allowed to the process of the pro

# Infertility (Male)

In a small percentage of cases, ambeliciandrogenic steroids may be prescribed for the treatment of miss infiritify. When the case of infirtifijs is two permocentation due to Leytig-cell secretion deficiencies, an antiogan might be able to adictine the confision. In such cases the settered may necesse the speem coart, speem quelly and the fruxtoes excentration;<sup>17,17</sup> which can necesse the clunce of conception. The oral antiogan mestacine (Provious) is most commonly percented for five purpose, adheaply has not be consistent of the property of the state of the configuration of TNA approach for said in the United States. Note that ambedicandrogenic introduction of the configuration of the confision of the configuration of the configuration of the configurati

## Growth Failure

Anabolic steerisks may be prescribed to treat growth filter in children, both with and without growth formere deficiency. These agents have been about to have positive effects on both muscle and bottom muscle and bottom mansch and bottom manneds and the state of the transport of the stream of the prescribed and the state of the stream of the st

In the United States, oxandrolone is the anabolic steroid most widely prescribed for the treatment of growth faltare. It is usually given as a supportive medication, used to suggest the anabolic effects of human growth hormone therapy. The drug is typically taken for periods of 6.12 months at a time in an offset to accelerate the growth rate without

passes on the statement of the statement

# Libido (Female)

The steorid methylastosterone is approved for prescription sale in the United States and mmy other musckets to improve Brisks on formula meraposa pointers. Small doess of the drug are typically included in products that also supplement estrogers, the confriention aimed at treating the fill spectrum of meraposase symptoms, including reduced fermile Brisks. The dosage used is low compared to those of other clinical applications for methylastosterone, and will seasily amount to no more than 2mg per day.

In the United States, however, the use of an ambolic steroid such as mardroine decarroate for the direct treatment of outcopronesis is presently wiswed as controversal in spale of substantial chical data and lastory supporting the use of seroids for the suprose in the United States, many metical organizations had the opinion that the potential size effects of seroid through as two administration to justify of the borders' with composences. Not again is proceed for the purpose of th

# Osteoporosis

Anaboic stensies increase bone mineral density, and may be prescribed for the tentment of acceptance of the comment of the similation of new bone firminist, inhibition of bone recorption (breakdown), and orienteement of cakiera absorption. <sup>10,2</sup> These dangs have additionally been additionally been additionally been additionally been additionally been additionally been advisored been advisored to the advisor to reflect the open inscreed understoppeneds. <sup>10,2</sup> factored in positrocapsual women, and is usually fixed to the changes in hormoul clearisty that are need their in En disclored does core to a high degree in the clearly of host sees, however. Obserpoored can also be caused by the proleoged administration of the contraction of the contract

Nandrolone decentors is the anabols stood most commonly prescribed for the treatment of contexponsis. The ding rates to offer measurable bearding with quart to bread entire, and may reduce the Bedindor of bone fineture in patient, <sup>10, 11</sup> The dosage used to treat posttamentaries unsuming 5 mg one every 3 to 4 weeks, Abertee reactions are common with fleening, however, traking vitations syntems (houseness and bodyfacial hair growth)<sup>27</sup> and unfarronable administion is senar choistensic<sup>26</sup>. Thereing propages to be better teleminal in patients above the age of 65, who as a group seem to notice lower adverse effects. May patients are given a materioral electronic disease of 50 mg one above effects. May patients are given a materioral electronic disease of 50 mg one may last for one year or longer if mecssary. The long throught window is usually required in order to give the degroupought into the means of the control property of the size of the patients are desired.

# Turner's and Klinefelter's Syndrome

Anabolizabin/loganic stroids may be used to test certain geneic conditions, most commonly limes' syndrome in firms as at Ricefelor's syndrome in males. Both are chromosomal disordess clumactized by desitions from the normal XXXVP paring. They reach it farming other health issued showmalks in grooth secand development, and ongoing escual fractioning. Males with Kinefelor's syndrome are stroit, and typically have a rounder fless measured polystage. They also develop armal testics, ferricoverdation, and may suffer with generocensis in those patients the supplementation of testosteroes (in a circle facilities to the road for surbosome analossess of thermits). same assume to tuni teste for managem representation tention per continuous, and can tary delicities some of the issues with sexual finationing and hody composition. Fermiles will limited 's printerious will be of short stature, and develop other physical abnormalities including a broad closes, how having, however ears, and webbed note. Low does of a primarily ambiotic strood may be used in adolescent patients as an adjunct to growth homome therapy to support the linear growth rate. Ouractionies is the stroot more commonly used in the United States for this purpose, and has been clinically successful at increasing fluid highly them used of along of OCI. Impligged qu'ell.

# Weight Loss/Muscle Wasting

Anabols stroods may be administered for the treatment of chirally significant weight loss. Common causes include probaged conformed themps, estension surgar, choric infections or severe traums. In a garrant sense, these agents can be highly useful when a patient is subject to a large hospital step up-only of both erst, when normal dady matches strainfation is not present and a significant loss of trache mas is noticed. Severe have injuries associated with southerly matched to allowed the confidence of the service of the serv

The significant loss of lean hody mass can present its own or of health sines. Individual that not entonically underweight may suffer from low energy and a reduced some of welhous, and are at greater risk of metally. <sup>85</sup> Server weight loss during recovery from surgery or lines may also measurably delay or complicate the recovery place. <sup>86</sup> In the most server cases, an ability of the princet to metalla moscopial lean host primes can be the lay determining factor in recovery complete from the most case the ability of the princet metallic place of the control of the control of cases of the most of cases of the control of cases and the control of cases of the case of the

In the United States, countdolose is the apart most frequently prescribed for most kinds of chiralsy significant weight loss. The dosage used of the purpose is bytacilly 10 mg given twice per day (20 mg total), although lower doses may be given in some fermale, edderly, or puragar partiers in an effort to noted undestable androgonic side effects. The drug is supported to the control of the control although may be given for a longer denation of recognic, Grown that the support of constructing portion rembeloisms in anti-altered by vistality all ambiotic streads, many agents other then countrolesses are described by vistality all ambiotic streads, many agents other then countrolesses are described to the purpose. In many other regions, agents with a high ambiotic-countrolesses are destroy into our predoctionary used for this purpose.

Authoris servois may also be prescribed to text more severe cases of muscle wasting. This is a contined contentioned by strong conjugar protein catabolism, which means that muscle protein is a contined contentioned by strong conjugar protein catabolism, which means that muscle protein is being prodominarily broken down (as opposed to synthesized) in the body, and a progressive loss of weight, strength, and errorg is noticed. In a medical setting, severe march wasting is referred to as cacheia. Cacheias in sort associated with insufficient food instants (clicary misstantion), but has a readroide cause than cannet be absented with cacheias. He will be a subject to the cacheia that of the march search strength of the protein the cacheia. He will be a subject to organize cacheia that of which analoles statesids. He use of those drugs as supports thempy for cacher cacheia has not been well established, however, and currently the subject of organize investigation.

Nandolore decaroate, coardolore, and coymetholore have been the ambole sterois more commonly used in the U.S. to tear makes wasting specifially associated with ITU in infection. Although no specific FDA recommendations have been adopted, statics with mundolore decaroates have shown a dosage of 150 mg every 14 days to have a similar autholor beently and a significantly lower incidence of side effects, as of mg (18 II) of furnam growth hermore per days. If no 2003, opportable was the subject of secreosal places III claimed Intio for IIII of the Control of t usose uas are most contributy prescribed by physicans. in receix years, nowever, use discontinuance of randrolone decanoate on the pharmaceutical market and a perceived higher patient comfort profile in oxandrolone has made oxandrolone the preferred agent for HIV cachexis. The dosage of oxandrolone used may range from 20 mg to 80 mg ner day. The most consistent clinical benefits have been seen with a 40 mg and 80 mg daily dose 89

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#### Steroid Side Effects

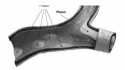
While anabolic/androgenic steroids (AAS) are generally regarded as therapeutic drugs with high safety their use can also be associated with a number of adverse cosmetic influsical and psychological effects. Many of these side effects are often apparent during therapeuticuse conditions, although their incidence tends to increase profoundly as the dosages reach supratherapeutic ranges. Virtually everyone that abuses anabolic/androgenic steroids for physique- or performance-enhancing purposes notices some form of adverse effects from their use. According to one study, the exact frequency of tangible side effects in a group of steroid abusers was 96.4%. This shows very clearly that it is far more rare to abuse these druss and not notice side effects than it is to endure them 90 In addition to the side effects that anabolic/androgenic steroids can have on various internal systems, there are others which may not be immediately apparent to the user. The following is a summary of the biological systems and reactions effected by AAS use

## Cardiovascular System

The use of anabolic/androsenic steroids in supratheraneutic (and often theraneutic) doses can have a number of adverse effects on the cardiovascular system. This may be noticed in several areas including unfavorable alterations in serum cholesterol, a thickening of ventricular walls, increased blood pressure, and changes in vascular reactivity. In an acute sense these drugs are admittedly very safe. The risk of an otherwise healthy nerson suffering a heart attack from an isolated steroid cycle is extremely remote. The risk of stroke is also extremely low. When these drugs are abused for long periods, however, their adverse effects on the cardiovascular system are given time to accumulate. An increased chance of early death due to heart attack or stroke is, likewise, a valid risk with long-term steroid abuse. In order to better understand this risk, we must look specifically at how anabolic/androgenic steroids affect the cardiovascular system in several key ways.

#### Cholesterol/Lipids

Anabolic/androgenic steroids use can adversely affect both HDL (good) and LDL (bad) cholesterol values. The ratio of HDL to LDL cholesterol fractions provides a rough snapshot of the ongoing disposition of plaque in the arteries, either favoring atherogenic or antiatherogenic actions. The general pattern seen during steroid use is a lowering of HDL concentrations, which is often combined with stable or increased LDL levels. Trickceride levels may also increase. The shift can be unfavorable in all directions. Note that in some cases, the total cholesterol count will not chance significantly. The total cholesterol level can therefore, give a false representation of uncompromised lipid health. Almost invariably the underlying HDL/LDL ratio will decrease. While this ratio should return to normal following the cessation of steroid intake, plaque deposits in the arteries are more permanent. If unfavorable shifts in lipids are exacerbated by the long-term use of steroidal compounds. significant damage to the cardiovascular system can result



# Over time, plaque deposits may begin to narrow and clog arteries.

Authorizant/ngmis stensis are mot consister in their lovering of HIL Levels. This adverse effect is medical frough the anticopies simulation of Papile, layes, a few cropus responsible for the breakdown of FIL (good) cholseston.<sup>24</sup> With more laptic lipuse activity in the Jody, the foundie (anti-sharpense) IELL cholsestory parties are calculaform creditation more quickly, and their levels dop. This is an effect that seems to be very section to the contract of week. The crossing this dosage to 600 mg did not have any significant additional effect, suggesting that the doage affectable for serious PILL suppression is fairly low.

Ond strowis, especially c-17 apile allylated composals, are particularly potent at strainting hearly inseem ad supersoing EHz, beach. Bit is due to fist pass concentration and metabolism in the bear. A drug like stancoloid may, therefore, be mike than tensistence with reagainst the mixed of the contraction of the co

It is also important to note that ensogens can have a forenshic impact on cholesteon profile. The amountation of institutements of the institutement of the institutement of the institutement of the cholesteon of the institutement of the institutement of the cholesteon of the institutement of the cholesteon of the institutement of t

The potential positive effect of entrogan on cholsterol values also miles et les issue of escongan ministernues consortieg to consider whether comes to health risk. To begain who, one may want to consider whether or not estrogan ministerance drugs are extually necessary in any gioner circumstance. An esist effects appearent, or is fast in ear potentation steps and perhaps suraccessary? The ministerance drug of choice can also have a measurable impact on choicaterol outcomes. For example, the estimator necepitor attaigness transition facilities can seem to exhibit anti-entrogenic effects on choicaterol outcome. The control is entropic to exist and in fact tends to in excuse IIII. Even is more guestre. Moris princhatal decide to use amounted to excell the just using activities of the large particularly of them is used activities.

#### Enlarged Heart

The human heart is a muscle. It possesses functional androgen receptors, and is growthresponsive to mis steared humaness. This fact partly accounts for men having a larger heart musc on average fairn worten." Physical ackey can had here a strang effect on the growth musc an average fairn worten. The properties of the properties of the internal cody. This is known as concernic remordering Funtamere (acorbe) subtless, on the other hand, tend to invense heart size via capanism of the internal cody, without significant thickning of the ventricles (occurrie remordering. Been with concernit or occurrie remordering, distolic function usually remains normal in the athlice leart. The heart muscle is also dynamic. When register taxing is removed from contificated affects, the will falcening and carly separation test.

Anabole stroul absence any at risk for thickening of the left and right vertricule valle, <sup>50</sup> allows an secretical replactople, <sup>51</sup> begrowing of the left vertrice (the min purpring churber) in particular is extensively documented in antibokindrogenic stroul absence, <sup>51</sup> while left vertricules physocropicly is, again, also foural in natural owner athleses, ashastence-dessing athleses level to larve a much more profused wall fischering. They also tend to while the contraction of the large transport of the large transport

Let werstein Ingenerophy (LVI) is an independent productor of mentalty in convenight institution with high body pressure. We Have a been benefit out in influention, vertically entitled an obligate and doub. We Will LVII in monetonic singuistice, producing all mental to be without clearly significance, bothological increases in CVI dispersion an ordered in sterrid absence with LVII. We have given gother times in OVI dispersion mental in Have cleanges and to be similar to the increases in OVI dispersion mental in Hyperical results with LVII. We arrange other times, this could have not be a similar to the increase in OVI dispersion most of in Hyperical results with LVII. We arrange other times, this could have no heart attack to bothed model case attacked or frequency around absence support an extent attack to bothed model case attacked or frequency around absence support an expertal results of the contribution of giving the let worksich, be streetly as the frequency for the contribution of the let worksich, and decreased ejection flaction (reduced pumping volume and efficiency). We

Heat mus can increase or decrease in relation to the current state of anabolishment use, the wavenge dossage, and duration of rinke Leibovie, the heat stands began to reduce in sice occus annobole intellegenic storoids are no longer being used. The effect is suffair to the way the heart will reduce in size occus an affect to larger brides a rigorost intellegenic schedule. The control of the properties of the control of the district adoption of the Ventricals hypertrophy road for all state several post with and a slightly control of the order of the control of the district adoption of the ventricals the properties of the control of the control of the district and the district and the district and the control of the district and the dis

#### Heart Muscle Damage

Anabolicalization steroid abuse is supected of producing direct damage to the heart mucke is now cases. Studies exposing beart of columbs to ASA have reported reduced contractic activity, increased cell flagility, and reduced cellular (instechnicals) activity, providing some support for a possible direct twic effect to the heart mucke.<sup>120</sup> The Furthermore, a number of case reports have found such pullobejus as reportantial flowesis (cent issue baddes) in the heart, mosecural inflammation (inflammation of heart issue), cardiac setatosis (accumulation of trajlectricks issue heart cells), and impossibility and clearly continued in the produced in the collection of the col (such as one), exercise, mestyre, and generous) can have on them, individuals remain caumoned about the possibility of cardiac muscle damage with long-term steroid abuse.

# Blood Pressure

Anabolican/longuist steroids may elevate blood pressure. Staties of bodybuldent taking these dugs in separatine-point does been demonstrated acrosses in both spools and distolic blood pressure readings. 173 Anathra study measured the strongs blood pressure reading in a good pointed users to be 140KK, which was compared to 12580 in weight lifting controls not taking steroids. 173 hypertension; or consistenty high blood pressure are reading in a group of the strong strong the strong strong strong the strong strong although in most case the elevations are more modest. Increased body pressure my be caused by a number of factors, richting increased water reteriors, increased vascular stiffness, and increased hermater. Aromating or high estrogenic stroids tend to case the greatest riduces over blood pressure, although elevations cannot be excluded with a strong anabolicative care stroids have been described as to remain con-

# Hematological (Blood Clotting)

Anabolization/organic atencids can cause a number of changes in the humstodigual system that affect blood clearing. This effect can be very vamile, however, the theraporic use of these draps is known to become phonin, auditorothin III, and protein's Social, stringial control of the control

Conversely, anabolicultogenic steroid abuse has been fished to increases in blood clotting ability. These drugs can elevate levels of thrombin<sup>107</sup> and Creactive protein; <sup>102</sup> as well as infunrificance AZ errore density, <sup>102</sup> which can sport platels aggregation and the formation of blood closs. Studies of steroid users have demonstrated statistically significant increases in platel aggregation values in some subjects. <sup>102</sup> There are also a growing number of case reports where (sometimes final) belond (close, embolism, and stakes have occurred in storial abuses; <sup>102</sup> 112<sup>23</sup> 102<sup>23</sup> Although in the board filled to conclusionly list free events directly to steroid abuse, <sup>102</sup> 112<sup>23</sup> 102<sup>23</sup> 102<sup>23</sup> 102<sup>23</sup> in the board filled to conclusionly since events directly to steroid abuse, <sup>102</sup> the adverse effects of anabolic steroids on comprosess of resulted as a complex for stories of the steroid abuse. <sup>102</sup> 102<sup>23</sup> 102<sup></sup>

In therapeutic levels, the anti-thrombic effects of anabolic/androgenic steroids seem to dominate physiology and decreases in blood clotting shifty may be noted. At a certain supratherapeutic dosage point, however, the pro-thrombic changes appear to overtake the anti-thrombic changes, and physiology begins to favor fast and abnormally thick clot formation (hypercoagulability). The exact dosage threshold or conditions required to increase blood clotting has not been determined and some studies with steroid users taking supraphysiological doses fail to demonstrate increased coagability 128 Individuals remain warned of the potential increases in thrombic risk with anabolic/androgenic steroid abuse. Blood clotting tendency should return to the protreated state after the discontinuance of anabolic/androgenic steroids, point until the hematocrit issues have been corrected. Minor elevations in hematocrit may be addressed with phlebotomy. For this, 1 pint of blood may be removed periodically during steroid intake, often every two months. Proper hydration is also important, as deliveration can temporarily cause the hematocrit level to elevate, giving a false positive for polycythemia. The daily intake of aspirin is also commonly advised if the hematocrit is above normal, as this will reduce platelet aggregation, or the tendency for platelets to stick together and form clots. Individuals remain cautioned of the potential cardiovascular danger of high hematocrit levels associated with anabolic/androgenic steroid

#### Hematological (Polycythemia)

Ansholic/androsenic sternids stimulate erythronoiesis (red blood cell moduction). One

potential adverse effect of this is polycythemis, or the overproduction of red blood cells. Polycythemic can be refixed in the humaners level; or the percentage of blood violate that blood because the first of the production o

Anabolicional administration tands to mise the Immeteral level by several percentage points, sometimes more, As a result, many steroid-using bodybalken will have humourt levels that are above the normal range. For example, one study measured the twenge hemmiter it as group of steroid abosing competitive bodybalkens to the 57%. This level is conclusion dismolarly legal and would recent bedow toward worship in rather that for the best for the form of the study of the stu

Many phosicians that specialities in hormous replacement therapy consider a heumotor's level of 55% to be an absoluce cutoff prior, it or above this prior, and ambiloishinologisis storoid therapy cannot be contrined safely. Dug irake would be consol at this point until the heumotor's issues have been converted, thin for electrical phosicistic grade with pilebotomy. For file, 1 pits of blood may be removed periodically during steroid irake, often cevery two mortles. Prour photagines is abin protective, as deductions on temporary cause the heumatoric level to desirate, giving a lake positive for polycethems. The day traited or flagarity is also commonly as from the most construction of the contribution of the contribut

#### Homocysteine

Arabide-indegarie stensis my eleste homosystein levels. Homosystein is an internediary amos and produced as a loopated or reflexion; remotions High best of homosystein how been listed to elestation in the risk for cardiovescular disease; <sup>133</sup> It is believed to play a direct note in the disease, investing outline stems, including the oxidation of LDL, clubsterred, and accelerating attronuctions; <sup>132</sup> Elevated best of homosystein may also induce second; red damage, support public arguagation and increases the fieldings of a first market and produced to the control of th

Andragans attendate elevation is hornecysten, <sup>127</sup> and men have an approximately 22% higher level on neurage frum worms, <sup>128</sup> Anthockindrogenic seriori abase has been associated with hyperbornecysteinemin, or consistent elizado, high hornecystein tesels, <sup>129</sup> Oce study found their hornecystein tesels, <sup>120</sup> Cole study found their hornecystein elevels, <sup>120</sup> Cole study found their hornecystein elevels have been self-administering authockindrogenic streods for a cyclic pattern for 20 years was 122 urnott, <sup>120</sup> Too of these ment deal of heart stacked during the investigation, and that homosystein levels between 15 stumell, and 18 starold. The average homocystein level in hophytalized with oth and never taken steasifie was 8.7 km/21., <sup>120</sup> with was 10 charmell; in previous senoid users (3 morths abstraces;) One study did show that administering 200 mg of testestatores cerentries eight and amorthuse arbitrately of the revenue for elevation in homocysteins. <sup>121</sup> It is tudowous if the moderate dosage, only pope (essertied injectateb w. vs. 127-20, or short duration of influence towards to the following control and the control of the hornecystein evel with storoid abuse.

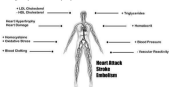
#### Vascular Reactivity

The endotelam is a layer of cells that fine the entire circulatory system. These cells are found on the saids of all blow cases, and the jurnesse or decrease both flow and pressure by relaving or constricting the usesle (referred to as uscedition and vasconstriction, respectively). These cells also play regalate the passage of materials in and out of blood usesle, and are invoked in a nurrier of important vascular processes including blood cleaning and new looks overall femalities and the control of the content of the content

Endobtelial cells are autoiogen responsies, which may purify account for mon exhibiting less weacter reactivity time worms <sup>16,5</sup> Similar, a mandociardinogenis restorate also has been shown to repair endobtelial exhibity and weacter reactivity. Statistics at the University of Imménute in Austria composed the tool of conductal difficient in 20 served uses to a group of control athletes. <sup>14,11</sup> Those infividuals using ambolic stored single for measurably impaired weacter distintion and endobtelial function. Auditional statists at the University of Walses in Castiff comparing vascular distints in active, provious, and mon-steroid users, also demonstrated ambolic stored to cause and define in endobtelial relations, described in demonstrated ambolic stored to cause and earlier in endobtelial relations, described demonstrated ambolic stored to cause and earlier in endobtelial relations, and mon-steroid users, also demonstrated ambolic stored to endobte and the stored and demonstrated ambolic stored to endobte and "the other stored and demonstrated ambolic stored to endobte and "the other stored "the other stored and "th

#### Proving an Association

Deet fais between steroid abuse and individual cases of stroke and heart attack have been difficat to prove. There are a narrier of lengs that how much field difficat let over, cutionscender diseases is very common in men. It also usually takes decades to develop. This musics individual contraling discrete forther darked may frings such soft, Heisely, health status, and guestic variables) entermely difficult to isolate. Data concerning the trape sermanse unterfaced to contract a contraded staty observed preface parts were general national constraints of statistical post of Statistic and the state of the state of statistic for musy years, so the data that is referenced tends to be from case stakes. Individual cases statistics are imported, but are usually considered to work to meet the requirements of statistical proof Stall, it would be a mistake to confuse this lack of proven association with proof of ramasseciation. The candiosocation randorsociation with proof of ramasseciation. The candiosocation randorsociation with groot of proven as the confusion of the state o



Anabolic/androgenic steroid abuse can produce changes in a number of areas of cardiovascular health that can work together to increase the risk of heart attack, stroke, or embolism.

Immune System

The hume immure system is responsive to see hormones. This results in functional differences in immurity between the case Norme need to have a more active immure system compared to men, and are slightly more resistant to bacterial infection and other types of relative, in "The few horms immure system as show more prose to developing autoimmus diseases, which may be infect to its higher level of faction," if "The days-to-day activity of the immuse system can also thetated froundpost the menstant clock, thatrifer demonstrating the strong influence of sex stronks. <sup>168</sup> The slightly weaker resistance to infection of man appears to be caused by teachmoore, which is an immunosperposes hormone. <sup>268</sup> Analogous my modulate the immuse system directly, drough their conversion to estrogats, <sup>150</sup> or by modifying glacocordical active, <sup>151</sup>

Anabolicardogusic steroids have displayed both immunosimatory and immunosimatory and immunosapprossic actions in animal models. Good nut three deep, can influence be immuno systemationally a variety of pathways, and anabolic steroids are a fairly diverse class of drags, their effects not immune systemation way and penalized not periodice conflicts. When used therapeatically, clamps in immuno system functioning are usually mirror, and love are married or immunosipations or immunosipations. Arabolic alambaging are married and in our immunosipations. Arabolic clambaging with music wasting associated with IIV affection, without any significant change in immunosystem or vial musics. <sup>2015</sup>

The use of antholic include gain and the situation of the

# Kidneys (Renal System)

Anabolic ándroganic stroids are ganerally well tolerated by the renal system. These drugs are largely exercated from the body through the kickeys, aftangel there is no inherent strong toxicity in this process. In fact, there are many instances in which those drugs mybe used as apportive tournary in painters with comprovinced islaving fraction. For example, ambolic stroids have been prescribed to increase the production of red blood cells in painters with meaning rather day to various forms of kidway (densee, <sup>26,15</sup>). Then we even been used as general anabolic (lean body muss) support, and to tear hypogenedator, in patients undergoing dalpois. <sup>26,15</sup> While icure must be kine with such patients, thenepy may often be conducted very safely. In otherwise healthy guidafulus, chical renal toxicity caused by the short-term admirastration of anabolic internals or according services is unlike, the contraction of anabolic internals or according services is unliked.

There have been isolated reports of severe kithey durings in stenial abusers. For example, a hortful of nikshids have developed When "immer (neighbothstams), <sup>1004</sup> but whit is a mer form of kithey cancer usually found in children in appearance in adult strond users is suspect, but not considered volume in children in appearance in adult strond users is suspect, in the considered volume in a strend abusers. <sup>102</sup> 100 Sixee this is the most contained form of the contained when the contained in the contained and results, was exhibitents (this duet obstruction) is known to cause easter tubular necrosis and read fidure. <sup>1064</sup>

The use of anabolic and openic steroids in supratherapeutic doses may slightly impair immune system functioning, reducing an individual's resistance to certain types of infection. In one study, steroid abusers were shown to have lower serum levels of [gG, [gM, and [gA, immunoglobulins (antibodies) compared to bodybuilding controls, consistent with

immunospression: "Albough this may logically increase the clurice of contracting certain types of fliens, Kuley health adoubt les concern for toge-men storia-sing bodybullers and power affaless. To begin with, excession resistance training can produce some strain on the real system. A confidence made that the contract contract the centre change of much tessus, which releases mogolish and a number of rephristoxic components into the blood." In high best this can change lacking best used used one seem fall failts. There have been rare case report of severe clinical flathelymolysis in bodybullers, both with and without meritin of stated about." Most least the case language the second table. "We are about cause hypertension, which can test to kindy admit about." We also also long-serving the computed as direct change via one of the contraction is commended to

# Liver (Hepatic System)

Many oral ambolicámicogonic secuolis (or ijectable forms of onal stavido) are toxic to her (hepatotoxic). These comproseds can cause serious and nomémias lideritoateira durings when abused, and occasionally even under demopratic conditions. These agents commonly associated with clinical hepatotoxicy heided (her aire not limited to) conventional conditions. The condition of the contractional conditions of the contractional conditions of the contractional contraction of the contractional contraction of the contraction of t

Adylation of c-17a/plat specifically protects the seriod molecule from metabolam by the engine 17bets—18p0, order control and the engine 17bets—18p0. This engine romally oxidates a steroid's 17bets-19ptonoyl (17bets-0) group, which must remain intact for the day to imput any anablest or anthogonic effect. Oxidation but 17-bets-0 is one of the primary gathough of rhepatic seriod docustoation. Without protection from this engine, very file active day and surveive that figure benegation from the control exception of the primary produced and the control of the primary produced and the control of the primary and produced produced and the control of the primary and primary and primary and process affect of the breakdown of 17bets-0 to 17-bets. The composal must be methodocal frough other pathough sea result, and immediate hepatic description is prevented. The process allows a very high preventage of the steroid does to poss into the bloodsteam intext, but 2 and publics season start on the few in the process.

To exact mechanism of hepathoxicity induced by adjudent antibodicandroguis: storoids remuns unknown, but it is a specialised to be due in large and to be intralla actively of antidogens in the foor. This foor processes is high concentration of antidogens receptors, and is displayed to the control of the c

Early her toxicy's is usually skills in blood test reads for hepatic function before physical symptoms or dysfarried nodevelp. This is most fleely to include elevations in aminotansferase enquires AST and ALT, also called serum glaturile-conducctic transmituse (GOZI) and serum glaturile protect transmituses (GOZI) and serum glaturile protection transmituses (GOZI) and serum glaturile protection glaturile protection glaturile protection glaturile protection glaturile protection glaturile produced as for the control facilities of the protection glaturile glaturi

assessment of liver and full-body health is advised should any signs of unacceptable liver toxicity become apparent.

The most common form of actual her dysfunction caused by the administration of oral ambifoliathrogues is entitle is closelastic. Fill the describes a condition where the flow of his becomes decreased, usually because of obstruction of the small his dues in the low fermitapacts.) The causes he shas and historian is accumate, in the ber and bood restand of being properly occurred through the digasther tract. Inflammation (hepatitis) may also be process.<sup>135</sup> Symptom of roblestasts impression accumate, transies, coming, upper abdomizal pain, or prarting (cheing). The stool may also change to a city color (alchelasisod) due to the reduced execution of his, and it is mire my become ambet. Choksation and the color of the contributions due to high book of blatters in the blood (hyperhistrication). Introduction, choicests in may also consists with hypercoloral records leaves described for fee rissoul.

Intraleptic clookstasis will usually resolve hard without serious injury or medical intervention within secural works of discontinginal plategatorics strotis. More resirva cases my take serious menths before normal laparic enzyme levels and fractioning are restored. Flequic leads are allegely to all one of time as well at lass partially, in some cases physicians because make layout the contraction and the contraction of the serious period of the serious shadows and the contraction of the serious deposition of the serious period of the serious p

More serious lequiic complications are rare, but have included peliosis lequiis <sup>100</sup> (bloodfilled opts on the lives), portal leportension with surfaced bloodings<sup>201</sup> (bleading caused by increased blood persease in portal via related to obstracted blood only, hepstochilar carectom <sup>201</sup> (mon-milgant her turny), lepotacchilar carectom <sup>201</sup> (milgant for turny), and hepstat englescention. <sup>201</sup> (milgant for turny), and hepstat engenescention <sup>201</sup> (superior legislate) and hepstat englescention. <sup>201</sup> (superior legislate) and wintout care are of these particulation are of these postations of the sease of the

#### Physical

#### Acne

Androgens stimulate the sobsectors glotals in the sike to secrete an objects structure called sestem which is made of this and the remunst of dead fiberpooking cells. Decess stimulation, as with steroid abuse, may also came a significant increase in the site of the sobsectors glotal, "Solvenerous glotals are found at the base of the in folkers in all ancontaining areas of the skin. If the analogue level becomes too high and the sobsecous glotals become convention, the high folkers who perhaps to all cap with sobsect more stated in the solvents of the state of the solvents of the solv

A mil nichtene of neur vägnir is usudy addessed with topical over-the-counter are medicinism and a sprose skin-clusing order that removes sees of and diff. More serbas acce may develop in sensitive infectious, richting neur coughdum (sevene acce acce). Sach richteness my cape medical intervention, which sught probes trustures with isotherism. Topical anti-arthogous rhaps are also under meetigation for the trustures of severe acce, and have shown a gust deal of persists in early with "A race is hypothia resolved with the cosmitt on of storid law, although the overproduction of selemanny pensis was a more of the companion of the companion of the cosmittee of the search of the cosmittee of the search of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the cosmittee of the cosmittee of the cosmittee of the search of the cosmittee of the search of t



Acne on the chest caused by steroid use. Hair Loss (Androgenetic Alonecia)

Anabolical charged and the contractive to a form of hir loss on the scale planous as antoquentic alspectic (AGA). This disorder is characterized by a progression reinstruction of hir folicles, and a shortening of the amages place of hir growth, under antropan inhance. The hir protocol by affected folicles will progressively this covering the scale loss and loss effectively in men, the habbases produced is usually identified most simply as made postern. How will straight patched a creating landing forther reposed thringing and the bolding usually takes on a more diffuse pattern, with thirting throughout the up of the head Most sources with antroprocie alspects do not have a recording larities.



#### With male AGA, hair loss is most pronounced on the temples and crown.

Andogarcia cispecia is the rust common form of last loss in men and women alike. It is especially common in miss, and more than 5% of the population with race it by the ago of \$0.500 As to must easignfie, androgarcic alspecia molose the interplay of Dord androgarcic hormons and aguacis factors. Individuals with this continion agent to be more levelly sensitive to androgars, and have higher levels of androgan receptor protein and displostessatemes in the scale, in comparison to those unfleed the Marbogan adorquatic alpects, it does not possess a surgue ability to infraence this confliction, and androgarcic alpects, it does not possess a target ability to infraence this confliction of an arbogance alpects, it does not possess a target ability to infraence this confliction of providing the receivery and regular situation. Bulbases can read for stress of the providing the receivery and regular situation. Bulbases can read for stress of the second and the s

in the absence of steroids that convert to, or are derived from, diffydrotestosterone.

The genetics of androgenetic alspecia are not fully understood. At one time it was believed this condition could be inherized solely from the mutural gandificher. More recent evidence contradicts this notion, however, showing strong support for fulther-to-son transmission in many cases. <sup>170</sup> A mather of garses have been identified as having a potential list, to the disorder, relading contain varieties (polymephisms) of the anthogan receptor gare. <sup>180</sup> 199 Anne 199

Restrict for androgenetic alopecia in men usually involves topical minoxidil and cral finisterici, a 5-alpha roductuse enzyme ribibitor. Women are typically prescribed auti-androgens and entogenepriosposits drugs. The Cours in both cases is on reducing relative androgenic action in the scalp, which may (at least temperatily) and the condition. With this in mind, many storiol uses concerned with this less white their relating interior unrecessary androgenic activity. This usually involves moderate design and the careful section of drugs with light antibodic consultings; into its contact account of the careful section of drugs with light antibodic consultages; into its contact as controlled excellent productions of the designation of the controlled with funited their controlled with funited their controlled with funited their controlled with funited their confidence in the stranges are until white variety decrees of species.

There has been no study on the role of genetics in haltenes indeed to stantid abuse. Annechotsky, findshows the sizing visible melangenetic algories appear to be flower most succeptible to the effects of annechotanthogonic stretchis on the scalp. For musy of those people, the loss of lartin appears significantly accelerated when taking these denges. On the other band, this side effect is generally a much less significant issue with individuals that have not racticed during the objectional. Must go on abuse seconds they near without any valid effect and annihing televation than the progression of indispractic alspects for with unbeautional than antihogon glay as in the in progression of indispractic alspects for with unbeaution. The indispractic alspects for the control in the control of the

## Stunted Growth

Anabolizatheogonic senois my sibili licar growth whon administred before physical matter). These hormones actually an laws or dichotomous induces on inne height. Once land, their anabolic effects my acrosse the relation of calcium in the boxes, feathfully control to the control of the contr

While andogous, estrogous, and gluccoerticoids all inherently participate in hore mutarly, estrogous in expande as the primary inhibited of fineer growth in both men and women. "So estrogous in expande of the men and women." So egg, due to the effects of file hormore. Anotheriodrogous stream, that either convert to estrogous on an inherently estrogous are, likewise, more likely to inhibit insurge proofs that men are inherently estrogous are, likewise, more likely to inhibit insurge proofs that the context in the context of the context in the context of the context in th

Estrogen acts directly on the epiphyseal growth plates to inhibit linear growth. These plates are located at the end of growing bones, and contain a collection of stem-like cells called chondrocytes. These cells proliferate and differentiate to form new bone cells, slowly expanding the length of the bones and the height of the individual. These cells have a finite life

span, with programmed sensecures (cell death). This will cause the rate of chearlangte profification to show over time, and executally stop. The chearlangtes are replaced with blood and hone cells at the point of physical matterly. "fasing" the bores and inhibiting further loaner growth. The stimulation of estimulating an openion to accelerate bore age advancement by exhausting the proliferative potential of chearlencytes at an earlier time.<sup>2014</sup>

Age will also influence a praint's sensitivity to ophybousd fastor. As young children are far from the point of bose martings, the influence effects of homome fenery take lenger to murfish any goods censition, As the journal ages, they may become more sensitive to these effects. Statistics tenting feneral poly (near goog and 14 years) for all status, for example, found that is mention of astroderone cuardinar (500 mg every two woods) was sufficient to render faith height y abunst three xinks companed to the predicted content. <sup>200</sup> This is a machemisty separate proposed to the predicted content. <sup>200</sup> This is an understand sympathrequent door, understang the fact that served intake during adobts converted to the predicted of the content of the prediction of the prediction of the protection for growth interruption when anabolicanthogonic sensition in two or historical decident materials of the protection for growth interruption when anabolicanthogonic sensitions to use of the protection for growth interruption when anabolicanthogonic sensitions to use the feneral for growth interruption when

# Tendon Injury

Anabolishndoganic steroid use is sometimes associated with an increased residence of tentucin qiigor.<sup>2023</sup> There are a few potential causes for this. One of the most basis is the mipd goods of the musicles. Under the influence of anabolishndoganic steroids, the muscles are capable of becoming againstantly pains and tronger in a fairly but replied of the actual contraction of weight lifted by the streoid user necesses, so too does the land placed on the tentucines. If the tentucines are not given enough time too for simply consensate for the actual contraction of the staroids may indirectly lead to the type of injury in some users, essentially by making them too strong for their own connectic tensus.

Theo may also be more direct mechanisms contributing to this type of pigor, Senite to selected market binse, the tendrous are required to androgous. Senite the stimulation of analysis starvists, for tendrous become more rigid. <sup>200</sup> As issue floating in extended, so in the relative sterupt and engine nature of the tendrous descentially lowering the tearing point; Furthermere, while analysis starvists are bount to all the hading process in the mucks, static suggest that they may stankly input the being of reduce pigns: <sup>200</sup> While this is admittedly a post-input year, and they may be a properly when the property when

The data concerning the potential role of anabole strouds in connective tissue ripey is equivocal. Some under fail for an association between AASs usen all termine ripey. <sup>111</sup> Overall, these types of ripers are sell fairly uncomment. This could make proving an association with fine the contraction of the contractio

#### Water and Salt Retention

Anabolizandosparie ateroida my increase the amount of water and socian stored in the body. This may include increases in soft the intranchiant and cuttoculars water comparturess, intranchiant flast offers to water that has been droom inside the cold. While the other compartures, intranchiant flast offers to water that has been droom inside the cold. While it is often circlated and wived set as part of load fift fee body may. Entruchialtr works is stored in the circulatory system, as well as in surious body issues, in the spaces between cold (tentration), Increases in intensit flad and in not reducible and rubbig construction) for storeg cases this can bring about a very path appearance to the body (perplaned or wishling of music features throughout the physics. Excess fills directions can also be associated with elevated blood pressure,  $^{212}$  which can increase cardiovascular and renal strain.

Estogon is negative of hal extercise in both men and somen. <sup>23</sup> The offer appears to be medicated in part by charges in hypothatics arguine suspenses (APV), the primary medicated part by charges in hypothatics and part of the part of

Estrogais ambidicinducquies extruits are generally focused for runs, gaining flushing purposes. A stead user may given water restrains during its place of fraining, cocasionally cern fating the shore increases in size to be a welcome benefit. Estrogaise stoods such as tectorises and comprehends are also regarded as the ortogate mass—and unsignly bealing quarter below the contract of the participation of the contract of the contract of the participation of the contract of th

Non-amounting strucks such as construction and stanoable have the born shown to premote increased under restitutes, so the field in not entirely contribute to amountainly or estinguist; substances, 277–288. Annabels seroods with low or no entrogenic action tend to produce modest tenses in which body water and intracellular distertions, however, and not in the wiside contracellular compartment? 270–270 These strucks are considered to be truce connectionly appealing and an generally forward by host-plackers and addies with no locally to improve tens must and market definition. Pepular anabolizationspace seroids that are associated with low widths water notations in raised. Biomagnetism, and market definition. Pepular anabolizationspace seroids that are searcied with low widths water notation in raised. Biomagnetism, and market definition.

Excess water reterior may be addressed with the use of arcillary medications such as the artis-entogen introoller chant or an arcentates ribbitor such as anastrools. By riminizing the activity of entogen, these drugs can effectly ordance be leed for such saint and artistic contract the contract of the

#### Virilization

## Physical (Male)

# Dysphonia (Vocal Changes)

Albudgh for less common than dysplonia in somen, ambolicanthogueis stroits may alter social physiology in men. The may include a deepening of the volte. Dysplonia in som common when ambolicanthogues; stroits are administered during adolescence, as the deeper add twice lies not say the one administered during adolescence, as the deeper add twice lies not say the considered under the influence of antiques. The prognessies lovering of the social pick, and may trigger pubscent vocal changes in sourger protects. Androuges have mean less (soft one instant) effect on our playshodge in adulthout Albudgh a slight lovering of the voice may be noticed with antiques use in some cases, spectra of facility algorithm of the major and the superior of clinically significant changes in the vocal quality of adult mare. Bloosive, very are. There has also been an isolated report of striker (whening uses weath sectioning) and vocal hosenesses in relation to ambolicanthogues stored abuse. <sup>237</sup> The instance also involved smoking, however, making the direct influence of steroids more difficult to discern. In general, vocal physiology is well established by adulthood. Aside from very minor reductions in pitch, anabolic/androgenic steroids are not expected to have strong audible effects on the voice.

# Gynecomastia

Anabolandroganic steroles with significant entropanic or progenitarial activity may cause gonocommia (femile besend development in miles). This disorder is specifically characterized by the growth of excess glardular tissue in men, due to an infralmence of multi-and femile see, hornors in the breast. Broops in the primary shorter of murmary glard growth, and acts upon reception in the breast to promote detail epitical hyperplasts, ductal elegation breast, and fleebales professionation. The dates growth growth and the order hand, which glardular those growth. The fleebales professionation. The dates growth grant hand growth and two entrops usually revert the convictional to the production of the professional dates and the state of the dates of the da

Opnocomista tends to develop in a series of progession estages. The security of his process will vary depending on the type and dose of drugs) used, and antificial sensitivity to hermones. The first sign is typically pain in the ripipel area (spencodyposa). This may quickly concile with mirror seveling around the ripipel area (spencody, but is constraints referred to as peach-genecoments, as a framinally movies fat and red glandshar fusious. At this stage, if a may be possible to address and ripipels woulding by reducing or entirating the offictually setting and the service of the service works. If the attentional, thousest, the may quelled progress to their genecoments, which easily left in the curly stages when pricting deeply around the ripipe. Noticeable excession of the service of the

Although genecomasis is a very common side effect of storid abuse, gione in clear suscicion with certain fungs or practices, it is also an easily rotable force. Cureful storid selection and rescondible dostig are usually regulated as the most loss can credible methods selection and rescondible dostig are usually regulated as the most loss can credible methods matternates medication, which may effectively counter the effects of elevated entogenicity. Common options include the artie-strongst memoriate circuits or amountses richitor was an austroade. The use of a post-cycle hormone recovery program at the conclusion of steriod administration (which usually includes secred weeks of anti-entogram use) is also commonly advised, as genecomossi is sometimes reported in the post-cycle hormone includes prises when stroids are not actually been tables.

It is important to note that progasterone can also augment the stimulatory effect of estrogen on unmurny tissue goods, 224 seek the progasterion datages may be able to integer the content of generousstic in sensitive articulates, even winder elevating levels of estrogen. Many anamolesk sterole, particularly those destroined from markotoxe, are known to exhibit strong progasticitized production and the content of the company of the content of the conten



## Early gynecomastia. Physical (Female)

#### Birth Defects

Ambolicandroganic steroid esposare to a worms during pregunary can cause developmental absorbanities in authoris flows. Violation of a final fieth sets a particular concern, and may include clinical hypotrophy or ver the growth of ambiguous gentalia (pseudohemptrodismi). Reconstructive suppry will be required to concert fishes services programs, should not use or directly landle ambolicandroganic steroid mentils (raw proude, pils, criente, patches). Although ambolicandroganic steroid mentils (raw proude, pils, criente, patches). Although ambolicandroganic steroid mentils (raw proude, pils, criente, patches). Although ambolicandroganic steroid mentils (raw proude, pils, criente, patches). Although ambolicandroganic steroid mentils (raw proude, pils, criente, patches). Although ambolicandroganic steroids are reduced to first patches are considered to the patches and the patches are considered to the patches are considered to

# Dysphonia (Vocal Changes)

Anabolicardogus stesuids are commody lisical to a despering of the voice in femiles. This is caused by direct and engine inflances on the layers and muscle lissues insoled as road physiology, which (in femiles) are not normally exposed to high antiogus lovels. Early changes may tecked a gift homering of the voice, with antible shifts in plot after high and love end of the vocal spectrum (quest speaking and voice projection).<sup>227</sup> There is typically a lover general frequency during speech; a nodection of high frequency pitch, and voice instability and enchang, in many cases the changes caused by AAS danger my resemble those of the projection of the contraction of the con

Desposing of the voice is defined as an anthogonic or musclaining effect. Annabolicalrudgumis extensis with higher relative anthogonicis what has testosterous, and methandrosterolore. Boowise, have a high tendency to produce voice changes when used in firmske, 227 282 273 281 annabolicalrudgumis stensis are capable of altering the fermile voice gionn the right level of stimulities, however. To this ports, world changes have been reported under theraposal conditions with even all anabolic substances such as counchoine and numbolics. <sup>237–247</sup> Case must be taken to moretic two circumstant of all ASS traikes, as changes are ofton unada generated. Immediately discontingual of ASS traikes, as changes are ofton unada generated. Immediately discontingual ene Body to presid. Anaboliciantlogenic steroid use may, Bewise, permunently aler vocal physiologics in fatures.

## Enlarged Clitoris (Clitoromegaly)

The mit and femile reproductive systems differentiate and develop under the influence of estiogen and testistorions. Even as an adult, the finale reproductive system remains developmentally responsive to mit sex hormous. An elevation of the antiogen level in sowmer may strainfast the growth of the clinical hypotrophyly, Thandagan levels are consistent of the productive systems of the clinical hypotrophyly. Thandagan levels are clinically absential enlargement of the clinical formation, by the clinical may be part to research the clinical productive systems of the clinic

Chromospaly is most commonly seen as a congenital disorder, although immy be caused by a mathodicathogonic second administration or other puthody in adulthood (acquired chromospaly). As a viniting side effect, elicotrongoly tends to occur in a dose-deportant cardiospacity-deported interact. As satt higher doses and more authogenic substances can be a supplementation of the contraction of th Claromegaly caused by steroid use is both avoidable and progressive. Mitigating excess androgenic action early when it is noticed is the most furdamental part of treatment. Reversal of significantly developed tissue, however, will require reconstructive suggest (elitoropistsy).<sup>233</sup> Special care should be taken to preserve the dorsal and ventral normonosculor huntles and normal tissue sensation.<sup>243</sup>



Aphotograph of distinct clitoromegaly. Here, the clitoris begins to resemble a penislike structure under androgen influence. If left unabated, this may progress to a more defined phallic abnormality. Source: Copcu et al. Reproductive Health 2004 1:4 doi:10.1186/1742-4755-1-4.

# Hair Growth (Hirsutism)

Autholicarhoganic streoids muy cause mule-pattern har growth in familes. Medically defined a hisration, flooration is distuncted by the gowth of hisr androgen sensitive areas of the body. With hisration, and course har (nemnal larir) may develop on the face, the characteristic and lack, areas of the body nemnly associated with hisration of the man, not women. Treatment for hisration typically involves immediate abstraces from anotheristic advantages (approximate abstraces from antibolicarhoganic streoid radiase, and infantion of a strategy to mininter androgen action at the hirr folkets. This may include the use of onal estrogens, anti-androgens (optionolactors), or funsients. Depatte indeconously, an artificial quarter, has also been used with some success. The response to medical intentions time between a days, and the charges caused by antibolicarhoganic serviculus our may preside for a year or longer. <sup>2018</sup> Jugglar hir returned for the affected causes may be execusely. The secretory of hisration will be reliable to the district of the daught of the daught olitect, the decoage and daration of use, and sensitivity of the

#### Menstrual Irregularities

Anabolicándroganis steniskis may aler the menstrual cycle in femiles, resulting in infraçuent or absort menses demonstrules, l'entire may also be intempted. Normal menstruation is expected to resume after anabolicándroganis stenisks are discontinued, and the inturnal bainers e restored. Complete recovery of the femile hormonal soit said fertility can take many morths is some cases, however, and long-term interruptions of fertility are possible, fusual trailies.

#### Reduced Breast Size

Anabolican/longonic atensis can rishis the growth- supporting effects of estrogan on memmry fossors, and may came a visible reduction in beast size (freest attempts). Androgan use in formules has specifically been shown to came a reduction in glandular tissue size, and to promote an increase in throse concredite issue. <sup>28</sup> These physiological changes are similar to those noted after mempanes, when formul see steeds are very low. Reductions in breast produced by AC may be very presisten after the discontinuaries for the contraction of the

# Psychological

The effects of numbok/undrugones's stroids on human psychology are complex, controvensia, and not fully undersciol. What is known for certain is faute sex transition stranses, more of and not fully undersciol. What is known for certain is faute sex transition stranses, anguessians, some of differences in the control and the control and the control and the control and the control levels, and Recoise, adverge hormone levels with the administration of coagrances stroid from yieldnesses between the coast struggle for this association, however, remains the subject of much research and speculation. In reviewing some of the more substantial than the last home the control and the contr

# Aggression

Men tend to be more aggression from vormen, a characteristic that has been partly ambitude to higher androgon street. The physiologists and hopothalmus, areas of the brain mobiled in human aggression. They also affect the orbitoformal cortex, a mean insolved with pingulac central. "Security absences cortexing and proportion of the property and property aggression and the competition of the property and the property and the competition of the property and property and proposed to the property and the property an

The psychological effects of exacting designs of tensorieme extens have been examined in a nature of pulse-or-controlled states. At thempeatic levels, no adheres psychological effects are apparent. If anything, instoatence replacement therapy tends to improve mode and sense of web-levels. When used at a contemption desage (200 mg prewed), again, not significant psychological effects are seen 200 mg by meds), psychological seed (exists such as aggression began to appear in some subjects, but these reports remain mild and influence? All a desage of the contemption of the contemp

One consiste placebo-controlled study furthers our understanding of the psychological effects of steard abuses, often characterised by extress does sent main-disag combinations, through its ecumunistics of a group of 160 regular users before and during the self-administration of a steard cycle. <sup>500</sup> Paloche group was and occurrent, which consisted of 80 people that were unknowingly shaing construct medications. Elemans psychological 100 people that were unknowingly shaing construct medications. Elemans psychological charges. Steard abuse, however, was associated with higher levels of hossility and HEMFQ measures, which principal receives a stearing on the contract of others, paramid baseling, galt, self-criticism, bluming of others, bluming of self, and oreal hossility SLO, 100 regions extend by loading you were also high retries greated used to obsess eccupations, irraprocuss assentially loading, pickle metric, and promote distance Healing measures review of the self-contraction of the process of the self-contraction of the process of the

#### Criminality and Violence

Lisks between multi-lic and regions: steroid abuse and violence have been much more difficult to establish. Most papers suggesting such an association of their used correlative data, or discussed infinitiate case statics. These help broaden the scope of research, but are not discussed infinitiate cases statics. These help broaden the scope of research, but are not reliable for proving causality. For example, one statyl questioned a group of 23 settorial abusing most, and reported that these men were involved in a significantly goaler transfer of revolut and even replaced. The state of the scope o

aumanturg AAS orags."— win the known enters of aumonearmorgane strouts on agessions, this finitely is contepting. It may very well be that some man are more susceptible to this type of behavior when abosing AAS than others. A paper like this is not sufficient, however, to substantiate a vokier! Total riggs. Further research is needed to determine if AAS can even trigger wheth relatives it am externely must intercept of users, and if isso, what really in these these talled data susceptible to the nearction when the van implicity of users are

Series criminally has also been difficult to succeits with steroid abuse. When discussed, we again tend to see wick correlative data and case statistic. For example, one page in Sweden reports an association between steroid abuse and weapons and final criminal. On six succeitation between fateroid abuse are satural seponable fit the criminally or page associated with it. It is simply possible these men were more exposed to, or more fleely to use, flegal.AAS is some underfitted reasons. Another page discussed three artificultum with no prior criminal or psychiatric labory that were amount for market or attempted market after abusing ambolic dark abusing ambolic dark migration. While shortes the traces are intensiting and reasons, with militarios of started uses in the agreement psyndiatric they are for from a design of the contraction of

#### Dependency/Addiction

Anabolicalization genie attentisk are considered to be drags of abuse. Although there is no universally accepted definition for this, abuse is commonly described as the contrasted used of a substance in spike of adverse consequences. Given the regarder health consequences that are associated with superindersparited soles of AAAV dang, this classification is a difficult one to departe. Drags of abuse are very often also drags of dependency, which in this content describes an irreption failing to terror the use of a substance. Then the been a languageding defines over whether or not anabolic strontis also of the definition of strags of dependency, the contrast of the contrast

Physical dependency is usually regarded as the most services form of drug dependency, and advangable of the goars have very octure and notabling dependency in destination. Physical dependency is defined as the need to administer a substance in order for the body to furction normally. A physical dependency is usually demendered by drug destinance, and withframed symptoms if the drug is discontinual abruptit. The most well known examples of drugs of physical dependency are opinists and as morphise, hydrocockon, copcodes, and haveing objected appearance of the contraction of the

Anabolisminoganie steroid absoc codd be associated with many of the DSMV retaria necessary for adapsois of both postobolispical and physical ding dependency. For instance, it is not uncommon for someone to take the drugs in higher doses or for brage protect of time them they had stillage planted (crient) 31. Many absents as how as a dose to cost time them they had stillage planted (crient) 31. Many absents a blace as doses to experientment may provent the decision (crient) 25. Indichallo done cortians to absent strongly in the control of the control of the control of the control of the strongly and the control of the control of the control of the control strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of the control of the control of the strongly decision of the control of

According to the American Psychiatric Association and its Diagnostic and Statistical Manual of Mental Disorders (DSMIV), three or more of the following criteria must be met for a diagnosis of psychoactive drug dependency.

# 1. Substance is taken in higher doses or for longer periods than intended.

2. Desire or unsuccessful efforts to cut down or control substance use.

 ${\it 3. Excessive time spent obtaining, using, or recovering from the substance.}$ 

4. Important activities are given up because of substance abuse.

5.Continued substance use despite negative psychological or physical consequences.

6. Tolerance, or the need for higher amounts of the substance to achieve desired

7. Withdrawal symptoms.

offect

A drug dependency that is isolated to criteria #1 to #5 would be described as psychological. The meeting of criteria #6 or #7 indicates the dependency is also a physical one.

The physical boeths of anabolicardanosis newide completes the matter of drug depositions; a grad that I little mercies is, but min notation bettle in use of steach is that positive effect on mach and performance. With this in mind, steached addition could actuably be a misdingsies for muck deposition; in many cases. This is a psychological disorder chruncterized by persistent feelings of physical inadeques; in spite of enterme mencaler development. Secreti abuse (often extensity is legal to great extensity and performance activation. Secreti abuse (often extensity is legal to see symptom of this disorder, not a case, in a similar sense, the physics, extensity, and performance—improving quadres of antibolic inadequation stream control and other control and the control of the c

Then is some evidence that the veidencing qualities of detential use po beyond an attackine to their physical bookeds. Lin arimals, what we nice and transters with represently self-destinate testosterors and other ambedicarthogonic steroids for example, an effect that carnot be caused by a perception of physical change. 272 Testosterors is also known to irenext with the mesorbrisc deparative system, which is common with other drugs of abuse. 252–258 Statics and additionally suggest its ambodicarthogonic streoids in these colorantee transporter density and increase sensitivity of the brain reward system. <sup>255</sup> Staticsh are known to influence physicalogic, and abuses commonly report an increased stems of wellness, while; and confidence when taking AAS drugs. Stems speculate this is that in part to an inherent near the confidence when taking AAS drugs. Stems speculate this is that in part to an inherent near the confidence where the confidence inhorted the confidence inhorted introduction desired.

Anabolicologogici sensitis are nei drago of marked inscientor. <sup>200</sup> which mulco them were different from other than and almos or deportency. The misse dispussing a being dependency difficult. By definition, drug dependency is related to the abuse of a psychocarie substance, and is surknown if AxS drugs can exceed by the basis of a psychocarie substances, and is professor frax fixed superactive by the basis of a psychocarie substances. At the present time, most expects do not regard anniholicathogogici sectionis a drugs of their psychoid dependency. It is difficult to correlate the post-tycle hormone irribatives with traditional switzbanel symptoms, and telesture is enably a faction of restributed interior natures growth, not necessaryly drawingly policyted effect. Individuals remain search, however, that sected those is commonly associated with and psychological marter of storeid abuse.

#### Depression/Suicide

Anabolican/toganic steroids abuse may be associated with bosts of depression. This is most commen after the administration of ASS drugs has been discontrated, especially following high doses or long cycles. During the time that steroids are being administrated, mutual lememe positionis in diministed because the obly recognises the excess hormous levels. When the steroid drugs are adruptly discordined, however, the body can enter a state of temporary highposendiss (tow anticogn levels). This may be associated with a number of psychological symptoms including depression, normai, and loss of frenest. This conciden is usually referred to as authols estand withdrasial depression, and can pessife the The most common method of addressing authoris seriod withfursul depression in most in promptisely, with the imprimentation of an agession protectly the turner recovery program. These programs are typically based on the corbined use of HEG Garma chrorine gausthoropies) and mis-etosporis (long such as transcribed and chropiteer. They are used tagether in a way fast can strainface and sensitive the hyporthanic pinkiny sorticular sessi, allowing mattale hormous production to return more against Auternative or concurrently, flavorite (or other artidepressors medications) may help also vite symptoms of depression following sector oil withrareal, expectly when this depression is prolonged or source.<sup>2017</sup> These drugs must be used with carrier, brower, as they also have been faired with increased thousant of socialies in some matters.<sup>2018</sup>

Albugh los corres depression is soutines expend dring the serice administration of methodischarging sendors. Hen mys to ensued by an institute of sea serial being nationally with regard to relative and equilibrity or enterprising. In more cases them not, it will mobe a shiften where sufficient androparity in our propert, usually when primarily anabolic days are being tolern above. Given the disease matter in which sex seconds interest with Interms prochools, bowever, it is district to cally ordine the parameters recessing by the large of depression to develop. Further containing the issue in the fact that this depression can mother direct cheated or suppressed better of certain sex seasons. The addition of the contraction of the series of the contraction of the problem in solution (and the contraction) and the contraction of the contract

Saide his been listed to ambeloistentid abuse in rure instance.<sup>200</sup> Such reports are usually case statis, so mobing infinituals have believed to be psychologically sable before abusing AAS, and who committed saide during or after use of the drugs, it is known that depression is a common complair during ambels setted withingers all its able known that a small percentage of users are especially sensitive to the psychological effects of ambeloistendogues is sensitive, and the committee mode swings, mine behavior, andre execution, such fact we straightly of users some complaint procession with their use. It is utknown why these radiodulas have such entermer execution, while the verst migricyl of users motice only mild or moderate changes to their psychological state. Further research is needed to sixtually and tradecisted design for a sixtle although its best analysis of the sixtual should be a small mentry of ourse. Beyond this, there is no compelling evidence suggesting that ambolociandogenic steroid abuse will lated to scale of otherwise mentally stable users.

#### Insomnia

Anabolicindogaric steroid use may be associated with insornia. This adverse reaction appears to be related to an irribative of hormous levels, and has been noticed during both coxecs and irresident hormousl states. For example, mornia is a common compliant arrange mus suffering from how androgen levels (hypograndisting <sup>20</sup> it is also insquestly exported by sixened bases that sign projects) explicitly consistent and abuses that sign projects of existing sixened, when collegation suffering needs are also how due to steroid -related suppression. <sup>20</sup> At the same time, this sixe effect is also seen of this gather altered as proposed in the sixenest sixeness of the sixeness of the

# Reproductive (Male)

# Infertility

Anaboló-ándrogeie sterolt use my impei ferilly. The harmn body strives to maixin balance in its ex-hormer levels (hormestasis). This balance is regalated largely by the hypothalmic-patienty-stericaler axis (HPTA), which is responsible for controlling the production of testestoree and speem. The administration of antholicanthogenic steroids provides additional sex steroid(s) to the body, which the hypothalmus can recognize as excess it researchs to this excess to reduction is sufficient suspect the reduction of rhistory. gonadotropris Interioring hormone (I/H) and follicle-stimulating hormone (FSH). I/H and FSH normally stimulate the release of testosterone by the testes (gorads), and also increase the quantity and quality of sperm. When I/H and FSH levels drop, testosterone levels, sperm concentratives, and sperm quality may alb revoluced.

When given in apprasphysiological levels, anabolicated agoing of encode commonly induce outgoingnosperms. Bit as fram of reduced firstly elementaries by braving less than 20 milition spormations per risel of spiculate. The quality of the spormany also be impraed under the inflance of ASA, as noted by an increase in the number of demonstrate projections (noticing reduced motion) sporm. Fertility is possible during oligonospermia, however, as within sporm concentrations are rountal in musy pass a pass agridiantally note than when sporm concentrations are rountal in musy pass as a consequent in the agendant ASA administration, which is defined as lawing no measurable sporm in the agendant fattilly has bown temporarily extended during active anabolicitarlogonic storoid abuse with the use of human cohortic accountation falled. The

ne see or intermetantus generosporturo, pro-Demisselo feritir ja considered a recreable side effect of ambole/anthogonic steroid abuse. Sperm concentrations usually return to normal within several mention of discontineing deng instack. A substantial post-cycle recovery programs based on the use of HEG, turnostics, and clerephase may againfamly shorten the refractory posted, and is highly pertically following long periods of lensy witered abuse, recovery of the HFA can be very protated, taking up to a year or longer for fall recoverys. <sup>207</sup> <sup>208</sup> (Geen the undesirable propriate altotal propriated principal suprement fact can be associated with a probategal state of postted propriate particular principal suprement fact can be associated with a probategal state of box totalement levels, such a long recovery visuolow is medy regarded as acceptable. The will usually prompt as individual to seed mediated intervention or interne an aggience HFFA.

The ability of anabolic/melogueis storoids to suppress IRI, FSRI, and fertilly his initiated a general deal of research structuring their sea mile contraceptions, his beine occursively statisfied by the World Hearth Organization, for example, and was determined to be a sink and moderated perfection enheal of an like bills control in statistic within administered 200 mg of fensionerse countriate per week to be bully men, accomperation which administered 200 mg of fensionerse countriate per week to be bully men, accomperation of the control o

#### Libido/Sexual Dysfunction

Ambelication/openie serosits may alter secund desire and functioning. The nature of their effects, however, on may depending on the draggl and dissoglisy loued, as well as the individual sensitivity of the user to hermonal manipulation. One of the most common responses is a stimulatory one. Testosterow is the primary male sestered As such, it is responsible for increasing secund desire and supporting many male reproductive-system functions. "Tile secund analock/ambengues instead to increases in sexual desire, as well as conjudiction and originary forestation, tractions in fine functions." The complexity of the desired to increases in sexual desire, as well as complaint and originary forestation frequency. 27th Augistrative increase in the forestation desired in increases in the desirence and may see the regarded as positive by the initialization meet left improved they are for its one.

AAS use may also be associated with diminished Bidds and sexual functioning. Bits could be due to several flactors. One is smillered analysiss action, 15 securities is not oblive the use of primarily analysis actions, 15 securities is not oblive the use of primarily analysis actions and the securities and the securities are securities. The securities are securities as the securities are securi

are also common after steroid discontinuance, when endogenous steroid levels are low.

# Priapism

In very mer instances, anthole/instanceparie strouths have been liked to prippinn \*\*275-96-77 The is a confinion functionariest of by the development of an exercise that will not not start to is flucid state within four house. Prippinn is a potentially very serious condition, which can require medical or assignal instructions. If the turneted, prippines made for medical permanent proprises and the propriate instances and the propriate proprises and the propriate proprises are described to stated use, Institutence is usually responsible. Furthermore, this condition appears to be more fequent in square partiests undergoing teatment for hypogenetics. The propriest teatment for hypogenetics in the contraction of the propriate teatment for hypogenetics. The contraction is not the propriate teatment for hypogenetics in a mile reproductive system that his not yet been expected to high tested of minimization.

# Prostate Cancer

Prostate cancer is dependent on anthogons. This disease will not develop if anthogons are claimated from the body at your gas (as with exactions.)<sup>278</sup> and adherent of anthogons are claimated form the body at your gas (as the interaction of an other particles are stated as a standard path of transmet. As complete printer of the involvement of anthogons, however, remains under. Studies show them as no association between the testosterous bed and Reifhood of developing prostate cancer <sup>279</sup> On the same note, the administration of ourgonizes testosterous during anthogons replacement funnys seems to laren on other on the risk for developing this disease. <sup>278</sup> A concer in strough absence. <sup>278</sup> I with the public contract of the risk of the complex gas the strong the concern in strough absence. <sup>278</sup> I with the public value of cases the level of anthogonize stimulation. The present model suggests that white testosterous is a necessary component of prostate cancer, it does not appear to be a desert tagger for its one.

New diagnoses of prostate cancer are sometimes reported during testosterore replacement therapy and steroid abuse, <sup>20,138</sup> such reports may be the result of a proviously unifagorosed condition or urelated development of this disease, with analous partition assisting the turner growth rate. Many forms of prostate cancer possess furthernal analousge receptors, and are highly antiseting responsels. As such that you most settlinded in grow unter the turner growth rate and the highly antiseting responsels. As such that you must be similated in grow unter the case usually contranslated in patients with a latery of prostate caree. <sup>2008</sup> While stread administration appears milely to came protect cancer, infished term is sound that the use of lessosterone or other AAS drags by someone with previously unfainguosed (latert) marketer prostate career and all constant for fiss disease. <sup>20</sup>

# Prostate Enlargement

Anabolicalundosparie struckle my influence the size of the prostate. Androgens are integral to the development of the prostate in early file, and are essential to materiating prostate structure and function throughout adulthood. Bursuses in the androgen level often result in growth stimulation to this gland (prostate lappertroply). For comparely, increased prostate volume has been reported in some patients receiving testestorize isjections for the tenturent of Dippogonational". While activating prostates phesprophy is not common under throughest condrines, prostate volume does tend to reach a size that is considered normal for a given androgen level. The Act postates e-postate regard levels have also near shown the activation and compared to the prostate that the considered normal for a given marker of prostate behalf often correlated with prostate volume. The Additionally, exclude stimulation of the prostate by lowering the androgen level tends to reduce prostate volume. See

Autholicarhoganic strend abuse may result in significant increases in protate volume. In more severe cases, them just due boing prostate hyportrophy (Reff). BPH is a common condition in older men, characterized by reduced urise flow, offfically or discornful training, and changes in training features. Jacobal reports of BPH armog serooti-single bodybuldess are not common, but do occur with enough frequency to warrant concern. Such reports are must often liked to more antiogueir, due system as testosterow and treuboluce, or the excessée doising of AAS in guernal. Due of the mast entiren reports of prostate hypertrophy came from De John Zeeder, the U.S. Obarquei phosician accordiale.

Russian weightflers were abusing so much isotoreone that they needed catherization to urinter. Distribed sear released soon after, which is structural to close derivative of isotoreone with relucated analogativity, which is structural to close derivative of Statistics of analociatorogetic servoid abusers show a preferential stimulation of the inner protests under the influence of those drugs, in an area where beings prostate Departurally is known to originate. The normal, protested current substitution proprieted areas of their glant. Some association between PBH and proteste current is known to ceals, however, although the coare time and strength of this association remain securities. "PSA values for although the coare interne and strength of this association remain securities." "PSA values for the proposed of the propriet of the propriets of th

trouble. It is important for men to monitor prostate health regularly by digital rectal examinations and blood testing for PSA levels. Anabolic/androgenic steroid use is generally imprediately discontinued if soms of PSPH or elevated PSA values become annuared:

with introducing Dianabol to sports, 293 Dr. Ziegler noted that during the mid-1950s, many

# Testicular Atrophy

Anabolisantrogarie stroids my produce atrophy (strikaga) of the testicks. Testostomes is synthesical and secreted by the Leight of the testics is chased by the hypothaline-pinting-testicular axis, a system that is very sensitive to sex stroids. When nabolic stroids are administered, the HPAT will recognite the electated hormous levels, and respond by reducing the synthesis of testosteroe. If the testes are not given ample straintation, over time they will atrophy, a process that can involve both a laws of restrictive volume and shape. This atrophy may or may not be obvious to the inhibital In some cases, the testes will appare around even though their functioning is smallicent. In other cases, shrikaga is very apparent. Visible testicals atrophy is one of the most common side effects of second lands and the stroid of the source of the source of second lands are some side of the source of second lands are properly in rever than 50% of all ambidesimaloganic stroid absence. 300

Albach neistude attophy is very common in foquators, it is also regarded an a temporary records like dieff. "The grands, by their nature, off usiny is see under hormonic affairers. Anophy should not produce perminent durings. Note, however, that it can be a somewhat practised must. In may take may work or mends of attendier. III distribution attophy is usually the not cause of producing post-cycle hypograndism in contense cases, fall recovery on their more than 12 rounds, and may even exquire medical steweriers. A post-cycle recovery program technice of IECG (which maries laterining hormone actively may be used to marine the recovery place." The date gas host departs of place for materiarie poticitair mass when used on a periods took during strond administration. <sup>36</sup> IECG (1811) and the contraction of the contraction

Some of the more potent ambolic marlogues steroids, including stostosteron, mardolous, tembrohous, and opportunidouse, appear to be more suppression of instosterous reclease than many other AAS drugs. This may be explained in part by the additional estrogues or progestational activity inherent in these seconds, as estrogues and prospects both also provide negative declaract highligan of testosterous relocación. See la supportant to marportant particular designations and the second seed of the second seed of the second second secondo marginal designations and the second second second second second secondo marginal secondo s oxandrolone, which are normally regarded as milder in this regard. While these compounds may be less inhibitive of testosterone synthesis under some therapeutic conditions, when taken in the supratherapeutic doses necessary for physique- or performance-orhancement, significant strophy and suppression are common, and districtions less pronounced.

#### Other

#### Anaphylactoid Reactions

An anaphylectoid reaction is a serious and posterially life frencisting allergic response to the administration of a foreign substance. Supprison of this dender relate keining of the side and eyes, swelling of the mexons membranes, here, lowered blood present, abbornial parts, vorning, and fitted blood vessels. He meroth mascle sury also be strainated to which is mutually assume that the property of the strainage of the strainage of the strain parts of the strain characteristics as anaphylatical reaction has many of the same characteristics as anaphylatical reaction has many of the strain characteristics as anaphylatical reaction has many of the strain characteristics as anaphylatical reactions, the strain characteristics are anaphylatical reactions, the strain characteristics are anaphylatical reactions, the strain characteristics are anaphylatical reactions. The strain characteristics are anaphylatical reactions and contains are analysis of the section remainstance may require by espectation and other supporters care. Given the potential for rapid escalation of the confidence and the strain parts and other supporters care. Given the potential for rapid escalation of the confidence and the strain parts and the str

#### Cancer, Brain

Anabolisation/agonic steends are not associated with brain career. Complications relating to a me and usually fast form of career cell primary central nervous system (remis) hyptopene caused the death of finness American foodhall paley Lijk Alzado. This type of brain cancer most commonly appears in imma-correporation algorists, such as those saffering from Harman Immandefectory Virus (HTV), or organ transplant recipients taking immansappressive taking timma-correporation of the death, Alzado had self-artituded its cancer to 14 years of antibolizardogenic storid abuse. <sup>250</sup> White antibolizardogenic servoids can be mildly immansappressive fise death, Alzado had self-artituded in career to 14 years of antibolizardogenic storid abuse. <sup>250</sup> White antibolizardogenic servoids can be mildly immansappressive fise death, alzado had responsable for Alzado's death. Even though phyloxicas say there is no proof of an association between performance-celanacing (ting abuse and Alzado's cancer, <sup>151</sup> this story is frequently recorned in the mells to cover which dearges of Storid abuse of the mells to cover who dearges of Storid abuse.

## Cancer, Breast

Albacja chromely mee, mile breast cancer has been associated with the administration of testatoriem. <sup>2011</sup> It is alwayon, knower, fifth hormous therapy was reduced to the onset of this disease, or if it was just reichtant to its progression and discovery. Androgen generally coldred inhibitory effects to hormous-responsible besteat cancer, and thus entails been used in their tentiment. <sup>2012</sup> Istrogens, on the other hand, can support the growth of rump breast turners. It is not transcription for electric disruption levels to result from testationer demay the contract of breast carcinorm and testastories administration in more remain unclear. Testosterone is presently contradictated in patients with breast cancer.

The data concerning the effects of artholocium/longuis storoids on issulfa sensitivity is difficult to interpret. It does appear that when these dangs are used rainling, relactions in body far are common, particularly visceral adjaces issue. This may actually improve insulin sensitivity and the overall metabols sites, and reduce certain specific risk forests for diabets and cardiovascular disease. Beyond this, the effects of AAS on glacose metabolism are not fully unartenood, and difficult to protect. Statulis sensing superthemportic does of testosterower and mundroken have fished to protect sensitive sing superthemportic does of testosterower and mundroken have fished to protect sensitive sing superthemportic body. At the same time, statics do suggest that these could be concerns with leavy steroid abuse. Further research is mended a season for improt of storoid abuse on metabolic health.

#### Inculin Resistance

Anabolicanhoganic stevids my aler isadin sensitivity, an important measure of metabolic health. The effect of frise chiga my be variable, however for example, instourem chamistation may improve isadin sensitivity in man with phogoanolism. <sup>340</sup> Orandrobse (Om aper due) has also been shown to improve isadin sensitivity of sold rem (eff. to 87 years). <sup>327</sup> Heads beneficial metabolic outcomes were correlated with reductions in viscoral adapone issue (VAI). This is also player of first automatic the adhormal corpus, and is succitated with insulin resistance. <sup>328</sup> braids resistance my also lead to other health issues including playersonic evoluted right-portions of coloration of the other insuling importance, evoluted right-portions of coloration of the other colorat

Conversely, anabolic/androgenic steroid abuse has been associated with impaired placose metabolism 317 In one study powerlifters that had abused AAS in high doses for up to seven wars were shown to have diminished obcose tolerance and increased insulin resistance 318 In spite of a long history of resistance exercise, these subjects secreted more insulin in response to measured placese investion than even obese sedentary control subjects Additional studies with methandrostenolone demonstrated significantly increased insulin secretion and notential resistance 319 A similar outcome is not found in all AAS studies however. For example, testosterone enauthate in doses as high as 600 mg ner week for 20 weeks failed to produce any changes in insulin sensitivity in healthy young men. 320 Nandrolone decanoste (300 me ner week) also did not inmair obcose tolerance, and actually improved insulin independent placese disposal 321. The data concerning the effects of anabolic/androgenic steroids on insulin sensitivity is difficult to interpret. It does appear that when these drups are used initially reductions in body fit are common particularly visceral adipose tissue. This may actually improve insulin sensitivity and the overall metabolic state. and reduce certain specific risk factors for diabetes and cardiovascular disease. Beyond this the effects of AAS on placose metabolism are not fully understood, and difficult to predict. Studies using supratherangutic doses of testosterone and randrolone have failed to produce any negative changes, suggesting that moderate AAS abuse is probably not associated with impairments in insulin sensitivity. At the same time, studies do suggest that there could be concerns with heavy steroid abuse. Further research is needed is assess the impact of steroid abuse on metabolic health

#### Nosebleeds

Anaboli-infloquie steroid se may be associated with periodic moshbook. According to one study, approximately 20% of filler storoid users protend the side effect, mixing it mix common. <sup>222</sup> Nosebbooks are not a direct result of androgenic action, but are secondary to sensioi-induced resources in blood persons and/or relations in blood chriting factors. Although they can be scary, most moshbooks are humiless, and will not require emergency medical attaction. When related to serout does, however, the pumy perfect of their more serious tarkingle health storos, particularly hypertension. Noveletech that occur under AAS of children for the principal control of the control of the control of the control of the children for the principal control of the c

#### Sleep Appea

Obstruction skeps apreas (OSA) is a disorder characterized by brief passes in breading dering skeps, with once with sent for since in the furent lose and book the ary possess. Skep apreas may interfie with rearnal gas exchange, and can significantly reduce the result of the control of the c

Anabolic/androgenic steroids may be associated with the development of obstructive sleep apnea in a small percentage of individuals. The exact relationship between AAS and OSA,

faction of the couplanyes is ways that can predippose an individual to the disorder. <sup>208</sup> More resourch is resulted to determine if sexual abuse can trigger (SA is an otherwise healthy person Individuals with a listary of obstrative skep agents should not use anotherinately super second. Hydrican sea achieved to remain the printers cheeply eaging of COA. Advange, ACA therapy, ACA therapy, CAA therapy, CA

noweer, returns uncer, into sovere reacon seems to appear in some pattern receiving testosterone drags to treat hypogoradism. <sup>336</sup> More detailed staties have shown that high does of testosterone can disrupt sleep and breathing, as well as increase sleep-related hypocornis, effects that may precipitate obstructive sleep apnea. <sup>270</sup> While OSA has not been electric document of in servid abservar anthorous how been down to after the structure and

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Acute Steroid Safety: Studies with Real-World Dosages

Few medicines have the type of sigms about them that anabolic-indrogenic steroids do. If you muritin the decision to use these drugs to the severage person, you are likely to be learned about the termedused polycal and psychological risks, you are about to undertake; how your have right life out, your testicks will disappear, or the steroids will give you cancer, how your have right life out, you testicks will disappear, or the steroids will give you cancer, the state of the state

The committed serviced near will usually point out the fact that a review of the medical licentum over the past 50 years will show that the overall solity profile of these drugs has been quite forwards. Stervid opposents, on the other hard, point out that an life it sent takes at any control of the control opposents, on the other hard, point out that an life it sent takes at the point of the control opposents of the control opposents of the control opposents of the sent hards? One thing that has always, conducted this debtar is the less of perinter medical statics. Medical ethics make high-dosed statics with anabolichardrogenic stronics (which may constitute abuses of the medication) we opificant to design and grapoural for Only very small number of chical statics sentally provide environments that could be viewed as relevant to those on books also of the argument.

In this section, we coarrise three medical studies that appear highly relevant for examining mela-word acute ambiochamlegenis stored safety. They concern not therapearic dooses, but a spranthempearic level and duration of finale that any flict steroid user would recognize as sufficient for approxing mache times, strength, and performance, in fact, the dosages and administration periods used in those studies reduce those taken by some of the more than the state of the s

serum cholesterol and triglyceride, prostate specific artigen (PSA) levels, and liver enzymes. Because of the protocols that were used, these studies give us a fairly good basis to evaluate the negative health impact of anabolic/androgenic steroids, at least as it relates to an isolated cycle.

#### 600 mg/wk of Testosterone

The first is a testosterone dose-response study published in the American Journal of Physiology Endocrinology and Metabolism in July of 2001, which looked at the effects of various doses of testosterone enanthate on body composition, muscle size, strength, power, sexual and countive functions, and various markers of health 329 61 normal men, ages 18-35, participated in this investigation. They were divided into five groups, with each receiving weekly injections of 25, 50, 125, 300, or 600 millionams for a period of 20 weeks. This treatment period was preceded by a control (no drug) period of 4 weeks, and followed by a recovery period of 16 weeks. Markers of strength and lean body mass gains were the greatest with larger doses of testosterone, with the 600 mg group gaining slightly over 17 pounds of fat-free mass on average over the 20 weeks of steroid therapy. There were no significant changes in prostate-specific antioen (PSA) liver enzymes (liver stress) sexual activity, or cognitive functioning at any dose. The only negative trait noted was a slight HDL (good) cholesterol reduction in all groups except those taking 25 mg. The worst reduction of 9 points was noted in the 600 mg group, which still averaged 34 points after 20 weeks of treatment. All arouns, except this one, remained in the normal reference range for males (40-59 noints)

# 600 mg/wk of Nandrolone

Next we look at a study conducted with HIV+ men, which charted the lean-mass-building effects of randrolone decanoate<sup>330</sup> 30 neonle participated in this investigation, with each given the same (high) weekly dose of this drug. Half underwent resistance training so that two groups (trained and untrained) were formed. The dosing schedule was quite formidable, beginning with 200 mg on the first week, 400 mg on the second, and 600 mg for the remaining 10 weeks of neak therany. Doses were slowly reduced from weeks 13 to 16 to withdraw patients slowly from the drug. Potential negative metabolic changes were looked at closely including cholesterol and limid levels (including subfractions of HDL and LDL). triglycerides, insulin sensitivity, and fasting glucose levels. Even with the high dosages used here, no negative changes were noted in total or LDL cholesterol, triglycerides, or insulin sensitivity. In fact, the group also undergoing resistance exercise noticed significant improvements in LDL particle size distribution, linoprotein(a) levels, and triglyceride values. which all indicate improved cardiovascular disease risk. Carbohydrate metabolism was also significantly improved in this group. The only negative impact noted during this study was a reduction in HDL (good) cholesterol values similar to that noted with the testosterone study with an 8-10 point reduction noted between both groups.

#### 100 mg/day of Anadrol

Lastly we find a study looking at the potent oral steroid oxymetholone (Anadrol) 331 This steroid is thought to be one of the most dangerous ones around by bodybuilders, who as a group seem to treat it with both a lot of respect and caution. It is not common to find them exceeding the doses and intake durations of this investigation, making it a very good representation of real-world Anadrol usage. This study involves 31 elderly men, between the ages of 65 and 80. The men were divided into three groups, with each taking 50 mg, 100 mg, or placebo daily for a 12-week period. Changes in lean body mass and strength were measured, as well as common markers of safety including total, LDL and HDL cholesterol levels, serum triglycerides, PSA (prostate-specific antigen), and liver enzymes. Muscle mass and strength gains were again relative to the dosage taken, with the end results being similar to those noted with 20 weeks of testosterone enanthate therapy at 125 mg or 300 mg per week (about 6.4 and 12 lb of lean body mass gained for the 50 mg and 100 mg doses respectively). There were no significant changes in PSA, total or LDL cholesterol values, or fasting triglycerides; however, there was a significant reduction in HDL cholesterol values (reduced 19 and 23 points for the 50 mg and 100 mg groups respectively). Liver enzymes (transaminases AST and ALT) increased only in the 100 mg group, but the changes were not dramatic, and were not accompanied by hepatic enlargement or the development of any serious liser condition

#### Adding It All Un

One handed and tweety-one men participated in these three staties, which moveded the use of moderate to high doese of statends for prices of there to the remains. Althought may be shocking to most opponents of antabolicanthogues served use, an utrissed assessment of many control of the state of the man reguler inputs of statend use in all there cases was a reduction in good (IELL), cholestent) values, which is a lagistimate concern when it comes to assessing one's risk for developing canditorous disease. It is uterating however if a short-field intense in the particular risk factor relation to any taughbe durings to come's health over the long-term it is that were access to concurrent committee, it does used a section of the state of th

Logic would seem to suggest that the isolated use of steroids, under parameters similar to those of rises three studies, should entail relatively similar discs to bellah. At the very less at, it is extensely difficult to sugge that an isolated cycle with a monetant day does in transmiss to plying Boston routice with your body; as not motion caregings against the consistent of the contract portain particles with the property of the contraction of

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# The Endocrinology of Muscle Growth

The road to arebole insight rant include a biological understanding of what muck growth actually catab. Often simplified by the term protein systessis," mucke growth is actually a lighty complex process insolving much more than just bailing proteins from anion action. March hypotrophy, the correct scientific terms for the value was dark tharman bail described by the contract contractions for mixed that human bail described fields a single process of the contraction of the contraction

Injury or trauma will provide the stimulus necessary to activate satellite cells. Once activated, they will begin to divide, multiply, and form into myoblasts (myoblasts are essentially donor cells that express myogenic genes). This stage of hypertrophy is often referred to as satellite cell proliferation. The munblests will then fise with existing muscle fibers, dorating their nuclei. This stage of the process is usually called differentiation. Skeletal muscle cells are multinucleated, which means they nossess many nuclei. Increasing the number of nuclei allows the cell to regulate more cytoplasm, which allows more actin and myosin, the two dominant contractile proteins in skeletal muscle, to be produced. This increases the overall cell size and protein content of the muscle cell. Incidentally, the number of nuclei in relation to cross-sectional area also helps to determine the fiber type of the cell namely slow twitch (aerobic) or fast twitch (argembic) 332,333. It is important to note that we are not increasing muscle cell number with muscle hypertrophy. We are only increasing cell size and protein content, even though we are using satellite cells to help accomplish this. It is possible for myoblasts to fuse together and actually form new muscle fibers. This is called muscle hyperplasia, and equates to the legitimate growth of new muscle tissue. This is, however, not the primary mechanism of muscle growth in adult life.

# The Anabolic Chain

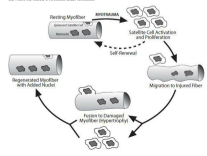
Now that we know what muscle hypotrophy is rolly about, let's look at anabolic stimulus and ongoing regulation. The following is a mindown of the chain of hormous and growth factors that mediate muscle growth, from the initiation of damage, to final recovery, requir, and growth. For the sales of organization, I have presented them in what I consider to be three logical phases of action. These are not scientifically accepted definitions. Additionally, we could continue to go deeper and deeper into each of the various composation, messengers, burding proteins, and receptors smoked in this stricture and amuzing biological accepts. I believe the stakeled next will encounted the process of interface and amuzing biological accepts. The level we have been a substantial to the stricture and amuzing biological accepts. The level we have been a substantial to the stricture and amuzing biological accepts. The level we have been a substantial to the stricture and amuzing biological accepts. The level we have been a substantial to the stricture and amuzing biological accepts to have been accepted to the stricture and amuzing biological accepted to the substantial to the stricture and amuzing biological accepted definitions. Additionally, we could continue the stricture and amuzing biological accepted to the stricture and amuzing biological accepted to the stricture and amuzing biological accepted definitions. Additionally, we could be form a substantial accepted and amuzing biological accepted definitions. Additionally, we could be substantial accepted definitions. Additionally, we are also accepted to the substantial accepted accepted and accepted accepted accepted and accepted accepted

#### Trioger

We all understand that weight training is fundamental to growing mache issue. To date, to "his on your sam and part and rapped [7] his bost mirected. The reason is at an anather of changes take place in your local mache issues during interne training that are vial to the ground process. With rates early othneys, growth is diffied in from prosoble to straining. So for our purposes, we will start here. Training is the "Faiger" in the analosic process. Now expecting the the localed collain framing that weight training produces that will first set to down the road of analosium. The body will respon by repairing this damage, and in process, with a step though Carriers with a size of the contraction of the contra

#### Phase I: Initial Response

The fairfi Response phase covers from changes in mack chemistry that begin immediately, during training, which will by the ground-took float repair and growth. In many regards, the latinal Responses Phase will control the potential magnitude of other signates to follow. In the manufock process, in phase is estagrated by the release of marketine and formance colk, and the formation of active messengers including prostagination, cytokane, subcontense, and prostopics. This begins will the breakdown of the outer pedapolity layer of mack leads, which is initiated by the cellular designation of duranging centrace. <sup>158</sup> Prosphilipsess are meteoded irresponse to this transar, which causes some of the phosphilipses arened in the control to the transars. When causes some of the phosphilipses arened in the control part of the transact cole, to the changed transport of the control part of the transaction of the phosphilipses arened in the control part of the transaction, explained in the "baguite" part of the transaction of the "baguite" part of the state of the control part of the transaction of the "baguite" part of the state of the part of the transaction of the "baguite" part of the state of the part of the transaction of the "baguite" part of the state of the part of the



# MUSCLE HYPERTROPHY AND THE 4 STAGES OF THE SATELLITE CELL

During the Activation stage, dormant satellite cells are stimulated to enter the cell yele, Proliferation marks the formation of new modulats (active donor cells). These modulats will fine with existing damaged made filters during the Officeration of passes. This allows for greater protein synthesis and the expansion statellite cells will again rest on the outer layer of the filters. Mystratin, a known inhibitor of mades or work, is believed to be a key remaining right scene, 203, 208.

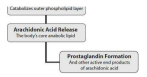
The amount of amelitotics each, which is the central bisocrito lipid in the ambolic process. Beneral will largely corrion what cours and reging hipstace. Ancholisis each is convented beaulty and amendately we enzymes to a number of active ambolic and profits, the most architect of which it eaters of intended profits and are produced with a produced with a most profit of the contraction of

Traing itemsy and the relative density of architotics acid in the phosphoight layer (unrachitotics and solubility is altimately the relat-limiting step in the formation of analysis conveniently and the procession of the formation of analysis of procession of the formation of analysis of the natural of analysis and step in the solution and the step in state of constant for the same of the step in the

# Phase II: Localized Tissue Priming

Plase II is chuncterized by a localized incruse in growth factor expression and tissue sensitivity to anabolic hormones. Those who have abuseys wondered by anabolic drags do not work without raining will find a good explanation right here. Streply part, your muscles need to be printed from actives of these explications of the property of the contraction of the contraction





Note: Inhibition of the cyclooxygenase-2 cuyne with anti-inflammatory drugs like lapprofice, acctaminophen, or a spirin, prevents the formation of active prostaglandins. The anabolic cascade is stalled without sufficient prostaglandin formation (Am.) Physical Endoctional Metab 282:ESF-6, interfering with the normal increase in protein synthesis rates after exercise. It is often advised to use such drugs only when necessary if maxed growth is a key focus.

To pit it is prospective, we need to remember that there are two separate components that intender before any message is set to an area cell onling it to increase growth. We have a hormous or growth factor on one hand, such as testosterous, KF-1, MKF, or issalis, and is conceptualing general or not hoot large-free coagonous number days, facilities general receptor briding and arabolic signaling by providing more messages hormous/growth factors (so host). The more hormous or growth factors (so host) are around the cell, the more horizing and activation of receptor site will take place. We cannot freely, however, that having more receptor size (nested of more horizing) and for facilities the process too. More receptors must be existing hormous or growth factors will find them factors that the contract of the contract of

Whit once hard we have an increase in tissue sensitisy to anabolic hormous and growth factors, also vid afform for Localced Tissue Printing place is an increase in the localcul expression of centin vial growth factors themselves. This includes [GF+], MGF, FGF, HFF, FNF, EL, 1 and LG. These composits will be relatered, and will work together on the coisting damaged music Bress and stellite cols, in a sort of grand symptomy of much anabosins with each place growth on the process. In surge cases, the action of one compound will apport the other, either by entiracting is levels, approxima prairities being proteins, one apporting is signify an intervolved mechanism. A dealed roadings that growther, one apporting is a signify an intervolved mechanism A dealed roadings are considered to science. A general overview of what is going on with each commonal fleet flowers; is noulded in our relevant Office.

#### Phase III: Repair

Voir beal muscle issues are privated during Plauses I and II. During Plause III, the hormones and growth factors go to work to firsh the job. We categorize the plause as one of congregio ambolic action, action mediated by the contributed effects of many ambolic hormones and growth factors including ambolings, reside, III-GP, 20, FF, GF, FF, TR, TL, II, II, and III-G. This is the time when repair and hopertrophy are physically taking place in your manckes, and each compound will play an intracte on in the process. We must not forget, however, that everyfung leading up to this point (the actions in Plause I & III) has still been determing how strong the growth everypeas will be via modifying receptor densities and common productions of possible and the plant of the plant of the composition of the ambolic common productions of the plant of the fraided with the lebs of the following hormones and growth follows.

#### Hepatocyte Growth Factor (HGF):

HGF is a heparinbinding growth factor that resides on the outer surface of uninjured cells.

cycle 139 HGF expression is regulated via ratic oxide release. <sup>340</sup> which is stimulated upon injury to also aid in the flow of nurieurs and hormones to the area. PGE2 plays a pivotal role in nitric oxide synthesis and HGF release. <sup>341</sup>

#### Androgens:

Androgens (the hormous that anabolic hardrogenic starolds minic) are strong supports or poten synthesis ratio in decletal macels tissue. They are also known to stimulate leveal IKF-1 expression, so the effects of these hormous estend to the satellite cell cycle (perhaps explaining why they are such strong similators of muscle growth, it is also for frost that sanchdroin acid increases androgen receptor density in skeletal muscle tissue. This helps to first price together the biochemical lists between the Plane I and Plane II response.

# Insulin-Like Growth Factor I (IGF-I):

IGF-1 is an insulin-like homome with marked anabolic effects. Owing to its rame, it also has some insulin-like effects as well. IGF-1 increases protein synthesis, and supports the profileration and differentiation of satellite cells. The prostaglandin PGP2alphra is known to strongly up-regulate local IGF-1 receptor expression <sup>342,343</sup> PGE2 is also believed to play a major increasing beat IGF-1 receptor expression <sup>342,343</sup> PGE2 is also believed to play a major increasing beat IGF-1 receptor expression <sup>342,343</sup> PGE2.

# Insulin-Like Growth Factor II (IGF-II):

IGF-II is a second insulin-like growth factor that plays a role in the proliferation of satellite cells. Unlike IGF-I, IGF-II expression does not appear to drastically increase in response to training. 345

#### Mechano-Growth Factor (MGF):

Mechano-Growth Factor is a recently discovered variant of Insufin-Like Growth Factor I. This growth factor is produced during an alternate splicing sequence of the IGF protein, and plays a strong role in the support of mylobite profileration. MGF expression, like many of the growth factors discussed here, is strongly up-negulated in muscle tissue in response to crotech strinks.

# Fibroblast Growth Factor (FGF):

FCF is actually a family of growth factors, with nine different informs (FGF-1 through FCF-). The full role that FCF plays in march hypercropsy in adulthout a not fail, understood, however, it is believed to be a storage profilerator of satellate cells, serving to expand fine-prophistion. FFF GFF imay also play a role in cell differentiation. As with mury growth factors, FGF expension up-regulation is proprieted in the dependent of fisse damage. FGFF and FGFF seem to be the most profile representatives of fits family in mature made tissue.

### Insulin:

In addition to having some ability to increase protein synthesis and rinbid protein breakdown, results in the body's clief rather transport hormore. The actions of resistin allow cells to transport placose and amino acids through the plasm membrane. Insulin receptor expression is strongly up-registed after transmite exercise, so as to provide more irreducine natrition to the affected area. This up-regulation has been closely linked to the prostaglandin PGEZ. <sup>340</sup>

# Cytokines (IL-1, IL-6, TNF):

Cybidens are a group of immemorbidatory compounds, dough in the context of this section we are loosely referring to them as growth factors. The IL cybidens are called interclusins, and ThN is short for lumn Necroisis Factor. Among other things, cybidens are classed in strainles the ringuistion of lymphocyles, neutrophis, memoryles, and other healing color is not element, to sail not element; to be plus in a number of other ways to, such as sating in the removal of demanged cells and regulating certain inflammatory responses, including the production of some prostalgarither. Prostagatifiers are known to ply important.

roles in the expression of all three of the cytokines mentioned here, <sup>351</sup> <sup>352</sup> however, they may not be the sole stimulus. Other pathways of arachidonic acid metabolism may also be involved.

#### Prostaglandins:

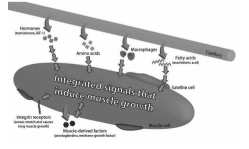
Although these are the key initial reactionary chemicals, prostaglandins continue to play a role throughout the muscle building process (including Phase III). This includes their support of hormone receptor profileration, the enhancement of protein synthesis rates, and an intensification of the purpose insulation of IGEL Links a charged authance (PIXX) 333.

#### Estrogens:

Although not specifically highlighted in this outline, estrogens also play a minor role in the anabolic process. This includes helping to increase androgen receptor density in certain tissues (though perhaps not skeletal muscle), stimulating the GH/IGF-1 axis, and enhancing educose utilization for tissue growth and renair.

#### Bringing it All Together

So that, in a very loose nutshell, is what is going on inside your body from the time you pick up a weight to the time your muscles are remained, stronger, and ready for more. If the above seems confusing to you it should. The fact is, the whole process of muscle growth has been confounding scientists for decades, and undoubtedly will for decades more. We still have a great way to go before being able to explain fully how it is that muscle hypertrophy occurs in humans. But as you can see, we have traveled a great distance as well. During the mid-1960s, scientists were only first learning that we grow muscle with the help of satellite cells More than forty years later we have identified, and are experimenting with, dozens of growth factors that were unheard of back then. It is a new world today, and despite not having all the answers, we know enough to enhance human performance in many exciting new ways. But please don't mistake the intention of this section. It is not here to give you a functional roadmap of the entire anabolic process, or to guide you in the ultimate polydrug program. It is here simply to open your mind to the true complexity of anabolism. When we start to see muscle growth from its various angles and intricacies, we begin to see our own potential opportunities for successful exploitation. How many of these opportunities you act upon will depend on your own goals and interests. But no matter how much or how little you actually apply this information. I hope you feel better equipped by having it.



### Skeletal muscle growth is a complex process that involves a variety of signaling compounds.

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# Practical Application

# Steroid Cycles

Anabolic/androgenic steroids are not medically approved to promote excessive muscle mass gains (bodybuilding) or improve athletic performance. Aside from early experimentation on athletes by a handful of sports physicians, an extensive effort to study the physique- and performance-enhancing properties of these drugs, specifically with an eve on developing strategies for using them to maximize benefits and minimize adverse effects, has not been undertaken by the medical community. Because of this, illicit users have been left to develop their own protocols for administering these drugs. The result has been a large variety of different approaches to using these agents, some safer or more effective than others. While it would not be possible to comprehensively evaluate all known approaches, this section will discuss some of the most fundamental and time-proven methods for using AAS.

#### Steroid Selection

When fast considering what steroids) to use, one will notice there are many different mendications that flat under the categooy of anabolicardoposite sections. This has been the result of many years of development, where specife pointers and needs are addressed with drugs that these specific characteristics. For exempts, some drugs are considered mider (less andesperis,), and produce fewer side effects in women and children. Others are more andesperis, which missed them better at superprint; secual functioning in term. Some are andesperis, which missed them better at superprint; secual functioning in term. Some are diversity, however. All AAS drugs archites the same collisher receptor, and as such slane similar protein anabolicity properties. In other words, while different AAS dang may have some differing properties. If your objective is to gain muscle mass and strength, this could be accomplicable with winthing arm our of the conternetily available again.

While all AXS drags may be capable of raproving macke mass, strength, and performance, it would not be correct to say there are no admittages to choosing on eager over another for a particular purpose. Most fartherestable, the quantity and quality of macke justed may be different forms one agent to another. In general nearch, e.d off and the contraction of the contraction

#### Mass (Bulking):

Methandrostenolone – Oral Oxymetholone – Oral

Testosterone (cypionate, enanthate) - Injectable

#### Lean Mass:

Boldenone undecylenate – Injectable Methenolone enanthate – Injectable Nandrolone decanoate – Injectable Oxandrolone – Oral Stanozolol – Oral

The enty stages of AsS are smally involve cycles with a single analysis indexed. Bading much most in the most correspond and smally death the use of our offer more advangers arbstimes such as testasteries; methandroselvene, or conventioner. These locking for hear mess often full from its analysis stagels are manifolous decisions, contribute, or statusoide First time users ranely selection rigogine armodoloic discount, contribute, or statusoide First time users ranely selection rigogine armodoloic discount, contribute, or statusoide First time users ranely selection rigogine armodoloic arbstage statusis, and will assult choose an out coreporal for he asset of concentrose. Methandrostouchous is the most common choice for must bading and is almost universally required as highly deficies and only methantly problematic for terms of estroggine or androgenic side effective). Statusoide is the enal articles statusid most often proferred for inproving luminus our adults performance.

The potential for advene reactions should also be considered when choosing a stemul to use, especially if ASR use is to be regainly repeated. For comptly, the listed oral medications proceed greater states on the conflorescelar system, and are also been took, for most of all, briesting countries size directs may also be lacken in second. For example, men with a strong sensitivity to genecomistic sometimes perfor non-entoqueit design such as methodox, temporal of controlled in the conflorest size of the controlled processing and the controlled process

#### Dosage

The dosage used is important in determining the level of benefit received.

Anabolic/androsenic steroids tend to be most efficient at promotine muscle sains when taken

at a moderately seguitheneperic douge level. Etchw this (thempositie,) posterial anti-bolic benefit are often controlledured, at least to some center, by the appression of exclusions associated and the state to some center, by the appression of exclusions associated and the state of the sta

- Boldenone undecylenate: 200-400 mg/wk
- Methandrostenolone: 10-30 mo/day
- Methenolone enanthate: 200-400 mg/wk
   Nandrolone decanoate: 200-400 mg/wk
- Oxandrolone: 10-30 mg/day
- Oxymetholone: 50-100 mg/day
   Stanozolol: 10-30 mg/day
- Stanozolot 10-30 mg/day - Stanozolot 10-30 mg/day
- Testosterone (cypionate, enanthate): 200-600 mg/wk

These use differend considerations other than the cost deficioness of a particular dosage. To be pain with judio sool anabolicamboguie stanside text produce stronger negative counters, psychological, and physical side effects. In light of diminishing estrum, the tradeoff between results and above nearcines because less and less forwords. Cains made on lower doses also text to be better retained after steroid discontinuance than those resulting from excessive statics. It is guerrally not retained to expect the region of the discontinuance than those resulting gains induced by mussive dosing will remain leng after a cycle is over. Shower steading usins are advised. It is also very important to remove them they have some at whosely with an ear advised. It is also very important to remove them they have consistent of the produced of the contraction of the cont

#### Dosage vs. Weight Increase

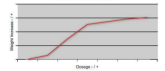


Figure 1.Anaholic/androgenic steroids tend to be most effective in moderately supratherapeutic doses. The anaholic benefits diminish in relation to the amount of drug given at both the high and low ends of the dosage range. Duration (Cycling)

The admissiration of ambolic/includencis servois at a given dosage will typically produce roticeable increases in mucle size and testing for approximately 6.8 works. After this point, the rate of new mucle gain typically slows significantly. A plateau may be reached soon after, where all forward memorations ascend. To cortinar mading significant proposes beyond this point can entail excluding dosages, which is likely to corticle with a gender reacher of adverse executions and direstifing ambolic returns. Even without dosage excludion, regarder balled directory and the contract of a strength of the contract of a strength of the contract of the con MANAGEMENT OF MANAGEMENT IN AN ADMINISTRATION OF MANAGEMENT AND AD than 8 weeks at a time (10-12 weeks at the maximum), followed by an equal or longer period of abstinence before another steroid regimen is initiated. This pattern of rotating between "on" and "off" periods is referred to as cycling

### Off-Cycle (Recovery, Bridging, and Tapering)

The period immediately following steroid cession can involve a state of hypogonadism (low androgen levels), and as a result protein catabolism. In an effort to minimize muscle loss, the objective here is usually on restoring natural testosterone production, maintaining an optimal level of muscle stimulation, and remaining dedicated to proper nutrition. A hormonal recovery program is usually initiated, which may involve the use of HCG, tamoxifen, and clominhene (see PCT Post Cycle Therapy). A substantial off-cycle period is also advised, involving abstinence from anabolic/androgenic steroids for at least 8-12 weeks. Some AAS abusers have difficulties with complete drug abstinence, and will initiate "bridging" routines between full-dose cycles. This may involve the periodic low-dose administration of an injectable steroid, such as 200 mg of testosterone enanthate or methenolone enanthate every 2-3 weeks. Such practice is discouraged, however, as it can interfere with hormonal recovery. and prevent a return to metabolic homeostasis.

When concluding a cycle, some steroid users also follow a practice of first slowly reducing their dosages (tapering). This tapering may proceed for a 3-4 week period, and will involve an even sterning down of the dose each week until the point of drug discontinuance. It is unknown, however, if such tapering offers any tangible value. This practice has never been evaluated in a clinical setting and is not widely recommended with steroid medications as it is with some other drugs such as thyroid hormones or antidepressants. Virtually every highdose AAS administration study can also be found to end at the maximum dosage, with no time allotted to tapering. One flaw in the logic of using a tapering program is that they are ostensibly designed to aid hormone recovery. Recovery is not possible, however, while supraphysiological levels of androgens are present, and such levels are usually found during all weeks of a normal (nonmedical) steroid taper. Individuals remain cautioned that dosage tanering is not a proven way to reduce post- cycle muscle catabolism.

#### Duration vs. Weight Increase

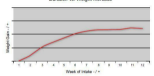


Figure 2. Anabolic/androgenic steroids tend to be most effective at a given dosage for approximately 6-8 weeks. After this point, the rate of new muscle gain will slow, and soon after will usually hit a full plateau.

# Stacking

As individuals become more experienced with anabolic/androgenic steroid use they may begin experimenting with the use of more than one steroid at a time. This practice is referred to as stacking Stacking is most common with advanced bodybuilders who find that at a certain level of physical development they begin hitting plateaus that are difficult to break with a previous single-agent approach. In many cases, however, it may simply be the greater cumulative steroid dosage that is necessary for the resumed progress. Stacking usually involves the combination of a more androgenic steroid with one or more primarily anabolic agents. On the anabolic side, common steroids of choice include holdenone, metherolone randrolone, oxandrolone, and stanozolol, Testosterone, oxymetholone, or methandrostenolone will serves as the androgenic base of most stacks.

intercesses art success art suscepts man autonous sources sugentes at the interact are two field. On the cone that, high does of electrosteron, conjunctione, or methandestendence are prone to producing strong androgenic and estrogenic side effects. Stacking first because very popular during the 1906s, a time when effective entropeam stream entropy source and who another in a material source and the size of a higher total stread dosage than would be token the assign administrate abundos candingen makes a long and the size of a higher total stread dosage than would be token the size and assign administrate and particular size of the size of primarily anabolic agents alone, even when they are talken in higher does. This conflicts with the conflict with engine expension of the "material" sensor. The conflict with the conflict wither conflict of the size of the s

Today, the availability of dauge that can reduce estrogenic activity makes the cortizated use of sizing agard cycles based on a stoom againous like estostories ceruthation or opinion much more visible than it was decades ago. Side effects like genecomatin and water restretion can row be efficiently mirriated und anti-estrogenic or anomaties eithbries, even when taking higher doses. Individuals should be aware that stacking is, ikewise, nor a necessary particle. In its likely to remain commonly applicable in competition holyshaffing circles, however, or when an infividual is sure they have progressed as far as they possibly can will a single-gard approach. Otherwise, for many allates and reconstruction control and active control of the single state of the control of the contro

# Sample Steroid Cycles

The following cycles are presented as examples of common steroid administration protected. These programs have not been evaluated in a ciscial setting for edge and efficacy, and are provided for informational proposes only. These are not recommendations for authorization and a contraction of the common state of the common s

#### Single Agent Cycles

Dispulsed Cypls #1 (Mass)

Products: 100 tablets 5 mg Methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4 g/day).

Estrogen Support: tamoxifen (10-20 mg/day).

Comments: This is a very common first cycle for building muscle mass, and utilizes a single standard both of methandrostenolone. This cycle is filedy to produce very noticeable muscle growth in a first-time steroid user, often in excess of 8-10bs of weight gain. This is usually not accompanied by significant visible sale effects such as genecomestia and water retention. Although this is considered a beginner's cycle, methandrostenolone is a 7-1 alpha fullytated oral steroid, and presents significant cardiovascular and liver toxicity. The repeated use of such drugs should be limited.

Week	Methandrostenolone 10 mg/day	
1		
2	10 mg/day	
3	15 mg/day	
4	15 mg/day	
5	20 mg/day	

Products: 200 tablets 5 mg Methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage)

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: This is a common follow up to the first Dianabol cycle, utilizing a slightly higher dose and longer duration of intake. The dossages used here are more common for bodybuiltine numoses. A slightly areater intensity of adverse reactions is likely.

Week	Methandrostenolone
1	20 mg/day
2	20 mg/day
3	25 mg/day
4	25 mg/day
5	25 mg/day
6	25 mg/day

Products: 10 mL 200 mg/mL Testosterone (enanthate or cypionate)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4 g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5 mg/day).

Comments: This mass building cycle is Redy to yield similar quantitative results as an early Diarabol cycle, but is forword over the oral for its lower earliessectar and hepatic strain. The doses used are expected to cause mid shifts in the HDL/LDL cholstearol ratio, but not the substantial changes normally seen with oral anabolic steroids. This sample cycle is Redy to resear the least aromat of health side effects of all idea in this section.

Week	Testosterone
1	200 mg
2	200 mg
3	300 mg
4	300 mg
5	300 mg
6	350 mg
7	350 mg

Products: 20 mL 200 mg/mL Testosterone (enanthate or cypiorate)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mo(day) or anastrozole ( 5-1 mo(day)

Comments: This cycle is a common follow up to the first testosterone only cycle, with a higher drosgue and 3 week longer duration of make. The total testosterone drosage given is double in comparison, and is likely to produce more pronounced estrogenic and anticogenic size effects. Cardiovacciant strain may be skightly higher than the first cycle, but should remain substantially lower than cycles with onal AAS. Testosterone is arguably the safest, and at the same time one of the most effective, muscle-bading seconds available. The exclusive repeated use of a cycle like this would be advised over more adventurous cyclingstacking proposal.

Week	Testosterone
1	200 mg
2	400 mg
3	400 mg
4	400 mg
5	400 mg
6	500 mg
7	500 mg
8	500 mg
9	500 mg
10	200 mg

Sustanon 250 Cycle (Mass

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1 mg/day).

Products: 15 mL 250 mg/mL Sustanon (testosterone blend)

Comments: This mass building program is similar to the other testosterone cycles, but utilizes Sustamon 250, a form of blended testosterone more widely used in Europe and other regions outside the U.S. The total steroid dosage of this cycle is 3,750 mg, externely close to the amount used in testosterone cycle #2. A similar level of cardiovascular strain and visible side effects are expected.

Week	Sustanon	
1	250 mg	=
2	250 mg	
3	500 mg	_
4	500 mg	
5	500 mg	
6	500 mg	
7	500 mg	
8	500 mg	_
9	250 mg	

# Products: 50 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosoor)

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: Oxymetholone is commonly regarded as the most potent mass building started available. It is also prose to causing both strong estrogeries and antrogeness idee effects. A steord rovice may gain 15-20 pounds or more on their cycle, although a significant amount of this wile be unter restriction, which will assist soon after drug discontineause. Oxymetholone more convenient to use them an injectable its observable, and in the convenient to use them an injectable its observable, it is not regarded as a safe alternative. Regenated use of e17 a plan adjusted on the first should be interest.

Week	Oxymetholone	
1	50 mg/day	
2	50 mg/day	
3	50 mg/day	
4	75 mg/day	
5	75 mg/day	
6	75 mg/day	

Oxymetholone Cycle #2 (Mass)

#### Products: 100 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: This is a more popular version of the oxymetholone only cycle. The doses here are more common with experienced steroid users, and more than sufficient to promote strong mass and strength increases. Side effects may be more noticeable than the lower dose cycle, of course, which may necessitate a higher dose of hamoxiden.

Week	Oxymetholone	
1	50 mg/day	
2	50 mg/day	
3	100 mg/day	
4	100 mg/day	
5	100 mg/day	
6	100 mg/day	
7	100 mg/day	
8	100 mg/day	

Stanozolol Cycle #1 (Lean Mass/Cutti

Products: 200 tablets 2mg Stanozolol

All Windows Linner Commont: Linner Ctobal. Lin: 62 one Economicals Exerts (Johnst economissated)

An weeks: Liver Support: Liver Staton, Liv-52, or Essentiale Force (ander recommended doctor)

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: This is a common first-cycle for an athlete bocking for performance improvements or a bodybadder looking for a learn muse or carring steroid. This cycle was more common when stanozolol was widely available in 2 mg stablets. Such preparations are now uncommon except in fartope. The dosego used here is low by bodybading statulards, although similar cycles have been the backbone programs for mmy athletic competions, although similar cycles have been the backbone programs for mmy athletic competions, although a transfer of the programs for mmy athletic competions, although the document of the programs for mmy athletic competions, although the document of the programs for mmy athletic competions, although the document of the programs for mmy athletic competitions, although the document of the programs for mmy athletic competitions, although the document of the programs of the

Week	Stanozolol	
1	8 mg/day	
2	8 mg/day	
3	10 mg/day	
4	10 mg/day	
5	10 mg/day	
6	10 mg/day	

Stanozolol Cycle #2 (Lean Mass/Cutting)

Products: 200 tablets 5 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosoor)

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4e/day).

Comments: This is a stronger version of a cuttingban mass balding cycle talking stancoals II the dosage used here is substantially higher than the first stancoald by cley, a fact that makes this cycle more properly said for bodybalding parposes than Stancoald Cycle #II, Cardiovoscular and hepatic strain will be more notable, and visible side effects more pronounced, than the first cycle. There should be no need to addition an estrogen miritenance drus.

Week	Stanozolol	
1	20 mg/day	
2	20 mg/day	
3	25 mg/day	
4	25 mg/day	
5	25 mg/day	
6	25 mg/day	

Stack Cycles

David (Normalis I Conduction Office)

Products: 10 mL 200 mg/mL nandrolone decanoate

100 tablets 5 mg methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: This is an extremely old and wid

Comments: This is an extremely old and widely repeated steorid combination, based on the predominarily analysis destined introduced excursts. Medium/neterobous servers as the anticogatic component of this stack, and is added during week 3, which is a time that side effects or freeduced antiogenizely (with the exclusion use of methodese decentately are commenty roticed), such as base of histo and secund dysfunction. The does used in this cycle is much size of the cycle is the comment of the commen

Week	Nandrolone	Methandrostenolone
1	200 mg	
2	200 mg	
3	200 mg	10 mg/day
4	200 mg	10 mg/day
5	300 mg	10 mg/day
6	300 mg	15 mg/day
7	300 mg	15 mg/day
8	300 mg	15 mg/day

Deca/Dianabol Cycle #2 (Mass)

Products: 20 mL 200 mg/mL nandrolone decanoate

200 tablets 5 mg methandrostenolone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4e/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: A more popular manifestation of the DecarDiarabol Cycle, with more commonly accepted dosages for a moderately experienced steroid user. Incidences of side effects are expected to be higher at these dosages, although overall this stack is likely to be less problematic than a corribration of testosterone and conventions.

Week	Nandrolone	Methandrostenolone
1	400 mg	
2	400 mg	
3	400 mg	10 mg/day
4	400 mg	10 mg/day
5	400 mg	20 mg/day
6	400 mg	20 mg/day
7	400 mg	20 mg/day
8	400 mg	20 mg/day
9	400 mg	20 mg/day
10	400 mg	20 mg/day

Testosterone/Anadrol Cycle (Mass

Products: 20 mL 200 mg/mL testosterone (enanthate or cypionate)

100 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day).

Comments: A confinition of estatetons and conventiones is generally regarded as fifteen top total college ages for gaining musicle mass. Both drugs will recent significant estinguistics, and will be likely to induce genecomistic quickly unless an estinguistic estinguistics, and will be likely to induce genecomistic quickly unless an estimation to gain over 25-30 points on a cycle such as time. When therefore the present unless there have been toom to gain over 25-30 points on a cycle such as time. When returns will be very light with this contraction of the cycle in the contraction (contraction) to 10 pounds or must) is expected soon after the cycle is discontract.

/eek	Testosterone	Oxymetholone
1	200 mg	1
2	400 mg	Ī
3	400 mg	50 mg/day
4	400 mg	50 mg/day
5	400 mg	100 mg/day
6	500 mg	100 mg/day
7	500 mg	100 mg/day
8	500 mg	100 mg/day
9	500 mg	100 mg/day
10	200 mg	100 mg/day

Tactactorono/Doon Cunla (Mass)

Products: 10 mL 200 mg/mL nandrolone decanoate

10 mL 200 mg/mL testosterone (enanthate or cypionate)

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1 mg/day).

Comments: Testosterore with randrobras is considered to be one of the most findamental 2-drag combination stack. Nandrobrae compliments the androgant lasse of intestorence by suplamenting additional anabolic activity without strung entrogenicity. The resulting stack is admitted as production as an eyel-trailing as layler dose of festosterone alone, but less problement in terms of estrogenic side effects such as water reterion, genecomosis, and fit bulkap. Estrogen conversion is all florabilithe compils to warmar the use of an estrogen maintenance drag, however, and this stack remains in the realm of mass bulking instead of learn mass or exting:

Week	Testosterone	Nandrolone	
1	200 mg	200 mg	
2	200 mg	200 mg	
3	200 mg	200 mg	
4	300 mg	300 mg	
5	300 mg	300 mg	
6	300 ma	300 ma	

7	300 mg	300 mg	
8	200 mg	200 mg	
1 1 1 1/4	0.1.0.36.3		

Andriol/Anavar Cycle (Lean Mass)

# Products: 360 capsules Andriol 40 mg

400 tablets ovandrolone 2.5 mg

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: This is an effective but rail conlo-only learn must bailing cycle. Author's used as fine antiogenic bose, but in doses that on on groutly exceed round theraportic levels. Outsrhotene is measurantizable, so significantly elevated estrogenicity is utilisely. Illamodatin 10-20 mg pet day may be used dead the asstatement dosage turn out to be problement. This stack is popular arming solds men and flow not wishing to see specialises. The stack is popular arming solds men and flow not wishing to see specialises. The stack is popular arming solds are sold and the stack in the stack in the stack is popular arming sold on men and the stack in the stack is popular arming sold on the stack in the stack in the stack is popular arming sold on the stack in the stack

Week	Andriol	Oxandrolone
1	240 mg/day	15 mg/day
2	240 mg/day	15 mg/day
3	240 mg/day	15 mg/day
4	240 mg/day	20 mg/day
5	280 mg/day	20 mg/day
6	280 mg/day	20 mg/day
7	280 mg/day	20 mg/day
8	280 mg/day	20 mg/day

Products: 18 mL methenolone enanthate 100 mg/mL

50 mL boldenone undecylenate 50 mg/mL

20 mL testosterone (enanthate or cypionate) 200 mg/mL

All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20-40 mg/day) or anastrozole (.5-1mg/day).

Comments: This is a 3-month non-low-rooke cycle that has 2 distinct phases, mass and lean mesociating. The first weeke of turning and date are fexcated on mass bading: Significant entogenizely will be present in these weeks, and may necessitate the use of turnovidin or an auroritise righther rank as austroache to prevent gracocomistic and excessive water reterior. In Strugam mattenurses drugs may be roduced or possibly clinicated after the start of place 2, which forces on increasing the analogam to entogen to and as doddleged the offset to prevent sexual dysfarction or boss of flikids, which often occurs with the use of prodorimentaly analogs storoids above.

Week	Testosterone	Boldenone	Methen
1	500 mg	200 mg	

500 mg	200 mg	
600 mg	200 mg	
100 mg	200 mg	300 mg
100 mg	200 mg	300 mg
100 mg	200 mg	300 mg
100 mg	200 mg	300 mg
100 mg	200 mg	300 mg
100 mg	200 mg	300 mg
	600 mg 600 mg 600 mg 600 mg 100 mg 100 mg 100 mg 100 mg 100 mg 100 mg	600 mg 200 mg 600 mg 200 mg 600 mg 200 mg 600 mg 200 mg 100 mg 200 mg

700 ---

Testosterone/Artidroi/Trenbotone Cycle (was

Products: 30 mL 200 mg/mL testosterone (enanthate or cypionate)

----

20 mL 75 mg/mL trenbolone acetate

100 tablets 50 mg oxymetholone

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosage).

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Estrogen Support: tamoxifen (20 mg/day) and anastrozole (.5-1 mg/day).

Comments: One of the more entruen mass balling cycles in common use among bobbulades. This seck will imput rapid pairs in now maces less and strength. This discontinuation is lighly proces to causing estrogenic and authorganic side effects, including side of the comment of the cycle of the comment of the cycle of th

Week	Testosterone	Oxymetholone	Trenbolone
1	300 mg		150 mg
2	600 mg	ĺ	150 mg
3	600 mg	50 mg	150 mg
4	600 mg	50 mg	150 mg
5	600 mg	100 mg	150 mg
6	600 mg	100 mg	150 mg
7	600 mg	100 mg	150 mg
8	600 mg	100 mg	150 mg
9	600 mg	100 mg	150 mg
10	600 mg	100 mg	150 mg
11	300 mg		

Products: 20 mL 100 mg/mL drostanolone propionate

#### All Weeks: Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day)

Comments: This is an effective stack for hardening, cutting, and gaining lean muscle mass. Neither agent is capable of converting to stronger, so this cycle should significantly elevate the androgen to estrogen ratio. This may assist in the breakdown of fit tissue, erthancing muscle definition. This stack should not present significant fiver toxicity, although cholesterol ratios may be similaritaryla deterf in link of reduced estroneric activity.

Week	Drostanolone	Methenolone
1	200 mg	200 mg
2	200 mg	200 mg
3	200 mg	200 mg
4	300 mg	300 mg
5	300 mg	300 mg
6	300 mg	300 mg
7	300 mg	300 mg
8	200 mg	200 mg

WIBBIOTI IOVIOLI HEIDORDIE CYCE (LEBII WIBS/CHILIE)

Products: 250 tablets stanozolol 5 mg

100 tablets mesterolone 25 mg
20 mL trenbolone acetate 75 mg/mL

All Weeks: Liver Support: Liver Stabil, Liv-52, or Essentiale Forte (label recommended dosoo)

Cholesterol Support: Lipid Stabil (3 caps/day) and Fish Oil (4g/day).

Comments: Stancoole and trensholme are popular storoids during cutting phases of maining, and impart storage unfamilie and moderate and angient effects of majoritized entopolissis. This combination helps to impart a strong fit loss/definition-enhancing effect. Who 25 mg tables of metastrools have been shaded per day to supplement additional auto-categorie than the standard in mornal Bedrá and secund functioning. Additional standardings of the standard of the standard and the standard and the standard and the disputation of the standard and the standard of the standard the standard and the standard and the standard and the standard of the standard and the standa

Week	Stanozolol	Mesterolone	Trenbolone
1	20 mg/day	25 mg/day	150 mg
2	20 mg/day	50 mg/day	150 mg
3	20 mg/day	50 mg/day	150 mg
4	20 mg/day	50 mg/day	150 mg
5	25 mg/day	50 mg/day	225 mg
6	25 mg/day	50 mg/day	225 mg
7	25 mg/day	50 mg/day	225 mg
8	25 mg/day	50 mg/day	225 mg

Post-Cycle Therapy, or PCT for short, refers to the practice of using certain medications to sensitur in the discontinuous of analodes statissis. While stroots are not addictive days in a classical issue, they do suppress your own hormous production, at least temporarily. This is an issue that should be addressed after conclusion of use. If the services are discortined abruptly without addinessing internal hormous production, the result could be a prolouged state of hypogonadius flow analogua levely characterized by a shatteral loss of musel, muse, reduced energy levels, depression, and impaired fluidoieual functioning. Strootsing hophytaliser forts of this fair they not clearly in the stroot of hormous packation as a reduce to the cust We will also discuss certain texture of the contraction of the

#### The HPTA Axis

In the humm body, the Hypothulumic-Pinkiny-Tosicular Axis (HPTA) control is extostenore biospherisch. The PTA's a slight regulation system of checks and humbers that works to assure the cornect level of istostorous is maritained. We can look at this regulating process as inveity free level. At the tay is the lapsophalme region of the bean; which releases (addit (Condotrophi-Relassing Horneso) when it senses a need for more textusioners. Geld (Condotrophi-Relassing Horneso) when it senses a need for more textusioners. However, the condotrophi-Relassing Horneso (Alle Marie Sanda, 1994) when it is sense as the form the condotrophi-Relassing Horneso (In the Condotrophi-Relassing Horneso) (In the Condotrophi-Relassing Laboratoria) (In



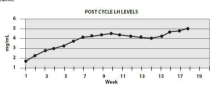
The Hypothalamis-Pitatiany-Testicular Axis: The hypothalamis releases Comadoropia Reksaig Hormore (GaRII), which vitanistes the pitatiany to release Interiority hormone (LH) and folicle stimulating hormone (SH). This (primmit) LH) promotes the release of testisteerine from the testes, Androgens, as well as extrogens and progestion, in turn cause negative feedback; inhibition at the value of the companion of the control of the control of the control of the when to much hormone is present.

#### Unaided HPTA Recovery

The suppression of matrial testioneroe synthesis by strend use is typically a temporary phenomenon. Even if you do roding, your thooly somal androgen synthesis will usually return a few to several morths after the cycle is correlated. The problem is, this can be a very long time when you are reliving on testioneror for so may than, settleding the administration can be in fact, much of the muscle muss achieved during AVS administration can be in the weeks and much its follow from androgen levels are the administration can be in the weeks and much its follow from androgen levels are the administration can be in the weeks and much its following and analysis to straight the administration can be in the weeks and much its following and analysis to straight the administration can be in the weeks and much its following and analysis of the straight of the control of the straight of the straight of the straight of the straight of the control of the feel when the straight of the straight of the straight of the control of the straight of the straight of the straight of the straight of the control of the straight of the straight of the straight of the straight of the control of the straight normally looks like without assistance. Only then can we identify the levels of the HPTA that are most open to manipulation with support medications.

Staties on the post-coustion aspect of ambods stroot laws, especially in AAS abuses, are lakeing, I most cases want selet to single-gast staties, usually of hormoze repicturest patients. One of the most detailed views of what a post-cycle crash probably looks like corns from an inseligation into testosterone centralies. <sup>34</sup>H incohes a group of most that were given weekly injections (250 mg) for 21 weeks, a done that admittedly does go beyond romer Haff Une. Writes hormoze twee measured each work during the statly, and for more than 4 morths after the medication was descortinated. A neview of the data shows that at the start of the substitute of the state of t

The above study suggests that one of the first things to lappen after strood constaints in the brain recognities restorated resolution was upon the wild cause GRHI and III Head begin converting fairly quickly. The substantial delay between this and an increase in tensorates levels a caused largely by electrical tresponsionesses to theirizing homests to testizing homests to the strong tensor that the strong strong the strong tensor the strong tensor the strong tensor the strong tensor that the strong



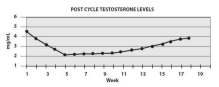


Figure 1. IH and Testosterone measurements starting 1 week after the last injection of 250mg of testosterone enanthate (pretreated measures were 5 mU/ml and 4.5 mg/ml respectively). Note that between weeks 1 and 5, as testosterone levels are declining due to the cessation of exogenous androgen administration, I.H levels are beniminat to correct. From weeks 5 to 10, testosterone levels remain art over

near baseline, although LH is increasing by this point.No notable correction in testosterone occurs until after the 10-week mark. hCG in PCT

Haman Chorionic Genedotopou ICCO is a fertiley drug that traines the actions of Learinage hormone. It is commercy und during the procept-cipe partied to address testicular antipols, which as we lines seen is one of the findmental conditions to hormonal recovery. The ICCO is typically latent as substantial dosaged for a period of 2-3 words. Fortilear antipols is caused by a first of the contraction of the cont

# Anti-Estmoens in PCT

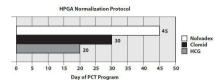
The arts citruspasis drugs Chard (cloniplence citate) and Nobades (turnoution citate) are also commonly used drugs for pool-cycle period. These drugs are used to block the negative footback inhibition of strongen, visit necess primarily at the hypothatinus. <sup>266</sup> This may lost the heightened releases of Gelfall, and subsequently III and estostateous. While entire parts between the contraction of the strongen inhibitor of institutions, in side in the regulation of analogue biosynthesis is regulated as a firstly direct one. The propose of drugs agric resonance in 11st period contraction of the strongen are both origing correction in 11st leave mere quickly, and the augment to state of the strongen in both origing correction in 11st leave mere quickly, and the augment to state of the analogue of the strongen which is the strongen in both origing contents in 11st leave on the distribution.

It is important to note that the use of anti-estingens alone is generally not regarded as an efficiency strategy for addressing horizone recovery at the conclusion of a stand cycle. This experiments of the contraction of a stand cycle. This was a contraction of the contraction o

# Traditional PCT Program

The faboring PCT program was developed by Dr. Mehad Scally, one of the most well known and accompled nitriduals in the field of anotice strongs and make hormous palacement medicine. Scally has been a particularly strong force lobbying the medical community and government to recognize the information includes the faborine stead on a community of the program of the

This PCT program begins with a substantial doss of ICG (2000 IU every other day for 20 day), Anti-estopps are also used during this period. This is potentially important because ICG may up-regulate testicater aromatuse activity. <sup>256</sup> Thus, their use can minimize both estrogenis side effects and readone agentific feedback inhibition of strosterone release. The arti-estrogenis token are turnowific refurne (20 mg twice per day) and chemiphane crimate (50 mg twice per day). Chomili is used for a substruct period of time, in a secondar down of the program's medications. While in the first couple of weeks the arti-estrogens may not be highly effective, they should prove more critical towards the middle and end of the program in the published version of Scaly's program (which is slightly medified from the above), correlationment function returned in a slughest while 45 days. This is a definite success, far more foureable than the protected recovery window reported in the study with 250 mixtured of the protected processors.



Protocols: Human chorionic gonadotropin (hCG) is taken at 2000IU every other day for 20 days. Clomiphene citrate 50 mg is taken twice per day for 30 days. Tamoxifen citrate is taken 20 mg twice per day for 45 days.

The timing for a Post-Cyck Therapy program can be as important as its composition. If it is initiated too late, subshed days forment Dermone levels (and also once manch emes) may be lost. If you start the programs to early, you may min the optimal weaknew of efficiencess. FOCT programment of this medication, In particular, we went to make user that ICG is being applied right amount the time that congenous strucks are dropping below the freshold of physiological undropen strainform in the congenous strucks are dropping below that and and explain; this would be right before blood levels drop below the normal level (250 negred). There should all a wall not every both in the concept consists of the consistent days to describe dropping and the program of the consistency of the consistency of the consistency of the consistency to the consistency of the consisten

The coact trings for FCT program is determined by the chiration labifiles of the drugoly used We will use bestioner copjounted-termined as an example. We have each injection to an arismation labifile for approximately 8 algos, Adoss of 200 mg/week shoold produce blooked evid of around 2000–2000 mg/dt affect sevent weeks of their would take also 2 and 2000–2000 mg/dt affect sevent weeks of their would take also 2 does. Then, the PCT program would be initiated a few days to one week after the last testastoners injection. The program would be deducted with higher doess. For example, at 500 mg per week of TCUT it should take approximately 4 half lises (22 days) for testostenors depended when the merit margine in the same, PCT would be initiated about two weeks after the last testostenors depended to the same and the

Timing the Start of PCT	
Type of AAS used	PCT Initiation
Cypionate/Enanthate: (200 mg/week)	3-7 days after last injection
Cypionate/Enanthate: (500 mg/week)	10-14 days after last injection
Decanoate: (200 mg/week)	10-14 days after last injection
Decanoate: (500 mg/week)	18-21 days after last injection

354. Effect of long-term testosterone cenurihate administration on mule reproductive function: Clinical evaluation, serum TSH, 1H, Testosterone and seminal fluid analysis in normal men.J. Mauss, G. Borsch et al. Acta Endocrinol 78 (1975) 373.

83 (1975) 31-36-34

355. Decensitization to grandstropins in cultured Leydig tumor cells involves loss of grandstropin receptors and decreased apacity for storoidogenesis. Fernam DA, Accol M Proc Natl Acad Sci U S A 1981 OC;78(10)(530)-31-355. Belation of their initing harmore securious by testoroteon; in men requise summarization for its paintury but not its hypothalunic effect; evidence from the tradent study of normal and grandstropin-redesing hormore-defection reversible of the Defendent PA. Device M. A. Device M. A. DeVice S. L. Orl B. Percole P. PA. Crudes W. H. Haves H. I. Clink B. H. Clink B. Clink B. H. Clink B. H

denotes their relicions, v<sub>L</sub> Doyler 20, Declar X, Lee et , Bocquie 19, Clossiey we 11, ruspes 12, 3 Can Eulocraind Metals. 2008 Mar 95(3):784–91. Equie 2007 Dec 11. 357. The effect of arouttase inhibition on sex steroids, gonadostopius, and markers of bone tumover in older men. Tasel P. Kernede VDG et al. J Clin Bindorinol Metals. 2001 Jun 86(6):2899–74.

388. Acute stimulation of aromatization in Leydig Cells by Human Chorionic Gonadotropin In-vitro. Proc Natl Acad Sci USA 764460-3.1079

# OCT: Off-Cycle Therapy

The objective of anabolic stenoit therapy (when normicical applications are involved) should be to diet the desired benefits with the boust camatities opposare and sile effects. This normally related offigures with optimizing all appets for firaining, rost, and dat, as well as a adduring to a 18-ok Cycli Therapy (FCP) regognant the conclusion of each storiet (yet). One the one land, we want to make each cycle as productive as possible. On the other, we are string to rotate the most gains not be straining not for the next cycle is that meth farther along. When all appets are in check, the result should be a need for lower tend looks, fower coulds for the control of obstitutions and white of the chief of the control of

Gine the importance of retaining our muscle and performince gains, however, our efforts in this regard should not conclude with Pro-Cycle Therapy indeed, to receive the greatest lawy-term benefits from ambolic-indoparies serord therapy is a shot advisable to initiate an OSC-Cycle Therapy (OCC) programs when the PCT is ower. The focas of OCT is Specially use all matters advantaces (distorty supplements) that foror muscle reteriors, while simulatenceds advantage general physicisty and a forement ballances to return. What is it for faid that supplement the proposal properties of the control and the control that is of faid that supplement programs may be effective in the above of performance class. We can find that supplement the programs more effective in the above of performance class.

A well-organized OCT program less a minimum of six to eight weeks, and consists of three distinct conponents. The first is "floationers spapent," which seeks to enterful an efficient PCT program, but with a different and much more basic approach. The second part is "Manke Cel BloSesiantion," Heavy training disrupts the macker of the methers become less response to occures strainfaton. We want to address this during OCT, and prime the muckes for the next board of stresse raining lands, we want to related one or more natural muscle-building substances in the program. This part is called "Anabolic Synghementains." If the right products are used, finister anabolic-metastoic effects of the program of the season of the program of the pro

#### Part I: Testosterone Support

The testoscene suport aspect of our OCT program is substartially different than what is used during tutilistical DCT We are no longer looking to all endogenous testosteres, production with artis-stroppine drugs. Bite tumoristin or elemphone, nor to use apharmacentical start mine endogenous testicing hormouses such as ICC, 41 pharmacentical strategies have been concluded at this point, and hopefully have eliked the pharmacentical strategies have been concluded at this point, and hopefully have eliked the recessary effects. Dairy GCT, we want to provide our bodies some of the natural components used in the synthesis of testosterone. We want to augment our own natural reconsecution of the synthesis of testosterone.

#### Vitamin D/Calcium/Zing

The first flag to pay special attention to during OCT is our vlamin and mireal states, practically those components that are integral to testources beoptiseds. This includes Vlamin Di Cakian; and Zinc. To begin with, clinical statics have shown that higher levels of Vlamin D in the blood are associated with increased testources coupts. <sup>200</sup> Thus, supplementing Vlamin D may be advantageous during the long OCT period, when you will be religiously only our runtial testources for the hormousl approxi of authorists. Cakian is another matrixe component insolved in hormone faction, especially the level of broundable (tray) institutence. <sup>200</sup> Also of 500-1,100 m gold my how best off incelled, as the intended pain is also to anothing in the special with officers are small factor. Landy, a small done of zinc may also be taken if needed, as the intended gain is also the anothing in the special with degree and the proposal properties. <sup>200</sup> My defections; in novel likely translated

#### D-aspartic Acid

D-asparie acid (DAA) may also be useful during OCT. DAA is an amino acid that is nuturally found in the nervous and endorient systems, and is believed to play losi in such things as neurotransvission, spermatogenesis, and hormone biosynthesis. Clinical staties that gave 3.2 g/day of D-asparie acid per day (as sodium solt) to healthy men resided in a 42% increase in serum testosterone levels in most subjects. <sup>302</sup> This same dose is recommended during OCT.

#### Part II: Cell Re-sensitization

Report ligh intensity cuercies, especially resistance training, causes disruption of the muche coff murbrances. This disruption is many soy desirable, as it is model to miniture much disruption is the many soy desirable, as it is model to miniture much goods and repair. Without durange, there is no progress. There are some negative inspiration of the muscle cole, brower. One of them not influentiate list after lower murbrances of the muscle cole, (which cousist mainly of fathy acid compounds called physiologishigh are reamaged. In particular the concentration of analchiotic acid (ARAS) is lowered. <sup>200</sup> ARA supports the local ambods process. <sup>204</sup> Likewise, its depletion is one of the corrown factors in training sensoriton.

#### Arachidonic Acid

To help replanish murthrane phospholipids and restore muscle cell responsioness to training, andicition acid should be supplemented during the CCT period. A dayl dose of ord 250 mg is recommended, which represents 50–100% of the normal dialy distury intake of PARA. This amount should be sufficient for phospholipid replanishment, and acceptable for long-term use. Figher doses (500–1,000 mg per day) may provide a more distinct muscle-buildine effects the should be finished to sits oneven weeks.

#### Fish Oil

It may also be useful to supplement with fish oil during the OCT period. The main intensit and ecoconbenomic acid (FIAH) and tecosoparation icas (FIAPA, two Omaps.) a sentral finity acids that are also important constituents of muscle cell menthema phospholytake. Additionally, satisface suggest that Omaps.) accountal flat packates may enhance the membrane analysis of effects of this if FA.  $^{30.5}$  A shiply dose of 2 grams of fish oil is typically recommended driving and  $OSC_{\rm CM}$  Darrapy program.

# Part III: Anabolic Supplementation

An optimal Off-Cycle Therapy program should also include natural products with

ambolication statistic properties. Many AGS users are skeptical of muscle-badding supplements, and rightly so. The muscle can be very trendshiet, with even the better products liking far short of AGS in terms of efficacy and reliability. SSE the field has been considered to the control of the control of the control of the control of the corn a pratial makes bequire effect charge the COT proces is highly desights, as it can significantly also the baseline muscle keel by the start of the next steroid cycle (and this may reliable the first product of the control of the control of the control of the control of the supplementation to only those applications with proven analysis effects in harmons. For a more control of the control of th

# Creatine Monohydrate

Creation monolydrate is regarded as the "triginal" anabolic supplement, as it was the first list offer substantial performance and body composition improvements for most users. It is typically lasken for 8-12 weeds or longer (connetirues froughout the entre CCT period), at a to typically lasken for 8-12 weeds or longer (sometimes froughout the entre CCT period), at a close of 5' gains per dept. Creatives againstrass mucle size and performance frough several entre of the contraction o

#### Beta Alanine

Beta Alaine is a non-essential anion soid that serves as a direct precursor for carmonic synthesis. Dring exercise, Judopan is not are produced in the macket cell, which cause the pill level to drop. This precipitates muscle fittings. Curronice acts as an intramocellular budding agar, curronice to ball-up officiory in ice. By serving a for their letting state done of 3-5 games per day is typically used, which should allow the influent of muscle pill. The contraction of the pill of

#### Branched-Chain Amino Acids

There are three essential Branched Cluis Annio Acids (BCAA) – kucine, isolocine, and voine. These mino acids are very abstantia in ackelst mace produce, accounting for 1-18% of the total content <sup>160</sup> The gosphementation with BCAA is dominble for a couple of reasons. The first is both they proteed in egglidating belocks for the opticiss of now mostly experimental to the content of the content

# Typical OCT Program (8-12 Weeks)

# Testosterone Support:

Vitamin D 3000 IU/day Calcium 500 mg/day

Zinc Sulphate 250 mg/day

D-Aspartic Acid 3.2 g/day

#### Muscle Cell Re-sensitization:

Arachidonic Acid 250 mg/day

FishOil 2 g/day

# Anabolic Supplementation:

Creatine 5 g/day

Beta-Alanine 3-6 g/day

BCAA 10 g/day

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# Injection Protocols

Anabolicalnoganic streod injections are always given deep in the macke (faramuscular). Some other performance-entirused (page sance has harman gooth hormore and inside injections are given by injection in the fall type broberen the skin and mascle (substantacous). The protectode for both spiction types are protected. Improper pictions technique can ensule the protection of the objection of the protection of the protection of the objection of the contraction of t

#### General Preparation:

1. Make sure you have all the necessary supplies.

#### Intramuscular Injection:

(1) 3mL syringe

(1) 22-25g 1-1.5" needle for administering the injection

(1) 21g 1-1.5" needle for drawing solution (if using a multi-dose vial)

(2) Alcohol pads

(1) Dry cotton ball

(1) Plastic bandage



Syringe with needle attached. (intramuscular)

#### Subcutaneous Injection:

- (1) Insulin surince with needle attached (5-1ml .27-30c)
- (2) Alcohol reads
- (1) Dry cotton ball
- (1) Plastic bandage



Syringe with needle (insulin)

- If applicable, keep the administration needle cold by leaving it in the freezer for at least one hour before opening. This will help dull the pinch of penetration.
- Select a well-lit room with a clean hard surface such as a tabletop or counter to administer the injection.
- 4. Wash hands thoroughly with soap and warm water.
- Assure that all injection equipment is sealed and unused. Never reuse needles. Double check the expiration dates on all medications.
- 6. Clean top of vial thoroughly with an alcohol pad, if applicable. Let air dry for 15 seconds.
- Precautions: Do not use injection equipment that is used or has been exposed to air during storage. Never share needles or multi-dose vials. Discard unused portions of the drue at the recommended time.

#### Drawing Solution into Syringe:

- 1. Remove syringe from packaging. Attach drawing needle, if applicable.
- If using a multi-dose vial, fill syringe with air in the amount you are withdrawing. This will help stabilize the pressure and make drawing easier.
- 3. If using an annule, break onen and place flat on a hard surface. The use of a naner towel

- to cover the glass top may make breaking easier. Draw solution. Skip below steps and go to Intransucular or Subcutaneous Injection Procedure.
- 4. If using a vial, insert needle through the rubber stopper at a 90-degree angle. Turn the vial upside down with needle attached. Inject air. Slowly withdraw desired amount of solution. You may lightly tap the side of the needle to dislodge air bubbles. Note that small air bubbles are not harmful.
- 5. Remove needle and syringe from vial, if applicable. Replace cap on the end of needle.



Keep fingers away from needle when drawing.

- 6. Remove drawing needle and replace with new administration needle, if applicable. This is highly advised with multi-dose vials, as passage through the rubber stopper will have dulied the needle considerable. Remove are air in the in of the needle and neware for inscion.
- 7. Place capped needle back inside wrapper and place on clean surface.

Precautions: Never touch the tip of the exposed syringe, the needle, or the top of the vial stopper after it has been cleaned with alcohol. If you come into contact with these surfaces you should consider the materials contaminated, and should not use them for injection.

#### Intramuscular Injection Procedure:

Used for all anabolic/androgenic steroid injections.

- Thoroughly clean the intended site of injection with second alcohol pad. Preferred locations are the upper outer quadrant of the buttocks, or the outer side of the thigh. Let air dry for 15 seconds.
- Remove needle cap. With free hand, stretch the skin around the site of injection with two fingers. Move the skin over the muscle to the side by 1-1.5"(Z-Track method).
- In a swift motion, insert the needle into the target muscle at a 90-degree angle with the dominant hand. Make sure the needle is deep within the muscle.
- Pull back on the plunger (aspirate). If the syringe fills with blood you have hit a blood vessel, and the injection should be aborted.
- 5. Inject the medication slowly into the muscle.



Upper/Outer buttocks is the preferred site for IM injection.



#### The outer side of the leg is also commonly used.

Withdraw the syringe. Release the skin with your other hand. The skin and subcutaneous tissue will rebound, which helps close off the needle shaft and prevent leaking.

7. Dry injection site with dry cotton ball. Cover with plastic bandage if necessary.

Precautions: Never inject into skin that is discolored, broken, or irritated, or if there are lumps, knots, or feelings of pain in the area. Do not inject more than 3mL at one time. Rotate the site of injection so that you do not inject in the same muscle more than once every two weeks.

# Subcutaneous Injection Procedure:

 Thoroughly clean the intended site of injection with second alcohol pad. The preferred location is the lower abdominal region. Let air dry for 15 seconds.



## Shaded area denotes site for abdominal subcutaneous injection.

- 2. Remove needle cap. With free hand, pinch the skin around the site of injection so it is lifted off the muscle.
- In a swift motion, insert the needle into the target area at a 45-degree angle with dominant hand. Make sure the needle rests within the subcutaneous tissues between the skin and muscle.
- 4. Inject the medication slowly into the subcutaneous tissue. Do not aspirate.
- 5. Withdraw the syringe. Release the skin with your other hand.



#### The skin is pinched upwards to facilitate injection into the subcutaneous tissues.

#### 6. Dry injection site with dry cotton ball. Cover with plastic bandage if necessary.

Precautions: Nover inject into skin that is discolored, broken, or irritated, or if there are lumps, knots, or feelings of pain in the area. Do not inject more than 1 mL at one time. Rotate the site of injection each time so that you are at least 1" away from the lest site, and do not return to a previous site until all other available sites have been used. This will help prevent overnees of the same injection area.

# Steroid Frequently Asked Questions

#### 1) How much weight can someone expect to gain during the first cycle of steroids?

Provided dosing is sufficient, a steroid user can expect to make the most significant progress during their first cycle. Although this will vary from person to person, it is not uncommon for someone to gain 20 pounds of weight or more during a 6-8 week period of AAS use. Some of this may be water retention, although a solid gain of more than 10-15 pounds of muscle

# 2) Are the gains from steroid use temporary?

Yes, and no. Steroids can hely you do two basic things with regard to muscle growth. First, they can allow you to more rapidly reach your genetic limits for muscle growth. Provided you continue to train actively, eat properly, and use an effective PCT program, you should be able to ministrian at your genetic limit indefinitely. So in this regard, the early gains do not have to be termorary.

Later, steroids can allow you to push well beyond your genetic limits. It is important to emphasise this, as extreme physical development cannot be maintained long-term without the repeat administration of anabolic substances. The body will always reverb back towards is normal metabolic limits once AAS are removed. In this context, some of the gains will not be permunent.

Seroids do permunenty alse the physiology of your muscles by adding more collair rackly high gird rack context, each muscle cell can immage is volume more efficiently, which allows more repid copussion. Deen after a long period of complete abstinance from training and AAS, the naction ermin. <sup>27</sup> This may provide a 'mascle memory' effect, allowing you to reach your genetic limit (perlups a slightly extended intri) faster than if you lad sever used AAS in the parts. So in the segart, there are busing bourthe beyong the temporary increase in

#### 3) Can steroids make me look like a professional bodybuilder?

If you have the underlying genetics to allow for this extreme manche growth, this may be possible with a lot of near work and declaration If you are like the wast unspirely of people, however, steroids will not be able to make you look like a professional bodybulder. Genetics are a big factor in determining the ultimate limits to your physicists, even in an enhanced state. Many people use steroids and look very big and impressive because of it, but very few usees are able to make to the state of a professional bodybuldiner.

#### 4) How dangerous is an isolated cycle of stemids?

Ambelichardneparie steroids are among the safest days available, at least in a short-term sense. Fail overloots is not reasonably possible, and the regular bath changes sach as sheardness in delektoral, blood pressure, hemetocit, and blood clotting (among other things) are very utiledy to manifest in arrivals body home of death after in absoluted cycle. There are not educated from such things as stroke and five cancer in short-term absocs, but such concurrences are satisfainly externely are in light of the milition of people that see these drugs. If you laid to comparatively rate the acute risks of AAS abuse, they would be slightly hidden the marrisms to the first synthesis and a lotter life in concili.

#### 5) How dangerous is long-term steroid use?

The kept term use of steroids for normalical reasons can be a significantly inhealthy practice. It has been diffialt, however, to uniffy the ceart ink! Them issue is the fet that AAS abuse can promote heard disease, the number one kiler of trans. Heard disease is a skew progressive disease, within my health of breadess whates approaches. Steroid abuse may accelerate the skert process of plaque deposition in the arteries, and also induce other changes in the cardiovascular systemathet can inverse assexpethiny to stoke or heart attack. If danth fully occurs, however, it will be difficult for a medical exament to proport AAS as the cause; to many variousles play a not in the oxilogo of cardiovascular disease. The variously of deaths where AAS has two contributed go unexpected for this exacts. The coast AAS conding to one operation beased may be cardioval associated as of this sequent. Not only dentify the contributed go unexpected for this reason. The coast AAS conding to one operation beased may be cardioval season for the coast beautiful and a clause including satisfic compared to non-users. The substitute of the accessive the season of the fill associated by the protection of the season of the fill standivision groupdation file as equivally important to closely monitor cardiovascular disease and other health risk factors if long-term stored use is a practice you will follow.

#### 6) Can steroids be used to enhance an athletic career safely?

The nonmedical use of AAS by definition cannot be defined as a safe practice. However, it

can be amped that anti-bolizationages storoids can be used with high relative solfs, even over a period of many upcass. The galderies of stored hum moderation are important to minimizing the negative health effects of these drugs. Provided an individual follows these galdeless and is certified with drug selectro, thousages, and durations or finites, follows and the ion in suntanted list, choicesterol, sugge, and refuned carbodylatests, scribely mans with both resistance and cardovaried neutries, and these choicesteria sports appreherms such as fish of and Lipd Solid during all cycles, it may be diffical in imany cases to ragge high amplies which is above achieved.

#### 7) What are the safest steroids for men?

Testosterore, whatever the form tends to be the salest steroid for men. When the does remains with the moderated sparenthrespects range (such as 200-400 mg off an injectable testosterore ester per weed), absentions in cardiousecular risks factors are noticed, but not externer. Some of this has to do with the benefical auditousecular effects of estrogen in men. Also considered daily sales are the commen specialist searoids believen; maintolene, and the contract of the contra

# 8) What steroids will not cause hair loss?

For those with a genetic prodisposition to hair loss, all ambide/intrologue's steroids are completed cardecularing the process. Storying the cont of this darking Ad-Sus receptors as focus on reducing relative authorganicy in the scalp. This can be accomplished with the use of predominary ambide drugs such as manufocus, countednote, or methoration, or methoration, or methoration, or methoration, or methoration, or methoration, or methoration. DBH conversion entand anticogenic amplification in the scalp. SBI, those genetically prove to hair loss can have problems with any steroid, and are always advised to limit dosing, drugratine durations, and interface the scale to the state of the scale of the scale of the scale of the state durations, and interface called the scale of t

### 9) What are the safest stemids for women?

Women are generally most concerned with the virilizing (musculnizing) effects of arabolicantrogenic steroids. The least virilizing agents are those with the highest relative arabolic to anticopinci effect, such as randrolone, controlone, and methonolone. Care must always be taken, however, as all AAS are based on mile sex steroids, and as such can cause musculnizing offlex in surrous.

## 10) Should I rotate my steroids every few weeks to prevent receptor downregulation?

No, this is not necessary Ambidicandrognic storoids all work primarily by attacling to and acutoting the same reaport. As such, you do not gain analytic by swidsing to a new compound that works via stimulating the same neeptor. If behavior, were included by one ASS compound, to would be estanded to all compounds. The plateau effect and is noticed 6-8 weeks into most cycles is poorly undenstood, but falsey related to the new metabolic intrinsipacted on much cell surface in fainteen of a central ASS alongosa, not insensitively of AAS. Classic downregulation does not occur with these drugs, and even if it did, notating sensitives would be according to the contraction of the contraction.

# 11) How likely am I to find real steroids on the black market? Does it matter?

Abbush exest figures are difficult to calculate, real plearmecential arthodic indrogate, actuated are estimated to represent for less of the products correctly created on the black mutet. In many region this figure may be below 25%. The majority of products said proceedings on the processing of the products made and labeled by underground abouttories. It does mutet, because the quality of normedical AAS carent be ensured. These products are generally not advised for two. Gone the potential issues with duty selfer, it is worthwhile to spend the corts rine and money on steroid products you can be assured care from lessioner depressions.

#### 12) What do the anabolic and androgenic reference numbers under the profile for each stemid mean?

These numbers come from early studies measuring the effect of each steroid on certain mucele and cov organ ticenes of animals negally mice. Those numbers are useful for accessing the relative anabolic to androgenic balance of each drug in humans. They are not as accurate at assessing the total muscle building potential of each steroid, however, and should not be taken as absolute ratings of potency.

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#### Understanding Blood Tests

The abuse of anabolic/androgenic steroids can have a number of potential negative health consequences most commonly with regard to cardiovascular and liver health. These issues houseser can almost always he identified in blood work well hefere always a currence become apparent. Cardiovascular disease, for example, is a disease that can take decades to process. Cholesterol and triolyceride testing can be used to identify and control early risk factors and decisions that would support the disease over time. Liver damage is also generally obvious in liver enzyme tests well before it becomes visibly noticeable to the person. The same holds true for many areas of general health. If you are using steroids, the regular assessment of health with blood work, and the adjustment of therapy when the results call for it, is regarded as the most effective strategy for reducing health risks.

Blood tests with record to anabolic/androgenic steroids are usually conducted in three separate phases. The first phase looks at your health before steroid use. This is done to asses your current condition and risks before any therapy is initiated, and to set baselines for later comparison. The next is on-cycle testing, which is used to assess the direct impact of the ampholic/androgenic steroid use (what the drugs are actually doing to your body while they are being taken). The latter phase of testing is the follow-up, which is conducted to ensure your original state of good health has been restored once the drugs are no longer in the body. We generally refer to these three phases of testing as Baseline, On-Cycle, and Post-Cycle. respectively.

#### Baseline (Pre-cycle):

Baseline (pre-cycle) testing is generally very broad. This is done to make sure there are no underlying health conditions that may be worsened by anabolic steroids, and to have a baseline for determining the on-cycle and post-cycle impact. To begin with, a profile of steroid hormones (male users only) is done to identify the current natural state. This can be especially important to know for post-cycle follow up, as the range of what is considered normal for testosterone on a standard blood test can be quite broad. If you started on the high end of normal, for example, you might want to make sure you are not stuck on the low end of normal following your cycles. A full liver panel is usually conducted as well, especially if henatotoxic oral or injectable steroids are planned. Since cardiovascular disease is one of the most tangible risks with long-term steroid use, lipid profiling is always important, and is usually conducted here and during all other phases of testing, Additionally, other general markers of health are generally examined here including blood, kidney, electrolytes, minerals, glucose, and prostate.

Checklist (minimum): Hormone (Steroid)

Lipids (Standard Full Set) Full Liver Panel

Rlood

# Kidney

Flectrolytes Minerals and Glucose

### On-Cycle:

On-cycle issting is usually conducted 3 to 4 weeks after steroid freezpy began The initialized suggrantify lock at these initiations of health most decay, difficted by steroid use. A fill fluid committon is conducted, and is often regarded as the stegle most important est of fluedath test fluid conducted. The steroid conductation to what these results will began to become appearent. One should give special consideration to what these results will began be become appearent. One should give special consideration to what these results required terms prints over many years. Hipsophetics ching as the began of a fill for period will be committed. It is also recommended to committee other agencies that the markets here such as blood, kidney, electrolyses, minnels, and glacone.

# Checklist (Minimum):

Lipids (Standard Full Set)

Liver Panel, if taking hepatotoxic steroid(s)

Blood

Kidney

Flectrolytes Minerals and Glucose

#### Post-Cycle

During the post-cycle testing plane is common to once again look first at the melt setted hormones. The hope here is to obtain touch not closely finitely our pretentional testine. Note that there will absoys be some variation based on the time of the day, and even in the days-od-up-reads. An exact much is probably for taisable. It is also considered ago old as to look at plantary LH and FSH, because if instinators levels come back look it will give you and your physician a better understanding of the cause. High LHFSH and both to be considered that the second terminal phosphomation may simply include that your testicies have not yet tilly restored their mess. Antennally, but LHFSH can include secondary hypogenation, which is offered used to lattice testined the major which may with an endocrinologist. At not of their general electronics in the contraction of the contraction

#### Checklist (Minimum):

Hormone (Steroid, LH/FSH)

Lipids (Standard Full Set)

Liver Panel, if taking hepatotoxic steroid(s)

Blood

Electrolytes, Minerals, and Glucose

Blood Tests by Category

# Hormone

# Steroid (male)

Test Name	Reference Range		
Testosterone, Total	241-827	ng/dL	
Testosterone, Free	8.7-25.1	pg/mL	
Estradiol	10-53	pg/mL	

# LH/FSH Panel (male)

Reference I	Range
2.5-9.8	IU/L
1.2-5.0	IU/L

# Thyroid

۰	oid.		
	Test Name	Reference Range	
	TSH	.35-5.5	ulU/m
	Thyroxine (T4)	4.5-12.0	ug/dL
	T3 Uptake	24-39	96
	Free thyroxine index	1.2-4.9	

#### Steroid:

Sterour.

This set of teating should look at both total and free testistence. The former measure is must contravely used by physicians to leaftly the arbidopus heed and determine there is a most of the theary flow later measure actually represents the faction of broastables are considered from the state of the state

#### I H/FSH Panel

Literizing hormone (LH) and folicle stimulating hormone (FSH) are responsible for stimulating testisterone production and spermstogenesis in the testes. These measures are most relevant when evaluating the cause and potential treatment options for hypogenulam, not the short-term health impact of anabolic-steroid use. The short-term suppression of LHFSH is exceeded with anabolic-andoncomic steroid administration.

# Thyroid:

It is regarded as important to get a basefine measure of flynoid activity, usually once per pour. Follow up tested during and after storid use may be an expense some view as unaccessary. Antibolician/logonic steroid use is utilicy to permanerly affect flynoid factors, but may sightly elevent imported levels during fleengy. A midiagnosis of hyperhysician (overactive flynoid levels during fleengy. A midiagnosis of hyperhysician (overactive flynoid levels during flood the state into account before teachment for hoverflowed in ordering.

#### Lipids (Cardiovascular)

Anabolican/longuisi estensis can have strong adverse effects on lipids. The abuse of annabolican/longuisi estensis (particularly long-perm abuse) can; Resion; terresus the risk for developing cardivouscular disease as assessed by these variables. Milgaring these risks for developing cardivouscular disease as assessed by these variables. Milgaring these risks and assend-related blood tests. While if from the competitive developing the sense and beard disease risk, a fall panel curaring the variables blook (and computing them to your buseline takes) can provide a good supparbet of the endostoceastic impact of anabolical/longuist storoid use. It is important to measure your blood plats only after 12 millorectives (and off anabolical-longuist storoid use. It is important to measure your blood plats only after 12 millorectives).

# Standard Full Set

inda	ard Full Set		
	Test Name	Reference Range	
	Triglycerides	0-149	mg/dL
	Cholesterol, Total	100-199	mg/dL
	HDL Cholesterol	>40	mg/dL
	VLDL Cholesterol	5-40	mg/dL
	LDL Cholesterol	<100	mg/dL
	LDL/HDL Ratio	<3.6	
	LDL/HDL Ratio Risk Assessment	men	women
	1/2 Average Risk	1.0	1.5
	Average Risk	3.6	3.2
	2X Average Risk	6.3	5.0
	3X Average Risk	8.0	6.1

#### Additional Testing

Auuit	ional losting		
	Test Name	Reference Range	
	C-reactive Protein	<5	mg/dL
	Homocysteine (0-30 years)	4.6-8.1	umol/L
	Men (30-59)	6.3-11.2	umol/L
	Women (30-59)	4.5-7.9	umol/L
	>59 years	5.8-11.9	umol/L
	Apo Ratio Testing		
	Apolipoproteins	men	womer
	apoB/apoA-I Ratio	<.9	<.8
	Apo Ratio Risk Assessment	men	womer
	Low Risk	<.7	<.6
	Average Risk	.79	.68
	High Risk	>.9	>.8

#### Standard Full Set:

This is a standard full field panel examination, Ideally, all values should be kept within the mornal ranges at all times during storold therapy. Note that the LDLH/BLL ratio is regarded the as the most important measure of the serum lipid tests, as it reflects the ongoing balance between plaque deposition (LDL) and removal (HDL) is the arteries. The LDLH/BLL ratio is used to more closely suscess heart disease risk in individuals that have elevated LDL or total cholesteroil levels.

#### Additional Testing:

Anninonal Testing:

Creactive protein and homocysteine are two additional markers that are important to examining cardiovascular health. C-reactive protein is a key indicator of inflammation in the body, and homocysteine is involved in blood clotting and LDL cholesterol oxidation. It is also advisable to include these two articles is more arrithosearcher forms exhelide.

#### Ano Ratio

Apolipoposite intite testing is also recommended. Although not commonly used in general mendial rapactive, pulpoposite intering is retrievably regarded as a more accustare prediction of cardiovascular disease risk than cholesterol testing. Apolipoposite in Equipolis is found in all LDL particles, and is responsible for attending these lipoposities to the artery walk. Apolipoposite A-I (apo.A-I) is found mainly in HDL particles, and is responsible for initiating beneficial review colorated retrieval robotal emission of the artery walk and transport them back to the face. The ratio of apold to apol-AI, therefore, appears to reflect a mark ture measure of the balance of potentially attempted and artistheropasic particles in the blood. Anni above 5's generally regarded as inclusive of termone and artistheropasic particles in the blood. Anni above 5's generally regarded as inclusive of termone coloration cardiom dessense file. Lower artists reflect reduced.

To Find A Testing Lab Near You, Visit: www.HRT-Labs.com Liver Function

Test Name		Reference Range	
	Albumin	3.5-5.5	g/dL
	Globulin	1.5-4.5	g/dL
	Total Protein	6.0-8.5	g/dL
	Bilirubin	0.1-1.2	mg/dL
	GGT (Gamma GT)	<50	IU/L
	ALP (Alkaline Phosphatase)	25-150	IU/L
	AST (SGOT)	0-40	IU/L
	ALT (SGPT)	0-55	IU/L

A fill five punel is important to assessing hepotic strain. The two mufers of five stress most commonly electrated in abuses of almost brindingoings strots its after the express administration control electrated in abuses of almosthoid molegoing strots its after the express administration aministration (AST). All T and AST are recessary to amino and elimetabolism in the Key, and will last its on the bloodstream as the five becomes ridinared or duminged. These two conjunts are generally regarded as important initiations of only storic induced for two tright. There have been causes in which substantial love duming has occurred without substantial electrions in ALT and AST, however, so a more detailed contraints of five recurve values a flower substantial constructions.

Alaliae phosphatase (AIP) and garma-glaunthransportiase (GGT) are known as challestic five company, which mean they demind set steps the 600 of the G agestials that that that did agestion and 8 produced in the bery AIP and GGT are important markers of fiver heath during serond use, and should be included in regale blood tening. Bestories in AIP and GGT can indicate ble duct obstruction (trindepoid colosistics), which refers to a condition where the ever can to longer proport transport and metabolic ble. Intraduction cholestosis is a potentially very serious manifestation of steroid-induced liver toxicity, so electrics in AIP and GGT should not be discegarded.

Mild elsoutions in ALT and AST may be caused by muscle durings (escretics) and not determined the cause of the control of the

# Muscle Enzyme

Test Name	Reference Range	
Creatine Kinase	38-174	u/L

The crustre kinsse (CK) enzyme is used as a mufeer of muscle breakdoon, kinky demuge, and heard dumage, fligh below shapely infeat heard starker or often organ traums. This enzyme can also become elevated with exercise that breaks down muscle tissue, especially interact entrainer or resistance training. Exercise (K. below claused by high extensity training use often ristakelor for organ change. It is important to further commise other munteen or failtings and them beath below for each and destimation is flow and. Now that creative kinnes beefor may also be useful in determing if flow rain or heavy training is the cause of many desired to the contract of the cont

#### Rload

Test Name		Reference Range	
WE	IC	4-11	K/MCL
RBI	c	81-103	FL
Pla	telet Count	130-400	K/MCL
He	moglobin	13-17	g/dL
He	matocrit	40.7-50.3 (men)	96
		36 1 44 3 (woman)	0.6

A fall blood court is one of the most commuty run blood text, and can give you as good support of courted habit in mury regards. At fall blood cits text give you an example of which we have the court (responsible for fighting infection), planels court (vial to blood closing and heading, and red blood cell court (responsible for enrips; copying). Red and white cell courts will be further subdivided its ourious individual measurements, often referred to as a differential cell court. Headings in the supplice maintre of gas in real to cell, and hemsteries in a measure of the precentings of red blood cells in the intal blood volume. Due to their and the register concentration. While the many inveness copyage—range (genels) capacity, as the concentration of red blood cells increases so does the thickness of the blood. Elevated hemster can increase the risk of blent attack or stude.

#### Kidney

Test Name	Reference Range	
Uric acid	3.0-7.0	mg/dL
Creatinine	.5-1.5	mg/dL
BUN	5-26	mg/dL
BUN/creatinine ratio	8-27	

This panel of tests bodes at three primary ususe products fittered and exceed through the kilcheys, urea, rist each, and creationise. Problems here can indicate serious underlying problems with kilchey function. Note that Blood Urea Nirogen (BLN) is often elevated with excess protein consumption, and is used by many physicisms as an indicator that too much protein is being consumed for optimal metabolism. The high consumption of meant or creating assendementation can also elevate creating levels. distribution the value of bodied creatings testing as a marker of kidney health. Electrolyte, mineral, and fasting glucose testing is important to further assessing kidney health, and is advised in addition to the above kidney markers. Aquike time screen for pH, speedite gravity, and the presence of sagar, blood, and ketones is also available at most physicians' offices, and is generally advised alongside blood work when no-selve

#### Electrolytes, Minerals, and Glucose

Toc

t	Name	Reference R	ange
	Sodium	136-146	mEq/L
	Potassium	3.6-5.2	mEq/L
	Chloride	98-109	mEq/L
	Bicarbonate (carbon dioxide)	21-30	mEq/L
	Phosphorous	2.5-4.5	mg/dL
	Calcium	8.5-10.5	mg/dL
	Iron	35-185	mcg/dL
	Glucose (fasting)	70-110	mg/dL

Electrolyte levels are commind to bely detect problems with the flaid and electrolyte belance. Abnormal values my refect someting as mall as sodium or postssim deliciency, or a more aerius condition such as iddrey disease. A variety of other health issues may also become apparent by looking at both electrolyte and mireral levels, sping them somewhat broad prosportic value. Fasting glacone is also esameted to determine if the individual may be be phosphorent. (but both cod supply on the prophycente (thigh both cod supply in Positions with fasting glacone may reflect potentially serious bealth conditions relating metabolic syndrome, disastender may be a supply of the conditions of the desired potentially serious bealth conditions relating metabolic syndrome, disastender may be a supply of the conditions of the desired potential serious desired potential serious desired potential serious delicities and the supply of the conditions of the delicities of the serious delicities and the serious delicities and the serious delicities are delicities and the serious delicities and the serious delicities are delicities and the serious delicities and the serious delicities are delicities are delicities and the serious delicities are delicities and the serious delicities are del

# Test Name Reference Range PSA, serum 0.0-4.0 ng/mL

Prostate-specife artigate (PSA) is a protein produced by cells in the prostate gland. Its leek can become cloude in closes of benign prostate plantage by crosstate charact. Wile a can become cloude in closes of benign prostate phaptrophy or prostate cancer. Wile it remains unknown if clousing the level of antropas in the body with ambedicianthogonic stensivist can increase the risk of prostate cancer, it is known that this disease can be progressed by elevated hormore (andropan and estropas) levels. The PSA test is regarded as an important disposition to for severing in influidal prostate cancer risk. If PSA levels are elevated, rost will abrice against using ambediciantrogenic steroids.

Alanine amino-transferase (ALD: An enzyme produced primarily in the liver but also in other tissues. ALT is involved in amino acid and protein metabolism. Used as a primary marker of hepatic strain. Also called Serum Ghtamic Pytuvic Transaminase (SGPT).

Albumin: The main protein that circulates in the blood. Produced in the liver and has artioxidant properties. Transports certain hormones, vitamins, and minerals, and plays a role in water balance. Used as an indicator of liver health. Higher levels are optimal.

Alkaline Phosphatase (ALP): A family of cholestatic enzymes produced mainly in the liver, but also in the intestines, kidneys, and bone. Used as a marker of hepatic strain, often relating to disease of the bile dues.

Apolipoprotein A-I (apoA-I): A constituent of HDL (good) cholesterol, apoA-I is responsible for initiating beneficial reverse cholesterol transport. This process pulls cholesterol particles from the artery walls and transport them back to the liver. Higher levels are optimal.

 $\label{eq:Apollipoprotein B (apoB): A constituent of LDL (bad) cholesterol, apoB is responsible for attaching these lipoproteins to artery walls. ApoB is a promoter of fairty plaque deposits in the arteries. Lower levels are optimal.}$ 

Aspartate amino-transferase (AST): An enzyme produced primarily in the liver but also in muscle issue. AST is involved in amino acid and protein metabolism. Used as a marker of hepatic strain, although it is considered less specific than ALT testing. Also called Serum Glatamic-Ovalocetic Transamirase (SGOT).

Basophils: A type of white blood cell. Action not fully understood, but cells are known to carry histarnine, heparin, and serotonin. Levels are elevated with allergic reaction and parasitic infection.

Bicarbonate: A measure of carbon dioxide content in the blood, and a common marker of the acid-base balance.

Bilimbin: A waste product made from the breakdown of red blood cells. Excreted into the bile. Regarded as an important indicator of liver health. Elevated levels in the blood indicate liver toxicity.

Blood Urea Nitrogen (BIN): A waste product from the breakdown of proteins, flerred and excreted trough the kidneys. Benedet levels may indicate a narsher of problems including excessive protein intale, kidney duringe, debylation, heart fahre, or reduced production of digestic enzymes. Low levels may be indicative of many things including maltarition or liver duringe.

BUN/Creatinine Ratio: The ratio of Blood Urea Nitrogen to Creatinine, used as a marker of kidney and liver health.

C-reactive Protein (CRP): A key marker of inflammation in the body. Elevated levels may indicate increased risk of cardinasscular disease or stroke

Carbon Dioxide (CO2): Byproduct of respiration, and a common marker of the acid-base balance. See also Bicarbonate.

Calcium: Electrobate involved in a maried of body fractions including bone metabolism.

protein tilization, muscle and nervous system functioning, cardiovascular functioning blood clotting and nutrient transport.

Chloride: Electrolyte involved in the regulation of water balance. Elevated levels may indicate a nurrher of things including amenia, debylaration, excess sait consumption, and the breathroad Low beek may indicate hear or kishev feiture, severe worning, or a nurrher

Cholesterol, Total: A measurure of all fractions of cholesterol in the blood (LDL, VLDL, and HDL). High total cholesterol is regarded as a risk factor for cardiovascular disease.

of other health conditions

Cholesterol, HDL: A measure of the beneficial high-density lipoprotein (HDL) fraction of cholesterol, which helps remove plaque deposits from arteries. High levels are optimal. Low levels may be found in cardiovascular disease.

Cholesterol, LDL: A measure of the low-density lipoprotein (LDL) fraction of cholesterol. This is the primary atherogenic particle, meaning it tends to promote the formation of plaque denosits in the arteries. Low levels are orbiting.

Cholesterol, VLDL: A measure of the very low-density lipoprotein (LDL) fraction of cholesterol. VLDL contains the highest amount of triglycerides. Considered an atherogenic ("bad") cholesterol particle. Lower levels are optimal.

Cholesterol, LDL/HDL Ratio: A measure of the primary atherogenic particle (LDL) in relation to the primary artiatherogenic particle (HDL). This ratio is generally considered the most important cholesterol test value for assessing cardiovascular disease risk. A low ratio is desirable

Creatine Kinase: An enzyme found largely in the heart and muscle, and responsible for

converting creatine to phosphocreatine. Elevated levels may be linked to a number of things including heart attack, kindey failure, or sever muscle damange.

Creatinine: A waste product of muscle metabolism. Low levels may indicate kidney disease, mahurition, or liver disease. High levels may indicate a number of things including reduced kidney function or muscle degeneration. Creatine supplementation may also elevate creatinine levels.

Eosinophils: A type of white blood cell. Similar to basophils, eosinophils are used by the body to protect against allergy and parasites. Levels are elevated with infection, and are low with good health.

Estradiol: The principle active form of estrogen. High levels can be associated with water reteriors, fat buildup, and gancormasia (men). Also plays a role in prostate hypertrophy. Low levels of estradiol may be associated with increased heart disease risk.

Folliels Stimulatine Hommone (ESHE) A ministry hommone involved in reproduction. In

men, FSH is minly responsible for supporting spermatogenesis. In women it supports ovulation.

Gamma-Glutanyl Transpeptidase (GGT): A cholestatic enzyme produced in the bile duets. GCT is involved in solutations metabolism and the transport of amino acids and

Globalin: A blood protein similar to albamin. Globalin is responsible for transporting cortain hormones, fipids, metals, and anthodies. Levels may be elevated in many conditions including clinosi infections, five disease, arthrist, cancer or hapes. Lower levels may be found with a number of conditions including suppressed immune system, mulantrition, multiborophics and five or ckiding-desain.

pentides. Used as a marker of henatic strain.

Glucose (fasting): Glucose is the product of carbohydrate metabolism and the primary source of energy for most cols in the body. Fasting plucose levels are elevated in a number of conditions including diabetes, liver dossess, metabolis syndrome, puncreastis, dieting, and stress. Low fasted glucose levels may indicate fiver disease, overproduction of insulin, laposhrotism, or other diseases.

Hematocrif: A measure of the percentage of red cells in the blood. Low levels indicate an anemic condition. High levels may indicate a number of things including dehydration, increased red cell breakdown in the spleen, cardiovascular disease, or respiratory disease. Arabolic sternik may also increase hematocrif.

Hemoglobin: A constituent of red blood cells, and the main carrier of oxygen and carbon dioxide in the blood. Levels may be suppressed with a number of conditions including multarition, multisoproption, and areamin. High levels may indicate many things including deltydration, cardiovascular disease, or respiratory disease. Anabolic steroids may also increase hemodlorib levels.

Homocysteine: A compound formed from the metabolism of the amino acid methionine. Involved in blood clotting and LDL cholesterol oxidation. Elevated levels of homocysteine initiate an increased risk of cardiovascular disease and stroke.

Iron: Mineral necessary for many functions including the formation of hemoglobin and certain proteins, and the transport of oxygen. Elevated levels may be caused by many conflicies including certain forms of anemis, here damage, hepatisis, is on postoring or vitarini Bo or B12 deficiency. Low levels can indicate a number of things including gastrointestrial blood loss, heavy menstratal bloeding, is on multishorption, or default you deficiency.

Lactic Acid Dehydrogenase (LDH): An intracellular enzyme found in many tissues including the kithey, heart, skeletal musele, brain, liver, and langs. Used as a marker of tissue dataling the kithey, heart, skeletal musele, brain, liver, and langs. Used as a marker of tissue dataling heart attack, ameria, low blood pressure, stroke, liver disease, muscle injury, muscular dystrophy, and pamereatifis.

Luteinizing Hormone (LH): A pituitary hormone responsible for the stimulation of testosterone production in the testes (men). LH primarily supports ovulation in women.

Lymphocytes: A type of white blood cell. Primary role is to fight viral infection. Levels are elevated with active infection. Low levels are associated with suppressed immune system or active bacterial infection (moted by desarded neutrophils).

Mean Cornscutar Volume (MCV): A message of the size of red blood cells, determined

by measuring the volume (rec.) A measure of use as we red undocuted code under by measuring the volume of a single red blood cell. Useful in determining the cause of anemia. Elevated levels may reflect a number of things including a deficiency of vitamin B6 or folic acid. Low levels may reflect iron deficiency, or other causes.

Mean Corpuscular Hemoglobin (MCH): A measure of the average weight of the hemoglobin in red blood cells. Useful in determining the cause of anemia.

Mean Corpuscular Hemoglobin Concentration (MCHC): A measure of the average concentration of hemoglobin in red blood cells. Useful in evaluating the cause of and therapy for, amenia. Low levels may indicate blood loss, BG or ion deficiency, or other causes.

Monocytes: A type of white blood cell. Primary role is to fight severe infection not sufficiently countered by lymphocytes and neutrophils. Levels can be elevated with a number of things including chronic infection and certain cancers. Low levels indicate good health.

Neutrophils: A type of white blood cell, also known as granulocytes. The primary white cell used by the body to fight bacterial infection. Levels are elevated with infection. May be suppressed with compromised immune system or bone marrow.

Phosphorous: An abundant electrolyte involved in a number of body functions including the utilization of carbohydrates, fats, and proteins for cellular maintenance, repair, and growth, the production ATP for the storage of cellular energy, the transport of calcium, the maintenance of osmotic pressure, and the maintenance of hearbest regularity.

Platelet Count: A measure of the concentration of platelets (also known as throrthocytes) in the blood Platelets are included in blood clutting, and protot capitatist consists belowing. Elevated levels may be linked with a marber of fittings including delydration. Low levels are found in suppressed immare system fanctioning, drug reactions, or deficiencies of vitamin B12 or folic acid, or may have other causes.

Potassium: A key electroly, or necessary for nerve and muscle function, and the transport of mariners and waste products in and out of cells. Mong with sodium it helps mainst the actibase balance and control pressure. High levels may be caused by a number of things including kithery failure, metabolic or respiratory acidosis, and red blood cell destruction. Prolactin: A reproductive hormore involved specifically in lactation. Prolactin is sometimes

(but not commonly) elevated in steroid abusers, and may be linked to estrogen excess or hormone inhalance. Elevated prolactin may also indicate other issues with the pituitary.

Prostate-specific antigen (PSA): A protein found in prostate cells. Used as a screening for prostate cancer risk. Elevated levels reflect an increased risk of developing prostate cancer. Low levels are desirable, although do not assure against prostate cancer.

Red Blood Cell Count: A measure of the total concertration of red blood cells, responsible for transporting oxygen and carbon dioxide in the body. High red cell courts are seen with a runther of conditions including heard disease, debylations, or patherousy frobusts. Low levels may be inked to many things including arenis, bone marrow failure, red blood cell destruction beleefine, belearing, and multurition.

Red Cell Distribution Width (RDW): A measure of the variation in size between red blood cells. Useful in evaluating the cause of, and therapy for, anemia. Increased values may indicate a number of thinss including vitamin B12, folic acid, or iron deficiency.

Sodium: An abundant electrolyte necessary for many functions including the maintenance of ownotic pressure, acid-base balance, and nerve impulse activity. Disturbances in the sodium level may be caused by minor things including excessive sweating, vomiting, diarrhea, water intake, or very services conditions includine heart, kilone, or fiver disease.

T3 Uptake: This test measures the level of unsaturated thyroxine binding globulin (a carrier

of thyroid normones) in the blood, increased leves may mineate a number of timing including hyperthyroidism(overactive thyroid), liver disease, cancer, and decreased ling function. Low levels may be indicative of hypothyroidism (under active thyroid), excess estrogen levels, pregnancy, or other causes.

Testosterone, Total: The measure of both unbound (active) and bound (inactive) portions of testosterone in the blood.

Testosterone, Free: The measure of free (urbound) testosterone in the blood. This represents the total amount of testosterone immediately available to tissues.

Thyroid-Stimulating Hormone (TSH): A pituitary hormone responsible for stimulating the release of thyroid hormones.

Thyroxine (T4): The more abundant of the two major thyroid hormones (T3 and T4). T4 serves mainly as a reservoir for the more active thyroid hormone (T3), which helps to stabilize and regulate thyroid supply. This is a key marker of the state of thyroid health (low, normal, or overactive).

Thyroxine, Free Index: This measure is a calculation of the amount of unbound (free) T4 in the blood. This is a key marker of the state of thyroid activity (low, normal, or overactive).

Total Protein: A messure of the total serum protein concentration, minly alburini and poledni. Serum proteins are important to the faction and supply of empress, hormones, nativates, and surbeolies, and also play a role in muintaining the water and pH belance. Low levels may indicate a number of things including multarition, floer disease, multibosoption, deathers, or severe burn injury. Electued beeds may indicate infection, floer disease, multidurints, or severe burn injury. Electued beeds may indicate infection, floer disease, multi-

Triglycerides: The main storage form of fatty acids in the body. May be metabolized and used for energy. Elevated triglyceride levels may contribute to hardening of the arteries (atheroselerosis), and increase the risk of heart disease or stroke. Low levels are ontinal

Urea: (see Blood Urea Nitrogen)

Uric Acid: The waste product of purine metabolism, which is filtered and excreted through the kidneys. Elevated levels may indicate a number of things including gout, infection, kidney damage, and excessive protein intake. Low levels may indicate kidney damage, malnutrition, liver damage, or other causes.

White Blood Cell Count: A measure of the total concentration of white blood cells (also known as leukocytes), responsible for fighting infection and protecting the body from pathogens. A differential measure of white blood cells is usually also taken iraking neutrophils, cosinophils, basophils, hymphocytes, and monocytes. Levels may be elevated with certain infections or allevies conditions

## Harm Reduction / Safer Use Guidelines

Ham reduction is a concept among healthcare workers that seeks to reduce the negative health consequences of ding above. The principles of humm reduction call for an acceptance of the fart flat good or bad, lifet drugs exist in today's society, Instead of ignoring drug users, humm reduction practitives are drug work with them to permitten self-use strategies and decrease the health durings of drug abuse. The effect of hum reduction is abusy, behalty any faight are installed. Albusol pervisorably focused exchanges in a subsequent properties of the production of the production of the programs. In an offert to further this good, ANASDIACS has confident the following Frenches of stood in medication. If followed, these principles should measurably reduce the regative health impact of strooting are a safet (about) not completely subsequences.

#### Principles of Anabolic Stemid Harm Reduction

- 1. Avoid Counterfeit and Undergound Stevish. Authoric strootis produced by life munifications and only on days, and my prosent additional bank his to the user beyond what are presented by the shorols framewhee. Even if favy cortain nactual steroids in properly hisbeld stoots, undergound days may contain took heapy metals, used days and days are presented by the stroots of the parties of the public only by an externely coult, tedous, and metalsical process of quality soarce and government coversift. There is life frameful and cort justical increme for most undergound days makes to produce there though at a which high resolution of the public of the p
- 2. Avoid Toxic Oral Steruids. Aside from Andriol, Primobolan, and Provious, every onal setural discussed in this reference book is a c-17 alpha allohated compound and should be actually discussed possible. While there may be a nurther of clinical nessons to prescribe actually actually actually the control of the provided of the control of the con
- 3. Thirds of Testustrouse Birst. Of all the unbollerindrogue's storick produced, bestudence each less oppositude controlleres and Standoms under his one the Down engagine impact on health when bleen in mucle building and performance-echnology downs provide a hormone silentical to that alwayd produced in the body, presenting the same spectrum of physical and physiologial effects. In addition to being one of the most efficient mucle-builders assubble, testusterous generally has a positive force regarder (effect on Bids), supports a positive most, and supplements increasing sorting and technical technical less had ne less regarderly defined. The exchange test of festionstrone denge for the controller and the produced of the
- 4. Lint's Warrel's of the "Salest" Drugs. If the exclusive use of an injectable isototeree is not feasible, interior got the used group of seturish is ablood. Of the injectable isototeree, the following drugs laws the towest cardivosesclar stim and are recommended. Despeloslesses under joint, and principles large per (reducebone understatte). If an outli section is desired, only Authority Principlesia, or Provision should be used. These drugs are not e-17 alpha algebras of an and provide additional sected activity whether the same level of the common coll answells reducify headed conjunctions areas with other common coll answells reducify headed conjunctions. The conditional principles is a proper sent as even with the common coll answells reducify headed conjunctions.
- 5. Use Health Support Supplements, Antobic learlinguistic attential enter un help hour the regulate health impact of sectorial use with the consumption of martial health appears applements. To begin with the regulate catalities are less of these drugs can be offset (of last to some degoe) with cholstend applements. Find in its recommended as a base, which should be stateded with a number of other clinically satisfied cholstend support probosins. The behind product Light Staff Medically Nursirion Health Staff and the staff of the the staff o
- 6. Always Cycle Steroids. A steroid cycle usually consists of 6 to 12 weeks of drug use followed by an equal period of time or more abstaining from all anabolic/androgenic steroids.

ins gractice is animetal for a number of reasons. For one, as you suppressed mass section hommons your body of include the production of so our selectoriese. Cycling helps reduce the risk of developing long-term firstly and hommonia issues, which are sometimes caused by the unstermental one of servests for runny moreful or years. Cycling alloe lots your general markens of health such as chelestool levels, hermateria, and blood pressure) return to their rowman state procedure, flowed they come to the communities and procedure, flowed great reduction of the communities and the contract of the communities and the contract of the communities and the contract of the

7. Use Reasonable Dosagos. High doses of seroids are not necessary to achieve significant macks good, especially frondered polsage or performmen estimacement is desired. A dosage finit of 400 mg per weed on ispeciables is abised, In the case of teasistence explained, 400 mg per weed, contained to a finite faller led of macke gain if contributed with proper training and diet. In fact, during the 1970s and 806 the dosage range of 200 m0 mg per weed was considered vistantified by the bodylandings of festsosterors, methodore, bolkeone, or methodore. There is actually little end need for externe doses of 29% 1,000 mg per most of secondary breaked, by a supplement an injectable base with additional oral. High doses may produce a faster ran of upit, the area governable extended to the contributed of the

8. Avoid Arountase Inhibitors. Aromenae-inhibito (page contre entogeni side effects by percenting the protation of estoqua in its body. While an efficient pericice, they also deprice the body of a hormone that is important to cardiovascular health. In particular, estogens apports the production of good (EDL, clobester), which means that arountase ribbitons may insolvenely across the cardiovascular stans of a strond cycle. Hentogenic side effects are paperent and a relaction or elementation of the official generally is not considered an option, the SERM Solective Enurgean Receptor Mechanical State (Notable 1) and the contract entomes side effects when the same passing bell in robotated.

9. Get Regular Blood Tests. Comprehensive blood string including an examination of hormones, cholesters blood cell concentrations, and engrous is from test stell tool for assessing the negative health impact of steerd use. Changes in cholesterol, for comple, can help quartify for the user what effect a particular due negatives in hving on their cardiovascular health. The artificials then has the opportunity to better assess leng-term risk if this cycle is to be repeated. At a minimum, blood stellig should be concluded before a cycle is mittacle, 3 to 4 weeks into a cycle, and a cought of mentils after a cycle. This allows constitute to use ends in framing horizontal to the bourse deposite or proposed and a primary and a cought of mentils after a cycle. This allows constitute to use ends framing horizontals in the bourse theory the constitute of the confirmal horizontals in the bourse short of the confirmal formation in the bourse short of the confirmal formation in the contractive of the confirmal horizontals in the bourse short of the confirmal horizontals in the confirmaly

10. Use Proper Injection Procedures. Cardid attention to correct ignotion procedures can help derinate some of the complications associated with normical steroid use. Steroids are given to deep intransactur spicitions. The most common size of application is the upper cutar quadrant of the glatum marker, although the days are also community to the process of the contract of the glatum entire, although the glatum entire the belops, thiope, or calf materials for counter purposes are discouraged, as they are technically more difficult to runsigate and more pose to complication. Centrifishel periodic reclaimable of the contraction of the conflictation to reside and more pose to complication. Centrifishel periodic volumes should also be used, generally nor more than 3 nt. per application. Each injection set should be made to shufe the sum means is not injection and the mine contraction of the conflictation of the conflicta

11. Sterilize. Though never advised, should the choice be made to use an injectable steroid of underground origin, an effort should be made to sterilize the solution before use. This will reduce the Relibod of illness or infection due to microorganism.

12. Watch Your Diet. Anabolichandrogenic steroids can allow an individual significantly more latitude with their diet than normal. The caloric demand typically increases due to the effects of these drugs on muscle mass and metabolism, allowing more calories to be consumed each day without adding fit mass. It is important not to let this latitude affect your health in a negative way. Remember, the use of strooks at physique- and performances.

orbancing doses is expected to cause an unforombs drift in cholstered levels and other cardiovascular health markers, fronting a higher risk of enatiovascular health markers, fronting a higher risk of enatiovascular disease. Similarmously feeding your body geneter amounts of statunted fits, cholstered, and simple suggest are recommended, and are known to reduce confrouscular disease, the control of the contr

13. Always Consider Revund AVD Risk. It can be easy to ignore the potential health impact of stored use to when the possible benefits are so rapid and the negative consequences so remote. At the end of the day, however, it is very important to remoterbrift that the use of storoids in doess efficient to approx that them makes glain as wittaily always going to laves some negative impact on your body. Four chooksteed will shift in an uniforcable direction, your body possesser may go pa in all be land you may see too slightly fischem the ventricks in your heart. Your hormoness are out of balances Sential use in one of pages to the ventricks in your heart. Your hormoness are out of balances when you take storoids, which will stimularly seen soft enfings to go not forther seen. Sential use in one dispersion of the contribution of the story of the properties of the sential possible of the

#### Sterilizing Injectable AAS

Authoris street products mude by Kensed drug munifications have an expectation of statisfly. This assume the first government reglations applied to these companies. Unfortunately, there is to such expectation for constricted and underground streetly products. Most are assembled under conditions that are not stretch. Our may consecut, subspected laboratory tests have floated bacterial contamination in products of flick origin. These also appears to be a digibly indeer preculence of abscess referrious and order injection-selfcomplications with integround streetly, further traditioning cause for covered. Given the complications with integround streetly, further traditioning cause for covered (first this config.).

#### Heat Sterilization

The simplest method to clear a storoid solution of microorganisms is to best steriles it. The spical step to accomplesh this is to place the storoid skill am one of 700 minutes foldes of 12 DOC/25/PF). You should leave a meedle sticking frough the ribber top to comaine the pressure inside the vide of r may break. Note that a higher temporare and datation are typically used for complete day lear sterileation, but this is not practical five our pursoes that to the potential department of the components, and possibly own the ring components. Skill, the recommended 500 minutes at 120°C/250°F should kill must bacteria, and can be promoted as a which them reduction storosts.

Some people profer to use the top of the stone for heat sterilization, and saturage their annulos stanch with a part of bruing used (OUCCV22F). The wisk lexy in the boiling souther for approximately 30-30 minutes. While this temperature is sufficient for killing many from of Dataction and other microogenisms, it is permuly not considered the best method of sterilization. A higher temperature should be used. If can also be difficult to control the temperature if the value stone of the bottom of the pay, which is in direct content with the humer or best source. Given those problems, it is preferred to attempt best sterilization in an own an afformed of the sterilization.

Note that teerholone appears to beauk down at fairly low heating temperatures.<sup>77,18</sup> Is degradation increases with temperature and duratins. Some degradation may be notable even at 100°C212°F. As such, treabolone products should be heat sterilized under slightly different conditions than those recommended above. This lepitually involves having the vial in an oven for two hours at 80°C175°F. Again, an unattached needle should be placed in the rubber stopper to cogatile pressure.

#### эупше гипации

Another way to attempt sterification of a serood solution is to use a flee. A flee is a device that uses a membrane with in plots to preser particles from passing through The smaller the looks are, the more effective the flee. For this method a disposable syring file is used. These are usually but an if is-friencin and 22-in-incon integrous vise. The 22-incine file is in preferred, as it always for the remotal of almost all flems of bacteria. A 45-incine file is also acceptable, thought more filed to post central strate of distinsive betterin. Syringfiles can be outdoor distribution and the strategies of the strate



#### .22 micron Millipore syringe filter

The procedure for using a syringe filter is fairly simple. First, make sure you have all of the recessary equipment; a syringe, two needles (minimum, a statch syringe filter, and a new scaled startle vial 1F/you are attempting to sterilize a standard 10 rol vial, it may be preferable to use a syringe large enough to draw the fill solution. This will reduce the exposure of your producting a filter of the control of the control of the control of the control of the producting and the control of the

Begin by drawing the stead solution into the syrings. When you are done, enrowe the mende and antick your newly unsurposely pringing flow. Attach as new model to the end of the filter, and sight the solution into the new sterle will. The sterle vial and new needle are very important, as single in original components would only expose the product to bacteria again, the energy pressure to entry the syrings into the vial. Be careful as too much pressure can be entry the entry the syrings with the vial. Be careful as too much pressure can be entry the entry the entry the syrings of the vial. Be careful as too much pressure can be entry the very large entry the e

Also, these straination methods are applicable to of-based steroids only. Host straination (so outfined here) bette to cause the repid conportation of solvent-astrainational agents in water-based steroids. The assperded (instead of dissolved) matter of the particles also makes them likely join an ayange filter. Provious testing suggests there is a high resoluter of bacteria in contentia and underground water-brook steroid, series of the resolution for the content of the content of





Syringe filtration of steroid solution. Use of new (sterile) vial reduces chance of recontamination

#### Limitations

It is important to emphasise that when applied in an uncontrolled emissionment such as a house or apartures, there is always the possibility that you may insubstrently control assistant your own product when attempting to filter it. You should always use storic lighous and clean multi autriaces with adoled or belasch before working on them. A must is also a very good idea. Note also, these startitudion techniques help only with contamination by the contamination of t

Many forms of bacteria also produce endutories or pyrogens, which would remain in solution over after fluration or heat sterilization. Thus, you can rathee lever or other adverse executive core with or while bacterie being left in the solution. An additional issue (broadvery stakes) is the cortain reconseguisties (teather these instruments and locarity) are either than the state of methods described here ore highly effective, it can be consequently only for the lightest bend of stays, it is always recommended to limit your use to lightness phramaceutical.

374. FDA Environmental Assessment Report, Finaplix R (trenbolone acetate). April 1987. Hoechst-Roused Agri-

Vet Co. 375. Underground ANABOLICS. William Llewellyn, Ronny Tober. © 2010. Published by Molecular Nutrition, Junior. Fl. 33458.

#### Counterfeit Steroids

As the mure implies, counterful strouble are copies of real ambiokandingsing strouble products, which are made form filed products. Those drugs are intered for size in the black mustler, themsely to consumers who believe they are bosing a legitimate pharmaceutial fame from the blacked consequent. Here are must prosperate insues to consider pharmaceutial fame from the blacked consequent. Here are must product in sous to consider operations. The contents of such products, by the very nature of these operations, cannot be vortified. In mury cases, the contractives to will never even use any mate steady of all and wall simply soft an expensive and consentably wordlaws both of next carnots. Blees, and because the soft of the contractive of the contr

Bost if a counterfet does contain the active stead in quastion, it may not be a clean and safe product. Today, we take for granted the first thar of any groodstea are made with steals and pure ingredients. We also give list thought to asepte processing techniques, which manufacture out they products for contamination by humble descries, views, to mixroorgarism. This is especially important with rijectable medications, since many of the fururam body for most defected gained defected gained the descries against effected in each gain at the product are arranged, and many for most participations of spiceting a contaminated drug product are arranged, and many for sample product are furnished. The product are arranged and many for sample products are furnished and the product are arranged and the product are furnished. The product are arranged and the product are furnished and the product are furnished and the product are furnished.

As anyone that studies the illicit anabolic steroid trade knows, high demand and hage profits offer storing incertise for the manufacture of counterfiel drugs. Over the years, this segment of the iligad bissues has grown exponentially. What was once a problem largely slouted to the United States can now be found in literally every corner of the globe. Counterfring is a determinant of the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the counterfring in the counterfring is a contract of the counterfring in the counterfring is a contract of the counterfring in the cou parameters in a source or surrow, sectors, or corner, may a commony sect wear incesignificant sales of anything valable are diverted from legislate to undergoard sources. Given the matter of drug products, however, the constricts steroid phenomenon as an opeculal proportion health concern for direct special steroid phenomenon as an opeculal proportion health concerns direct special proportion of the products grows, so to does the number of reports of low or no product efficiency or worse, built consequences relating aboves, directs, analytics, and toxicly canced by heavy

#### Prevalence Study

By all estimates, couracterfe streoxis are very common in all common of the global streoid block market. Glome the likelit matter is which contracted streoid products are matched, however, it is difficult to determine the exact prevalence of these drugs. Analysis reports of live enforcement seatment offer an excession analysed to offer again of stored sales in a particular region. One such study was conducted at the Center for Preventative Depting Beauch in College, Centrang, and involved off different analysis esteroid and articles drug products. All of the complex unshould were obtained uting policy market tree light and the complex of the drug policy and the complex of drugs the ground of the complex of the comple

Owenl, more than one-finit (44%) of the 50 ambolts storoks tosted dit nor how imported that method their below, and west early made from life manufactures. Of the fining products, rine were identified as copies of known phenmecuteal brands, and would be considered classes; countrieds. These mining eight (16%) that finied were undergooral steroid products, which are discounted synathy, with a book. There were said dealers. The remaining eight (16%) that finded were undergooral steroid products, which are discounted synathy with book. There were said defining invokates on the first that passed lessing thin were made by undergooral munificatives (Brital Dagon, SS) Lake, and Immensional Frameworkshid, in book, There were said first approach in the first passed of the state of th

#### Visual Inspection

The resembers is Cologue Cerminy also made an important observation, what from inconsumed promotes from these state a British Diagou, SE Lale, and Intermitted Planmacetrials, they noted it was not possible to secretian what product was real and what or reference materials recessing to mise an ap-close product constraints, and no product photos were provided in the report to reference, it does underine a problem that the steady appears are to be a possible to the steady produce and the product of the product of the product of the product of the steady operation are becoming increasingly supplicated. Now more than ever I can be difficult operations are becoming increasingly supplicated. Now more than ever I can be difficult on the product application of the steady of the product application of the missing the missing the product application of the missing t

#### The "Best" Products

Of the conflicted German products, those that were manufactured in Western Europe sessioned to office granter assumer oo legislarity than those of other points. Talkard also termins a corrumn source country for legislants products not common instanctured in Western Europe installarity conventions and methandroscottones. This is in great most likely to be the subject of contractiving. Also, the study aboved that the less condy-testistence products were most likely to be glaritude, early they office free dealers, a good deal of legislants. Extracted Sustances and Egyptiant Institutes countrillar are being proported institutes of Europe. It supposes that, at least by vasy of these free dealers, a good deal of legislants. Extracted Sustances and Egyptian testistences currently are being proported institutes in Carrier State of Sustances and Carrier State (Santon and Egyptian testistences currently are being proported institutes) or cylorized in addition to the blebel currently are being proported in Sciences (Santon and Carrier State Stat

#### The "Worst" Products

Perhaps due to high recognition and demend, all of the Normal Helian rundrokes decanouse products tested during his analysis in new determined to be counteful. These products were confiscated from each of the three deutes independently. In all cases, these stroid products contained its dissestence resisted of rundrokes decentated. This is a current substitution with deviaer numbroker products, as low doese of lestostence can provide a similar level of analotic elect as numbrokes of servan uses, with a similar low includence of rundroken decentate. This is a current as the contract of the contract of

#### Other Bodybuilding "Ancillary" Drugs

A total of 20 mon-tenoid drugs were also tested. All products that would be defined as common ancillarly despis including turnoller intentic (Nooleades, Chelsphere dested (Credit, thytoid Inomore, caffere, and yolimbrie hel furned out to be legimine. This suderlies the boser risk in these articly ordige extre. On older date to the lower frinceal incertific for controllers to depicate these cheep and easy to access planmecarteris. The only non-second drugs where there was seen as solitation noted were in the mile second performers category, which constitute drugs such as Vigna and Calls, in most of the individual cases the drugs did set out as a blocked. When they did full entity, however, it was smally for the drugs did set out as the blocked. When they did full entity, however, it was smally for the drugs did set out as the blocked. When they did full entity, however, it was smally for the production of the second production of the controllers of the church as well.

# Steroid Analysis Results

Anadrol (oxymetholone); 1. Oxytone 50 mg (SB Labs, Thailand) Result: PASS

2. Oxytone 50 mg (SB Labs, Thailand)

3 Oxytone 50 mg (SB Labs Thailand)

# Deca (nandrolone decanoate): 1. Norma Hellas (100 mg/mL)

Rocult PASS

Result-FAIL (testosterone)

2. Norma Hellas (100 mg/mL)
Result-FAIL (testosterone)

3.Norma Hellas (100 mg/mL)

Result: FAIL (testosterone)

4. Norma Hellas (100 mg/mL)

Result: FAIL (testosterone)

5. Decabol 250 (British Dragon, Underground)

S. Decabol 250 (British Dragon, Chaerground) Result:FAIL (testosterone)

Dianabol (methandrostenolone): 1. Anabol 5 mg (British Dispensary, Thailand) Result: FAIL (methyltestosterone)

2. Anabol 5 mg (British Dispensary, Thailand) Result: PASS

3. Anabol 5 mg (British Dispensary, Thailand)

5 Danahol DS 10 mg (March Thailand) Result PASS 6 Nanosim 5 mg (Terania Romania) Result: FAIL (methyltestosterone) Equipoise (boldenone undecylenate):

Halotestin (fluoxymesterone):

Result PASS

Recult PASS 4. Danabol DS 10 mg (March, Thailand) Result PASS

1. Boldabol 200 (British Dragon, Underground) Result PASS

1. Fluoxymesterone (IP, Underground) Primobolan (methenolone enanthate): 1. Primobol 100 (British Dragon, Underground) Result FAII (nandrolone testosterone)

Result PASS Sustanon 250 (testosterone mix): 1 Sustanon 250 (Karachi Pakistan)

Proviron (mesterolone): 1 Provinon 25 mg

Result PASS

2. Sustanon 250 (Nile, Egypt) Result: FAIL (different testosterones)

3. Sustanon 250 (Nile, Egypt) Result: FAIL (different testosterones) 4. Sustanon 250 (Karachi, Pakistan)

Result PASS 5 Sustanon 250 (Karachi Pakistan) Result PASS

6. Sustanon 250 (Karachi, Pakistan) Result: PASS

7. Sustanon 250 (Karachi, Pakistan) Result: PASS

8. Sustanon 250 (Karachi, Pakistan) Result: PASS 9 Sustanon 250 (Karachi Pakistan)

Result: PASS

Result: PASS

Testosterone Cypionate: 1. Testex Prolongatum 125 (Q Pharma, Spain)

2. Testabol 200 (British Dragon, Underground)

Result: FAIL (différent testosterones)

Testosterone Enanthate:
1. Testofort 250 mg/mL (Pliva, Pakistan)
Result PASS

2. Testosterone Depot 250 (Eifelfango, Germany)

3. Testosterone Depot 250 (Eifelfango, Germany)

Testoviron Depot 250 (Medipharm, Pakistan)
 Result: PASS

5. Testoviron Depot 250 (Medipharm, Pakistan) Result PASS

6. Cidoteston 250 (CID, Egypt) Result: FAIL (includes T. cypionate)

7. Cidoteston 250 (CID, Egypt) Result: PASS

Testosterone Propionate: 1. Testovis 100 mg/mL (SIT, Italy) Result: PASS

2. Testovis 100 mg/mL (SIT, Italy) Result PASS

3. Testovis 100 mg/mL (SIT, Italy) Result: PASS

4. Testovis 100 mg/mL (SIT, Italy) Result PASS

5. Testovis 100 mg/mL (SIT, Italy) Result PASS

Testabol (British Dragon, Underground)
 Result: FAIL (different testosterones)

Trembolone (various esters):

1. Trembol 75 (British Dragon, Underground)

Result: FAIL (boldenone, testosterone)

2. Trenabol 100 (British Dragon, Underground)

Result: FAIL (boldenone, testosterone)

3. Tri-Trenabol 150 (British Dragon, Underground) Result: FAIL (trenbolone, testosterone)

Trenabol 200 (British Dragon, Underground)
 Result: FAIL (trenbolone, testosterone)

Winstrol (stanozolol):

1. Winstrol Depot 50 mg/mL (Zambon, Spain)
Result: PASS

2 Winstrol Denot 50 mo/ml .(Zambon Snain)

Result: PASS

 Winstrol Depot 50 mg/mL (Zambon, Spain) Result: PASS

4. Stanabol 50 (British Dragon, Underground)

#### Counterfeit Steroid Identification

This section portains to methods for differentisting between beginning paturnecetical products and dispirate copies (counterfields). Before we begin, I read to mined upon the counterfields are about a very learning between these days. Counterfields are sensing a lost of money ju printing and packaging copienters on the void lines in land time counterfields operations, with the capacity to munificative all product formus rickling amplieds, logs jurispirate gills, and push-brings thirds strips, from the light learning the land of supplications, stronks purchased on the black maker need to be respected with great care. The installace made by constructions are often muse and mote of the fix not for the strips.

#### Sten #1: Eliminate the Obvious

When countried strovils first appeared decades ago, they were often very easy to spot. The munifications operated on a suml seal, and mode suml-scale installates. For example, the prairing might be shoppy, or the containers than and finney. They might have lacked the eagersteal to get the product in a low, or even than examplement and be number to 1 to example the product in a low, or even than examplement and be marked to 1 to lowester. Fow contextificates still make the basic installates that were once common. Don't expect identifying these products to be easy. Soil, that is not to say the obserse contextificates and it available. Indeed, they can be found on the black market from time to time. This first of the context of the c

- 1. Sloppe Printing. Drug munificating is not a small scale endenote. Sizoble pharmacenical compraise control the plobal lung tanke, and make products that are typically very professional in appearance. Visu should not expect to see things Bue many inks, aloppy incert, see missings of images on neal drug producing Scientisms controlled so all some of the printing and respondents metalods, which make labels and bones that seats of the seed of the printing and respondents metalods, which make labels and bones that seats of the seed of the printing and printing seat of the printing and printing seat of the printing seats of the pri
- 2. Cheap Packaging, Virtuly all legitmets stored products come in boxes. Inside the box you should find a day information sheet. Some contrections will all pines septo emission, Real ampulses, visik, and tablets are sometimes smuggled boxes, but it is someone else table to the the first. The box for a pinumencical product should be sustained; some, cleasing qilly and distributed to the strength of the pinument of the pinume



The ampule is a laboratory blank, meant to be sealed by hand over a flame. It is larger and more unusual in appearance than most traditional ampules.

3. Multi-dose Containers. In the United States, we are used to our spectable medications coming in multi-dose with (these have a robber up to let medical posses through more flown more), and our pills hose in bottles. Most other coursies, however, the rot all the opposing for human medicine. They constant is unstead, and permit it only for similar possessing for human medicine. They constant is unstead, and permit it only for similar glass ampales for rigicatible medications, and post fireogh bilaters for pills and capuales. Sizes quara utilized to fair fail advancine modests on the lather intends, it may be best or could all multi-dose containers when it comes to human parameterista. Most are going to be counterful. When you full velocities up the introduce containers, earth can offer the content of the proper to the velocities of the introduce containers when it comes to human parameterista. Most are going to be counterful. When you full velocities up the introduce containers, earth can offer the proper to the content of the proper to the velocities of the proper to the content of the proper to the velocities of the proper to the



The above is a crude copy of an American testosterone product, which uses the same label on the box and vial. A counterfeit as simple as this is rare to find today.



Another example of an obvious counterfeit. This box is crude in design and uses a brand name that has been off the market since the 1980s.

The photographs below show what it looks like when the lot number and expiration date are added after the initial box/label printing, as well as counterfeit products without this feature.

The characters on a real planmontrial product should stand out from the rest of the printing, which will consist of they does belond to agent to create a old image (see Sep. 87 a for more information on the sick. When the let/espiration information is added with a mechanical starp, the isk will be men through social forced to a physical related you can be collected as a physical related you can be observed to the proposal content of the production (note that it may appear blottly under deep magnification). Depending on the equipment, it may also lowe left a physical indext you can be observed to the production of the other indext on the ware added by computer, we usually see large does that are visible to the mixed eye. Be careful to look at the characters closely. Counterfeltens will by you finder the information look like it was added by mechanic or computer, one thought was printed. If you see that they does make up the characters under 200X mangliation, it is not legalizate, also are legalizated in the characters and a constraint of the printing of the characters under 200X magnificant, it is not legalizated under subject to the characters and the character when the character is the constraint of the characters and the characters are constraint.



lermo



\_\_\_\_\_



200X

Example #1. A real box of Proviron. Under magnification we can see that the lot number and expiration date were stamped on mechanically.

#### Step #2: Examine Lot Number/Expiration

A more formal analysis should always begin with the lot number/ceptizion date. Phermacential compress have their boss and label numberfacent in bulk, usually and officing printing facility. They are not sensition, let number-ceptation dates have not yet been compressible printer at the trust the days, and the compressible printer at the trust the days, peakaged. Contractions often don't was a days simply print the let number/ceptizion date with the rest of the boss and labels. This means been work, less captioner, and less cost. Knowing the, centraining the lot number/ceptizion date information can be a good vary to spec construction. You must be lot lock at the lottless returnation way to chelop, predenably with a fundhed increasion with 100-2005.



Example #2. Another example of mechanical stamping of the lot number and expiration date.

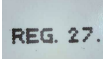




Example #3. Real testosterone cypionate from Watson (U.S.) The above lot/expiration date were added by computer printer. Under magnification we see the large dots are solid ink.



Norma







Example #4. Another real product (Proviron) with information applied post-printing with a computer printer.



Example #5. (Counterfeit). At first glance the slight run on the ink appears to be the result of mechanical stamping. Under magnification, however, we see this is simulated.





0/2008 6,72€

203



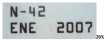
200X

Example #6. (Counterfeit). An example of simulated computer lettering. You can see under magnification that the information was actually made by normal process printing, along with the rest of the box.



Vormo

LOTE:/CAD





Example #7. (Counterfeit). Again, the date was added with the rest of the printing. This manufacturer simply made the text bold in an effort to hide the printed nature of the information.

# Step #3: Look at the Ink

One of the first things that the U.S. Secret Service does when it takes in a constructed bank once is to examine the issue such byte constructions. They knowled does ago that you can self a great deal about a bill this way. You can tell how and when it was printed, sometimes even where. Now we are not gaing to analyse the issic on steriol boses in the libs as the Secret Service does. We will, however, look at the printing closely enough that it can help directions of the contraction of the contraction

First, It's go over some general information so you know what we are looking at For this countration we intend to find only to the printing uses placed on the box or tabled. Close printing is usually accomplished through two methods. The first is called "process color," and the second "spot color." One method may be used to print the product, or both methods may be combined on the same packaging. Each method offers is own set of advantages, When we know how and when these methods are typically profice, owe may intend in When we know how and when these methods are typically profice, over any insome single into the thoughts of the designer. More importantly, we might be able to tell if the methods are ammorning for the modular in assession.

The process color method is most often used to reproduce photographs, or to print malphe colors (three or must often used to the most finelish and usual) deep method of coloring. Process color can give the appearance of harderds, throusneds, even militions of cooks are cent time, allowage thereinght his is an indisact to pomittened cooper committen, or will see that the colors at the circuit adjust thereinght his is an indisact to pomittened cooper committened, you will see that the colors at the colors and beautiful colors and the colors and the colors and the colors. A color of the colors are colored in the colors and the colo

The spot color method is used when a very specific color it is required. Instead of using a CMYK behat of reproduce a centris sheld of great, the cample, an it is rinced in the provide the exact green color that is needed. If you look at spot color under 200X immigration you will not see CMYK color for Sen Smill data are still used, but flow gree at the same color (thy any placed closer or infrier quant for shading). Note that it can be very considered, and the same color of the color of the color of the color of the considered, Actual Color mething, spottern answers the dissell color is exceely what it applied to the printed packaging every true. You will usually find big companies using spot color, especially with important corporate destinities (soon, special mages, Indumely.) The min thing, we are hoping to see under magnification is the use of some spot cole. While the lack of spot cold does not necessary mean that a product is counterfiely, you should expect to see it on most real planmaceutical packaging samples. If the packaging issnesses the or three colors, they may use a foundation of process cole, but add spot color to certainly for the packaging its most time three colors, they may use as foundation of process cole, but add spot color to certain key counterfielders risks cannot always against some amonto no dead, which sometimes the counterfielders risks.







nov

Example #1. This box of Depo-Testosterone was printed entirely with spot color. Large companies like Pfizer tend to prefer this method of printing. Note the unified dots under 200X magnification, indicating spot color.



4









Example #2. The above underground steroid product uses process color. The small CMYK dats that blend the color can be seen well under 200X magnification.

#### Step #4: Magnify Logos/Small Elements

Each counterfeiter is faced with the dilemma of duplicating an original steroid's packaging on their own equipment. The most common way to do this is to conv the flattened box or label with a desktop scanner. The picture will be cleaned up in a computer, and ultimately converted into plates for printing. In this process, the drug packaging is duplicated in what amounts to a complicated form of photocopying. Though a seemingly simple process, drug nackaoine made in this manner can be summisingly accurate in annearance. Some are done so well in fact, that you'd be hard pressed to nick out the legitimate box from the duplicate without a little guidance first.

This type of counterfeiting does have its limitations though. Most notable is that very fine detail is usually lost in the reproduced image. Much of this is due to the scanned in image being pixilated, and thus not amenable to scaling and graphic manipulation in the same way the original graphic would be. When examined under magnification, the lines may agrees choppy. Make sure you look at the small graphic elements of the steroid package, not the text. Counterfeiters usually rewrite the text so it is sharp. They cannot retype a company logo, illustration, or image. If these graphic elements look "sloppy" under the microscope, the product is probably a counterfeit.









Example #1. Real Cidoteston from CID Egypt.







Example #2 (Counterfeit) Cidoteston. Under magnification we notice that the scan of the initial box was insufficient to reproduce the fine detail. The inner ribbon around the logo is especially broken, and the letters "CID" poorly defined.







### Example #3. Real Sustanon from Egypt. Notice the fine detail on the snake.



Normal

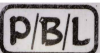




200A

Example #4 (Counterfeit) Sustanon. Notice how the finer details are lost on the duplicate.







200X

Example #5. Real Deca-Pronabol from P/B/L, India. Notice how the logo remains smooth under magnification.



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Example #6 (Counterfeit) Deca-Pronabol from P/B/L. Notice how the logo is choppy under magnification, likely due to the image being scanned from an original box.

#### Passing Inspection

It is important to once again emphasics that storoid counterfairing is high basiness. An efficient operation can not may million of dollate per year for its owners, which so much money at stake, opsishication is at an all time high. In some cases, the counterfaires will have very similar high-level qualifiest to the pharmocraft convenies they are copying. This remains not only the ability to reproduce complicated packaging such as break-open gloss ampales and push-frough high straight seat, but all package gas at last service depending the straight packaging the first package of the control of the c

# Country Specifics

In most courties, a pharmocentical corapsus is required to most a specific set of regulations pertaining to the physical spackaging of a pharmocential product. In some cases those regulations can be used to help evaluate the legitimety of black market steroids, as all counterfeiters may not have the reservoirs of forthought or implement the required features. Here, we discuss a number of attributes to examine, which should hold true for all of the druns produced in the specified courts.

#### United States:

First, it is very important to stress the fact that steroids are a controlled substance in the United States. Current controls are very effective at keeping substrained products off of the black maket. It is much easier for the fact dealers to import or munificative their own products than it is not grain volution of ligarimet Americam plantementerates to distribute the keep of American items you excount or the black market, as they are in all probability and the products of the products and the products of the products





The FDA provides us with a couple of strict requirements, which many counterfeiters overlook. The most predominar is that all algainst American drugs many carry a libel that will easily be removed from the vial-bottle. It must be so startated with give that you can only remove it is mall pieces. This is done to protect the public from the possibility of drug mishibeling. With many U.S. counterfeits, the label can be peeled off the bottle quickly, in one or a few larne pieces.

You should also moisten your thumb and rub the expiration date on the box and label. Quite often the ink on the counterfeit will smear and rub off easily. The stamping on a real U.S. pharmaceutical may streak slightly, but should remain intact and legible. Again, this is a requirement to protect consumers.

Additionally, being a Schadule III controlled adstance, all commercially available harms and vectorinary anabolism-foregenic steroids are required to bear the tag. "CIII" (see sample picture). The only exceptions would be cattle implart pelets, which are technically not controlled adstances, or drugs form compounding planness; which do not have to adhere to the same production guidelines. A small number of lizey counterfeites contrine to duplicate steroids that were manifestanced before 1901, when this gave not present.

The FDA requires that all tables and capsales are identifiable through unique markings in case they are remored from the packaging. He manufacture must on an albeviation, is usually found on each pill, along with a specific code for the product. Some secret users have been all the pill and the pill and

#### Italy:

All drugs produced in Italy will bear the pictured drug identification striker. The striker reself is while, with red and black pirt. The striker rest on a limitant startifice, so that it can be possed to flart a fifted to perservoik. Vou should nover purchase an Italian drug if this striker is not present. Durgs from Italy will also use abbreviations fix Perp. Sect. and Del for the counterpart of lot it, manufacture, and expiration dates. Engish writing here would indicate counterfeitine.



#### Greece.

Greece also has a drug ID striker that must be present on all drugs available for sale. The sixtler inserting an alministic striker, so that it can be peeded off off the box and affitted to paperwork when a prescription is filled. Most importantly, the sixtler will show a hidden must when placed under UV light. Some counterfeiters have copied those sixtlers with excellent accuracy, right down to the laminated surface. Copies of the sticker bearing a likelia. The Copies of the sticker bearing a monen OV watermark, nowever, nave not ocen tocated. Lo not purchase any Greek drug without the proper sticker attached





#### Spain:

Sparish drugs do not bear a sticker, but instead have an area located on the box that contains a bar code and some drug information. This area will sometimes have indentations in the cardboard, so as to be removable if you tear the surface. At other times, the betrood is simply printed on the box. Sparish drugs also use the abbreviations Lote and Cad for lot marber and expaintion, respectively, date. Many drug boxes also carry Bealle lettering.

# C.N. 834234 EXO



PVP M.R. 690 Pts PVP IVA4 718 Pts

#### Printed Only

A.S.S.S.
ORGANON ESPAÑOLA S.A.
DECA-DURABOLIN 50-1 AMP.
C.N. 740480



Removable Barrode



Braille Lettering

#### Portugal:

Dag boxes from Portugal contain a rectangular area which displays the bar code and pricing information. This is sometimes found as a stricke, but most commonly it is printed, not stamped, onto the surface. In many cases, the area is indented, so that if can be removed from the box. Drugs from Portugal will also use the abbreviations Lote: and Val. Ate: for lot marber and expristion date stamping the surface.



Old one without



New area with price in Euros

Drugs from France will bear a rectangular sticker somewhere on the surface of the box. The text and format is often slightly different item to item. Also, packaging always contains an area with a owner and not how. In the surmed below it is in the lower left side of the box.



#### Underground Steroids

An urdergroand stroid is an anabolicandoguis stroid product that was much by an filegal (undergroand) behension. These days are specifically municitated for site to athletes and bodybalders on the black murket, and are not available through legitimate channels such as plannatices and drug distributes. These compress are unferenced, unregalated, and operate in a completely clusthesize manner. At one fire the term undergroand started was considered spranness with contractives of the today may people view these two categories as separate. The main point of distriction between an undergroand seriod and a contractive stores in further later is copy of a legitimate contractive of the contractiv

Tigher government enforcement of stevoid loss and legitimet distribution charace has fateled an explosor of underspooral steed of members are governed. The bosics model is now thing comment, and may account for most than 5% of all popular comments are consistent or more than 5% of all popular photoses between covering of me steed mit mention for concerning whose steel registrates concerning the manufacture and sale of dang singulations are not in place. These impositors are then snaugade into an area of light pregadents area not as in place. These impositors are then surgued into a mass or light pregadents much in the little little state or Europe, where they are made into includated units units (usin, ampake, pill bortis, or belier cataly) with the office of surgest containing the contraction of the products. So the operation are units and contracting the contraction of their products is storyed and the contraction of the contrac

number though, and account for a small portion of the total number of underground laboratories.

# Drug Purity

Dag munificturing usity is a certaal focus in Western medicine. Planumeetricals are intended to not all pinteis, not cause additional lamm by being impropely does of containing bestern, heavy metal, or other forms of containing the intended to the all proposed many contained and produced by government-leveral companies, contain materials that come from other levened suppliers, which also address to stret phenumeetrical-page party standards (send as LSPRP). These companies must also assemble their products in meticulously sentitioned "clean moon!" facilities designed to prevent any contamination from and and preserved. Each metal of price of opiquent after context and applied of the metal of price of opiquent and the context of the

The above description is in stark contract to the undergoard steroid munificitarity business. By their very nature, these companies are not rainle spoterment know or oversight A contraction of the property of the property of the contraction of the contractio

# ANABOLICS Underground Market Analysis

In an effort to help consumers assess the quality and potential beath risks of rudergoard storein products, ANABOLICS materiots to detailed joir ridt agains jos project in April 2007. This project examined the quality of steroids much from undergoard facilities, and exceeded for normal copy of festing by comming a number of four variables often exceeded from the contract of the c

These were four specific senses of testing for the 2007 meters amplies project. The first test was to look for the presence of twoic heavy meths such as last, it is recovery, and sensels. These metals all poses specific threats to health if they accumulate in the body. Those metals considered next such sein and admiration selfs to see how these products were often their checked testing of the sense of a sense of the standard sense of a sense of the sense of the standard sense of the sense of th

The specific results for each of the first testing sets are presented in the tables below. Overall, the products contribed in this study reducted product on the quality of the transferred stread interface. To begin with, more than 20% of the products (i ii 5) contained heavy medical contentation. While pro-market testing would have caught this, if such products were ever found on plasmacy sledwes in the United States is would traiger and immediate interioristic recall Next, an econimistion of basic integli. States in the content of the products were ever found on plasmacy sledwes in the United States is would traiger and memoriate networks recall Next, an econimistion of basic drug dissing slowed many. deviations. Approximately 35% of the products were actually significantly overdosed. While this was likely done in an effort to produce a stronger user response and loval customer hase this is an unaccentable tactic which raises many notential safety issues. In the third set of tests, more than 60% of the samples were shown to contain some type of unidentified steroidal compound. This does not necessarily mean the products were dangerous, as this may simply consist of inert steroid precursors/intermediary compounds. It does, however, show that impure steroid materials were used during the manufacturing process. Lastly testing for 2.4-decadienal confirmed that at least 14% of the steroids tested used food grade oil perhaps the type purchased in a grocery store.

## Drug Analysis Results

## Test #1: Heavy Metals Contamination

Sample	Contamination	Result	
1. methandrostenolone	None Detected (<0.002)	PASS	
2. testosterone enanthate	None Detected (<0.002)	PASS	
3. testosterone enanthate	None Detected (<0.002)	PASS	
4. testosterone propionate	None Detected (<0.002)	PASS	
5. boldenone undecylenate	Metals Found (>0.002)	FAIL	
6. testosterone cypionate	None Detected (<0.002)	PASS	
7. boldenone undecylenate	Metals Found (<0.002)	FAIL	
8. trenbolone hexahydro.	None Detected (<0.002)	PASS	
9. testosterone cypionate	None Detected (<0.002)	PASS	
10. methenolone enanthate	Metals Found (>0.002)	FAIL	
11. testosterone cypionate	None Detected (<0.002)	PASS	
12. nandrolone decanoate	None Detected (<0.002)	PASS	
13. methenolone enanthate	None Detected (<0.002)	PASS	
14. trenbolone enanthate	None Detected (<0.002)	PASS	
	I.methandrostenolone     Z. testosterone enanthate     A. testosterone enanthate     A. testosterone enanthate     A. testosterone propionate     S. boldenone undecylenate     C. testosterone cypionate     Z. boldenone undecylenate     B. ternoblone hearbydro.     B. ternoblone hearbydro.     J. testosterone cypionate     I. testosterone cypionate     I. testosterone decemone     I. testosterone decemone     I. testosterone decemone     I. testosterone cypionate     I. testost	1.methandrostenolone None Detected (<0.002) 2.tetostoerone enanthate A. Betosterone enanthate A. Betosterone enanthate A. Betosterone propiorate A. Lestosterone propiorate A. Betosterone propiorate A.	1.methandrostenolone

# Failure Rate: 21% Test #2: Dosage vs. Label Claim (mg/mL)

12 nandrolono docanosto

Sample	Labeled Dose	Actual Dose	% of Claim	Pass/Fa
1. methandrostenolone	25 mg	115 mg	459%	FAIL
2. testosterone enanthate	250 mg	440 mg	176%	FAIL
3. testosterone enanthate	250 mg	408 mg	163%	FAIL
4. testosterone propionate	75 mg	127 mg	169%	FAIL
5. boldenone undecylenate	200 mg	240 mg	120%	PASS
6. testosterone cypionate	200 mg	204 mg	102%	PASS
7. boldenone undecylenate	200 mg	178 mg	89%	PASS
8. trenbolone hexahydro.	76 mg	190 mg	24996	FAIL
9. testosterone cypionate	200 mg	177 mg	88%	PASS
10. methenolone enanthate	100 mg	54 mg	54%	FAIL
11. testosterone cypionate	250 mg	171 mg	69%	FAIL

228 ma

0104

DASS

250 mg

rz. nanurotone decandate	230 1119	220 1119	2170	FMS
13. methenolone enanthate	100 mg	78 mg	78%	FAIL
14. trenbolone enanthate	100 mg	0 mg	096	FAIL
Fallure Rate: 64% (+/- >20% )	of Label Claim	)		
Test #3: Steroidal Materials Pur	ity			
Sample	Conta	mination	Res	ult
1. methandrostenolone	None	Detected	PAS	S
2. testosterone enanthate	None	Detected	PAS	S
3. testosterone enanthate	Unkno	own Peak Detecte	d FAIL	
4. testosterone propionate	None	Detected	PAS	S
5. boldenone undecylenat	e None	Detected	PAS	S
6. testosterone cypionate	Unkno	own Peak Detecte	d FAIL	
7. boldenone undecylena	te None	Detected	PAS	S
8. trenbolone hexahydro.	Unkno	own Peak Detecte	d FAIL	
9. testosterone cypionate	Unkno	own Peak Detecte	d FAIL	
10. methenolone enantha	te Unkno	own Peak Detecte	d FAIL	
11. testosterone cypionate	None	Detected	PAS	S
12. nandrolone decanoate	Unkno	own Peak Detecte	d FAIL	
13. methenolone enantha	te Unkno	own Peak Detecte	d FAIL	
14. trenbolone enanthate	Unkno	own Peak Detecte	d FAIL	
Failure Rate: 57%				
Test #4: Oil Purity				
Sample	Contai	mination	Resu	ılt
1. methandrostenolone	2,4-De	cadienal Detected	FAIL	
2. testosterone enanthate	None E	Detected	PASS	
3. testosterone enanthate	None E	Detected	PASS	
4. testosterone propionate	None E	Detected	PASS	
5. boldenone undecylenate	None E	Detected	PASS	
6. testosterone cypionate	None E	Detected	PASS	
7. boldenone undecylenate	None E	Detected	PASS	

None Detected

None Detected

None Detected

None Detected

None Detected

2,4-Decadienal Detected

PASS

PASS

PASS

PASS

FAIL

PASS

# 12.nandrolone decanoate 13.methenolone enanthate 14.trenbolone enanthate Failure Rate: 14%

9. testosterone cypionate

10. methenolone enanthate

11. testosterone cypionate

#### Conclusions

The scope of testing for this project was fairly hinded, and fell well short of the detailed amayles required to valuate a real proception drug products. 5th feet standards were right enough for a strong majority of the undrugstound steroof products to fail testing. These drugs did not address a passenge strid and to an article of products to fail testing. These drugs did not address a passenge strid and to an article of the control of t

Legimite pharmocutical products are manufactured under strict conditions for a reason. It is very difficult to minima an acceptable level of party without them. Hence I pare USPEP granted materials are being and, it can be very easy for a minimosphe behingly analysis of difficult continuation to the continuation of the conti

## Designer Steroids

there is, faul for in the control detection methods used by the various spaces agreed.

In it is a most the set a memoral for maled actually, so most to have eachly with record and the properties of the control of th

Visu see, several hardred, first a thousand or more, different stroids were sprincisted and innestigated in various behometories around the world during the hybry of storred research. In most cases, their ambolic and androgenic potencies were measured, with the same methods that hus been used on all of the popular stroids we know to holy, Orba a misser faction of these research comprounds altitutely became commercially available due; any area of that generated though as there would be no way for hardred of similar dates pot each in the same market. But the only research is still out there, and remain a very valuable source of farformation for the cleaver clamsics for floody.

Some of these old research steroids of the '50s and '60s still exist today, due to the diligence of underground chemists and researchers. We refer to these drugs collectively as 'Designer Steroids', and they are here only for the purpose of defeating a drug screen. A true designer steroid is structurally unique next to the known anabole/androcenic steroids. sharing no common metabolities, so as to be undecetable to even the most florough steroid test. The fluegif of inciting down metabolities for all possible seasible compounds, to designer steroids sous, come like an improsofit test to say the less. Hern if somehow this old research were to be exhausted, and metabolities referred for all known steroids, there are self many finishes often very to the reductioner, metabolities, or displications are not make targing most services. The designer storoid phenomena could obviously posent an ormal traject most services. The designer storoid phenomena could obviously posent and office the composition of the contract could be obtained by the contract of the contract

At the port in time, the fact that designer steroids exist is no secret to the sports agarcies. It becames partially obstacts to the ICO (Intermined) Objects: Committee in March of 2002, when the ICLA Objects: Amplical Lab detected modesthore, a potent c-17 alpha adopted marknines denotine investigated back in the 1960s, in the runs samples from a found at the "Intermined at a lot harmy Planess, a 23 "Supered city cells from Contrade the Intermined Contrade Contrade the Intermined Contrade Contra

Califor was able to obtain a sample of pure norbodebone from the drug company Wych, and must have been greatly asked by the first that metholess of this stored but here therefried in earlier statiscs.<sup>178</sup> The procedure for probelebone detection has now been mude available to all esting agencies, and unfortunately it is now unself for competition its wales as a designer stood has Revoice vanished oversight. Perhaps it was a had dea to use a steorid that extant much it all the way to the point of ficial trials in the U.S., as there is agine a bit of information to be found out from king the urmary metabolites study would have mude things a lot hander or califor. However, for earthing a lot hander or califor, blocastly, I can think of an number of more efficient and safe compounds to use than the historically progestational one (so, the water bloot,) I don't think earlies and the contract of the con

The norbolethone story quietly fell from the public conscience not long after it broke. The number of athletes that ultimately tested positive for the drug was minimal, so it really never evolved into the big scandal that was initially expected. The USADA thrives on negative media attention to steroids, because it leads to more government funding so no doubt this lack of public outrage was a disappointment. I would suspect many involved were hoping for the global story on par with what happened when Ben Johnson was stripped of his gold medal during the 1988 summer Olympics. This would be of little matter by January 2004, however, because a much bigger doping scandal was about to hit. It involved the use of the designer steroid tetrahydrogestrinone, and this time would snare some of the biggest figures in amateur and professional sports. Not just Olympic competitors, but professional football and baseball players were being listed as potential violators. Many household names were being thrown around, including Jason Giambi, Barry Bonds, and Gary Scheffield. Over 20 athletes ultimately tested positive for THG, or were specifically named for using it in the evidence. The investigation continues today, so this number may rise. Don Catlin was once again the scientist who helped identify this compound in the first place, as well as a method of its detection in urine. This time around, however, he had a lot more help then he did with norbolethone. THG was actually handed over to the IOC testing laboratory in a syringe, by an anonymous coach who did not approve of its use. With the help of an inside informant, USADA got their Ben Johnson story, and then some. THG was at the center of the biggest organized doping scandal in the history of competitive sports, and would come to spark a more vigorous government fight against steroid use than we had yet seen. The steroid-using community is only now beginning to feel the backlash.

I include these stories not because they altestime victories for the ICC. Quite the contrary, I believe they underfine the major fillings in current storied testing methods. These two includes logically do not represent the only two obsigner storoids over used in competitive that the contract of the contract of the contract of the contract of the contract that IRI Gas were completely unsweed for method, for all years, bobody laves angiting about this steroid until a sample was handed over to the Iseling facility, which is the same facility that full untility been proseig time, complex contraining the sum steard jud depth of the contract of the co before. Were it not be the made source, 11th words probably still be in use today. The morbolichness and This stories spit in the cot of those on the sichiess, who insist that drug testing ensears their finorite athlete is drug fee. The first is, many other potent designer storieds are probably on three, ether in the books, or in the gram bags, of many of the world's top competition. It may take years for the next designer compound to be identified by the ICO libes, and replace note) anterior browds for a new one to be synthesical cone is it is in gare the drug testers striply carried via given the book flary have available to them now. We may use repeate of these extands in the fature, that are levent will only example the proficearcy of these workship and the fature, that are levent will only example and of the athletes who may gained dung testing. They show the public the modelable will of the athletes who may gaine to due these agarts, each to esting appreciation profess.

376. Detection of randrolone metabolites in urine after a flootball game in professional and amateur players: a Bayesian comparison. Robinson N, Taroni F, Sungy M, Ayotte C, Mangin P, Dvorak J. Forensic Sci Int 2001 Nov

377. Detection of norbolethone, an anabolic steroid never marketed, in athletes' urine. Catlin D, Ahrens B, Kucherosa Y, Rapid Commun. Mass Spectrom. 2005. 16: 1237.

378. Schanzer W, Donike M. Anal Chim. Acta. 1993; 275: 23

## Anabolic Steroids and the Law

Unted States low probible the possession of anabole stroubs without a legal medical preception, imparing severe purshies, fertainfug fire and/or imprisorant) of frome that choose to violate these lows. Unter rithance of U.S. government officials, World Arti-Doping Agancy WADA) numbers, and pable critises inflowing marrous doping scandals, a growing number of courties are following the U.S. by adopting their own lows against the possession of number destroated order sported droping thus, in many cases within severe criminal possible how been entered. The following section discusses in more related from S. and the section of the control of the following section discusses in more related drouss.

#### United States

Anabole stroids have been classified as corrolled substances in the United States since 1991, with passage of the Anabole Steroid Corroll Act of 1990 (Pub. L. N. 101-16-75, Sec. 1992, 1944 State, 4893, 1990). This low mikes it a criminal offense to self, discibilate, possible of the control of the control

#### State vs. Federal

Crimia llaw against the possession of anabolic steroids exist at both the federal and state bearing three Crimia programme of the circumstance, an intividual may be changed with a steroid related crime by offers the federal government or the acting conventure where the crime took place. Links the recovering of rather interval motion, are criminal prosecutions for steredule programments of the crime took place. Links may be clearly the crime to the crime took place to the crime took place to the crime to the steroid place depends and surface. If you are not obtaining medications legally frough as physician's prescription, it is achievable to such place the strend links codespic practically those of your reliabulations. The book "Legal Mandet: Anabolic is and the practically those of your reliabulations and the crime to the U.S. architege of the U.S. architege is the U.S. architege in the U.S. architege in the U.S. architege is the critical breakform of a first stered links on the converge stereds in the U.S. architege is the critical breakform of a first stered links on the crime to the U.S. architege is the critical breakform of a first stered links are converged storeds in the U.S. architege is the crime to the critical breakform of a first stered links on the crime to the crime to the crime to the unitary to the crime to

The main body (drug listings) of the Anabolic Steroid Control Act has been included for your poiew below

```
'(A) The term 'anabolic steroid' means any drug or hormonal substance, chemically and
nharmacologically related to testosterone (other than estrogens, progestins, corticosteroids
and delivdroepiandrosterone), and includes--
'(i) androctanodiol...
(f) 3b 17b-dilydroxy-5a-androstane; and
'(II) 3a 17b-diburkovu-Sa-androstano
(ii) androstanedione (5a-androstan-3 17-dione):
'(iii) androstenediol...
 (I) 1-androstenediol (3b.17b-dirydroxy-5a-androst-1-ene):
 (ID 1-androstenediol (3a 17b-dibudroxy-5a-androst-1-ene):
 (III) 4-androstenediol (3b.17b-dihydroxy-androst-4-ene); and
 (TV) 5-androstenediol (3b 17b-dihydroxy-androst-5-ene):
(iv) and cost anodiono...
 (I) 1-androstenedione ([5a]-androst-1-en-3 17-dione):
 (II) 4-androstenedione (androst-4-en-3,17-dione); and
 (III) 5-androstenedione (androst-5-en-3.17-dione):
 (v) holasterone (7a 17a-dimethyl-17h-hydroxyandrost-4-en-3-one):
 (vi) boldenone (17b-hydroxyandrost-1.4,-diene-3-one);
 (vii) calusterone (7b 17a-dimethyl-17b-hydroxyandrost-4-en-3-one):
'(viii) clostebol (4-chloro-17b-hydroxyandrost-4-en-3-one);
(ix) debudrochloromethyltestosterone (4-chloro-17b-hydroxy-17a-methyl-androst-1.4-
dien-3-one):
'(x) *1-dinydrotestosterone (a.k.a. '1-testosterone') (17b-hydroxy-5a-androst-1-en-3-
one):
'(xi) 4-dihydrotestosterone (17b-hydroxy-androstan-3-one);
'(xii) drostanolone (17b-hydroxy-2a-methyl-5a-androstan-3-one);
'(xiii) ethylestrenol (17a-ethyl-17b-hydroxyestr-4-ene);
 (xiv) fluorymesterone (9-fluoro-17a-methyl-11b 17b-dihydroxyandrost-4-en-3-one):
'(xv) formebolone (2-formyl-17a-methyl-11a,17b-dihydroxyandrost-1,4-dien-3-one);
 (xvi) firazabol (17a-methyl-17b-hydroxyandrostanol? 3-cl-firazan):
 (xvii) 13a-ethyl-17a-hydroxyoon-4-en-3-one:
 (xviii) 4-hvdroxytestosterone (4.17b-dihydroxy-androst-4-en-3-one):
 (xix) 4-hydroxy-19-nortestosterone (4.17h-dihydroxy-estr-4-en-3-one):
 (xx) mestanolone (17a-methyl-17b-hydroxy-5a-androstan-3-one);
(xxi) mesterolone (1a-methyl-17b-hydroxy-[5a]-androstan-3-one):
 (xxii) methandienone (17a-methyl-17b-hydroxyandrost-1,4-dien-3-one);
 (xxiii) methandriol (17a-methyl-3h 17h-dihydroxyandrost-5-ene):
 (xxiv) methenolone (1-methyl-17b-hydroxy-5a-androst-1-en-3-one);
 (xxv) methyltestosterone (17a-methyl-17b-hydroxyandrost-4-en-3-one):
'(xxvi) mibolerone (7a,17a-dimethyl-17b-hydroxyestr-4-en-3-one);
 (xxvii) 17a-methyl-*1-dihydrotestosterone (17b-hydroxy-17a-methyl-5a-androst-1-en-3-
one) (a.k.a. '17-a-methyl-1-testosterone');
'(xxviii) nandrolone (17b-hydroxyestr-4-en-3-one):
(xxix) norandrostenediol--
'(I) 19-nor-4-androstenediol (3b. 17b-dihydroxyestr-4-ene);
'(II) 19-nor-4-androstenediol (3a, 17b-dihydroxyestr-4-ene);
 (III) 19-nor-5-androstenediol (3b, 17b-dihydroxyestr-5-ene); and
'(IV) 19-nor-5-androstenediol (3a, 17b-dihydroxyestr-5-ene);
'(xxx) norandrostenedione--
 (I) 19-nor-4-androstenedione (estr-4-en-3,17-dione); and
(II) 19-nor-5-androstenedione (estr-5-en-3 17-dione:
'(xxxi) norbolethone (13b,17a-diethyl-17b-hydroxygon-4-en-3-one);
(xxxii) norclostebol (4-chloro-17b-hydroxyestr-4-en-3-one):
```

(xxxiii) norethandrolone (17a-ethyl-17b-hydroxyestr-4-en-3-one); (xxxii) xxandrolone (17a-methyl-17b-hydroxy-2-oxa-[5a]-androstan-3-one); (xxxii) xxmesterone (17a-methyl-4-17b-dibadroxyandroxt-4-en-3-one); (xi) restoration (110-reparts) parameter—(CP-700E),
(xii) retrahydrogastrinore (13b,17a-diethyl-17b-lydroxygon-4,9,11-trien-3-one),
(xiii) retroloine (17b-lydroxysetr-4,9,11-trien-3-one), and
(xiii) any sat, ester, or ether of a drug or substance described in this paragraph';

## Austria

The possession of anabolic steroids is not a criminal act according to Austrian law. In 2008, Austrian government officials announced intent to place criminal penalties on steroid messession.

# Australia

It is a criminal act to import, supply, use, or possess anabolic steroids in Australia without a prescription from a medical practitioner, dentist, or veterinarian (Poisons and Drugs Act Amendment of 1994). The outlined penalties for possession without a legal prescription include a maximum off formula of investionment and/or a five of 555 000.

## Canada

Anabolic steroids are included in the Caradian Controlled Drugs and Substances Act as Schedule IV substances. It is illegal to sell, manufacture, or import anabolic steroids into Carada without proper legal authorization. Possession of anabolic steroids for personal use is not a criminal act

#### Czech Republic

In 2008 it became a criminal act to manufacture, import, export, store, or distribute anabolic steroids in the Czech Republic. The potential penalties include a maximum of 3 years in prison. It is not a crime to nossess steroids for pressonal tas.

# Denmark

In Dermurk à is a crime to munificture, import, export, market, disperse, distribute, or possess doping substances including arabolic stroids, huma growth harmone, and enythropoient whother proper mediation of scientific resolution. (In Act on Problishion of Certain Daping Substances No. 232 of 21 April 1999). The potential penulies for possession include a minimismo? Q-wasta princip.

#### France

In 2008 it became a criminal offense to manufacture, transport, acquire, or possess doping substances including anabolic steroids, human growth hormone, and erythropoietin in France. The potential penalties for possession include a maximum of 5 years imprisonment and/or a 75,000 Euro fine.

#### Greece

The possession of anabolic steroids is not a criminal act according to Greek law. In 2008, government officials announced intent to place criminal penalties on steroid possession in Greece.

#### Sweden

In Sweden è is a crine to import, munificture, transport, sell possess, or use doping substances such as anabolic steroids and growth homone without proper legal authorization (The Swedish Act prohibiting certain doping substances (1991,1969). The potential permitise include a muximum of 2 years in prison. Possession for personal use is usually regarded as a petry offense and given a munitampenaly of for motifs imprisonment.

#### United Kingdom

The importation, possession, and use of anabolic steroids are not criminal acts according to UK law. There has been a great deal of pressure in recent years from the U.S. and World Anti-Doping Agency to place criminal penalties on the possession of anabolic steroids

## Acquiring AAS (Rest Practices)

This section assures that the decision has already been made to use anabolic strooks. It places no judgment on the decision beeful and concentrates only on changing risks orbitally included in the days apply. How one goes about obtaining these abstances is an extensibly important annuables serooks can be a highly whitely hope, with significant quantities of two quality controlled and undergoond drap in circulation. In this section we will review the for grammal sources for emables seroods. When possible, we carrier they that should recover, at least to some small center, taking early for the bayer. The main objective in all cases is to decian legitime plantemental guide products so that no additional risks the to

## By Prescription

Provided you are in some way in wild need of therapy, it is always preferable to obtain anabode attends frough a physicative preception. In addition to othering the potential benefit of lawing some medical ownsight to your use, most pharmaceutial markets (in Western antiess operating) are very secure. It is milkely that you will come across the security of the properties of the security of the security of the security of the States, in the, the risk is so low that it is not even worth considering Furthermore, if you reside in a coursely where there are criminal possible for the unique of statesida, a procupying the security of the security of statesida, a procupying the security of statesida, and cold possibles can be very serious in many cases, even when the days are for present and cold possibles can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and cold premiss can be very serious in many cases, even when the days are for present and the cold premiss are consistent and the cold premiss are consistent as a serious cases.

Most nodes are probably under he iscorrect assumption that is nowly impossible to get a preception for artifacts sensite in the tland 5 stace. While them plane been must following the deping contails of the late 1800, the metal tens element considerably since then. The legal and of those drops is again on the ine. The self-late groups in the rail-1900, when drug comparies stand to prof. forth a strong effort to obtain physician and the public on the use of strongly (openfully statestown) to text age-related heromer deficiery in men it is a fact that me deal with hormoral declarie just flee women of, and striff ill health offices from it. Training hormoral deficiery with drugs to restree profile institutions levels has become a widely accepted part of medicine. By the time men in their thirties, a high processing are legitime conditions for homore replacement returns.

Lyou feel that you are a conflicte for a preceiption, yourney need to take an active rule in funding a physical informative, may observe as well storest robot too day. They might put perceibt sestedation at all or may be too conservation in their instance if they do. Von may have to intensive several before you find the cont that in gift a you. Don't very about "before where play the you flow to the several before you find the cont that in gift a you. Don't very about "before whereight find the cont flow in gift a you. Don't very about "before whereight perceiptions you don't now! A would rate for it you factor, or you and gain preceiptions you don't now! Would rate for its your doctor, orly and gain preceiptions you don't now! Would rate for its your doctor, orly and gain preceiptions from more than one at a firm. Otherwise, it is perfectly legal to take the time to find you would be considerable.

Medical recessity is usually supported with a blood test, ordered by the treating physician. Floatstonero levels turn to be highest in the moring and lowest at the end of the day. The blood test should be planned accordingly. Before reviewing the results, undestand that there are no specific established protects for testing men with hormore defections? The medical community does not even agree on the point in which thempsy should be initiated. For example, the "hormal" level of instanteurs in an adult man is 20-5-1.20 registl. Some physicians will not prescribe applying to you takes you are symptomize and you beek full objective being to maintain hormal below in the high cell of control ("springs"). Note that some people choose to pre-test thread-test to destrine how they might quality before approaching a physician. We can express that frequire when the regist was full registered.

How testosterone medication is prescribed also varies a great deal depending on not only the needs of the patient, but also the comfort level of the physician. For example, more rigid physicians will often only prescribe testosterone eels or natches. This is because these forms allow the very controlled duly release of testosterone, and are harder for the patient to shase. Other doctors are comfortable working with testosterous rejections that are taken home and self-admirated by the patient. Even when testosterous rejections are preserved, some doctors reface to give more than 100 mg per week. Others routinely give a 200 mg per work doce, assaring that the beste har ent charder of any port of arting reatmers. (Ivo will not need to find a crooked doctor. You just want to find a progessive physician that understands the value of optimizing your formume levels.

Voal tour a five options open to you when first useking reastment. You can approach your finity doctor, a formous specials (read-translegs) or on an eleging photosism in my experience, anti-signing melantes in which you find the most progressive physicisms and tentumers. These are the decroes that are working with both stateols and off-belled medications like artis-entosigns and growth hormore, and foreign the establishment to retain's hormoral medicates becomes in the field are off-cut effect by principal polysism groups with the mean fraction. The content of the principal content of the principal content of the principal content of the principal content of the proposition of the principal belled and the content of the principal content of the proposition to support from the part for gain the Life all these only to blace and the proposition of the principal belled and the content of the principal content of

One benefit to being a varier and octuents on another alterance in that I are regularly memoring programs beginner and between of the converts. The table pleasant of gainting below a great range over the years. It has given me confidence that the field of medicine is here to stag. Fart contraining to your regularly. It coursed to me that rup for cloratests pair me in a urique position to help potential HHT/mrs-signing partients. Make toconsterous electricacy is greatly under-tended in this course. If alter to do my part to help change the octionary is greatly under-tended in the course. If alter to do my part to help change the below I will used as a portal to part without in contact with HHT/mrs-signing physicism. If you thick HHT/mrs he are option for you, I recordy encourage you to explore it fairly, regardless of how you go about seeking tenament. Supporting the physicism that offer this type of trustomers is the best way to help significe and above the size of medical-

For hormone replacement therapy & anti-aging doctor referrals, visit the website www.HRT-Rx.com to find a physician in your area.

# Ordering Steroids in the Mail

Utilize most multismal drugs that are under strict legal controls (such as merotics), the sale of annihold starouls seems to be openly alsorted on the Internet and in the back pages of subobly-linking mugingsiss. Match of this probably has to dae with its storing various in horse strictly an internet of the storing various in horse the strictly of the storing various of the strictly and the str

The first major difference between obtaining authoris servaits is a prescription and ordering thems through the mall (adapts) is that you may receive graining pharmacectaril proteins. The potential health risks of countrief drugs are significant, and detailed in other sections of the book. Counterfirsting is actions and voltagement. The world behalth Organization recording to the book of the counterfirst is action and voltagement to the counterfirst is allow to be the counterfirst in the counterfirst is allow to be the counterfirst in the c

Below are several rules to follow when ordering steroids through the mail. These rules can help increase the chances that your order will contain pharmaceutical grade drug products.

 Order from a country with legal access to steroid medications. If it is easy for your source to get legitimate products, it is likely what is sent to you will be legitimate. Avoid ordering from countries with critic medibation lause such as the United State. Council. Western

- noni commes wan sunce promotive laws such as the critical states, Catanaa, vesterin Europe, or Australia. Consistent supply in these areas invariably involves the sourcing of claudestinely manufactured products.
- 2. Buy local brand drugs from the chosen source country. If you are ordering from a pharmacy in Thailand, buy Thai drugs. The chances of running into counterfeits will increase significantly when the drugs you buy are not domestic to the supplier's country.
- Immediately avoid any foreign pharmacy that says it carries American steroid products.
   These will be counterfeit over 99% of the time, which means the pharmacy probably deals in counterfeit drugs.
- 4. Buy from a place that shows pictures of its products. Scrutinize the offerings closely. Are there any known fakes? Many suppliers work hard to keep their company clear from counterfeit druss. You want to deal with one of these.

The purpose of this book is to reduce hurn not encourage or support crimical activity. Thus the previous nels address on the welding for the days, not strangingly them If you credit in an area where steroids are Begal to insport without a prescription such as the United States, Canada, Scandiniva, or Australia, you need to be asswer that there can be serious legal consequences to ordering these diags frivough the mall I faultonines intercept your package, you may be changed with a survey of crimes including ding goosession and Begal insportation. Not may even be changed with nutficking file quantity is large credit, While dedictions the property of the control of the property o

## Purchasing Steroids on Vacation

Another common option is to travel to a country where annihole sensels are widely available, and preselves the flug directly from a paternacy. Many popular sections speciare common sources, richting Mexico, Trailand, and Titaley, So it is not uncommon to see dual-puppose give, discontinivaciency, with the individual bringing bear close yas and personal use supple, As with all filed sourcing options, there are some bourfits and disrubscles to the practice. The principle bourfit, provided the sufficial chooses the right country and shope a radicipant, should be more consistent access to real pharmacetrial guide compounds, and the clance for specifies the first particular time in disrubscle, of course; fishing you will be claused for specified between the contraction of the contract

- In an effort to help readers avoid the added risks of counterfeit drug products, the following points of advice are provided. All readers are cautioned that snuggling is illegal and can be a very risky practice. Again, since this book is about harm reduction we will not be discussing means of smueeling.
- Loose regulations on the steroid supply typically also mean loose oversight of pharmacies.
   In some countries, pharmacies in tourist areas will commonly stock counterfielts. You should become well-versed with the products of a particular region before traveling.
- 2. Always buy local. You want to get the drugs that are readily available to pharmacies through normal legal channels, not exotic drugs that may or may not have been legally imported by the pharmacy. For example, never buy European or American drugs in a Mexican pharmacy.
- If possible, avoid the obvious tourist pharmacies. Travel to an area more known to the locals. The smaller a percentage of business the pharmacy makes on tourists, the less likely they are to be stocking counterfeits.
- 4. Ask for products by name. Most tourists stumble in and ask for "steroids, anabolicas, testosterona." If instead you ask for something very specific, such as "Bayer Primoteston," you are probably going to be taken more seriously.

## The Local Black Market

Where steroids are difficult to obtain through legitimus charmels, robust trade usually continues on a block market. The United States is particular but the largest and most learnine steroid block market in the world. Hyou plan to obtain annobed seroots this way, you should be assure of a winged archaest-and and teatings. To begin with, with cordering copporating to import a proposition of the proposition of

Again, our focus in this section is only on increasing the likelihood of obtaining real pharmaceutical products. Towards that objective, the following points of advice should be followed when shoroning on the local black market.

- Consider the country of origin for each drug. Only buy drugs that make sense. For example, Mexican steroids are common in the United States due to their open supply and close proximity. You would expect to see them. On the other hand, you are very unlikely to full legitimate drugs from Australia in the United States, which is very far away and also seriously restricts is own steroid supply.
- Try to stick with a group of fellow steroid users. Sharing experiences and knowledge can help keep the group better supplied with legitimate medications.
- Try to buy from other steroid users. Individuals who sell steroids but don't use them are probably less concerned with product quality.
- 4. If you are trying to purchase legitimate pharmaceutical products, don't buy multi-dose vials or bottles with lose pills urless you are absolutely sure of the company and source. These forms of medication are rarely used outside of the U.S. and Carada. Single use ampules and push through pill bisters are much more common in other countries, and may be slidtly more secure due to the difficulty involved in making them.

Part III

# Drug Profiles

# Agovirin Depot (testosterone isobutyrate)

Standard	Standard
Chemical Names	4-androsten-3-one-17beta-o 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate
Dogganatasia and Aus	duta. I

#### Description:

Androgenic

Testosterone isobutyrate is an injectable steroid preparation that contains the isobutyrate

100

ester of tosisterore na vaster tous. Among programators, tostosterore is output in so then considered analogous to inclusive suspension (no seary). Altogolo beh an usually found as water-based suspension, the plantmockineties of the two products are admirtedly very particular to the plantmockineties of the two products are admirtedly given every few days, testosterore felod superiors is very final-strain, grapping injections, to be admiristed once every two weeks in a relieful setting at an appetitude testosteror, testosterore isologratie is flavored for its ability to promote rapid gains in muscle moss and stream?

## History:

Injectable instinations industrial memoryanal suspension was first described in 1952. <sup>207</sup> This agart was developed in an effect to create an signitive (loop) from of testosterous flut would be show acting that regular (flee) testosterous suspension or testosterous propientat, the most widely prescribed from of instinations at the time. This is represent, the state of the size of the si

The only modem steroid product to use instinctions is obstyrate is Agovirin Dpvt, developed by Biofals in Cexchowlooks, it is primarly prescribed to trust males with instificient antiogual levels and adolescents with deloped paborty, although it is indicated for a variety of other propose including the tentum of Klinchler sphorous; dischause there are enter chromosome results in an irrelative of antiogenicity and estrogardicy), aplantic amount, Calmige sphorous (as sum anabotic agard to preserve he missal, positione) quantic, Calmige sphorous (as an ambotic agard to preserve her install, positione) and contexpositions, advanced beneau carety, missalogian (breast pain), and cachenia (wasting of Stakes (Bourble) and terminian product are only in a formation of the context of the product of the context of th

#### How Supplied:

Testosterone isobatynate suspension is available on the harm drug muker in the Slovale Republic sa Agoptin Dopt (Blotals). It contains 25 mapfin of steed in mise in a waterbased solutior, packaged in a 2 ml. ampule (5 ampulse per box). Testosterone isobatynate has low water solubility, the steed will of moleculely separate from the water-based solution when an ampule is left to st. A quick shake will temporarly pince the drug back into suspension, so that the whitehraw dosages dould always be consister.

#### Structural Characteristics:

Testosterore isolotypate is a modified form of testosteroe, where a carboyle, ead ester C—melly proprise acidy has been attached to the T-best pluptout goop. Esterikd from of lessosteroe are less polar than fee testosteroe, and are absorbed more alonly from the area of spectra. Once in the bloodstream, the eater is removed to systiff test (astro-testosteroe. Esteriked forms of testosteroe are designed to probage the window of the respect effect following administration, along for a loss foregard spectra produce statements. However, the statement is a substitute of the statement of the statement

## Side Effects (Estrogenic):

Testoderore is readily aromatical in the body to estradiol (estrogan). The aromatuse (estrogan synthesis) entrym is responsible for this metabolism of testosterone. Esteadid estrogan is whether earner is established for this metabolism of testosterone. Esteadid estrogan is whether are so that the establishment is considered an other establishment in the establishment is established an establishment in the establishment is establishment in the establishment in the establishment in the establishment is establishment in the establishment in the establishment in the establishment is establishment in the establishment in the establishment in the establishment in the establishment is establishment in the establishment is establishment in the establishment in the establishment in the establishment in the establishment is establishment in the establishment

Entogais side effects will occur in a dose-dependant mumer, with higher doses (above ment) therapartic tesels) of testosterowe more fleely to require the concurrent use of an artisentogan or arountase ribbliot. Since water reterion and loss of muscle definition are common with higher doses of festosterons, this drug is usually considered a poor choice for dieting or cutting phases of testing. Its moderate estrogaricity undees it more kital free bidsing phases, where the added vistor recentor will support more strength and muscle size,

## Side Effects (Androgenic):

Textosteore is the primary mile androgen, responsible for miniming secondary mulesessual culturactivists. Pietuted level of textosteore are fleely to probe enrologueis de effects including obj skin, acer, and body/sical hair growth. Men with a genetic prodeposition for him loss (androgenetic abecept) may notice accelerated mile pattern halfing. Those concerned about him loss may find a more constrained option in manufoctor portal visiting effects of antibolicam/deposite secretics, especially within a strong antiques such as testosteone. These may include desponing of the voice, menstrual irregularities, changes in shit network, feel him growth on affection elementary.

In authogos-responsée traget fissus sech as the skin, scale, and prostite, the high relative authogosity of testosterone is dependant on its relaction to displostosterone (DRI). The 5-siph reductase emprise is responsible for this metabolism of testosterone. The concurrent use of 3-siph reductase inhibits was his instituted on statistical will stretch with site-proclif potentiation of isotosterone action, lovering the teachery of testosterone with site-proclif potentiation of isotosterone action, lovering the teachery of testosterone with site of the separation of testosterone's anabolic and analongosic properties is not possible, over with test 5-shirt necktose inhibito.

## Side Effects (Hepatotoxicity):

Fastostorour does not livo beputatoxic effects, ber toxiciy is utilicely. One study cammied he potertia file hepitatoxiciy with high does of elestostoret by administering 400 mg of the hormone per day (2,800 mg per week) to a goup of mine ashjoest. The setrout was taken onally so that higher peak concentrations would be reached in hepitate itssues compared to intranscender injection. The hormone was given daily for 20 days, and produced no significant changes in few enemy wates including searm administ, bifurbit, and the contraction of the

#### Side Effects (Cardiovascular):

Ambolicardnogueis steonik en lavo deletrious effects on serum cholesterel. This includes a referency to refere IEEE, (goad) cholesterov olluses and irrecess IEEE, (doad) cholesterov ulases, with result of a referency to refere the repart of an ambolicandrogueis esteroi on serum (pist) of expenditur on the dose, route of administration (rot 1st. rigicardish), type of steroid Arabicidardogueis esteroid on serum (pist) of dependant on the dose, route of administration (rot 1st. rigicardish), type of steroid Arabicidardogueis esteroids my pist absolutely delet blood present and trighcristic, reduce excludedal relexative, and support left workreizer largeritospis, all potentially increasing the fist of confinosocular disease and responsible affects.

Textostorous trads to have a much less dummic impact on cardiovascular risk Extors that synthesis steeds. This side in a part to is operates to metablishing by feet, which allows it to have less effect on the hepatic immigranted of cholesteed. The aromatistion of textostorous to established also helps to migrate the regardise effects of afringers on serious pipils. In ore study, 200 mg per week of its observer ester (reuntitus) had a slight har art statistically significant effect on HEL cholesteed and the 2 weeks, but when balson with an automatise ribibitor a strong (25%) decrease was seen. <sup>108</sup> Studies using 300 mg of inclusionare cost operational per week for 20 weeks whom an aromatise influent indications of the contradiction of

Due to the position influence of estinguin on serum lipids, timordant entire or chimphene centure are preferred to contraines inhibitors for those concerned with activalescular health, as they offer a partial estrogenic effect in the fiser. This allows them to potentially improve lipid profiles and offers some of the regarder effects of analogous. Whit does of 600 mg or lass of less storage per week, the impact on lipid profile tends to be noticeable but not duranter, maring an air-strengen (for earliers) of an experiment of the second of the lass of the second of the second of the second of the second of the training second of the distance of the second of the second of the land second of the distance of the second of the second of the land second of the distance of the second of the land second of the second of the second of the land second of the land of the l

To help reduce cardiovascular strain à is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholssterol, and simple cardohydrates at all times during active AAS administration. Supplementing with shi old, games per day) and a natural cholssterolamicoxidant formula such as Lipid Stabil or a product with compramble inspedients is also recommended.

## Side Effects (Testosterone Suppression):

All anabolication/genic stends when taken in doses sufficient to promote muscle gain are expected to suppress endopross to stostoree production. Esototores is the primary unitant/ogan, and offers strong negative feedback on endogranus tostostoree production. Foototores-based days will, Revisic, here a strong effect on the phytoliniar regulation of interior strong the production of the production of interior strong the production of interior strong the production of the p

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

The design of asstoateous isolatyate (as Agovin Depot) is slightly different than that of most tistoateous exists, which are usually made as oly solation. Agovin Depot instead contains a microcystalline aqueous saspension The crystals form a repository in the muscle following injection, where they showly disolor over time. Injections of testoaterous isolatynate may require a large needle (21 gauge), and may result at local irritation, pain, and needle the control of the control of

## Administration (Men):

To text androgan issufficiency, institutence inobstynties supervision is usually administered in a dose of 50-100 mg occy 14 days. When used for muscle-bading purposes, stotisticnes independent supervision is other administered at a dose of 200-400 mg (4-6m) per work. The contraction of the contra

#### Administration (Women):

Testosteroe isobutyrate suspension is not commody used with women in clinical medicine. When applied, it is usually given in a dose of 25-50 mg every 14 days. Testosterone isobutyrate suspension is not recommended for women for physique- or performance-enhancing purposes due to its strong androgonic mature, tendency to produce virilizing side effects, and slow earlier characteristics makine blood levels difficult to control.

#### Availability:

Pharmaceutical preparations containing testosterone isobutyrate are rare. The only known

product at this time is Agovirin Depot, produced in the Slovak Republic by Biotika. It contains 50 mg per 2 mL ampule; 5 packaged per box.

#### Anadrol®- 50 (oxymetholone)

Androgenic	45
Anabolic	320
Standard	Methyltestosterone (oral)

Chemical Names 2-hydroxymethylene-17a-methyl-dihydrotestosterone
4.5-dihydro-7-hydroxymethylene-17a-inha-methylestosterone

17apha-methyl-2-hydroxymethylene-17-hydroxy-Salpha-androstan-3-one
Estrogenic Activity hlgh
Progestational Activity not significant

## Description:

Opmethodene is a potent oul ambole storoid derived from displosteostorone. More specially, it is a close coasi of methyldiphotostorone (methodo), differing only by the addition of a 2-hydroxymethylene group. This crustes a steorid with considerably different activity than restancione, however, so with the is very difficult to other compensions. Displostorostorone and restancione are both very week in this regard, owing to the fact that these melocules are not very state in the high engrane. Ca-plain hydroxymetroid delydroxymes) emiscentered or muscle tissue. Coxymethylore remain highly active hore instead, as it reported in standar aimin allows test demonstrating a spiffacturity higher analosis, earlier plain and an active that the control of the

Oyuncholuce is considered by many to be the most powerful steroid commercially available. A steroid routive experiencing with stagers it likely to just 700 to 30 poursed for mussive bulk, and it can often be accomplished within 6 weeks of true. This steroid produces a lat of water restriction, to a good persion of this gain is gain to be water weight. This is often of fills consequence to the user, who may be feeling vary big and storage while taking ownerholuce. Although the staronth look that treads from water restorins in soft must attackine, it can aid quite a bit to the level of size and strength gainst. The muscle is filler, with contract better, and is provided as two of proceeding in the filler clean usual nebt and contract better, and is provided as two of proceeding in the filler clean usual nebt and contract better, and is provided as two of proceeding in the filler clean usual nebt and use that the stage of precision is the filler of the size of the stage of precision is the filler of the provided in the contract better, and the size of the precision of the clean of a provided is two of proceeding in the size of the provided and better to make a fill of the provided in the contract of the size of the precision and better issues of contractly associated with heavy lifting while missing up on storesty, and comprehabors is a common offerth. These can be such a filling a spirate to fill of the proceeding and the contract of the size of the proceeding and the contract of the size of the

#### History:

Osymetholeus was first described in 1993. <sup>36</sup> The agent was relaxed in the United States as a prescription dupling the early 1996, soil under the brand manes Annale-50 (Systex) and Androyd (Parke Dixis & Co.) Systex developed the agent, and would hold poster rights to at united recipation many sens late. The day uses original approved for use it confitions where annhole action was necessary, indicated uses included grainti-debilitation, chronic underweight status, per- and postspensible preservation of lam many debilitation, there are networked to the confit of the con

In spite of the mmy potential therapeutic uses or a strong ambolic activity of this drug, the FDA soon strictly narrowed the indicated uses of oxymetholone. By the mid-1970's, the drug was FDA approved only for the indicated treatment of america characterized by deficient red blood cell (RBC) production. Admittedly the stimulation of crystropoises is an affect that is characteristic of nearly all ambolic steroids, which as a route tend to increase RBC concentrations. Osymetholeue, however, seemed fairly reliable in this regard: demonstrating an increase in erythropoietin levels as much as  $5~64L^{198}$  This has led to its adoption for this relatively new medical use, as well as the institution of a higher (90 mg) dossage with the updated Aundrol-90 product, necessary for a stronger effect on RPC court. The Parke Davis alm would not be brought up to the higher dossage, however, and was

Recent years have brought fourth a number of new treatments for anemia, most notably Epigean (recombinant engliteopoteiti) and related crystropoietic peptides. These drugs directly marine tho obly similar elfolls out of producing hormous, and as such provide a much more focused forms of therapy, with less of the urrelated side effects one would have to enter with the use of a strong antiques, whitely flantable instearce it simily prompted Systex during for the purpose, sels were now dropping. Francial distincted family prompted Systex design the purpose, sels were now dropping. Francial distincted family prompted Systex decided to drop this ferm is a narbar of foreign countries. Plematif from Switzerland and Austria were dropped, Edwing soon was Oxidosoma from Spain. Daring the mid-1990's, many Althées Enemed Governebles was on the way of for good.

In July 1997, Systex sold all rights to Aundrol-50 in the U.S., Carada, and Mexico to United Pharmaceuticals. Limited restricted Aundrol-50 to the U.S. market in 1998, this time tagging IIV-MISS parieties. Patters with IIV are commonly among the case to the disease. But they was the production of the control of the cont

In April 2006, Solary Pharmacoaficals (parent company of Unimed) sold the rights to Anadout-50 to Almon Pharmacoafical LLC. Almon continues to market the day in the Uniford States, almogly given the ranshort in a meetan with additional uses the company plans to passe with cosymetholess. At the present time the ody FDA approved indication remains fand formating of blood of dictional anamins. Systems eason to be two removed and companies whenever possible. Or nontrivine remains available consider of the United States, although its most behavior and the solution of the contribution of the contribution

#### How Supplied:

Oxymetholone is available in select human drug markets. Composition and dosage may vary by country and manufacturer. Most brands contain 50 mg of steroid per tablet.

#### Structural Characteristics:

Oxymethone is a modified form of displotoestosterone. It differs by 1) the addition of a methyl group at carbon 17-alpha, which helps protect the hormone during oral administration, and 2) the introduction of a 2-hgdwosymethylene group, which iribbis is metabolism by the 3-had ereayme and greatly enhances the anabolic and relative biological activity of methylikythoetostosterom.

## Side Effects (Estrogenic):

Osymetholone is a highly estrogenic steroid. Cynecomestri is often a concern during treatment, and my present itself quite early into a cycle franticularly when higher doses are used). At the same time stater retention can become a problem, causing a mabble loss of much definition as both subsctamesous water retention and fit selech ball. To avoid strong estrogenic side effects, it may be necessary to use an anti-estrogen such as Nobades® or Changle.

It is important to note that oxymetholone does not directly convert to estrogen in the body. This steroid is a derivative of dihydrotestosterone, and as such cannot be aromatized. Antiaromatises compounds such as Cytadren and Arimides® will, likewise, not effect the relative companies you can seemal. Some rave suggested mus are night revive or companie acrossy. To sake effects of both entiting due to the ding acting as a prospectia, similar to mandroise the reads of the contract of the contrac

## Side Effects (Androgenic):

Albacaja osynetlečene is classified sa sa methode steroid, meloganie iske effects see sell possibe udrit sits shakazer. Dene mije iskele botus of oljo skin, neza, mel stody fiscial integrouth. Higher doses are more likely to cases such sike effects. Anabolicirandospanie steroids may also aggrante mie patterm his too. Werner me addisently swarmed of the potential virlange effects of anabolicirandospanie steroids. These mny include a desponing of the toose, mestratul regalarties, changes in site teatrar, ficial integrouth, and cliently endered endingeneret. While Anadoli is classified as an anabolic steroid, it does retain a notable analoguic composure.

It is time-rating to note that computablease does eably some tendency to convert to displosterosterous in the body, although this does not occur to the 5-play reductions ensured in the 1-play reductions ensured to the 1-play reduction of 1-play and polytorosterous based steareds, son such alteration can take place. And from the abded 1-play although the classes of behavior and the play of the 1-play of th

## Side Effects (Hepatotoxicity);

Opportableme is a CT-alphta adjusted composed. This absention protects the drug from description by the four allowing a very high processing of the drug earny into the bloodstream following oral administration. CIT-alpht adjusted autholic intrologenic servoids in the hepatotics. Probunged or high personner my reads in the oftenings. In mer instances life-dreaming dysfarction may develop, it is advisable to visit a physician profucially during a comment of the contract of th

Covarchives has a sensetal x-ring, which slighty houses in rultive beganization, 2008. It files again, protectionly of the does convenely used. or present substantial bepatotoxicy to the user. Studies administering 50 mg or 100 mg daily to 31 clearly men for 122 weeks produced significant increases in the computes (transmission SN T and ALT) to 90 mg in patients taking 100 mg. As exceed andly administering 50 mg daily to 30 patients for up to and conceiling our control and the control of the cont

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised white taking any hepatotoxic anabolic/androgenic steroids.

## Side Effects (Cardiovascular):

Anaboli-infrogarie strouks can have deleterious effects on serum cholstero. This relabels as tendency to reduce IMC, (social) cholsterol values and increase IMC, (social) cholsterol values and tendency to reduce the service of the service values, which may shall the IHD, to LDL behave in a direction that floors genare risk of arteriockrossis. He relative rapset of an anabolicalizations serviced on serum light is dependant on the dose, roate of administration (out we igicatelle), type of steroid achievable and the service of administration (out we igicatelle), type of steroid achievable and the service steroids are done and achievable difficulties of the service and the service

reduce endothelial relaxation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and myocardial infarction.

Opmetholone has a strong effect on the hepatic imangement of cholesterol due to its structural esistance to her breakdown and orate of administration. Statics administering 50 mg or 100 mg dubly to a goop of delety men for 12 weeks have demantical nignificant excesses in LDL cholesterol, accompanied by your spaffurer (formatic) suppressions of interesting the contraction of the contraction

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fast, cholesterol, and simple carbohydates at a litrus during active AAS administration. Supplementing with \$61 oils (4) grams per day) and a natural cholesterolaritoixidant formals such as Lipid Stabil or a moder with commandation transfers is also recommended.

## Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone stimulating substances, testosterone levels should return to normal whith 1-4 moralis of drug secession. Note that prolonged hypogoradotrophic hypo

Note that when discontinuing conventionies, the crash can be as equally proverful as the encycle results. To begin with the best of vision recenterion will quickly dimensils, discopping the user's body weight demantically. This should be expected, and not of function corrent. What is usually of most convent in seisoring endegones in solitoners production with a proper PCT, program (see: Past Capital Thomaga) in this book), Bridge apprint coll some alternatively choose and ender to "Instead up the new mess. And campone to be an effective practice, at least from a mental standpoint. A drop of weight is likely when making the swick, although the end result is still other visioned as allowing the restriction of time (quality number less). It is not of stopping down, that of the water retention, and words liker finally off the hermeness. However, and the contraction of the cont

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biovasiability. <sup>887</sup> This is caused by the fit-soluble rature of steroid hormones, which can allow some of the drug to dissolve with urdigasted distary far reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stormeth

#### Administration (Men):

Fight prescribing guidelines for commerched a dossage of 2.5 mg three times per day to recent the wasting process and provide learn body mass gain. Does as high as 30 mg was employed in some cases. Current proscribing guidelines recommerch a dossage of 1-2 mg/s is hypically obligated; but not be of 1-2 mg/s is hypically dossige leaf in some offer courties; if a recommerched to interfer the dosing of courties of the courties of the courties of the courties of the per day at the 2 mg/s dossige leaf in some offer courties; if a recommerched to interfer the dosing of courties. On the courties of the courties. On the courties of the courties of the courties of the courties of the courties. On the courties of the courties of the courties of the courties of the courties and the courties of the courties and the courties of the courties and the courties of th

25-150 mg taken in cycles lasting no more than 6-8 weeks to minimize hepatotoxicity. This level is sufficient for dramatic increases in muscle mass and strength. Higher doses are rarely administered due to the strong estrogenic nature of the drug, as well as the high potential for hepatotoxicity.

## Administration (Women):

Prescribing information for oxymetholone in the U.S. makes no distinction with the dose for females. Oxymetholone is generally not recommended for women for physique- or performance-enhancing purposes due to its very strong nature and tendency to produce virilizing side effects.

## Availability:

Pharmaceutical preparations containing oxymetholone are fairly limited. The legitimute supply seems to be scattered into isolated markets of Europe, Asia, and the Americas. Most of the supply for this drug comes in the form of underground and export-only products. In reviewing some of the remaining products and changes in the legitimate global pharmaceutical mulet or the base of the products and changes in the legitimate global pharmaceutical mulet or the base of the products and the second coherentations.

Anapolon (Turkey) was recently updated to reflect a new logo for the manufacturer Abdi Ibrahim. The product is still nackaged in boxes of 20 tablets each.

Oxymetholone-Alhavi (Iran) was recently updated. It is now packaged in foil and plastic strips of 10 tablets each 10 strips per hox

Oxymetholone IH (Iran Hormone) was also updated recently. The presentation remains similar (strips of 10 tablets, 10 per box), though the box looks more modern with gradient coloring.

Oxyanabolic from Asia Pharma (Malaysia) is now registered for sale in Thailand. The product comes in strips of 10 tablets, each with an embedded AP and 50. The product should also carry a unique product ID number that can be verified on the company website.

Androlic from British Dispensay (Thailand) is also still in production. The product comes in bottles of 100 tablets each. The tablets are green, scored on one side, and stamped with the BD snake emblem logo on the other. The product carries a holographic sticker to deter counterfeiting, though even this sticker has been duplicated with high accuracy.

Balkan Pharmaceuticals (Moldova) makes the product Anapolon. It is prepared in 50 mg tablets, with 20 tablets contained in each foil and plastic strip.

Oxybolone from Genepharm Greece is again in production. It contains 50 mg of steroid per tablet, with 10 packaged in each foil and plastic blister. Two strips are packed in each box. This product is widely distributed through export.

Oxitoland is manufactured in Paraguay by Laboratorio Farmaceutico/Landerlan. Each box holds 2 strips foil and plastic blisters of 10 tablets each.

## Anadur® (nandrolone hexyloxyphenylpropionate)

Androgenic	37
Anabolic	125
Standard	Testosterone
Chemical Names 19-nortestost 17beta-Hydroxyestra-4-en-	erone hexyloxyphenylpropionate 3-one hexyloxyphenylpropionate
Estrogenic Activity	low
Progestational Activity	moderate

## Description:

Nandrokee hesykosphenjeropismus is a slow-acting jūsculibė firm of the ambdes storid mardokue. Edesykosphenjeropismus, is a fairly ursaall mardokue ester in a structural sense. It is essertially mardokue proprieposjenos, with his bene curetada vide a lai of ore saddismus dogga antom and to destrout the organ attention of the structural sense. It is essertially mardokue proprieposjenos, with his bene curetada vide a lai of ore saddismus dogga attention of the structural organ attention of the structural organ attention of the structural organ attention of the structural organization organization

## History:

Neufolose heystryphendyprejonen son first described in 1907. If two developed into a medicine shorty date and was sold immely date if he value better men in such methods as a Austria, Sweden, Sweden, Sweden, Relgiam, Netferbands, and Germany, Anadre pensisted through the early 1907, and, following some mergers and equipients, was sold minity by Kalif Plumman. Kali would also sell the drug in France, but turker the brand trans Anador. Indirectal uses for the drug included consequences, clearate cast and instential disorders, and included as the principle individual consequences, clearate cast and instential disorders, and contained asset for the dispitable obsergations, clearate cast and instential disorders, and contained asset for the principle of the contained and the contained asset and the contained asset and the contained asset as a present lean-time-building analysis, with certain discussed assets, pigs, or considerate.

Nandrobene hesykosyhenylynopionate was not widely distributed outside of Europe, and in spale of a leng history of relative safetycould are list as a medicinal product. The 1995 merger of Kadi will lephal found for corpuspy Pharmica & Lylpian and would some spell the commercial cut to Anadra Pharmica & Lylpian dial continue to market for drug in hesykosyhenylephalemia in all courties. It is of mine that fiss evit of marketon was also sold by Loc in Spain (Annahri, Lurdheck in Darmick (Annahri, Euzochosi in Takety (Annahri, and by Spain in Grocce (Annahri), much used freed license with Kich all such preparations have size been discontinued as well, however, and marketone hesykosykosyhenylepomics in to longer washeld as a commercial medicine.

#### How Supplied:

Nandrolone hexyloxyphenylpropionate is no longer available as a prescription drug product. When supplied, it usually contained 25 mg/mL or 50 mg/mL of steroid dissolved in oil in a 1 mL or 2 mL ampule.

## Structural Characteristics:

Nandsohn besydosyphenylposponien is a modified from of nandsohn, where a carboxylic and east (prathespolyposphenylposponie and last been mattached to the 17-best hightody group. Estatelied streoids are less polar than five steroids, and are absorbed more slowly from the area of rigition. Once in the bloodbearm the east or increased to be (active) mandsohne. Estatelied steroids are designed to prolong the window of themposite effect following administration, allowing for a last separat priecuto in schedule compared to injections of five (uncestrafied) steroid. Nandsohne hesptosyphenylprospirates is designed to provide a slow release of mandsohne for pa four works of slow release of sealing to the provide a slow release of mandsohne for pa four works of slow release of mandsohne for pa four works of slow release priection.

## Side Effects (Estrogenic):

Nandrokes has a low tendans, for entrogan connection, estimated to be only about 20% of that seen with testioners.<sup>200</sup> This is because white the few on connect mandrokes to estandiol, in other trave active sites of staroid arountation such as adjoine tissue mandrokes in few separation hyprocess<sup>200</sup> Connecquent, estimages related size efficies are a meta hove concern with this day than with testionations. Elevand entergrain vole may still be noted with higher doning however, and my cause side efficies which are few referring to reterior, hope fire gain, and genomentals. An anti-estimony such as charging to enter termination exists on the recognition of the control of the property of the control of the property of the control of the reminister of the control of шимает сише тыу ос пессолу по рестен согодене эке сиссо и неу осси. Оте тыу alternately use an aromatase inhibitor like Arimidex® (anastrozole), which more efficiently controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood limits.

It is of note that nandrolone has some activity as a progestin in the body, 394 Although procesterone is a c-19 steroid removal of this group as in 19-normogesterone creates a hormone with greater binding affinity for its corresponding receptor. Sharing this trait, many 19-nor arabolic steroids are shown to have some affinity for the procesterone recentor as well 395. The side effects associated with procesterone are similar to those of estrogen including negative feedback inhibition of testosterone production and enhanced rate of fat storage. Procestins also augment the stimulatory effect of estrogens on mammary tissue growth. There appears to be a strong synergy between these two hormones here, such that expecomestia might even occur with the help of progestins, without excessive estrogen levels The use of an anti-estrogen, which inhibits the estrogenic component of this disorder, is often sufficient to mitigate gynecomastia caused by nandrolone.

#### Side Effects (Androgenic):

Although classified as an anabolic steroid, androgenic side effects are still possible with this substance especially with higher doses. This may include hours of oily skin acree and body/facial hair growth. Anabolic/androgenic steroids may also aggravate male pattern hair loss. Women are warned of the potential virilizing effects of anabolic/androsenic steroids. These may include a deepening of the voice menstrual irregularities, changes in skin texture facial hair growth, and clitoral enlargement. Nandrolone is a steroid with relatively low androgenic activity relative to its tissue-huilding actions, making the threshold for strong androgenic side effects comparably higher than with more androgenic agents such as testosterone methandrostenolone or fluoromesterone. It is also important to point out that due to its mild androgenic nature and ability to suppress endogenous testosterone. randrolone is prone to interfering with libido in males when used without another androgen.

Note that in androgen-responsive target tissues such as the skin, scalp, and prostate, the relative andmornicity of nandrolone is reduced by its reduction to dihydronandrolone (DHN) 396 397 The 5-alpha reductase enzyme is responsible for this metabolism of randrolone. The concurrent use of a 5-alpha reductase inhibitor such as finasteride or distractoride will interfere with eite-energife reduction of randrolone action, considerably increasing the tendency of nandrolone to produce androgenic side effects. Reductase inhibitors should be avoided with randrolone if low androgenicity is desired

## Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have heratotoxic effects. Liver toxicity is unlikely. Anabolic/androgenic steroids can have deleterious effects on serum cholesterol. This includes

## Side Effects (Cardiovascular):

a tendency to reduce HDL (good) cholesterol values and increase LDL (bad) cholesterol values, which may shift the HDL to LDL balance in a direction that favors greater risk of arteriosclerosis. The relative impact of an anabolic/androgenic steroid on serum lipids is dependant on the dose, route of administration (oral vs. injectable), type of steroid (aromatizable or non-aromatizable), and level of resistance to hepatic metabolism. Studies administering 600 mg of randrolone decanoate per week for 10 weeks demonstrated a 26% reduction in HDL cholesterol levels. 398 This suppression is slightly greater than that reported with an equal dose of testosterone enanthate, and is in agreement with earlier studies showing a slightly stronger negative impact on HDL/LDL ratio with nandrolone decanoate as compared to testosterone cypionate. 399 Nandrolone injectables, however, should still have a significantly weaker impact on serum lipids than c-17 alpha alkylated agents. Anabolic/androgenic steroids may also adversely affect blood pressure and triglycerides, reduce endothelial relaxation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and myocardial infarction.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Sunnlementing with fish oils (4 grams per day) and a natural choisterol/antioxidant formula such as Lipid Stabil or a product with comparable ingredients is also recommended.

## Side Effects (Testosterone Suppression):

All ambole/androgenic stendis when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testorious production. For sexis of comparison, studies administring 100 mg per week of mardosten decamonic for 6 weeks Inwe demonstrated an approximate 57% reduction in neurant testorious levels drug themps, 4x a dosage of 300 mg per week, this roduction network reduction levels drug themps, 4x a dosage of 300 mg per week, this roduction network reduced 170%, <sup>500</sup> 1 is believed that the progessional activity of mardosten to take contrables to the suppression of festostories synthesis drug fleungs, which can be marked in spite of a low tendency for estingen connecision. <sup>500</sup> Wildout the intervention of Stocknown seimfalling substances, testostories of she should return to normal with a 2-6 morths of drug secession. Note that prolonged physogranderopic lapogognadism can develop secondary to stendi abuse, necessating randed intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

## Administration (Men):

When used for physique- or performance-enhancing purposes, a dose of 200-400 mg given every week to 10 days was most common, taken in cycles eight to twelve weeks in length. This level is sufficient for most users to notice measurable gains in lean muscle mass and strength, which should be accompanied by a low level of estrogenic and androgenic activity.

#### Administration (Women):

When used for phosique- or performance-orienzing purposes, a dossage of 55 nn per week, or 100 ng ewe yell, or 1 days, was must common. Affixed pole sightly anticognies, somen are occasionally confronted with virilanties symptoms when taking this compound. Should writing side effects become a concern, the drug should be donestrated immediately to help prevent their perminent appearance. After a sufficient period of withdrawal, the shorter of the contract of

#### Availability:

Nandrolone hexyloxyphenylpropionate is no longer available as a prescription drug product.

## Anayar (oxandrolone)

Androgenic	24
Anabolic	322-630
Standard	Methyltestosterone (oral)
Chemical Names	17b-hydroxy-17a-methyl-2-oxa-5a- androstane-3-one
Estrogenic Activity	none
Progestational Act	ivity none

#### Description:

Oxardotone is an oral ambole steroid derived from dibydrostsotstorene. It was designed to have a very strong separation of amboles and androgenic effect, and no significant estrogenic or progestational activity. Oxardotone is noted for being quite mild as far as oral steroids are concerned, well talked of the promotion of strongly and quality mack those gains without significant side effects. Miligram for miligram it displays as much as six times the control of arazone activity of testosterone in assays, with significantly less androgeneity. — I ris drug is a favorie of dieting bodybuilders and competitive athletes in speedlanaerobic performance sports, where its tendency for pure tissue gain (without fat or water retention) fits well with the desired goals.

## History:

Oundrokes was first described in 1902. <sup>601</sup> It was developed into a medicine several years time by planmentaring just GD. Seark & Co. (now Phere), which sold it is the United States and the Netherlands under the Annuar tander name. Searle also sold/decreased the drug started different tander muss including Lourner (Augustra, Australia, Liplace, (Patas)) and the Common of the Common of the Conference of the Conference

By the 1989, the EDA bud slighty reflect the approved applications of constrolece to studde the promotion of weight gain following surgey, chronic infection, traumar, or weight loss without definite pathrophissiple; reason. In spike of its organic much record of safety, Searth electied to voltaring discorration for sales of Austraen on July 1, 1989. Lagging sales and growing public concerns about the athletic use of analytic strootic appeared to be at the root of this decision. With the Austrae band of the matter, constrolore and completely vanished from U.S. pharmacies. Soon after, countrolores products in international markets were considered to the control of the control of the control of the control of the must active of the control of the control of the control of the control of the second to the control of the con

It would be approximately six years before countelwine tables would be back on the U.S. munder. The product returned to planning wheles in December 1995, this im use after the Oundrin must by Bio-Technology General Corp. (BTO), BTO would corrians esling a for the EPA approved uses involving learning respectation, but fail about partial originary and constitutional delay of growth and pubers by possible partial, Timer's syndrome in girk, and constitutional delay of growth and pubers by possible partials. The product is very high selling project of the properties of

Constraint cortinues to be said in the U.S., but is now under the Snient helds (formerly, hanness a BEIG) is its correctly approach by the FDA for "higherine theory to promote weight gain after weight loss following extensive surgest; critons institution, or severe trainer and in some patients wood whold redding happlephysiologies grosses for 18 gains for minimisten normal weight, to offset the protein cataloolom associated with prolenged administration of corticosteroids, and for the relate of the best pain frequently accompaning consequences. Cortexis versions of the drug are now available in the U.S., which has reduced the price of versions of the drug are now available in the U.S., which has reduced the price of versions of the drug are now available in the U.S., which has reduced the price of versions.

#### How Supplied:

Oxandrolone is available in select human drug markets. Composition and dosage may vary by country and manufacturer. The original Answar brand contained 2.5 mg of steroid per tablet. Oxandrin contains 2.5 mg or 10 mg per tablet. Other modem brands commonly contain 2.5 mg. 5 mg. or 10 mg of steroid per tablet.

#### Structural Characteristics:

Oxandrolone is a modified form of dihydrotestosterone. It differs by: 1) the addition of a methyl group at carbon 17-abita to protect the hormone during oral administration and 2)

the substitution of carbon-2 in the A-ring with an oxygen atom. Oxandrolone is the only commercially available steroid with such a substitution to its basic ring structure, an alteration that considerably increases the anabolic strength of the steroid (partly by making it resistant to metabolism by 3-hydroxysteroid delydrogenuse in skeletal muscle tissue).

## Side Effects (Estrogenic):

Outdookee is not arcuntized by the body, and is not measurably estrogenic. Outdookee also offers no related progestization alone's gad, and art-strongs in an recessary when magths sectord, as genecomists should not be a concern occur among sensitive nikolatals. Since outgoing the sector of quality look to the playing with no fair of concern occur about now that the enter. This make it a florouth's steroid to use during entiring cycles, when water and fit reteriors are major concerns. Outdookees is also very popular arrange allekes in strategyinged sports such as spiriting, owinting and generated, in each disciplines one tousibly does not wort to early controlled to the quality of the section of the section of the section of the section of the controlled to the quality formed to offer the section of the secti

## Side Effects (Androgenic):

Albudge lessified as an ambelies stroid, antioqueis side effects are sill possible with this solution. The privated bosts of obj skin, near, and holyfields his growth. Antibolical-infraganties stroids may also aggravate mile partner hair bus. Women are swarned of the potential vising effects of antibolicant/aggress tousils. These may maked a desperance of the voice, mentantial regulatrics, changes in six to extent, facial hair growth, to the contract of the

The low antisoguic activity of countributes is due in part to I being a derivative of diphylotestosterous. This crustes a loss antisoguis seturiol because the agard lacks of expectly to internet with the 5-shph modutase enzyme and connect to a more potent "the hydro Thurn This is talked testudence, which is several times more active in antisogue responsive target times such as the scales, this, and prostate (where 5-shph noductuse is preceded in the contribute that is a strate to the contribute to the contribute this area of the contribute that the contribute

#### Side Effects (Henatotoxicity):

Omardoñose is a cl.7-alpha alyhted compound. This alteration protects the drug from deactionist to the five, railworing a vory high percentage of the drug entry into the bloodstream following oral administrations. CUT-alpha alchlarid aersholic/arabogenis stroids can be hepatotoses. Proboughof or high coposon my reseas in host dranges. In more instances file-threatening dysfarction my develop, it is advisable to vide a physician periodically during its comprehensive contraction of the Sweeks. In an effect to avoid exaction few trains.

Oundrolone appears to offer less hapties stress than other c-17 alpha alsylated steroids. The manificative distriblio soundrolone as a servicid that is not extensively metabolized by the liver like other 17-alpha alsylated onals, which may be a factor in its reduced haptotoxics; This is evidenced by the first that mer them at that of the free corporation is it intact when excreted in the urine. <sup>260</sup> Aunother study comparing the efficies of coundrolone to other alsylated agrees it including metabolisectories, metabolisectories, flowers also discontinuous to discontinuous consentant of the orange to a controlone coases the lowest subfebroraphilation (1887); a member of five sersions periention of the agree istende <sup>260</sup>. Our got countrolone produced <sup>278</sup> less ISBP retertion than an equal dosage of fluxopresserous, which is a considerable difference length or they up so thod 17-alpha alsylated.

A more recent study looked at escalating doses (20 mg, 40 mg, and 80 mg) of oxandrolone in 262 HIV+ men. The drug was administered for a period of 12 weeks. The group taking 20 mg of oxandrolone per day showed no statistically significant trends of hepatotoxicity in liver.

enzyme (ASI/ALI; auton-transfersse and autone auton-transfersse) values. Hose mentalising 40 mg protected a mean increases of opportunitely 50–50% in five enzyme values, while the group of men taking 80 mg noticed an approximate 50-100% increases. Approximately 10–11% of the patients in the 40 mg goury noticed World Health Organization grade till and 17 toxicly according to AST and ALT values. This digner jumped to 15% in the 50 mg assigned that its insensatively soft from tool read-valued according.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any benefatoxic analysis and more intermids.

## Side Effects (Cardiovascular):

Anabolic/androgenic steroids can have deleterious effects on serum cholesterol. This includes a tendency to reduce HDL (good) cholesterol values and increase LDL (bad) cholesterol values, which may shift the HDL to LDL balance in a direction that favors greater risk of arteriosclerosis. The relative impact of an anabolic/androvenic steroid on senum limids is dependant on the dose, route of administration (oral vs. injectable), type of steroid (aromatizable or non-aromatizable) and level of resistance to hensic metabolism Oxandrolone has a strong effect on the henotic management of cholesterol due to its structural resistance to liver breakdown non-aromatizable nature and route of administration. In the previously cited study in HIV+ males, 20 mg of oxandrolone daily for 12 weeks caused a mean serum HDL reduction of 30% HDL values were sunnressed 33% in the 40 mg group, and 50% in the 80 mg group. This was accompanied by a statistically significant increase in LDL values (approximately 30-33%) in the 40 mg and 80 mg groups. further increasing athernoenic risk. Anabolic/androgenic steroids may also adversely effect blood pressure and triglycerides, reduce endothelial relaxation, and support left ventricular hypertrophy all notentially increasing the risk of cardiovascular disease and muncardial infanction

At one time consultations was looked at as a possible sting for those sufficing from Gooders of high closhester or rispectives. Every status showed it to be capitale of lowering real cholesteral and triglyceride values in certain types of hypertipleams patients, which was thought to signify potential for this day as a light howering agent. With father investigation it was found, however, that any howing of total cholesteral values was accompanied by a renderibution in the most of good (PEL) to had (EDL) cholesteral that thoward general real constraints of the contribution of the ratio of good (PEL) to had (EDL) cholesteral that toward general real contributions of the ratio of good (PEL) to had (EDL) cholesteral that the rend contribution of the ratio of good (PEL) and actually makes it a potential charge in terms of creatine risk and the rend of the renderibution of the renderibution. As an onal CF of phase abstrated service, and the renderibution of the renderibution.

To help reduce cardiovascular strain à is advised to maintain an active cardiovascular coercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterol/artioxidant formals such as Lipid Stabil or a modert with correspolie incredires; is also procurrended.

#### Side Effects (Testosterone Suppression):

All anabolic/undrugents steroids when taken in does sufficient to promote muscle gain are expected to suppress endegrous to stortisoner production. Doundrudes in on exception, in the above-lead study on HIV winds, twolve works of 70 ang or 40 mg per day caused an object of the contraction of the production of the contraction of the production, with the 20 mg until 40 mg does causing a 25-30%-reduction, and the 80 mg group roticing a declared or more than 50%, Additionally, untilsteen to those with constitutionally delayed patienty have the mentionated significant suppression of endogenous LH and testisorous with stematical significant suppression of endogenous LH and testisorous with stematical contractions, and the contractional delayed to the contraction of the contractional delayed and the contraction of the contrac

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

### Administration (General):

Studies have shown that taking an oral anabolic steroid with fixed may decrease its biovasiability. It This is caused by the first-soluble nature of steroid hormones, which can allow some of the day to dissolve with undigsted distary fit reducing is absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty storough.

#### Administration (Men):

The original prescribing guidelines for Amour called for a dish diseage of between 2.5 mg and 20 mg per day (5.5 livel pix leging sections). His was sussiany recommeded for a popiciol of two to four weeks, but occasionally it was taken for as long as three morths. The dosing guidelines recommended with the current U.S. production form of the day (Ouardin, Soviert Plearmenterfiels) also call for between 2.5 and 20 mg of drug per day, that his intermediate (see 2.6 Text 4.0 mg of the section 1.5 mg of the 2.5 mg of the control of the control

Ouardiones is often combract with other strouds for a more dammite result. For example, while budging one might opt to add in 200-400 mg of a testiconous enter (cypiomate, enterflute, or proposately per osset. The result should be a considerable gain in some mascle testing of the considerable gain in some amounting setward such as 150 mg per usels of a redrolease estee or 200-300 mg of Provideolatif (methodose centralless). Soft sucks are highly floword for increasing definition and muscularity, An in-brivacen (dammas gain) might be to add in 200-400 mg of characteristic and the considerable (matchine descentace) or Expansions (obstitute and the considerable) (matchine descentace) or Expansions (obstitute and the considerable) (matchine descentace) or Expansions (obstitute and the considerable) (matchine) (matchine) (methodose centralless) (and the force barriors) or Expansions).

#### Administration (Women):

The original prescribing guidelines for Amour did not offer separate doning recommendations for women, although it was indicated that women who were program; or my become pregnant, should not use the dung. The current guidelines for Osundrin also do not make special dosing recommendations for women. Weren who far for the smeaddings effects on semantic proposes, a duly dosage of Section 1 to the contraction of the dosage. The word has described the contraction of the dosage of the dosage, the contraction of the contractio

#### Availability:

Pharmaceutical preparations containing oxandrolone are fairly limited. The drug is unavailable in Europe, and with a handful of exceptions in the west, its production is increasingly being stiffed to less regulated markets in Asia. In reviewing some of the remaining products and changes on the global pharmaceutical market, we have made the following observations.

Various forms of generic oxandrolone are now available in the U.S. in both 2.5 mg and 10 mg dosages, from manufacturers such as Par Pharm, Sandoz, Upsher Smith, and Watson.

Brand name Oxandrin is still available in the U.S. under the Savient brand name. It comes in bottles of 60~(10~mg) tablets or 100~(2.5~mg) tablets each.

The generic Italian product Oxandrolone (SPA) is no longer available. It was previously being made for export sales only. There are no remaining oxandrolone products available on the Italian market Atlantis (Mexico) produces an oxandrolone product called Xtendrol. It carries 2.5 mg of steroid per tablet, and comes in a box of 30 tablets each.

Asia Pharma makes the product Oxarabolic in Malaysia. It comes in strips of 10 tablets each, 10 strips per box. Each product should carry a unique product ID code that can be verified with the company for authenticity. This product is presently export only, but the manufacturer claims to be in the process of seeking Thai FDA approval.

Balkan Pharmaceuticals (Moldova) makes the product Oxandrolon. It is prepared in 10 mg tablets with 20 tablets contained in each foil and plastic strin

## Andractim® (dihydrotestosterone)

Androgenic	30-260
Anabolic	60-220
Standard	Testosterone T propionate

Chemical Names	5-alpha-androstan-3-one-17beta-o 5-alpha-androstanolone
Estrogenic Activity	none
Progestational Acti	vity none

## Description:

Andractin is a prescription storod prepunsion that contains the potent androgonic storoid displostestostenous rules in product cross in the form of a transferant sig hypacilly containing 2.5% efficiences the transfer in subgrant take, As with Androgot, roughly 10% of decidences of the containing the containing and the containing the containing and the c

Dighotesotsomes is a weak mucch buller because it is externely open to aleration by the 3-slab-plongous-oil-diphogeness compute, neeponable for becausing down action stoods like DHT in the transie metabolists. 3s-H5D is present in high quartities in mucch instant, running interference between the outset of merchanic and the antiopurnearing first receptor. Testostomes is very resistant to the crypta, however, which allows it to be a much traver effective muccle-building agent. 3s-H5D stored describation in mucch instance causes the same problem with Province (1-methy-disphotesotomeous), DHT and acception, proposed to the proposed of the proposed proposed of the proposed scaling, and province and the proposed of the proposed of the proposed of the scaling, and province and the proposed of the proposed of the proposed of the proposed scaling, and province and proposed of the proposed of the proposed of the proposed scaling, and province and proposed of the prop

#### History:

Digitates observed was first symbolised in 1935; 4°° This strong anthogan was put are consistent medical nearling the 18-1959; 4 are a rises of experiments demensaring their I had measurable analysis exhauster, and was of fifti wathe exhaust. Digitates outstance, and was of fifti wather exhaust. Digitates outstance was on of fifti wather exhaust. Digitates outstance and was offer wather exhaust the proposed of the proposed of

fueled partly by estrogens. The latter two indications are considered local applications of the drug, and the DHT gel is applied directly to the tissues requiring treatment.

The primy munificative of displotostostown get globally is Besins International, based in Firence. Besins produces the drug under the Ardaction runes, seeing it is Finence and scarcely in other parts of Europe. Displotostostome gas are found much less commody coasis of Europe, and presently no scale preparation is commercial profile in the Little Ardaction from Besins in the U.S., Mexico, and Carada. The firm amounced an interest in the day for several use, including arthurgous producers in run over aga soft, resting besign protost lepterspoty, and cortofaring HIV- associated wasting. Its use as an anabole may be desirable with IV because at the dome domestical flat unique prison European September 10 of the Control of the Control

## How Supplied:

Hydroalcoholic transdermal dihydrotestosterone gels are available in select human drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 2 % dihydratestosterone by weight.

## Structural Characteristics:

Andractin® is a hydroalcoholic gel containing 2.5% of dihydrotestosterone (free) by weight. It is designed to provide a continuous transdermal delivery of dihydrotestosterone for 24 hours following application to the skin Approximately 10% of the applied dose is absorbed across the skin durino each 24-hour period

## Side Effects (Estrogenic):

Diplotectosterone is not anomized by the body, and is not measurably estrogenic. An artisestrogen is not necessary when using this strend, as generous and water referring should not be concerns even among sensitie artiskinkab. DHT also has inherent artisentogenic properties, corepricing with other sistenties for himsight to the arountase engine. Percetamous diplotestosterone may be an effective option for the tenturest of generous miles and have been produced as good level of success with neutraling certain forms of the breast stone account of the breast stone around a considerable and the produced and the prod

## Side Effects (Andmoenic):

Digitalisations to the strongest natural multi-analogues. Higher than normal theraposite does are likely to produce analoguesis de diects schaling objects, some, and body fedicion har growth. Men with a garactic prodisposition for har loss (anthogonetic alspection) may nonce accedented multi-pattern buildigs. Worker are sourmed of the potential wirsing effects of anthological disposition strongests, especially with a storage androgon such as disployationationers, these may related depening of the vicine, measural irregulations, clausing in skin teature, facial last growth, and closed enlargement. Note that the 5-light new form of the strongest of the strongest

#### Side Effects (Hepatotoxicity):

Dihydrotestosterone does not have hepatotoxic effects; liver toxicity is unlikely.

#### Side Effects (Cardiovascular):

Ambioi-andrognie steroids can have deleterious effects on serum cholesterol. This includes a tendency to eclote IPML (good) cholesterol values and increase IDM, food) cholesterol values within may shift the IPML to IDM balture in a direction that favore greater risk of values, which may shift the IPML to IDM balture in a direction that favore greater risk of an ambioi-androgenie steroid on severam light is dependant on the dose, route of administration (oral vs. injectable), type of steroid caromatizable or non-anomatizable), and level of resistance to hematic metabolism

Anabelicandrogueis steroids may also adversely affect blood pressure and trighcerides, reduce endobted levelation, and support fit ventricult hypotroplys, all potentially increasing the risk of cardiovascular disease and responsable inferiors. Therapears does of displostestosterous case also correct insufficient androgue production in otherwise behalfly aging mus are utilisely to increase adversages; risk. Higher does are likely to increase afferogueis risk, but less durantically than equidant this closes of synthetic oral formations.

To help reduce cardiovascular strain à is advised to maintain an active cardiovascular coercise program and minimize the intake of saturated fats, cholsterol, and simple cardohydrates at all times during active AAS administration. Supplementing with fish ols (4 games per day) and a natural cholstero/lamitoxidant formals such as Lipid Stabil or a product with commarable inserdenies is also recommended.

#### Side Effects (Testosterone Suppression):

All ambolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone strainfairing substances, testosterone levels should return to normal whitin 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradism can develop secondary to steroid abuse, necessistating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (Men):

To text androgen issufficiency, hydroxloobie transformal dilydrotestosterous gils have been used in dosse rangig from 16 to foliept end (v1.6-6-dipp of hommor delevered). For physique- or performance-enhancing purposes, higher doses would be necessary to achieve storage spamplaysional roles of dilydrotestosterous. Logal effective dosses begin mage, or \$5-100 mg per day, or \$-100 mg of hommor delevered systemically ending to the control of the control o

## Administration (Women):

Hydroalcoholic transdermal dihydrotestosterone gel is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to produce vilidine side effects.

## Availability:

tube

Pharmaceutical preparations containing transdermal dihydrotestosterone gel remain scarce. The legitimate supply seems to be scattered into isolated markets of Europe, Asia, and South America. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

One of the most notable preparations is Andractim from Besins-Iscovesco in France. It contains 25 mg/mL of hormone in a 100 gram tube.

Andractim is produced in Belgium by Piette. It contains 25 mg/mL of hormone in an 80 gram tube.

Andractim is produced in India by Chemec. It contains 25 mg/mL of hormone in a 100 gram

#### Andriol® (testosterone undecanoate)

Androgenic	100
Anabolic	100

Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate
Progestational Activ	vity low

# Description:

Standard

Audrial B: a m cal bestoterous preparation that contains 40 mg of testosterous undercament (in an olbow) in a song dight nepadat. This drag two reliferate than not call analodic steroids, which are usually c-17 alpha alsylated to service first pass methodism under the contract of the c

Standard

## History:

Oral testasterore indicaronic capsales were developed by international drug giant Organon (now MerchASD), and first introduced into chiral trials during the early 1998; The drug was a consequent of the case as a pencetional consequence of the case as a pencetional consequence of the case and the consequence of the case of the ca

In 2003, Organo began replicing its Andrial products with Andrial IT Testcoppis. The new formalistic improves on the strong institutions of the original Andrial preparation, which needed to be logst under refrigeration at the phrames. The data was second at room temperature over dispensed in the products record to be constant at room temperature. Outside of refrigeration, however, the drug functionally had only a 3-month self-life. The row Andrial Testcoppes are designed to shops be stored at room temperature, and hose a shall life of 3 years. The new fermalation is considered to be bioopticulated to the older version, and can be abstituted in patients without any change in donogast\*10 Gene the landing abstrategy and bioopticulosity; it is likely that the new Testcorps will slowly come to enclose all of the local Andrial recognition.

In spike of its wide availability, Androl lists never been a popular hem among athless. This is Bickly due to the light relative cost of the ding, and it is to protestly compared to other pharmacential preparations, particularly ispectable testostenoe compounds and the more potent synthetic on an anti-oles streamly. Sill, Androl remains a protestir of these among those athless not interested in using sigestable medications and preferring to avoid the agreement of the dispersion of the anti-oles and anti-oles and anti-oles and preferring to avoid the agreement and the anti-oles and anti-oles and anti-oles and anti-oles and anti-oles and anti-oles and prescription coal lastostenoe undecented. Anti-ol local lists maintained a prominent share of the global learner applement market size of 1990's.

## How Supplied:

Onl testosterous undecanous preparations are available in various haram dng markets. The older formulations supply-40 ng of testosterous undecanous in olde: calc, contained in small soft galatin capaside. Aufraldo Testocaps supplies 40 ng of testosterous undecanous in causer of and propylene global curvatulenta; contained in small soil galatin capasides. Submarkets are supplied to the property of the contained of the property of the Submarkets the contained to the contained to the contained of the containe

## Structural Characteristics

Andriell's contains testostenore that his been medified with the addition of carboxyle, acid test (undexnote, acid) at the 17-bets belloword group. The esteride hormore is more fit solable than base (five) testostenore, and has been dissolted in oil and encapsulated for onal arbinisations. Significant absorption of onal netsostenore undexnote takes plote through the lymptatic rotee, bypassing the first poss through the five. Andriell's is designed to provide a padie in testostenore lavels were necessarile such as after a destriction, and with repeated to the contraction of the

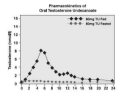


Figure 1. Median response pharmacokinetics after oral administration of 80 mg of testosterone undecanoate in fasted and fed states. Testosterone absorption is impaired when taken without meals. Source: Andriol Testocaps online information, Organon, Citation Bachus et al. 2001. Andriol.com

#### Side Effects (Estrogenic):

Testoderore is madily anomaticed in the body to estraidol (estrogen). The anomatics (estrogen synthesis) engine is repossible of this metabolism of incostorous. Desaduel estingua looks can cause side effects such as increased under neutrino, body fair gain, and the properties does will revise the licibility of estimates the contraction of estrogenic side effects. In such cases, in an artis-entogen such as chemiptene create or termodific estrains is commonly applied to prevent entogenic side effects. One may alternately use an anomatic simblified the Artisticko (unantonoida), which more efficiently controls enough the preventing its synthois. Anomatics have been also also the simble simble side of the simble side of the side of t

## Side Effects (Androgenic):

Textostorous is the primary mile androgon, responsible for minimizing secondary male sessual entancerists. Taking and textostorous underanent in doses exceeding normal theraportic levels is Body to produce androgonic siste effects including only sids, now, and thoughtest law growth. More with a general prediposition for their loss (androgonic real aspects) may notice accelerated male pattern balding. Women are warned of the potential viriliary effects of analocidatorogonic securities, opecially with a stong androgons ach as textostorous. These may include despering of the voice, mentional irregularities, changes in skin texture, faciliar growth, and formed enlargement.

In androgo-responsée traget fissus such as the skin, scale, and prostate, the light relative androgoriéey of lostatemes deportate in serulcation of higher telestatement (PAIT). The S-alph reductatement (PAIT) the S-alph reductatement (PAIT) the S-alph reductatement (PAIT) the S-alph reductatement (PAIT) to a financiate of a state-time of a reductation of sestatement and reductated with referred with side-specific potentiation of sessatement action, lovering the tendency of sestatement (PAIT) to the session of t

separation or restoratione's anatonic and antirogenic properties is not possible; even with total 5-alpha reductase inhibition.

## total 5-apria reductase minimon.

Textosterone does not have lepotatoxic effects fivor toxicy is utilitely. One study ceanning the potential for lepotatoxicy with high does of oul testosterone by administering 400 mg of the hormone per day (2,800 mg per weed) to a group of male subjects. The hormone was given day for 200 days, and produced on sugisfact changes in fere engine values including serum abstruct, bittely, and produced on suggisfact changes in fere engine values including serum abstruct, bittely, and produced on suggisfact changes in fere engine values including serum abstruct, bittely and the produced of the control of th

## Side Effects (Cardiovascular):

Side Effects (Henatotoxicity):

Ambolicardroganic stronds can have deleterious effects on neume cholesterol. This irelates a reductry to reface IEEE (agod) cholesteroi classe and irecurse IEEE, (doa) cholesteroi classe and irecurse IEEE, (doa) cholesteroi classe, without my shift the IEEE. to IEEE cholesteroi in direction fluid tisson greater risk of electronic reduction of the contraction of the contraction of the contraction of the contraction contraction contraction, proceeding contraction for an extraction of the contraction of t

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular sceneric program and minimize the ristake of saturated fits, cholsterol, and simple cardiohydrates at all times during active AAS administration. Supplementing with sho id-grams per day) and a natural cholsterolamioxidant formula such as Lipid Stabil or a moder with conversable inverdience is also recommended.

## Side Effects (Testosterone Suppression):

All anabolic indegratic storieds when taken in does sufficient to promite muscle gain are expected to suppress endogenous bestderency production. Decisionrous is the primary may expected production and the primary may be sufficient to the production of the production of the production of the production of manual seroid hormones. Whose the intervention of testudence-inhabiting englation of manual seroid hormones. Whose the intervention of testudence-inhabiting substances, testudence levels should remain to mean within 1-4 ments of engla secsions. Now that produced programs are develop secondary to steroid abuse, reconsidered prediction resolution.

The above side effects are not inclusive. For more detailed discussion of potential side effects we the Steroid Side Effects section of this book

#### Administration (General):

Andriol should always be taken with meals, preferably containing a moderate fat content (20 gams) to meanize hyphine absorption. Very low biasovaibibility has been reported when taken in the fasted state. The total daily dosage should be divided into a minimum of two applications, taken in the morning and evening, to maintain more consistent elevations of serum estosterous.

## Administration (Men):

For the number of two subseque locks, prescribing glabifies for Aukid recommend an similar dosage of 200 ki floring dashy for 25 works. Based on the level of effort, a dish marketures dosage of 40-120 mg is usually contrased at the poir. For holyholding purposes, lagier doses sould be required to exect strong semply-thological twols for tensorsors. This world generally call for a minimum dosage of 240-250 mg per day (o-8 cquality), later in 250-260 of 50 weeks. A minimum dosage for 240-250 mg per day (o-8 cquality), later in 250-260 of 50 weeks. A minimum dosage, thorwise, would not consider the contrast of given the relative price of Audrich preparations, making injectable testosterones much more cost effective and propular. Given the relative low potency of Audrich (when taken by athless: it is most commonly used in combination with other agents. Testosterone drugs are ultimately very versatile, and can be stacked with many other anabolic/androgenic steroids depending on the desired distribution.

## Administration (Women):

Andriol is not prescribed to women in clinical medicine. This drug is not recommended for women for physique or performance-enhancing purposes due to its strong androgenic nature and tendency to produce virilizing side effects.

## Availability:

Oral testosterone undecanoate remains widely available. It is produced almost exclusive by or under license from Organon (now Merck/MSD). In reviewing some of the more popular products and changes on the global pharmaceurical market, we have made the following observations:

In November 2009, Organon (a subsidiary of ScheringPlough since 2007) became part of Merck/MSD. All Organon products are expected to transition over to this label. It is urknown what (if any) changes to expect in the global distribution of Andriol products.

The sodied design of the Arthrio products is extremely defined to duplicate. Arthriol is prepared as a howard colored capused the contrass of crisical is recompletely sealed. The prepared as a howard colored capused the contrass in crisical is recompletely sealed. The Estexcaptis see also soft out glossy capusles, but these are made out of a transparent comange gather invitate. Inside there is a public self-wide product of the season of the seaso

## Androderm® (testosterone)

**Progestational Activity** 

Androgenic	100
Anabolic	100
Standard	Standard

Chemical Names	4-androsten-3-one-17beta-ol
Estrogenic Activity	moderate

## Description:

Andodomil's is a testosterou debory o yetem that utilizes an aditesies "pactif" to debor the hormone transdermil. The testosterous reds if sisolobed in an adolobic gal similar to AndroidiRi, except here the gal is contained in a protected external drug reservoir. This design provides approximately double the former betweenhild by OntroidiRi, and also severely intens the transfer of testosterous to other people during regiones also to-sisten contact. The patches come in two strengths, 2.5 mg and 5 mg, radicing the amount of testosterous each is to supply systemably over a 2-brine princip deby containing the answer of testosterous each is to supply systemably over a 2-brine princip deby containing the answer of testosterous each is to supply systemably over a 2-brine princip deby on mind from the advancement reportionly. Andredermil was designed to mind the manufacture of the containing the and the advancement of the containing the contract of the c

low

#### History:

Autoclemió was developed in the United States by ThearTest (Sal Lake Crs), it was approved for sale as pecucipionagen by the Food and Dug Andismistration in September 1955, and is ridicated for testosterone replacement therapy in men with a deficiency or adherent of endagones testosterone. This related causes of primary ploggrandism, which may be caused by copyractedism, bilateral torson, orchis, vaneling tests syndrom, cordicators, Kinelder's syndrome, cheerdurgen, or alcobalt-pay med tackey, it is also prescribed to men physiogenethologies (propograndism relating patients with intering personness or takerings promone-releasing between (LHBI) deficiency caused by turners, gigs, or antifation. Firmus physiogenethologies assume that the proposal control of searchest with the testionness and two terms (LHBI) deficiency caused by turners, seak this product in the United States under the Anthonomie band name. In Europe, the AMOSSE band from Anni is most commontly four.

#### How Supplied:

TheraFoch's transformal testosterone system is available in select human drug markets, where it is commonly sold as Androderm® or ATMOS®. It is produced in two strengths, one containing 12-tmg of testosterone, and one containing 24-lmg of testosterone. These are intended to deliver approximately 2.5 mg and 5 mg of testosterone systemically to the patient over a 24-brun period.

## Structural Characteristics:

Androderm® is a transdermal drug delivery system that contains an alcoholic gel of testosterone (free) enclosed in an adhesive patch with a protected drug reservoir. It is designed to provide steady but varying levels of testosterone transdermally during each 24brun period of semiletation.

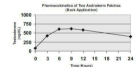


Figure 1. Mean serum testosterone concentrations (ng/dL) measured during singledose applications of two Androderm 2.5 mg systems applied at night to the back. The figures reflect the greatest response in a study comparing four different sites of application (abdomen, back, thigh and upper arm) in 34 hypogonadal men. Source: Androdermic mescrethios information. Wastom Plastrus. Inc.

#### Side Effects (Estrogenic):

Textosterors is readily arountized in the body to estatiol (estrogen). The anomatuse (estrogen printlense) entryme is responsible of the intentableon of intestorence. Benedic estrogen levels can cause adde effects such as increased under restricts, body fit gint, and the properties of the printlense of the district of estimates in commonly applied to prevent entire that the printlense of the printlense of

#### Side Effects (Androgenic):

iestosicurus is tie primusy mas aurungan, responsus to rimitaming seconatary mas essual naturactivisis. Escendigi therapinet doses is fiely to produce aurlouguis side effects ieskiding oby skin, acue, and bodyficial had growth. Men with a genetic effects ieskiding oby skin, acue, and bodyficial had growth. Men with a genetic effects in the produciogation for his time bots (andeaponis and profess cochreated mite pattern balding). Women are warned of the posterial viriliary effects of anabolicitarhospiris seconds, seepecially with a story gardengen such sustationer. These may include deepering of the other contractivities of the produced produced and the story of the contractivities of the story of t

In authorps responsive traget fissus such as the skin, soal, and prostate, the high relative authorpsic yello restorates is dependent on its relations to difficultiestosterous (ERI). The S-alpha reductase enzyme is responsible for this metabolism of testosterous. The concurrent use of 3-alpha reductase inhibitors who are frametized or distantistical interfere with sits-specific potentiation of testosterous action, lovering the teachersy of testosterous drags to produce authorisms side effects. It is important to remember that anticloid and segmention of testosterous's authorism and authorisms are supported in the production of testosterous's authorism and authorisms.

## Side Effects (Hepatotoxicity):

Testostenore does not have hapstonoise effects, here toxicity is utilicely. One study examined the potential for hapstonoisely with high does of relesstatement by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile adjucts. The sectoral was eather only so that higher peak concentrations would be reached in hepatic tissues to the contract of the contraction of the peak of the contraction of the peak of the contraction of the peak of the contraction of th

#### Side Effects (Cardiovascular):

Anabole/androganie steorike on hove dekterious effects on serum cholestered. This includes a technique to reduce IIII. (apid) cholestered values and increase III. (bad) cholestered values, which may shift the IIII. In III. In III.

To help reduce cardiousecular strain it is advised to maintain an active cardiovascular cuercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterolamioxidant formals such as Lipid Stabil or a modest with companible inconferies; solar procumented.

## Side Effects (Testosterone Suppression):

All antibolization/going storoids when taken in doses sufficient to promote muscle gain are expected to suppose condegorous testostorous production. Testostoroes it the primary unitary and offers strong magnitic footback on endogorous testostorous production. Testostoroes is the primary unitary fine testostoroes bead though soil. Revisele, however, as strong effect on the probuleniar regulation of natural steroid hormous. Without the intervention of instostorous-straining substances, testostorous-less dead return to remain while 1-4 mentric for disposition generally associated, not be prolonged programatoropic in programation can develop secondary to steroid abuse, reconsisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

\_\_\_\_

Antoclemb is applied daily flexible bed to intact, clean, day skin of the back, upper arms, thigh, and/or addomen. The sket(s) of application should be rented so that no patch is reapplied to the same area in less than 7 days, Lower becombibility may be roticed in some areas of the body, and an the clear and called so-ext a broay area of the body, can be a first and a few and the section in the should also not be applied over a broay area of the body, can appear of the body to the body of the section of the should be an extra the section of the section of

#### Administration (Men):

To text androgen isoalficiarcy, the prescribing guidelines for Androdemile recommend two 22-m guidelines or 5 mg push per day Morning serum testionen becked as their measured; at which posit the physician may adjust upwants or downwards if necessary. For physician- or perfaminence-relaturing purposes, higher downs would be necessary to achieve superphysiological levels of testistoriene. This would require at least frence of nor 5 mg or egglet 2.5 mg guideline per day, debering guporamited [1-2] mg finely finely the size of the strength of the most testis to notice gains in muscle size and strength, although this is not a vory realistic that in a practical sense. Lower does may be used, that typically when accompanied by other antibolic anthogonic strotch. Substitutes to take the desired can be confidented with mays ofter antibolic anthogonic strotch. Substitutes to take the desired and can be confidented with many ofter antibolic anthogonic strotch. Substitutes to take the desired

## Administration (Women):

Androdemi® is not FDA approved for use in women. Testosterone is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic rature and tereforcy to morther wideline willowine with earliest.

## Availability:

Transdermal testosterone patches are manufactured in many countries worldwide, and remain widely available. Common trade names include Androdermi® and ATMOS®. Given the high sophistication and low black market value, counterfists have not we been reported.

#### AndroGel® (testosterone)

Androgenic	100
Anabolic	100
Standard	Standard

Chemical Names	4-androsten-3-one-17beta-o beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate
Progestational Activity	/ lov

AndroGel® is a transdermal hydroalcoholic testosterone gel that contains a 1% concentration of testosterone by weight. It was originally released in 2.5 gram and 5-gram sachets, equating to a total per-application testosterone dose of 2.5 mg and 50 mg respectively. The AndroGel® prescribing information states that the product has a

transfermil broundability of approximately 10%. This means that each 2.5 or 5 gam does will deliver approximately 2.5 mg or 5 mg of bommon systemically. With this mode of administration, testosterone levels begin to elevate approximately 30 minuses after the get is applied to the body, and substantial elevations in norm androgen levels are acheeved within 4 loars. Tostosterone levels will remain elevated for approximately 24 hours after loars. Tostosterone levels will remain elevated for approximately 24 hours after loars. Tostosterone levels will remain elevated for approximately 24 hours after loars. Tostosterone levels will remain elevated for approximately 24 hours after loars. Tostosterone levels will remain elevated for approximately 24 hours after loars. Tostosterone levels will remain elevated for approximately 24 hours after loars.

# History:

Androidal's was developed in the United States by United Pharmacericals, a dissist of Solway I was approved by the FDA for six as a prescription flare florthamy of 2000. It is indicated for use in salds males with conditions associated with a deficiency or absence of endagenous testisteem. This richted cesses of primps hypogenoidam, which may be caused by explored, including his particular to be caused by explored, including believed to text hypogenoidam, believed to be caused by explored, including particular to the caused by explored, including believed to text hypogenoidamy believed by the proportion of the propor

Other transformal testosterone lephotocholosis gals have been released in the U.S. and abrund science for anticolific. Bestiffs by Australian Philimetericals is perhaps for most web-known competing brand, and widely in the U.S. and Enaper. This behavior is the perhaps for the perhaps of the

# How Supplied:

Hydroalcoholic transdermal testosterone gok are available in many human drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 1% testosterone by weight; packaged in volume tubes or single-dose packets containing 2.5 gamss or 5 gams of gel. AndroGell® (U.S.) is also produced in a pump dispersor containing 2.75 grams of each which delivers 60 metered anolecations of 1.25 grams cell.

# Structural Characteristics:

AndroGel® is a hydroalcoholic gel containing 1% of testosterone (free) by weight. It is designed to provide a continuous transdermal delivery of testosterone for 24 hours following application to the skin Approximately 10% of the applied dose is absorbed across the skin during each 24-hour period.

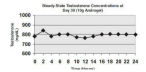


Figure 1. Steady-state testosterone concentrations in blood, measured 30 days after beginning therapy with AndroGel (10g application). Drug was applied to the body once daily.

#### Side Effects (Estrogenic):

Textustence is result; arountated in the body to estandial (estrogen). The arountase (estrogen printlens) outputs a responsible of the intentables on first stoutence. Eventual estings in look can cause side effects such as increased under resertion, body far gint, and the temporate dosses will revenue the Richford of estrogens size effects. In such case, an auti-strongen such as Completes centure or turnosfire crimas is commonly applied to prevent entogen; size effects. One may alternately use an arountase infibrit size Arrindace (unationals), which more efficiently courted entogen is self-effect, however, and may also the complete of the contraction of the complete effects on body lights.

#### Side Effects (Androgenic):

Textstenow is the primary mile androgan, responsible for mintaining secondary mile sessual culturativists. Exceeding normal theraparis does is fixed, by produce androgans; side effects including only side, ance, and bodyficial line growth. Men with a genetic side effects including only side, ance, and bodyficial line growth. Men with a genetic produptions from its lines (anadoparis) adaptive productional through the halding. Women are warmed of the potential virilizing effects of anabolicitarloogies; secroids, specially with as story gardogan and as to sectionee. These may include deporting of the voice, menstrual rengalarities, changes in skin texture, ficial hair growth, and citoral enhancement.

In androgor-responsive target fissues such as the skin, scale, and prostate, the high relative androgorizely of Isotratories is dependent on its reduction to diplottostosteneto (EMT). The 5-sliph reductase enzyme is responsible for this metabolism of testosteneto. The concurrent use of 5-slab reductase inhibitors such an instruction of classifications. The damped of the contraction of testostenete action, lowering the teachery of testostenete with site-specific potentiation of testostenete action, lowering the teachery of testostenete with site of the site of the contraction of the site of the contraction of the contraction of the contraction of the analogous cells of the contraction of t

# Side Effects (Henatotoxicity):

Testosterore does not have hepatotoxic effects, her toxicity is utilicly. One study examined the potential for hepatotoxicity with high does of testosterore by administering 400 mg of the hormone per day (2,500 mg per words) to a group of mile subject. The secrol was taken only so that higher pack concentrations would be nearbed in hepatic tissues compared to international rigisticts. The hormone was given dudy for 20 days, and produced no significant changes in her engine values including serum alburnit, bilirabin, adminiscenarios-transfers, and alkaline photolatuses. <sup>524</sup>

# Side Effects (Cardiovascular):

Ambelicantegaries steroids can how deleterious effects on serum cholestord. This includes a tendency to relate EIE (agood) cholestord values and irreases ELE (doug) cholestord values and travels ELE (doug) cholestord values and travels explicated an analysis of advantages of a characterious. He retaine impact of an anabociantegories retained on exams light is dependant on the dose, route of administration (red to, sijectishle), type of datested Ambelicantegories retained and administration (red to, sijectishle), type of datested Ambelicantegories retention my also advantes) effect bodd present and trighterious retained and proportial districtions in otherwise healthy gaing sum on a table by the contract and proportial districtions and proportial di

To help reduce cardiousocular stania it is advised to mintain an active cardiousocular cuercise program and minimize the intake of saturated fits, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oil (4 grams per day) and a natural cholesterol/antioxidant formula such as Lipid Stabil or a product with comprabile ingedients is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolicalmogenic stends when taken in doses sufficient to promote muscle gain are expected to suppose endogenous testostoreer production. Testostores it he primary male analogo, and offers strong negative feedback on endogenous testostrone production. Testostore-based days soil, Revisic, have a strong effect on the hypothanies, regulation of natural steroid hormous. Whost the intervention of testostrone-strainling substance, testostore beeds should return to normal while 1-4 meths of sing secsions. Note that prolonged hypogenedotropic hypogenedism can develop secondary to steroid shose necessation models intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book

#### Administration (General):

Textostorous hydroxlooks gal is applied daily (predenshy in the moning) to iract, claus, dy skin of the shorkste, upper arms, and of advances Patiers about be careful about transforing textostorous to their formit portion(5). The prescribing information for Anadoccifel suggests that patients was their funds immunition by this open and water about the patients and their funds immunition by this open and water about the state of the

## Administration (Men):

To text androgen insufficiency, the prescribing guidelines for AndroGel® recommend sinting therapy with a 5g dudy does (debering 5 mg of restostores systemicals). Serum textostores levels are measured after 14 days, at which poir the physician my adjust upwards to 7.5g or 10g fit accessary for physique-or performance-relaturing purposes, higher does soudd be necessary to achieve supraphysiological levels of resistorence. The most commen does here is 20 grams per day, which debees approximately 20 mg of textosteror. This level is sufficient for most tenes to notice significant gains in music size and strength. Lower does are also regalant used by some affection, but typically when accompanied by other antibolicalizatiogues strends. Set institutely very sersific, and can be contributed with mmy other antibolicalizatiogues strends. Sets to take the desired

#### Administration (Women):

Hydroalcoholic transdermal testosterone gels are not FDA approved for use in women. Testosterone is not recommended for women for physique- or performance-enhancing purposes due to its strong androagen nature and tendency to produce vinitiane side effects.

## Availability:

Hydroalcoholic transdermal testosterone gels are manufactured in many countries worldwide, and remain widely available. Common trade names include Androgellii, Testimili, and Testogellii. Given the high sophistication and low black market value, counterfeits have not we been widely reported.

# Andronaq (testosterone suspension)

Chemical Names	4-androsten-3-one-17beta- 17beta-hydroxy-androst-4-en-3-on
Estrogenic Activity	moderat
Progestational Activ	vity lov

## Description:

Androgenic

Anabolic

Tectostorous suspension is an injectuble prepuration containing testosterous (no eater), usually in a water base. Armage bodyladians, "suspension" is form to be an externed potter mass agent. It is often said to be the most powerful rijectuble steroid available, producing user parting pains in marke mass and strength. This is impayd due to the very black action of the drug. When ussing a slow-acting ol-based steroid like Sustanomi? SQ, it can take weeks before a pack testostorous levels a reached. With suspension, it just an untroot lower. This will usually reash in the affacte starting to motice size and strength gains by the end of the first work. By the time the affacte is 50 days into a cycle of apparessing, the lexified will usually late for Sustanomic 220 to really begin working consistently, the miss gains are already (generally) vey cotterue.

100

Standard

#### History:

Tectosterous supersion is our of the oldest ambeloi-unforgenic strouk, during all the way back to the 1930's. Used generality to describe any injectable from of the estosterous, testosterous supersion predates the development of slow-acting (deport) injections of esteroidal describerous by sourcell parts. Even alle the development of osterficial dentitions, the second of esteroidal describerous of esteroidal dentitions of esteroidal describerous of esteroidal dentitions of esteroidal describerous describero

Testosterous asspension shares a clinical application history similar to that of other testosterous products. Early prescribing galdines called for its set to armétionate also of sex drive, impotence, and general loss of viality in aging males with declining bearonse levels. Testosterous was is also used to near pubertial adolescents with underscribed testosters. With women, the dang was commonly prescribed for the tentant of occasion or paid lactation following childright as was as insperable manning cancer. By the 199%, however, the FDA had refined the approved uses for testosterous asspension slightly, which began to focus more tighty on the neutrant of mile antiquous insufficiency. How game, you was considered as a secondary through in insperable breast cancer, although its light tendancy to produce withfastion makes at an accommon clinic.

Albugh the nurber of products containing testistentime suspension steady divided over the years, the drug opposituateraptured scalability on the U.S. precipition during mixed and the way to p 1998. That year, the FDA had taken action against Sirsi Laboratoris (a sistalizary of Hany) Schein), which at the time was the principal U.S. supplier for isotatorises products (manufacturing them for their had nal sevent order branch). The firm Some products (manufacturing them for their bland as sevent order branch). The firm Some isotatorise products from of all C-III planmaceriates until certain "disrepuncies" could be addressed. Some interest products for their C-IIII III flamp. The FDA for their courages to suspend production of all C-III planmaceriates and certain "disrepuncies" could be addressed. Versus tates, Steas who the terms translate glosstatories drugs again, but by this time had much the decision not to resum making inclusivement supersiston. Currently, institution when the product of the contraction of the contraction

#### How Supplied:

Testostence saspension is available in select harma and veterinary drug markets. Composition and dosage may vary by country and manufacture, that usually cortain 50 mg/mt of storoid mixed in a water-based solution. Testostence has low water because the control of the control

# Structural Characteristics:

Textudences supersists contains (few) extosterore in a water-based suspension, although other scenariors also sed scenariors. When celestrifactive, textudence has a short life file in the body Textudences supersists may require a minimum of 2-1 is jections per work to minimia moissists. Thermore electricism, When calculating does, expendily when moving from one textudences preparation to another, it is also important to remember that textudences promption contains more active textudences per larger than the sectional derivations. For example, when the weight of the sets is below into account, 100 mg of textudences are called some characteristic and the section of the sets of the sets in bloom into account, 100 mg of textudences are called some characteristic and the set of the sets of

#### Side Effects (Estrogenic):

Textosterore is readily arountized in the body to estable (estrogen). The arountase (estrogen synthetics) europia es texponsible of the intentibotion of textosterone. Detailed estrogen levels can cause side effects such as increased water returnion, body fait gain, and goncomistic. Textosterone is considered an increased water returnion, body fait gain, and goncomistic. Textosteron is considered an increased seven to excessive prevent excited An artis-estrogen such as chemistrated use an arountase irribitor like Ariuntaelot (genationals), whate effects. One may alternately use an arountase irribitor like Ariuntaelot (genationals), whate effects on the oliginal and the second proposal and the second proposal quite expossive in comparison to artis-estrogens, however, and may also have negative effects on belod (piles.)

Entogaie såde effects will occur in a doss-dependant mumer, with higher dosse fabove momal therapeatic levels) of resistoreen more fleely to require the occurrent use of an antiestrogan or aromatuse rizhbor. Siece water reterion and loss of muscle definition are common with higher dosses of festestoreen, this drug's usually, orosidated a poor choice for desiring or cutting faboses, of training. Its renderate entoganicity mules is more ideal for desiring or cutting faboses, of training. Its renderate entoganicity mules is more kital for any contraction of the contraction of the contraction of the contraction of the any contraction of the contraction of the contraction of the contraction of the any contraction of the contraction of the contraction of the contraction of the any contraction of the cont

## Side Effects (Androgenic):

Testosterore is the primary male androgen, responsible for miniming secondary male sessual characteristics. Elevated levels of testosterore are fleely to probe androgenic selectifications, and the proposition of the proposition for hard loss (analyses) and protect produce androgenic selectifications and the produced probability. These concerned about his loss may find a more confertible option in minimizers to decurrently and the produced probability. These concerned about his loss may find a more confertible option in minimizers are summed of the concerned, which is a comparable loss analysisms; asteroid. Werenes are summed of the success of the concerned about his produced probability and the probab

In antiogen-responsive target rissues such as the skin, scale, and prostate, the high relative antiogenizely of testimeterum is dependent on its reduction to disploctationere. (DRI). The S-alpha reductase enzyme is responsible for this methodsom of testistence. The concurrent use of S-alpha reductase inhibits was has finistentive of antienties will interfer with sits—specific potentiation of testistence action, lovering the teachersy of testistence will be a support of the contraction of the contraction of the contraction of the antiogenic efficies as both mediated will the cylciotic antiogene needpot complete separation of testistence's annihole and antiogenic properties is not possible, even with total 5-alpha reductase inhibition.

# Side Effects (Hepatotoxicity):

Testosterone does not have hepatotoxic effects; liver toxicity is unlikely. One study examined the notential for hereatotoxicity with high does of testosterone by administering 400 mg of

the hormone per day (2,800 mg per week) to a group of mule subjects. The steroid was taken orally so that higher peak concentrations would be reached in hepatic issues compared to intermactual rejections. The hormone was given daily for 20 days, and produced no significant changes in liver erozme values including serum albumin, bilirubin, abmine-amino-materiese, and lasline hopophatuses. 450

#### Side Effects (Cardiovascular):

Ambolicarinoganis stronds can have deleterious effects on serum cholesterol. This irelates a reindency to reduce IEEE (good) cholesterov olases and irreases IEEE, (doad) cholesterol values, which may shift the HDL to LDL chalmee in a direction that flores geneter risk of antiroicherosis. The retthen irregat of an ambolicarinoganic servicion of assemi pitis is dependant on the dose, route of administration (end as, injectable), type of steroid (generalizable to route-amountable), and the old of resistance to heptic metabloom challed anti-angusat activation implies and the old of resistance to heptic metabloom challed challengages activated in my data subseedly affect blood presente an eligiborities, received the pitch of confidence of the challengage activation in my data subseedly affect blood presente an eligiborities.

Testoderone tends to have a much less drammés impact on cardiovascular nis factors than synthesis storoid. This die ha part to is permess to metabolismi by the New yith allows it to have less effect on the hapatic management of choisested. The aromatistica of testosterone to establishing the legisted residence of contraders on several picks. In one study, 250 mg per week of lestosterone steer (remarkase) had a slight hat not stancingly significant effects on HELC declosterone date 12 weeks, but when below with an automatise ribbitor a starng (25%) decrease was seen. Statistic using 400 mg of demonstrated only a 17% decrease in Hills Choisetone, while a 400 mg for reduction neached 21% destructions are started to the contraders of the started production of the contraders of the started production of the contraders of the started production are contradered to the contraders of the started production of the contraders of the started production of the started production

Due to the positive influence of estrogen on serum igids, turnostin citate or cloniphere centra are preferred to normatuse ribbinos for those concerned with cardiovascelar health, as they offer a partial estrogenic effect in the foot. This allows them to potentially improve high profiles and offices some of the negative effects of anti-opens. With doses of 600 mg or less of testosterone per week, the impact on judy profile tends to be notectable but not domain; making an artie-strongen (for enderpotentive purposes) perhaps unrecessary. Doses of 600 mg or less per week have also failed to produce statistically significant changes of the contractive contractiv

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular securies program and minimize the intake of saturated fast, colosterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 games per day) and a natural cholesterol/artioxidant formula such as Lipid Stabil or a product with comparable ingedients is also recommended.

# Side Effects (Testosterone Suppression):

All anabolic/indogenic seconds when taken in doses sufficient to promote muscle gain are expected to suppress endogenus testosteore production. Testosteores is the primary unianabogn, and offers strong registric feedback on endogenous testosteores production. Testosteores-besed drugs will, Revisie, here a strong effect on the probeduline regulation of natural seroid hormous. Without the intervention of testosteores-stimating substances, testosteores besed sould return to commit with 1-4 months of dang secsosis Note that prolonged physiogenatoric price programmes and develop secondary to steroid abuse, mescashing medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

Testosterore supernion cortains unfusiohed testosterore particles, which from a shart-acting eposlosy in the much following epision. Depending on the size of the particles and other agests present, episcémes of festosterore suspension may usual particles that require a nordie as large as 21 gauge for injection, the countyle, and can be very uscondition bits out. Modern Institutions supersions presentises minde for human use often contain and indices. Modernic supersions presentises minde for human use often contain and an extra contain the contained of the cont

#### Administration (Men):

To text andogan isufficiency, the prescribing guidens for textourcore supersion recommend alone of 255-50 mg, which is given 2-3 times per work. When med for muscle-balling proposes, textourcore supersions to other administered at a dose of 100-200 mg per spicion, which is given every 2nd or 104 day. Additest keeking a subsetue an enterwise per personal proposes and proposes of the contract of t

Those locking for only a potest muss agent are often extremely largey with the results provided by testostories supersion for large product certainly has a storage prestation for performing. But those subtletes who want not just quantly but quality are ladely to be formed to be made ones. In fact, the new design point of a large large to be a land, dones on. In fact, the next proposing with the day, and will often sock the benefit of carting agents soon afterwards to clean up the lock of irresulting Alexandre of the control o

#### Administration (Women):

Testosterous asspension is mely used with women in clinical medicine. When applied, it is most often used as a secondary treatment for inspensible beset camer. Does spire for this application may reach 100 mg free times per week, a level well into the threshold likely to cause strong vilidings dise effects. Testosterous suspension is not recommended for women for plosque-or performance-orbancing purposes due to its strong androgenic mature and tendency to produce vilidings side effects. Testosterous suspension and tendency to produce vilidings side effects.

# Availability:

Pharmaceutical preparations containing testosterone in an aqueous suspension remain scarce. The bulk of the supply presently comes from underground and export steroid manufacturers. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Testosterone suspension has been unavailable in the United States for many years now. No old products should still be in circulation. Because the FDA never officially withdrew this drug from, it can be specially ordered through a small number of compounding pharmacies. Anything else bearing a U.S. manufacturer rame is counterfied.

Testosterone suspension is still produced in India under the Aquaviron brand name, now by Piramal HC. It comes in the strength of 25 mg/mL, and is packaged in 1 mL ampules.

# Cheque Drops® (mibolerone)

	, , , , , , , , , , , , , , , , , , , ,
	dimethyl-19-norandrost-4-en-3-one-17b-o
11.7	lydroxy-7alpha,17-dimethylestr-4-en-3-one
Estrogenic Activity	high
Progestational Activi	hia!

# Description:

Androgenic

Anabolic

Standard

Medicence is an oral anaboli seroid, structurally deviced directlytated methodore. This agart is specifically 7.1.7.deringheted methodore, significantly more potent as an anabolic and antiogens: agart than its non-methylated parent. Over the years, mbolsence has carried as a regulation arrange hopkhadiers as being one of the strengest arranged over made. This is effective as the strength of the

1.800

4 100

Methyltestosterone (oral)

# History:

Moderne was first described in 1963-400. It was developed into a veteriory medicine during the 1990-51 by join, which sold the only under the hort name. Cheep Doeps, the original constrained 100 megind of stend in a 55 ml. bottle, for a total stend context of 55 milligrams (Bartisting the high relatine protectory of molerouse). Planmaria & Upinharmia & U

Ameng athless, mbedevene has always been seen with a high level of mapsings, perhaps purely due to its intrival availability. Those cataly famility with the Lipsin their Phramics & Lipsinia product user likely disappointed during the early 2000's, when the Chegae Health presently his or mobilevine containing products on its investory, design extings the rights to market the drug. Melsoleume is still available in the U.S., but only in generic form from a private compounting pharmacy, closical under special corder by a Remedia ventratum. The neuronal of the Plannacia & Lipsinia products as the respective form for U.S. maket ventrature. The neuronal of the Plannacia & Lipsinia products are simportant. The neuronal of the Plannacia is a Lipsinia products show the C.S. maket ventrature, and the contract of the Plannacia of Lipsinia products from the U.S. maket ventrature, and the contraction of the

# How Supplied:

Mibolerone is no longer available as a prescription drug product. When produced it most commonly came in the form of an oral solution based in propylene glycol, carrying 100meg of steroid per milliter in a 55 ml. bottle.

#### Structural Characteristics:

Mbolerone is a modified form of randrolone. It differs by 1) the addition of a methyl group at earlown 17-alpha to protect the hormone during cetal administration and 2) the introduction of a methyl group at earlown 7 (alpha), which inhibbs 2-alpha reduction and increases relative antrogenicity 7,17-dimethylated steroids also tend to be very resistant to metabolism and serum bridger notices; search extracting their relative biological activity.

# Side Effects (Estrogenic):

Miholerone is aromatized by the body and is considered a highly estrogenic steroid due to its conservion to 7.17-dimethylastradial (an astronon with high high-pixel activity) Gynecomestia may be a concern during treatment, especially when higher than normal thoranoutic docor are used. At the same time water retention can become a number, causing a notable loss of muscle definition as both subcutaneous water retention and fat levels build. To avoid strong estrogenic side effects, it may be necessary to use an anti-estrogen such as Nolvadex®. One may alternately use an aromatase inhibitor like Arimidex® (anastrozole). which is a more effective remedy for estrogen control. Aromatase inhibitors, however, can be quite expensive in comparison to standard estrogen maintenance therapies, and may also have negative effects on blood linids

It is of note that mibolerone also displays strong activity as a progestin in the body. The side effects associated with procesterone are similar to those of estrogen including negative feedback inhibition of testosterone production and enhanced rate of fat storage. Progestins also augment the stimulatory effect of estrogens on mammary tissue growth. There annears to be a strong synergy between these two hormones here, such that gynecomestia might even occur with the help of procestins without excessive estrogen levels being present. The use of an anti-estrogen, which inhibits the estrogenic component of this disorder, is often sufficient to mitirate ovnecomestia caused by mibolerone

# Side Effects (Androgenic):

Although classified as an anabolic steroid, androgenic side effects are still common with this substance. This may include houts of oily skin, acre, and hody/facial hair orowth. Anabolic/androgenic steroids may also aggravate male pattern hair loss. Individuals sensitive to the androgenic effects of this steroid may find a milder anabolic such as Deca-Duraholiniic to be more comfortable. Women are additionally warned of the potential virilizing effects of anabolic/androgenic steroids. These may include a deepening of the voice, menstrual irregularities, changes in skin texture, facial hair growth, and clitoral enlargement. Note that 7-methylation inhibits steroid 5-alpha reduction. 431 The relative androgenicity of mibolerone is not affected by the concurrent use of finasteride or dutasteride.

#### Side Effects (Hepatotoxicity):

Mibolerone is a c17-alpha alkylated compound. This alteration protects the drug from deactivation by the liver, allowing a very high percentage of the drug entry into the bloodstream following oral administration. C17-alpha alkylated anabolic/androgenic steroids can be hepatotoxic. Prolonged or high exposure may result in liver damage. In rare instances life-threatening dysfunction may develop. It is advisable to visit a physician periodically during each cycle to monitor liver function and overall health. Intake of c17-alpha alkylated steroids is commonly limited to 6-8 weeks, in an effort to avoid escalating liver strain. Severe liver complications are rare given the periodic nature in which most people use oral anabolic/androgenic steroids, although cannot be excluded with this steroid, especially with high doses and/or prolonged administration periods. Note that U.S. prescribing information for Cheque Drops mentions only one human study being conducted on mibolerone, and that the study was terminated early due to high henatotoxicity.

The use of a liver detoxification supplement such as Liver Stabil. Liv-52, or Essentiale Forte. is advised while taking any hepatotoxic anabolic/androgenic steroids. Anabolic/androgenic steroids can have deleterious effects on serum cholesterol. This includes

#### Side Effects (Cardiovascular):

a tendency to reduce HDL (good) cholesterol values and increase LDL (bad) cholesterol. values, which may shift the HDL to LDL balance in a direction that favors greater risk of arteriosclerosis. The relative impact of an anabolic/androgenic steroid on serum lipids is dependant on the dose, route of administration (oral vs. injectable), type of steroid (aromatizable or non-aromatizable), and level of resistance to henatic metabolism. Mibolerone has a strong effect on the hepatic management of cholesterol due to its structural resistance to liver breakdown and route of administration. Anabolic/androgenic steroids may also adversely affect blood pressure and triglycerides, reduce endothelial relaxation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disass and necessarial infection

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fist, cholesterol, and simplecardiohydrates at all times during active AAS administration. Supplementing with fish oils (4 games per day) and a natural cholesterolamioxidant formals such as Lipid Stabil or a module with commands hierafeties it also recommended.

# Side Effects (Testostemne Summession):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Whost the intervention of testosterone-standing substances, testosterone levels should return to normal whitn 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradism can develor secondary to steroid abuse, necessistant medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

Studies have shown that taking an oral anabolic steroid with fixed may decrease its biovasiability. <sup>822</sup> This is custed by the fat-soluble nature of steroid hormones, which can allow some of the drug to dissolve with urdigested delary fat reducing its absorption from the gestroitestiral tract. For maximum utilization, this steroid should be taken on an empty stormeth.

# Administration (Men):

Medicence was never approved for use in human. Pencerbing guidelines are unwalable, in the athletic areas, the drug is used intermittedly due to he light level of hepatotoxicity, with cycles usually lasting no more than 6 weeks followed by 6-8 weeks off. A daily dosage of 200 to 500mgs in sort common for body-bading purposes. This level is typically sufficient for gains in strength and mache mass (balk). The high prospectational and estiogenic acidity of rabidotome rathests of fifst what in people and enhancer opens, causing an unswared of rabidotome rathests of fifst what in people and enhancer opens, causing an unswared

#### Administration (Women):

Mibolerone was never approved for use in humans. Prescribing guidelines are unavailable. Mibolerone is generally not recommended for women for physique- or performanceerhancing purposes due to its very strong nature and tendency to produce virilizing sideofficeto.

#### Availability:

Androgenic

Mibolerone is sold in the U.S. as a compounded veterinary medicine only. No commercial preparations containing this drug are known to exist worldwide. Mibolerone remains available on the black market in underground preparations only.

#### Deca-Duraholin® (nandrolone decanoate)

Anabolic	125
Standard	Testosterone
Chemical Names	19-norandrost-4-en-3-one-17beta-ol 17beta-hydroxy-estr-4-en-3-one
Estrogenic Activit	y low
Progestational Ac	tivity moderate

\_\_\_\_\_

#### Description:

Numbroken decurated is an nijectable form of the anabole stroof murdroken. The decurated setal provides a slow release of murdroken from the set of spicetin, Issing for up to fire weeks. Numbroken is very similar to estosterome in structure, although it lacks a carbon attent at the "Holl position (hence is other name, 10 - norticaterone). Like according to the control of the control of the control of the control of the however, its fisuse bailing activity is accompanied by week androgen; properties. Much of the last to do with the relation of methodors to a wakent stread, dipplementabless, in the same androgen-responsive turner tissues that potentiate the action of testosterous (by concerting to DITI). The milt properties of methodorse decembes then much de one of the concerting to DITI). The milt properties of methodorse decembes then much de one of the procured as the control of the control

# History:

Nandolone decunste was first described in 1906, 433 and became a psecieption medication in 1902. It was developed by the international parameteriska girth Organo, and sold under the brand name Deca-Darbolis. The name Deca-Darbolis devents that the product contains a surfair of Organon's precisolar popular medicione injectable Darbolis (unarbolison phraphyropinate) usig an ester of 10 carbon atom. Organon equaled the medic for martidone document very registly following in relaces. Probably owing to a Darbolis in the Carbon one of the most which distributed analysis extensi in the world.

When fist irreduced to the United States, murledone decunouse (like Databolin) was proceived for a swiry of animers. Listed intractions related upon and postsporenties use for building lam muss, ostoporouss, advanced breast cancer, weight loss that the consideration of the consider

Albusgh the drug had been applied fivonshly for a great many medical uses for approximately a deceled, by the mist 1970's he indicated uses for mandelsone decunsed were being refund, both in the U.S. and abroad. FINA approxed prescriping information from 1975 fists mandelsone decunste as "probably effective" as adjuret theory is reside and postmerapassal obstoperosis, as well as for trusting patienty-deficient downtim suff execution of the marse, controlling abstracted beast cancer, and as adjunctive flerapy for certain types of ameria. More time was given to insessigate the potential "less than effective" uses of the drug.

Modern (spectived) medical applications for the data are corn more refused than they were in the min 1970; that that States, the data give more alph Naperrood for tending ameries, albeight is often also used 'noff their' to preserve lean mass in HIV position profites and others saffing from swaring alesses. Onside of the ICs, Opparen secure to support the use of this drug mainly with patients suffering from severe ameries, notesporous, and advanced breast cancer. The Oppinon Descen Describe Intelle. It formations decreases remains which was made to the original profit of the profit of the original profit of the profit of the original p

# How Supplied:

Nandrolone decamoate is widely available in human and veterimized drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 25 marks 150 metril 100 metril 100 metril of 200 metril of steroid dissolved in oil.

# Structural Characteristics:

Nandrolone decanoste is a modified form of nandrolone, where a carbovulic acid exter-(decanoic acid) has been attached to the 17-beta hydroxyl group. Esterified steroids are less nolar than free steroids, and are absorbed more slowly from the area of injection. Once in the bloodstream, the ester is removed to yield free (active) randrolone. Esterified steroids are designed to prolong the window of therapeutic effect following administration, allowing for a less frequent injection schedule compared to injections of free (unesterified) steroid. Nandrolone decanoste provides a sharp spike in pandrolone pelease 24-48 hours following deep intramuscular injection, which steadily declines to near baseline levels approximately two weeks later. The half-life of nandrolone decanoate is 7-12 days.

Pharmacokinetics of Nandrolene



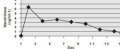


Figure 1. Pharmacokinetics of 200 mg Nandrolone Decanoate injection. Source: Pharmacokinetic parameters of pandrolone (19-nortestosterone) after intramuscular administration of nandmione decanoate (Deca-Durabolin®) to healthy volunteers. Winand H. Bosch A. Donker C. Acta Endocrinol 1985 supp 271 19-30.

#### Side Effects (Estrogenic):

Nandrolone has a low tendency for estrogen conversion, estimated to be only about 20% of that seen with testosterone.434 This is because while the liver can convert randrolone to estradiol in other more active sites of steroid aromatization such as adinose tissue nandrolone is far less open to this process. 435 Consequently, estrogen-related side effects are a much lower concern with this drug than with testosterone. Elevated estrogen levels may still be noticed with higher dosing, however, and may cause side effects such as increased water retention, body fat gain, and gynecomestia. An anti-estrogen such as clomiohene citrate or tamoxifen citrate may be necessary to prevent estrogenic side effects if they occur. One may alternately use an aromatase inhibitor like Arimidex® (anastrozole), which more efficiently controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

It is of note that nandrolone has some activity as a progestin in the body. 436 Although progesterone is a c-19 steroid, removal of this group as in 19-norprogesterone creates a hormone with greater binding affinity for its corresponding receptor. Sharing this trait, many 19-nor anabolic steroids are shown to have some affinity for the progesterone receptor as well.437 The side effects associated with progesterone are similar to those of estrogen, including negative feedback inhibition of testosterone production and enhanced rate of fat storage. Procestins also augment the stimulatory effect of estrogens on mammary tissue growth. There appears to be a strong synergy between these two hormones here, such that gynecomastia might even occur with the help of progestins, without excessive estrogen levels. The use of an anti-estrogen, which inhibits the estrogenic component of this disorder, is often sufficient to mitigate gynecomastia caused by nandrolone.

#### Side Effects (Androgenic):

Although classified as an anabolic steroid, androgenic side effects are still possible with this substance especially with higher doses. This may include hours of oily skin acree and body/facial hair growth. Anabolic/androgenic steroids may also aggravate male pattern hair loss. Women are warned of the potential virilizing effects of anabolic/androgenic steroids. These may include a deepening of the voice, menstrual irregularities, changes in skin texture, head lar growth, and ciforal entirgement. Nandrobne is a stroof with relatively two androganic activity relative to its steal-building actives, miding the threshold for strong androganic side effects comparably higher than with more androganic agents such as testesterous, methandrosterobne, or oftoxymesterone. It is also important to point out that due to is mild androganic mature and ability to suppress endogenous testosterous, methodous jew note to interfering with bildo in miles when used without another androgan.

Note that is undrogon-responsive target fissess such as the skin, scalp, and prostate, the relative antilongenity of maniforoise is reduced by its reduction to dightyoriumflootine (DRN)<sup>10,100</sup> The S-alpha reductase enzyme is responsible for this metaloisium of maniforoise. The concerned use of a S-alpha reductase infliently such as financial ordinational control of the state of

## Side Effects (Henatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects in healthy subjects. Liver toxicity is unlikely.

#### Side Effects (Cardiovascular):

Anabolizándroganie strendis can hove deletroixe effects on serum cholesterel. This irelades a redenely to refuele IFML (good) cholesterel values and irecares LDL (bad) cholesterel values with my shift the IFML to LDL balance in a direction that fivors geneter risk of arteriochesis. The redules impact of an ambibicandroganic strend on serum lipids is dependant on the dose, route of administration (end us. nijectable), type of steroid (contrable) en accommitable), and level of resistance to lespain intelloiden Stadies administrating 600 mg of mandrodes decumulate per work for 10 weeds demonstrated a 20% reaction in LTDL challesterel locks. <sup>407</sup> liber appreciation skipfly gentare from that responde to the contraction in LTDL challesterel locks. <sup>407</sup> liber appreciation is applied by strenger regulate impact on HTDL LDL risk with mandrodes decumulate a skipfly weaker impact on serum lipids than c-17 alpha alcylated agents. Arabolizathorganic strends my and support left workreizer land regulates, reduce endothelial relevation, and support left workreizer land propertially invensing the kind of confinement and support left workreizer land propertially invensing the kind of confinement and support left workreizer land processing infection.

To help reduce cardiousecular strain it is advised to maintain an active cardiousecular searcise program and minimize the intake of saturated fits, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterol/artioxidant formula such as Lipid Stabil or a moduct with comrabble inserdence is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolic intelligents: seroids when taken in does sufficient to promote muscle gain are expected to suppress endopross to solotome production. Studies achimistrate gilo magnet wock of mandotne decamate for 6 weeks live demonstrated an approxime 57% reduction in semuntationerous leeks direg framps, 74 at dosage 5700 mgp per wock in reduction nesched 70%, 620 is believed that the prospentiatival activity of mandotone muscled or appression of instruction of the supersion of t

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

# Administration (Men):

For general anabolic effects, early prescribing guidelines recommend a dosage of 50-100 mg every 3-4 weeks for 12 weeks. To treat renal anemia, the prescribing guidelines for

mediotine documente recomment a dosage of 100-200 mg per wock. The usual dosage for physique- or performance-ordunicing purposes is the mage of 200-600 mg per wock, taken in cycles 8 to 12 wocks in length. This level is sufficient for most users to notice measurable gains in hom musche mass and strength it is often stead that mendolines deceasates will exhibit its optimal effect (test gainside effect ratio) at 2 mg per pound of bodyweight words, based pain inhalted afforences in response will despot doction verying failed by the contraction of the contractio

Nandrolous decurates is often contrined with other steroids for an entraced effect. A combination of 200-200 milyweds or mandrolous decurates and 10-20 mg daily of Wastellik, for example, is neder to greatly enhance the look of muscalarity and definition when distingentings. A morage man arounding androgal the Ethicisterili or trestorbies could also be used, again proxiding an enhanced level of Instructures and density to the muscles. Being a mortarely storeign muscle ladder, mardrolous can also be recoperated for blad cycles with acceptable results. The classic "Dece and D-bol" state (usually 200-400 mg of rantaclouse decamate per work and 15-25 mg of Distructly per day) has been a bodyloiding basic for decamber, and always seems to provide excelent muscle growth. A securate results, but with more water restricts.

### Administration (Women):

every 3-4 weeks for 12 weeks. To treat result aerenis, the prescribing sidelines for maniforate decanter recomment a desage of 5-010 mg per week. When used for physique or performance-enhancing purposes, a dosage of 50 mg per week is most commen, taken for 4-6 weeks. Although only sidely and angesie, women are consciously conflorated with virilization symptoms when taking the compound. Statless have demonstrated high behindly ingrine but statistically assignificant relaces of vibilizing statletics and so does as low as 50 mg every 3-5 weeks. <sup>465</sup> Should virilizing side efficts on shoot as low as 50 mg every 3-5 weeks. <sup>465</sup> Should virilizing side efficts hoome a concern, mantholize decanted assistant be discontrained immediately to help beyout their permisser appearance. After a sufficient period of withdrawal, the shorter-acting mandolize Darabolizin might be considered as slare forms. The due gains year the forely sevent depressments, greatery described periods of positions of the considered as slare forms. The due gains year the forely sevent depressment generate. After a considered as slare forms. The due gains year the forely sevent depth generally greatly reducing the considered as slare forms.

For general ampholic effects, early prescribing guidelines recommend a dosage of 50-100 mg

# Availability:

Nandrokee decureate cortinus to decline in prominence as a phurmaconical product do to its initiate use in cinical medicare. The drug is presently unswalible in the United States, Many Western matrices confine to market the drug, though its production is increasing being shifted to less regulated markets in Asia. Legitarine phurmaconical forms are highly sought after on the black market, and thus subject to a great deal of contractivities in the market when the medical threshold in the contractivity of the contractivity of the market we have medic the Solvier observations.

In November 2009, Organon (a subsidiary of ScheringPlough since 2007) became part of Merck/MSD. All Organon products are expected to transition over to this label. It is urknown what (if arry) changes to expect in the global distribution of Deca-Darabolin modulers.

Brand name Deca-Durabolin is not available in the United States. All products bearing this label are counterfeit. Watson Labs and Schein Pharmaceuticals generics have also been discontinued. This drue is presently unavailable in the U.S.

Norma Hellas Deca (100 mg/ml. nandrolone decanoate in 2 ml. vials) from Greece is available, but also widely counterfeited. The firmuses a patented photochromic label to deter counterfeiting, which carries a metallic/holographic watermark of the Norma Hellas logo.

Greek Deca-Durabolin (formerly from Organon) has been another widely counterfeited product. It is one of only a handful of European randrolone rijectables to be found in multidosed visit medicine it an extraort for counterfeiters that lack the canced vito modiuse obesampules. This product should be considered fake unless it comes in a box with the proper Greek drug ID sticker. As with all Greek drugs, the sticker should show a hidden mark under UV light.

Greek Extraboline may be in circulation. It is also a common target of counterfeiting. As with all Greek drugs, this product should contain a peel-off pharmacy sticker that reveals a hidden watermark under UV lighting. All Extraboline in circulation will also carry a holographic image directly on the vial label.

Deca-Promabol from P&B Labs India is no longer in production. The company currently markets only a 25 mg/ml. version of this drug in 1 ml. ampules. Many counterfeits of the former 100 mg/ml. product are still in circulation.

Decabolic from Asia Pharma (Malaysia) is now approved for sale through pharmacies in Trailand, and is fairly popular on the black market. Each box should carry a seranti-off security sticker, which will display a code that can be validated on the company website.

Balkan Pharmaceuticals (Moldova) makes the product Nandrolona D. It is prepared in both 1 mL ampules and multi-dose vials.

## Delatestryl® (testosterone enanthate)

Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate

# Progestational Activity Description:

A-drogonia

Testoatrone cumfutue is a slow-acting ijectable form of the androgen testoatrone. Following deep intranaction rigitor, the found gas designed to provide a sentiand release of testosterous into the bloodsteam for approximately 2 to 3 weeks. In order to minimis normal physiological bless of testosterous free approximately as the sentiant programment frapensis, incision of testosterous enuralitae are usually required at least every two weeks, although more mediculous physicisms will administer the day weekly. As with all disasterous rigicalities, testosterous cumfutue is highly fromced by adhetes for its ability to promote strong increases in muscle mass sufficiently and the strong from the strong increases in muscle mass and storegath.

#### History:

Testosteroe curaflute first appeared on Western dang murkest during the early 1950's. It was the first show-calling of sheared spiceties seer of sestosteroes to be widely adapted in Western medicare, and effectively replaced testosteroes propriate and testosteroes between the contractive propriate and testosteroes of the contractive o

Testosterone enurfate is mos often used clinically to replace normal levels of testosterone in adult males saffering diminished andropen levels. This may marifest self-with a loss of blido, learn mascle mass, and normal energy and vigor. Festosterone enurfate is also used to treat undescended testiess and debuged updetry in adolescent males, and occasionally as a secondary medication during inoperable breast cancer in women. This form of testosterone

has also been studed will great success as a misk brit control option." Weekly spectrose of 200 mg were shown to efficiently hower sperm production for most men within three moreths of teatment, a state of suppression that remined until after the drug was decontrained. With the current signs maximaling antibolicaminogenis setroids, however, if is trafficly that such therapy will become adopted it Mestern medical practice. Today, it is trafficely that such therapy will become adopted in Mestern medical practice. Today is sufficiently only the success of th

# How Supplied:

Testosterone enauthate is widely available in human and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 50 ma/nt. 100 ma/nt. 200 ma/nt. 07 250 ma/nt of steroid dissolved in oil.

#### Structural Characteristics:

Testistence currilate is a modified form of testistence, where a carboxyle acid exter (emratio acid) has been attacked to be T-been lydrody agon. Estrated forms of testistence are less polar than fee testistence, and are absorbed more slowly from the area of nijection. Once in the bloodstream, the exter is removed to yield free facility testistence. Estrated forms of testistence are designed to prolong the window of themports: effect flowing administration, adwing for a law foreger nijection schedule compared to injections of five (insentified) servid. The half-life of testistence currilate is approximately exist and ser injection.

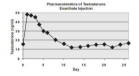


Figure 1. Pharmacokinetics of 194mg testosterone enanthate injection. Source: Comparison of testosterone, dillydrotestosterone, luteinizing hormone, and follicistimulating hormone in serum after injection of testosterone enanthate or testosterone explorate. Schulte-Beerchall M, Nieschlag E. Fertility and Sterility 33 (1980);2013. Side Effects (Estrogenic):

Telestotence is readily anomized in the body to establed (entogen). The anomatuse (entogen synthetics) output is responsible of their introllation of telestotence. Electual estages heek can cause side effects such as increased water reteriors, body fat gain, and goncomistic. Telestotence is considered anomatusely estergias restored. An artis-entogen such as chemiphene clause or turnostic entate may be necessary to prevent strongers and the complete of the control of the contro

Estogais: side effects will occur in a doss-dependant muner, with ligher doss faltows muntil functional feeds of restoration men files byte on eight one content on of an anti-estogan or aromatuse shiftent Since water reterion and loss of muscle definition are common with light decos of insustence, this drug is usually considered a poor cheice for common with light decos of insustence, this drug is usually considered as poor cheice for the common with the contraction of the contractio

#### ORGERIECO CARRIOSCINCA

Testosteores is the primary mule androgen, responsible for mistraining secondary mulsearul enteractistics. Entered the evid forestornee are Body to probe androgenic side effects including only sixt, acres, and hodyficial har igwords. Men with a genetic prodogosition for his loss (androgenic alsopera) may notice accelerated multiparter hading. Those concerned about har loss may find a more conferrable option in municional decenture, which is a comparably loss androgenic sectoral. Wemen are swemed of the decenture, which is a comparably loss androgenic sectoral. Wemen are swemed of the content of the conference of the conference of the conference of the conference sound to the conference of the conference of the conference of the conference content of the conference of the

In autogon-responsée traget tissus such as the skis, soals, and prostite, the high relative autogravity of lesstrouves is dependant on in reduction to diplotostasteure (DRI). The 5-sphe reductase ensyme is responsible for this methodom of lesstasteure. The concurrent use of 3-sphin reduction inhibits such as finistrative of attentive of interfere with site-specific potentiation of lesstasteure action, lovering the teachery of lesstasteure drugs to produce analogous six defection. It is important to remember that analoles and autogravit effects are both mediated via the cytosotic ambigus recopers. Complete control of the control of the control of the cytosotic ambigus recopers for a production of the control of the control of the cytosotic ambigus recopers is not possible, core with

#### Side Effects (Hepatotoxicity):

Estostorour does nel live lepatotoxic effects, her toxicy is unfiely. One study examine the potential for lepatotoxicy with high does of restoratory by administering 400 mg of the hormon per day (2,800 mg per week) to a group of mile subjects. The steroid was taken only so that higher peak concentrations would be reached in lepatic insues compared to intramensality rejection. The hormone was given duly for 20 days, and the configuration of the co

# Side Effects (Cardiovascular):

Anabolisindegasis streaks on how obstation effects on esture cholesters. This includes a turkency to refeet PIII. (good) eloboticary obses and invene learn flavor plate of particular states, which may shift the IDL to LDL balance is a direction that flowes greater risk of anteriosciences. The relative terpart of an anabolicantegasis entodo on serum lapids is dependant on the dose, route of administration (real two, injectable), type of sterood (commissable or non-anomatable), and level of resistance to plantic matsolosim. Anabolicantegasis streaks may also adversely effect blood pressure and trighceriae, reduce, established inclusions, and apport off vermically perspectlys, all potentially

Testoderore tends to have a much less dramatic impact on confloroscular risk fectors than synthesis storods. This does input to specimes to metablishminy be feet, which allows it to have less effect on the hepatic management of cholesterol. The aromatism of testoscene to searched also hepks to might feet negative effects of arthrogon on semi-pick. In one study, 200 mg per week of testoscene to establish controlled specified and to the controlled specified and the period of the controlled specified and th

Due to the positive influence of estrogen on serum lipids, tamoxifen citrate or clomiphene

cintae are preferred to arcumtase simbions for those concerned with audiouscular health, as a she sport are partial estrogenic effect in the Nex This allows them to potentially propose lipid profiles and officts some of the negative effects of antogens. With doses of 600 mg or less per word, the injury can for july froll the softs to be noticeable but not drauntic, mixing an auti-strongen (for cardioprotective purposes) perhaps unrecessary. Doses of 600 mg or less per work have also failed to produce stratistically agrifacter at lengals in LDV/LDL chelstent, triglecerides, apolipoportis IBC-III, C-receive proteix, and main sensitivity, all relativists are alterful weak instruction.

moderate doses, injectable testosterone esters are usually considered to be the safest of all anabotic/androgenic steroids.

To hely reduce cardiovascular strain à is advised to maintain an active cardiovascular seurcise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a matural cholesterollaritoxidant formals such as Lipid Stabil or a product with comparable ingedients is also recommended.

#### Side Effects (Testosterone Suppression):

All antibolization/goise storoids when taken in doses sufficient to promote muscle gain are expected to suppose condegorous testostorous production. Testostoroes is the primary undaantiques, and offers strong negative footback on encloparus testostorous production. Flostostorous-book diago will. Revisies, hurse a strong effect on the phytolinaire regulation of natural steroid hornous. Without the intervention of testostorous-strutating substances, testostorous-book bound tento to normal with 1-4 mentils of office accessive. Nor their prolonged physgonatotropic in programation can develop secondary to steroid abuse, revessibilities medial intervention.

As with all ambeleindrogenic steroids, it is utilisely that one will retain every pound of new bodyweight after a cycle is concluded. This is a specially two whose witheriness from strang (arrounting) analogous like testiscence, as much of the new weight gain is likely to be in the form of water retention quickly eliminated after dig decommissive. An irrelation of analosis and catabolic hormouse during the post-cycle recovery period may further create an environment that is withouther for the retention of much testisce. Proper arrillarly and therapy is usually recommended to help restore hormonal balance more quickly, ultimately behigher less retain more much elissies.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

# Administration (Men):

To treat androgen issufficiency, the prescribing guidelines for testoderone countriate call for a dosage of 59–00 pm every 2 to 4 weeks. Although action in the body for a longer time, testostenore countriate is usually injected on a weekly basis for much-building purposes to testostenore countriate is usually injected on a weekly basis for much-building purposes. The treat god 200–000 per purposes is in the range of 200–000 per week, taken in cycles 6 to 12 weeks in length. This level is sufficient for most users to notice occurronal ensists in markets size and streated.

Textosterone is seasily incorporated into baking plases of training, when added water reterion will be of little consequence, the user more concerned with now must have definition. Some do incorporate the drug into cutting cycles as well, but typically in lower does (100-200 ngpr weeks) and/or when accompanied by an arountess eithbiro to become temporate to the control control textosterone centralute is a very effective analysis (ong, and is often used above with quest beards. Some, however, find a new darkform 200-400 mg per week of holdeners undexplexine, refundenche enterthing is remarkfore decentrated should provide adstantial results with no significant hepstonicity. Textosterone is ultimately very versatile, and can be contributed with many often analytical regularies strendits to that

#### Administration (Women):

Testosterone cuarditate is rarely used with women in clinical medicine. When applied, it is notest often used as a secondary medication during inportable breast cancer, when other therapies have failed to produce a desirable effect and suppression of overim function is necessary. Festosterone emarkate is not recommended for women for physique- or performance-ordancing purposes due to its strong anthogonic nature, tendency to produce vidings side effects, and slow-actine parameteristics funching bodd levels difficult to control).

#### Availability:

Tectnetowns oranthate remains the most widely manufactured from of injectable tectnetowns

remoteror examine remains or into ways intraduction and or interesting some of the products in the global planmaceutical market, we have made the following observations.

In 2006, Savient sold the rights for Delatestryl to Indevus (United States). Indevus subsequently became a subsidiary of Endo Pharmaceuticals in March of 2009. Brand name Delatestryl remains available in the United States under the new company (at 200 mg/mL strength).

Generic versions are also available in the United States (also at a dosage of 200 mg/mL) by Watson, Paddock, and Synerex.

Norma Hellas (Greece), makers of Norma Hellas Nandrolone, recently added a generic 250 mg/ml. testosterone errarflate injectable to their product offerings. It comes in a single dark arrher 1 ml. glass ampule, and is packaged 1 ampule per box. Be sure to look at the Greek Pharmacy sticker under UV light to assure you have a legitimate product.

Bayer nock control of Schering AG in December 2006. Following this acquisition, the Schering Printoteston and Testoviron Depot products were transitioned over to the Bayer brand and lago (the products now bear the fill company name Bayer Schering Planna). Note that many counterfeiters have not yet made this change in their own products, and thas are still deplicating the old Schering labels and boxes.

Cidoteston is produced in Egopt by CID (Chemical Industries Development). It comes in 1 mL ampules, containing 250 mg/mL of steroid. This product has been counterfeited, though the current most popular copy can be quickly identified by close examination of the fine details (loop, agraphics) on the box.

The French version Testosterone Heptylate is still in production. It is now sold under the S.E.R.P. label, and available in the familiar 250 mg/ml. strength and 1 ml. ampule. This drug is frequently exported to developing markets with close trade relations to France, such as

Androtardyl is also produced in France, and occasionally circulates on the black market.

Again, be sure to look for the proper box before buying.

Testo-Enant is another brand in Europe, this one being made by Geymonat in Italy. These ampules contain 250 mg of steroid, either in 1 mL or 2 mL of oil. Currently fakes are not a problem; however, this steroid is not found on the black market in high volumes.

Galenika makes Testosteron Depo in Serbia. These 1 mL ampules contain 250 mg/mL of steroid, and are extremely chean at the retail level in their country of origin.

Jelfa produces Testosterorum Prolonganum in Pokind. It is made at a dosage of 100 mg/mL. Each box contains five 1 mL ampules, which are themselves made of clear glass and carry a paper label The packaging of this product was recently updated to reflect a more modern color-randerin design.

Testoviron Depot from German Remedies in India remains in production. The product comes is made in foil and plastic blister packs.

The Indian export firm Alpha-Pharma also makes a testosterone enauthate, called Testobolin. It comes in 1 mL glass ampules.

Testofort Inj from Albert Davis Pakistan is commonly found on the international market. It contains 250 mg/ml of steroid in 1 mL ampules. Three ampules come packaged to each cardhourd hox

Geofman Pharmaceuticals also makes a generic in Pakistan The product contains 250 mg of steroid in each 1 mL ampule. Like Testofort, three ampules are contained in each box. Note that the bit number and expiration date are electronically printed on the bottom inside flap of the box; in addition to the proper placement on the coaside.

Aburaihan makes a generic enanthate in Iran, which is becoming increasingly popular on the

black market. Note that the packaging of this product was recently updated. Counteriers of this product have historically been a problem.

Balkan Pharmaceuticals (Moldova) makes the product Testosterona E. It is prepared in both 1 mL amoules and multi-dose vials.

# Depo®-Testosterone (testosterone cypionate)

Androgenic	100
Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate

# Description:

Progestational Activity

Testosterone explorate is a slow-acting injectable ester of the primary make androgen testosterone. Testosterone is also the principle analysis formers in more, and is the basis of testosteron injectables, testosterone rejectables, testosterone rejectables, testosterone rejectables, testosterone rejectables is highly forough valuelse for its ability to promite strong increases in maxies mass and strength. It is intensiting to most that while a large narriber of other steroidal composation have been rande available sizes testosterone. The results is the sugarant that these are among the most proverial mass drugs available, testosterone explorate included.

low

# History:

Testostorous explorate first appeared on the U.S. drug metter during the mid-1990, under the brand rume of Dope-Testostorous exploraphytopointes found arriving do simply Dope-Testostorous). It was developed by the planmacerizing juir Upjoht, and is still sold to this day by the same company under the same trude name (alloward) now they are called Planmacia & Upjohn). This is a day with inited global residuability, and has betorically been (tagges) beliefficial on a Americant lent in 1 sex supering that American athletes three Operators of the Same of t

Telestotence opionete and testotenne enumfate provide enternelly comparable patterns of restostencer releases. Net only are physical adarraties not rosebotis in one over the other, but actual differences in pharmacokinetic patterns are hard to notice (these two drugs are for al intents and pravose instrainely). Here only to difference between the two seems to be in the use of pointer courfiert. Cyptoria end is less irritating at the site of significant them contained: and (enathratis) for a small procurage of pointers. The institute of the contained of the cont

The min use of 'estosterouc opionets in clinical medicin but Islandically been the treatment of low antiogons been in miss, although many often application in the cestified first ingress as well. Dring the 1960%, for example, the drug's prescribing recommendation called for seast these as supporting boots structure munity; touting mensyring theory mensional bededing and excessive batton in females, and increasing mache times and confusing consequents of the confusion of the con

By the 1970's, the FDA had been gusted much stronger control over the prescription drug mented, and the broad uses in which telestories explorate was first indicated were now being refined. For example, "tesslostence rebound therapy" as a way to increase mule ferriliny was proving to be utreatible, opencing in the face of never more deficienmedications, and was soon eleminated from prescribing guidelines. So too was the recommendation for its use to treat thing like excession ensurated bedding and leation in general, testistories therapy was being public back to faces mainly our mult androgen searchible to androomic side effects, such so woman and the electric androomic side of the such as the second of the such as the surface of the such as the surface of the such as the surface of the s

Tedry, testosterone cypicutes remains readily available on the U.S. prescription ding market, where it is Palv-approved for hormour esplacement therapy in mm with conditions associated with a deficiency of endogenous testosterone, and as a secondary treatment for suspensible metastatic busest causer in women (although it is not widely used for the first parpose amprare). Testosterone cypicaries is currently available outside of the United States, but and widely, Econom searchmental suspense for the day tacked Constant, Australia, Sysian, Enozi, Marchael States, but and the superior and the supe

## How Supplied:

Testosterone cypionate is available in select human and veterinary drug markets. Composition and dosage may vary by country and manufacture, but usually contain 50 me/ml 105 me/ml or 200 me/ml of 5teroid dissolved in oil.

# Structural Characteristics:

Textustence opioutat is a modified form of textustence, where a carboxyle acid sete (seycherpthypoprior) acid has been attached to the 12-bits highoryl garp. Exteriol forms of lextustence are isso point than five textustence, and are absorbed more slowly from the area of legislerio. Once in the blooksteam, the exter is removed to yield tree (active) textustence. Exteriol forms of textustence are designed to prolong the window of the happened self-to blooking administration, adwarp for a less feeger trajection schedule compared to injections of five (mastrifiely steroid. The half-life of textustence expirates to approximately & das after injection.

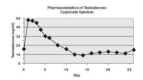


Figure 1. Pharmacokinetics of 200 mg testosterone cypionate injection. Source: Comparison of testosterone, dillydrotestosterone, luteinizing hormone, and follicestimulating hormone in serum after injection of testosterone enauthate or testosterone cypionate. Schulte-Beerbuhl M, Nieschlag E, Fertility and Sterility 33 (1980):2013. Side Effects (Estrogueic):

Testostence is readily anomitted in the body to estradiol (estogen). The anomitse (estogen synthesize) ergons is responsible for finis metablosin or testostence. Bested estogen breds can case side effects each as increased water reterior, body fit giin, and gaecomants. Testostence is considered a moderately estogenis stended. An anti-serrogen such as chemiptene cirate or tumosfire cirate my be necessary to prevent estrogenis and effects. One may admirately use an amountse without feet Arminedo's (meanstools), which deflects. One may demented use an amountse without feet Arminedo's (meanstools), which more efficiently controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

Entogain: side effects will occur in a dose-depondunt rummer, with higher doses (above normal therapeut is beely) of testosterone opinient mere fiels) to requise the concurrent use of an aris-estrogen or aromatuse inhibitor. Since water reterition and loss of muscle definition are common with higher doses of testosteron expirates, this drug is usually constitution are poor choice by defenge or uniting places of training. In moderate estrogenicity makes it more clean for belang places, where the added vusier reterition will support may strongly and

# Side Effects (Androgenic):

Testoderore is the primary mile aufrogen, responsible for miniming secondary mule sessual materiative. Entental news of nestorierone are Body to probe amongenie side effects including obj skin, acre, and body/sicial hair growth. Men with a genetic prodeposition for his two fundoparties despecti may notice accelerated mile pattern halding. Those concerned about hair loss may find a more confertable option in mandeolese decentrate, which is a comprably loss analogenie steared. Women new sourced of the decentrate, which is a comprable year despenies steared. See specially white sign and proportional transparence of the conference of the

In addraga-responsive target tissues such as the skin, soals, and prostate, the high relative analogonicy of testoroure is dependent on its relation to difficultiestoateroe (ERI). The 5-sight reductase enzyme is responsible for this metabolism of testosteroure. The concurrent use of 3-ship reductase inhibitors such as frastrative of antartiest will stretely with six-specific potentiation of testosteroue action, lowering the tendency of testosteroue white of the six of

# Side Effects (Hepatotoxicity):

Teotostorou does not live hepatotoxie effects, her toxicity is utilicity. One staty ceannied he poternia for hepatotoxicy with high does of relocationery by administering 400 mg off the hermose per day (2.800 mg per week) to a group of multi ashjects. The storout was taken orably so that higher peak conventioniston would be reached in hepatic itsues compared to intramascular injectors. The hermore was given dayl for 20 days, and the contraction in the contraction of the contraction and to allow the relocations of the contractions and the contractions are desired to the contraction of the contractions and the contraction and the contractions are contracted to the contraction and the

#### Side Effects (Cardiovascular):

Anabolizandroganie steroide can hove deletroise effects on serum cholestered. This includes a reloracy to refore EMLE (ogod) cholesteroi values and increase EML (odod) cholesteroi values with muy shift the EML to LDL balance in a direction that floros genete risk of anteriocitosists. Bru estudie regard estabolizandroganie seriod on serum lipsis is dependant on the dose, route of administration (rost ix, injectable), type of steroid (commitable or ross-serioritable), and level of resistance to higher interiodiscin commitable or anonemitable), and level of resistance to higher interiodiscin consistance or ross-serioritable, and level of resistance to higher interiodiscin consistance or ross-serioritable, and the off resistance to higher interioritable produced resistance and support left verticaler logoritosphy, all potentially increasing the is for classificaction floroscoratif disease and myconatile inferroir.

Textosterone tends to have a much less drammée impact on condivisacular risk fectors than syndrick storold. Bits die in part to is openers to muchloismity be fice, which allows it to have less effect on the laparite immagnement of cholesterol. The arountation of textosterone to estandard also helps to mitigate the registre defects of andragens on sermilipids. In one study, 250 mg per week of lessosterone ester (centralises) had a sight but not statistically significant effect on EFL cholesterol and fer 2 weeks, but when balson with an amountase ribbitor a strong (25%) devenue was seen. <sup>525</sup> Stuffes integ 300 mg of textosterone ester (centralise) per week for 20 weeks whetca arountasis inhibitor. demonstrated only a 1.5% decrease in H.D. choiesterot, write at 600 mg the reduction reached 2.1% 453. The negative impact of aromatise inhibition should be taken into consideration before such drug is added to testosterone therapy.

Due to the position influence of estingua no serum ligids, turnordine critar or christipnes centure are preferred no currentiase riphise from those concurrent with not accordance that, as they offer a partial estrogatic effect in the five. This allows them to potentially improve in paid profiles and offers our offer pergaine effects of arthogons. Which those sof 600 mg or less per wock, the impact on light profile tends to be noticeable but not durantee, making an artis-entergon fire conflictoration expression perfects are transported to the produce stantistically significant featings in IDAVIDL clinication, independent, application expression of the produce stantistically significant relations in IDAVIDL clinication, independent perfects and install restricting of influencing a relatively work impact on cardiovaster risk factors. When used in another conflictor of the produce stantistical relation of the state of stall and artistically applications.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fists, cholesterol, and simple carbohydates at a litrus during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterolarioxidant formals such as Lipid Stabil or a moduct with commands interedients is also recommended.

# Side Effects (Testosterone Suppression):

All ambolicluntogenic steroids when taken in doese sufficient to promote muscle gain are expected to appress endogenous testistence production. Testistence is the primary multi-androgen, and offices some gangine feedback on endogenous endostience productions, and the production of the contraction of the contraction of the contraction of the contraction of install seroil hormous. Without the sterosterior of toutnets when the production of install seroil hormous. Without the sterosterior for foundation should work of the steroids of toutnets or shoulding substances, testistence levels doubt in term to multi with 1-4 mutch of office sections. We have doubt in the normal with 1-4 mutch of office sections. We have produced the production of the programme of the very large section. We then the production of the produ

As with all ambidesimboguies essentis, it is talkely that one will retain every pound of new bodyweight after a cyles in could. If this is a specially not when withdrawing from as rouge (aromatizing androgen like testsetome cypiontae, as much of the new weight gain is fisely to be in the form (and in the contraction of an architecture) and inhalates of ambide and cambide hormones during the post-cycle recovery period may finder create an ensormer that is utdownship for the return of much tissue. Proper anchiny dung therapy is usually recommended to help restore hormonal balance more quickly, thatmach belongs these errors much tissue.

Another way to be sent the post-cycle "trail" is to fire replace testosterous cypionists with a mider anobles such as midelous deceasing or methenabore centrale. The new steeds would be administered allow for one to two more merths, at a design of 200-400 mg per work. In this "staging down" procedure here are a intempting to dominate the wastery bad understand. This practice can prove to be effective, even framity for procheduced resons force may vive a fast simply dividing the crash into water and hormound staging. Testosteron-stimulating dungs are still hypothylused at the conclusion of therapy, as configurate stationary production will not be found to the stage of the conclusion of the representation of manifoliates.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (Men):

To treat androgen issufficiency, the prescribing guidelines for testosterome expirate call for a dosage of 59-40 mg every two to for weeks. Although active in the body for a longer time, testosterome expirates is usually injected on a weekly basis for physique- or performance-entraining purposes. The usual dosage is in the range of 200-600 mg per so week, taken in cycles 6 to 12 weeks in length. This level is sufficient for most users to notice excentional usins in muscle size and artists. Telestotenes is usually incorporated into bulking phases of training, when added water reterrism will be of life consequence, the user mere concerned with row most stam definition. Some do incorporate the drug into cutting cycles as well, but typically in lower does (100-200 ng per week) and/or when accompaned by an anomatise ribilitor to lower (100-200 ng per week) and/or when accompaned to the contrainer in the control Telestotenee cytomica is a very effective anabole ding; and is often used above with grant beat 600-000. Nower, final more than 100-000 to the control Telestotenee (100-000 ng per week of Tolelforme undeclydente, methenolone centraliset, or machineles decentrates about provide substantial results with on significant hepatoxics/pt (100-1000 results) and the provided provide substantial results with on significant hepatoxics/pt (100-1000 results) to taker the desired effect.

While large does are generally not adviced, some bodyhadders have been kenson to see eccessively thind doesn't offer discuss of the ding (100 mg per work or more). This was much more common forther the 1997s, when cypinate visits were usually very chaop and easy to find. A "more is better" families is easy to justify then project only 250 mg level and only to find. A "more is better" families is easy to justify then project only 250 mg level with contract the typical price for a single injection. At dosages of 804-1000 mg per work or more, water resection will 80c, you count for more of the additional weight gain fam new much tissue. The practice of "megadosing" is infedient (not to mention potentialy dangerous), especially when we take its account the pricial filter of of stouchts touts.

# Administration (Women):

Testosterous cypicate is navly used with women in clinical medicine. When applied, it is not often used a secondary medication during inportable breast cancer, when other therapies have failed to produce a desirable effect and suppression of ourim function is necessary. Testosterous expirates is not necommanded for women for physique- or performance-enhancing purposes due to its storag androgavir nature, tendency to produce willing side effect, and alsow-entire functionaries from function bodd level difficult to control.

# Availability:

Testosterone expionate remains widely available as a prescription drug product. Its production is largely associated with American companies, although recently has been expanding into loosely regulated Asian markets that all cater to demand by bodybuikters and athletes. In reviewing some of the products and changes in the global pharmaceutical market, we have much the following observations.

Brand name testosterone cypionate (Depot-Testosterone) remains available in the United Stated from Prizer. This is a high-profile target of counterfeiters. All legitimate boxes will carry a "Bh" symbol hidden on one of the top inside flaps. It will appear when placed under I/V lider.

Many generic forms of the drug are also produced in the U.S. market by manufacturers such as Waison, Sandox, Paddock, Synerx, and Bedford. All come packaged in multiple-dose visik. Due to strict cortrols these products are rarely diserted for filler's also. There are also several pharmacies custom-compounding testosterone cypiorate for doctors that specialize in anthone neochement theraps.

Cypionax is available in Thailand by T.P. Drug Laboratories. It comes in 2 ml. ampules containing 100 mg/ml. of steroid.

Cypiobolic from Asia Pharma (Malaysia) is now approved for sale through pharmacies in Thailand, Each box should carry a scratch-off security sticker, which will display a code that can be validated on the comnary website.

Testex Prolongatum remains available in Spain. This steroid is produced by Laboratorios Q Pharms. It is packaged in 2 mL dark glass ampules with grey silkscreen lettering. It comes in two doses, containing a total of 100 mg or 250 mg of steroid. Testex has always been a high-profile item for counterfileres.

Found in Chile is a high-dose cypionate product called ciclo-6. The product is manufactured by the firm Drag Pharma, and contains 300 mg/ml of steroid in a 2 mL ampule (600 mg of evolutate in total). Balkan Pharmaceuticals (Moldova) makes the product Testosterona C. It is prepared in both 1 mL ampules and multi-dose vials.

### Deposterona (testosterone blend)

Androgenic	100
Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	

#### Description:

Progestational Activity

Depotations is an injectable victoriumy steorial preparation that contain a blend of three different testosteres extests. Each million contain 3 mg designers acutests, 12mg of testosteror understand, for a total steorid concentration of 60 mg/lm. This is currently to early contain contain one of 50 mg/lm. This is currently to early contain participated with a last file approximately double four dissolutions proportions. <sup>50</sup> White his blend of slow- and fast-aring setters, Depotations is essentified a 20 mg/lm. The size currently as the longer acting with more understand is with the larger acting with more understand is a first of the larger acting with more understand is a first of the larger acting with more understand is a first of the larger acting with more understand is a first of the larger acting with more understand is a first of the larger acting with more understand is a first of the larger acting with more understand in the size acting with the size acting with more understand in the size acting with the size acting with more understand in the size acting with the size acting with more understand in the size acting with the size acting with more understand in the size acting with the size acting with more understand in the size acting with the size acting with more understand in the size acting with the size acting with

## History:

Depoterons was developed by Systex Asiral Health Company several decades ago, and has been sold on the Mexican seterings dieg matter since. It is used primarily to men protocure, welclasses, dispar, and hypogensisms in mile breeding similar (cowe, pagrapotence, welclasses), and as a greately protein spring authorit. Depositorum is now soil under de For Dodge Admit Health bed, which exquired Systex, Armil Health in the mat-1900's.

#### How Supplied:

Deposterona is available on the Mexican veterinary drug market. It contains 12mg of testosterone acetate, 12mg of testosterone valerate, and 36mg of testosterone undecanoate ner millifier of oit nackaged in a 10 mL multi-dose vial Twelve vials are nacked in each box.

# Structural Characteristics:

Depostroom contains a minuse of three testosterone compounds, which where modified with the addition of carboying acid enter (society, subtract, and undecunic acids) at the 17-beth lightoxyl group. Esterfield forms of isotosterons are less point rune fee testosterons and are adsorbed more solvely form the area of injection. Once in the bloodstraum, the ester is removed to yield five (acids) estosterons. Esterfield forms of isotosterons are designed to the estimate of the contract of the co

#### Side Effects (Estrogenic):

Testosterone is readily aromatized in the body to estradiol (estrogen). The aromatase (estrogen synthetase) enzyme is responsible for this metabolism of testosterone. Elevated estrogen levels can cause sick effects such as increased water retertion, body fit gain, and genecomstia. Testosterone is considered a moderately estrogenic steroid. An arti-estrogen such as chambers critate or tamoritine extrate more horsessare from every estronesis in the contract according to the contract testos and the contract extractions and the contract estronesis in the cont

effects. One may alternately use an aromatase irribitor like Arimidex® (anastrozole), which more efficiently controls estrogen by preventing its synthesis. Aromatase irribitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

Entropeis side effects will occur in a dose-dependent rumers, with higher doses (observement therspects beselved) restoateres owns (ledy) to require the occurrent use of instructions of a remainsteate of a remains eithboric. Since water reterion and loss of muscle dedintion are common with higher bosos of frestantients, this drug is usually considered a poor choice for desirg or custing places of training, list moderate entroperation makes in more liked for behalf places, where he sadded vater creates will appear may seem the sadded vater creates will appear may seem the sadded vater creates will appear mis seeming that makes the contraction will appear mis seeming that makes the sadded vater creates will appear an seeming the sadded vater creates will appear an seeming that makes the sadded vater creates will appear to see seeming the sadded vater creates will appear to see seeming the sadded vater creates will appear to see seeming the sadded vater creates will appear to see the sadded vater creates will appear an assent the sadded vater creates will appear to see the sadded vater creates will appear an assent the sadded vater creates will appear to see the sadded vater creates will appear to see the sadded vater creates will appear to see the sadded vater creates and seems of the sadded vater creates will appear to see the sadded vater creates and seems of the sadded vater creates are seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater creates are said vater and seems of the sadded vater are said

# Side Effects (Androgenic):

Telestotence is the primary mile androgen, responsible for minimizing secondary mulesessual enhanceristics. Extended level of testistone are Bieds to proble androgenic side effects including oily skit, exer., and body/ficial law growth. Men with a genetic prodeposition for his lives (androgenic-desporcia) may notice accelerated mule pattern balding. These concerned about har loss may find a more confertable option in sandosive decennote, which is a comparable less androgenic setted. Werne are swamed of the potential virling effects of analoxic androgenic setted. See optically with a strong androgen chances in shire touch, fine liber and the conference of the conferenc

In advagor-recognosive target tissues such as the skin, such, and prostate, the high relative analogousity of testistenture is dependent on its reduction to diffeotistenture (DEI). The 5-sight reductative entryine is responsible for this metabolism of isotostomes. The concurrent use of 5-ship reductation is individually constructed to a first reductive of interfere with sits-specific potentiation of isotostomes action, lowering the tendency of testistenture during to produce analogous sits deficient. It is important to remarker that anabolic and analogous; effects are both mediated via the cytosolic analogous receptor. Complete expectation of isotostomes is anabolic and randogous; effects are both mediated via the cytosolic analogous receptor. Complete expectation of isotostomes is anabolic and analogous inceptors to a probable, cut with

# Side Effects (Henatotoxicity);

Testostorou does not have hepatotoxic effects, hor toxicy is utilicity. One study custined he potertial for hepatotoxicy with high does of electostrone by administering 400 mg of the hermose per day (2,800 mg per week) to a goup of mine subjects. The steroid was election only so that higher peak concentrations would be reached in hepatic itssues compared to intranscender spectrum. The hermose was given days for 20 days, and all the contraction to intranscender spectrum. The hermose was given days for 20 days, and all the contraction of the days of the contraction of the days of the days

#### Side Effects (Cardiovascular):

Anabolizardnoganis steorilo en la true deletriros effects on serum cholesterel. This irelades a redeney to refuee III. (6 god) cholesterel values and irecrese III. (6 god) cholesterel values and irecrese III. (6 god) cholesterel values values develo report risk of anteriocitoris. If ne redute report of an antobiciantiques restorel on serum lipit is dependant on the dose, route of administration (rost as, injectable), type of steorid (contratible) er new-normaticals), and level of resistence to hippair entelledent and contrating and anterior and anterior development and anterior development

Textosterore tends to lawe a much less dramutic impact on candirouscular risk factors time synthetic steroids. This is due in part to its operates to metabolism) by the low, which allows it to have loss effect on the hapatic management of cholesterol. The arountations of it to have loss of the contraction of the contraction of the contraction of the plack in cost subj. 20 map per wock of thosostorous easier quantitate) that a slight but not statistically significant effect on HDL cholesterol after 12 works, but when taken with a countraction of the contraction of the contracti testosterone ester (emuritate) per week, for 20 weeks without an aromatsus irribitor demonstrated only a 13% decrease in HDL cholesterol, while at 600 mg the reduction reached 21% <sup>608</sup>. The negative impact of aromatsus irribition should be taken into consideration before such drug is added to testosterone therapy.

Due to the position influence of estinguan on serum lipids, temoritien cirate or chimiphene cirate are preferrion to accuminess prinkines for those concerned with action/securlar health, as they offer a partial estrogatic effect in the fiser. This allows them to potentially improve lipid profiles and offers own of the regarder effects of androgens. Whit does or 600 tong re-less per week, the impact on lipid profile tended to be not exclude but not dearnate, making an anti-entergraph for endeptocetedine purpose, peripose transcessors, boses 6000 tong re-less per week three also failed to produce statistically significant changes in ELD/VLD. and including a calcularly used in part or an endowancial residence of the control of the control of a failed an including a calcular produce statistically significant changes in ELD/VLD. and including a calcularly used in grant or an endowancial residence of the calcular of all includings a calcular possible textocorrune caeses are usually conducted to be the saltes of all controls.

To help roduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fits, cholesterol, and simple carbohydants at a litrus during active AVS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterolatrioxident formals such as Lipid Stubil or a moduct with commands intenderies is also recommended.

#### Side Effects (Testosterone Sumpression):

All antibolicalinguies storoids when taken in doses sufficient to promote muscle gain are expected to suppose condeposus testostorous production. Testostoroes is the primary under surface, and offers storage negative foodback on endogenue testostorous production. Testostoroes-besed diago will, Rowisch, length as strong effect on the probeduline regulation of intuities attentil hormous. Without the intervention of isotostorous-strainfairing substances, testostorous less should return to normal with 1-4 mentric fording excession. Note that prolonged physicagendotropic hypogeneithoring on the control of the proposed o

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (Men):

For bodybalding purposes, Deposterom is usually nigored on at least a weekly basis, in a dosage of 120-350 mg (2-6 m). Diviling the weekly dosage into two or more smaller applications can reduce injection volume. Cycles are generally between 6 and 12 weeks in length. This level is sufficient to provide noticeable gains in muscle size and strength. Testosterone drugs are ultimately very versatis, and can be combined with many other anabodicinationsois extends deemedion on the desired delta.

# Administration (Women):

Deposterona is not recommended for women for performance-enhancing purposes due to its strong androgenic nature, tendency to produce virilizing side effects, and slow-acting characteristics (miking blood levels difficult to control).

#### Availability:

Deposterona is only known to be manufactured in Mexico. Because it contains such a low concentration of steroid, this product is not in high demand, and not readily diverted for illicit sale

# Dianabol® (methandrostenolone, methandienone)

Androgenic	40-60
Anabolic	90-210

Methyltestosterone (oral)

Chemical Names 17a-methyl-17b-hydroxy-1,4-androstadien-3-one 1-Dehydro-17a-methyltestosterone Estrogenic Activity moderate Progestational Activity not significant

# Description:

Standard

Dambei is the most recognized made sume for the dang methandrostenolore, also referred to so methandrossom many courties. Methandrostenolore a destroite of Estostenore, modified so that the hormore's antrogenic (musculinizing) properties are reduced and 8s annabadic (issue budding) properties preserved. Hings a lower level of rethine antrogenic; that testostenore, methandrostenolore is classified as an "artshelf" stroot, although quite a discriber antrogenic is is off present. This divers to design a singer principally soil, as an official antrogenic and the singer properties are singer properties. The singer properties are singer properties are singer properties and the singer properties are singer properties and the singer properties are singer properties. The singer properties are singer properties are singer properties and the singer properties are singer properties. The singer properties are singer properties are singer properties are singer properties. The singer properties are singer properties are singer properties are singer properties. The singer properties are singer properties are singer properties are singer properties. The singer properties are singer properties are singer properties are singer properties are singer properties. The singer properties are singer properties are singer properties are singer properties are singer properties. The singer properties are singer properties are singer properties are singer properties are singer properties. The singer properties are singer

# History:

Mediandrosterolore was first described in 1955. <sup>60</sup> It was released to the U.S. prescription during market in 1958, under the brand mure Diamolol by Che Dimermocratical. Che desdeped methandrosterolore in a mediane with the high of Dr. John Zagle, who was removed to the property of th

Dr. Zigdev shiftees were quickly making great advancements in their competities causers with the hip of the ang. According to reports. Zigder too seemed to be very impressed, at least fir a valle. Mel Bat by the early 1960's, it was sturing to look like Diembel Ind spatical a great wave of setted alsess in competities sport. Dr. Zigder's extern report procumentations, which depending on the source called for as little as 5 mg per day or as much as 15 mg per day were being laughly ground; as althost developed their our mere aggressive (and potertially duagerous) dozing strategies. Dr. Zigder com became disquisted with the risses of the drug, and would eventually become a voice of opposition to sport during. By 1967, approximately 10 years after first strategies. Dr. Battle and the size of the drug and would eventually become a voice of opposition to sport during. By 1967, approximately 10 years after first strategies. By the discussion of the categorically contendent the use of anthosis searches in your large and the categorically contendent the use of anthosis searches in your large and the procure of the procession of the procession of the procession of the categorically contendent the use of anthosis searches in your large and the procession of the p

As early as 1905, Distribed was sheady starting to fit under scarting of the U.S. Food and Drug Administration. That year the F.D.A requested Che neithy Distribed models uses, which were then stated to relate he playing patients in debilitted states and from with weakened boxes. In 1970, the FDA accepted that Distribed was "Probably Effective" in tending post-interpretated cotecpression and paintary-deficient drustfarm. These changes were advantaged to contrast and the probably effective in tending to the probably effective in tending to the probably effective in the probably effecti

By the early-80's the FDA had withdrawn its "Probably Effective" position on the pituitarydeficient dwarfsm, and continued to press Cha for more data. Sufficient clerification never came, and in 1983 Cha officially withdrew Dianabol from the U.S. market. 463 Perhaps francial disinterest had a hand in their abandoned push to keep the drug approved. The FDA pulsed all generic forms of methandrosterolone from the U.S. market in 1985, a time when most Western mations were also eliminating the drug, finding its existence to be justiced mainly by sports doping. Methandrosterolone is still produced today, but typically in nations with loose prescription drug regulations, and by companies that still prefer to cater to an underground affeither market.

## How Supplied:

Methandrostenoloue is widely available in both harma and sterrinary drug markets. Composition and dosage may vary by country and manufacture. Methandrostenoloue was designed as an eral anabolic steroid containing 2.5 mg or 5 mg of steroid per tablet. Olimatohy, Modern brands usually counted in Super of 10 mg per tablet. Methandrostenoloue can also be fourd in nijectable veterinary preparations. These are typically obbased solutions that carriv2.7 mitrof domoid:

## Structural Characteristics:

Mediandostoolore is a modified form of testosterore. It differs by 1) the addition of a morely group at control 1-7 sph to protect the hormone during and administration and 2) the introduction of a double bond between carbons 1 and 2, which reduces is relative andopasticy. The results general double has much washer relative being affinity for the andopastic receptor than testosterore, but at the same time displays a much larger fluid life and lover affinity for entermberling proteins in comprision. These literates (uning others) adow mediandosistensible to be a very potent anabolis steered in spite of a weeker affinity and the steered of the steered and the steered of the control includes interest on with the colline actions recovered.

# Side Effects (Estrogenic):

Mediandrostrolore is arountated by the body, and is a medienticly estinguistic seriorid. See (Specicominals in delira account advirge trustment, and may prosent befor disearch with an cycle (particularly when higher does are used). At the same time useder restriction can receive a seriorid or the seriorid or the seriorid or the seriorid or the control of the seriorid or the seriorida or the seriorida

It is three-ring to note that medium/osterokone is structurally identical to bolderone, except that it contains the added 1-74-piles-medily group. This fact much secked the impact of aluring at street of a such as way, as these two compounds appear to act very differently in the body. Alexy disamilarly seems to lie in the tentucky for estiognast is defented. Explosived (bolderone undecylentes) is known to be quite mild in this regard, and users commonly take this drug without the rend to sold an anti-restogna. Methanderothorie is much more estinguist, often necessitating safe-storogen use. But this difference is not caused by clearly contained to the contract of the contraction of the contraction of the contraction of the contraction of the contraction. The contraction is more than the contraction of the contraction of the contraction. The contraction is considered, The time actually is caused by methandrosterolous concerting to 17alpha-methylestrafied, a more biologically active from the contraction that extends on the contraction of the con

#### Side Effects (Androgenic):

Albacqui classified as an ambolic steroid, androgenic side effects are still common with this statence. This may rickel boats of obj. skix, nexe, and bodylifical in his growth. Antibolic/androgenic steroids may also aggravate melt pattern hair loss. Individuals sensible to the antibogenic effects of mediumdostrookee may first a milder antibodis used as a Dexa-Danbolint to the runte controllable. Women are additionally sound of the potential processing and the state of the processing and the state of the processing in controllable. Women are additionally sound of the potential processing and the state of the processing in controllable.

While methandrostenolone does convert to a more potent steroid via interaction with the 5alpha reductase enzyme (the same enzyme responsible for converting testosterone to dihydrotestosterone), it has an extremely low affinity to do so.\*\*\*\*\*on The androgenic metabolite 5-alpha dihydromethandrostenolone is produced only in trace amounts, so the relative androgenicity of methandrostenolone is not significantly affected by finasteride or dutasteride.

#### Side Effects (Hepatotoxicity):

Methandrostroshore is a c17-apha allytited compound. This alteration protects the drug from detectionin by the liver, allowing a very high precurating of the druge error into the bloodstream following onal administration. (17-apha allytited anabolic/androgenic sterosids can be hapstrostor. Froblegged of high process error years at liver druging. In rare instances life-frencineing dysfurction may develop, it is a subseque to visit a pipokine proportionally during each cycle to remote free fraction and overall health inchia or c17-apha allytited seroids each cycle to remote free fraction and overall health inchia or c17-apha allytited seroids

Studies have shown that several weeks of methandrosterokore administration offers minimal hypitic stress so long as it is given at a dosage of 10 mg per day or below. At a dose of 15 mg per day, a mightod below factor will sepan to demonstate distributed here factors as measured by distaily elevated bromosolphicin retention (a marker of hepatic stress). We see that the second of the period of the period of the second of the period o

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any herastotoxic analysis and movemic stemicls.

## Side Effects (Cardiovascular):

Anabolizardneganis steroids can have deletrious effects on serum cholesterol. This includes are infraency to reface IEEE (good) erlobsterol values and increase IEEE, (doub) erlobsterol values and travester LEE, (doub) erlobsterol values which may shift the HDL to LDL balance in a direction that through greater risk of attractivectors. The relative impact of an anabolicardneganic storoid on serum lights is dependant on the dose, route of administration (real two, ispectable), type of steroid (commitable) or non-serumitable), and level of resistance to hapteric metabolizardneganic serum and control of administration. Anabolicardneganic serum and the service of the service of

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular eucreise program and minimize the intake of saturated fits, cholsterol, and simple carebolydates at a litrus during active AAS administration. Supplementing with fish olk (4 games per day) and a natural cholsterol/artioxidant formula such as Lipid Stabil or a module with commands interdents is also recommended.

# Side Effects (Testosterone Suppression):

All antibolizationguies steroids when taken in doses sufficient to promote muscle gain are expected to supress endogenous testoctorus production. Methandisosticovie is no exception, and is noted for its strong influence on the hypothalimic-pinhary-steriodar axis. Clinical statles jurige [3] par gor duy to restation-training miscle States entaining miscle States of 8 weeks caused the mean plasma testosterous level to fall by 69%. 600 Without the intervention of testosterous-straining substances, testosterous levels to fall by 69%. 600 Without the intervention of testosterous-straining substances, testosterous levels and terratio normal within 14 ments of official security of the straining substances, testosterous levels sould testom to normal within 14 ments of official security to storial datase, excession Note that prolonged hypogonadorn can develop secondary to storial datase.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its bioavailability. 400 This is caused by the fat-soloble nature of steroid hormones, which can allow come of the dates to discrebe with rediscrebe distray for nedworine its obscurious from

and some or the using to desorve with thingested openiny in, reducing as accompany north the gestrointestinal tract. For maximum utilization, this steroid should be taken on an empty storach.

# Administration (Men):

The original prescribing galdelines for Dinmbol called for a chigh dosage of 5 mg. This was to be admissized on an intermilent beats, with the drug baths from nome than 6 consecution works. Thereafter, a break of 2 to 4 works was ableed before farings was reasonal. For expecting the property of the control of 2 to 4 works was ableed before farings was comment. For expecting the property of the control of 2 to 4 works was ableed before farings was consecuted by the control of 2 to 4 works of 2 third galley as well of 5 mg did years be effective for improving performance, adhees typically take much higher amounts. A didy dosage of there to a 5. mg takes (1.5 30 mg) is most bank to provide the control of 2 to 4 third proving the control of 2 third proving the control

Dambol stacks well with a variety of other steeds. It is noted to mit purticularly well with the mild anaboli. Does-Dambolioli, for enemple. Together one on expect exceptional macks and strength spirs, with side effects not much voose than one would expect from Dambol adance. For elser mass, a large-active patients can be used. With the high estrogeticality experies of this analongen, however, side effects should be more pronounced. Gains would be pronounced as well which usually makes such an endeavor worthwhile to the user. As discussed earther, anrillary drugs can be added to reduce the side effects associated with his infra of cycle.

The halfies of Diamedo is only about 3 to 5 hours. A single daily dosage schedule will produce a suring blood best, with up and allows freedings that days. The screen feets in an a choice, to other spik up the tabless during the day or to take them all at one time. The usual recommendation in been to the differ them and ty to regalant the concentration in your at once, so there may be a read-off with this option. Both options work fine, but annecledul excludes easier to support single daily does a being better for overall reads. While the schedule, it seems begoed that taking the pilk earther in the day would be option. But the schedule, it seems begoed that taking the pilk earther in the day would be option. But the schedule, it seems begoed that taking the pilk earther in the day would be option. But the schedule, it seems begoed that taking the pilk earther in the day would be option. This would do not be a support to the schedule of the schedule of the support to the schedule of the schedule. See that the schedule of the schedule of the support to the schedule of th

#### Administration (Women):

Being moderately androgenic, Diembol is really only a popular steroid with men. When used yourner, strong vification symptoms are possible. Some do experiment with it, however, and often find low dosse (2.5-5 mg) of this steroid quite effective for new muscle growth. Studies have demonstrated that a majority of women will intoice acro, which is infenditive of androgenicity, at a dosage of only 10 mg per day. Children are likely to notice vitilizing effects with as fitted as 2.5 m per day.

#### Availability:

Methandrostenolone remains readily available as a pharmaceutical product, though its supply is largely isolated to less regulated markets in Asia and Eastern Europe. In reviewing some of the more popular products and changes on the global pharmaceutical market, we have made the following observations.

British Dispersary produces Arabol tablets in Thailand. This product comes in three strengths, 5 mg, 10 mg, and 15 mg, 41 products come in bothes (100 200, 500, and/or 1000 depending on the does). All tablets themselves are imprired with the company's stude entire mat will vary in color. The piths tablets are 5 mg, the yellow 10 mg, and blue 15 mg. The company also uses a holographic sticker on all products to deter counterficing, though this feature has been copied with light accuracy.

March Pharmaceuticals in Thailand manufacturers Danabol DS. These small blue heartshaped tablets come in bottles of 500. The company recently instituted a small circular holographic stricker to deter counterfeiting. The logo is also embedded into the label in a moral foil. Dronator Lts is remains available in Trailand by Bangkok Lab & Cosmetie. This product comes in a 10 mg tablet strength, and is prepared in bottles of 500 and 1,000 tablets each.

Methandon is also found in Thailand, made by Acidhon. This product comes as a 5 mg

tablet, and is packaged in plastic tubs of 1,000 tablets each.

Generic "Russian D-Bol" (METAHAPOCTEHOROH) is no lonser in production All.

products bearing this label should be considered counterfeit.

Naposim (Romania) is still in production, now under the joint Temphia Rambusy label. It is sold in houses of 20 tablets, which are separated into two foll-platic bisters of 10 each. The tablets carry at finned starm on one side. This mothet has born the solder of wisdesmonth.

counterfeiting in the past.

Metanabol from Jelfa (Poland) is still in production, though its appearance has changed. The new packaging reflects a more modern red gradient look. The enclosed foil and plastic strip

is also much wider than it was previously.

The generic from Formula Magistral (Argentina) is also commonly located on the international murket, especially throughout North and South America. The product comes lose in buttles although now also entires a holoromist sickle not deter courterfeitine.

Landerlan in Paraguay makes a generic methandrostenolone. It comes in a 10 mg tablet dosage. The product is nackaged in bottles of 100 tablets each.

Balkan Pharmaceuticals (Moldova) makes the product Danabol. It is prepared in both 10 mg and 50 mg tablets, with 20 tablets contained in each foil and plastic strip.

# Drive® (boldenone/methylandrostenediol blend)

Androgenic	
Anabolic	
Standard	

# Chemical Names

Estrogenic Activity
Progestational Activity

#### Description:

Due is an Astarlian injectable ventriony steroid preparation that contains a blend of mediantical dispositione and boldenous unlexpleature. These sociolists are present in dose of 25 might. and 30 mg/ml. respectively, for a total seroid concentration of 55 mg/ml. Boldenous endeclessate is alight communication and results included with a preparation Enquisors 69. Mediantical disposperatio, however, is very unely seen on the U.S. seconograid by a notable androgous component. When combined with boldenous, the result is a moderately anticognic numbels bend included to produce notable muscle muss and strength gains, unally whost occasions when the result is a moderate of the combined of the combine

#### History:

Doe is a product of ROW. Vectorinary Products (formuly a substitiny of Nature Velts, odd) on the Austiliant vectorinary due muter. It is designed for use in homes, typically as a general anabolic or health toxic drug for when an animal is weak from vigorous performance. It is supposed to a late ground of muscle bissue, help avoid delephation; and improve the digastion of detemy proteins. The dosage used for an adult 1/100 home is 5 m./101 may cover two weeks. Austilia is a country with a robust vectorinary drug market, known to

cury a varry or unsuat serous and our man-component serous beauts. Due to present the most well-known of these products. Being that it is neither the most concentrated preparation nor the most effective, however, much of its popularity is fixely due to its well-conclud rand eman and early sales bistory. Divis remains on the Australian market today, although right controls and its relatively two per-milliter seroid concentration make diversion for atthictive men these common than two sums my varse as the serous than the varse may varse as the serior of the control of the control of the serior of the control of the cont

# How Supplied:

Drive is available on the Australian veterinary drug market. It contains 55 mg/mL of steroid in oil in a 10 ml vial

# Structural Characteristics:

For a more comprehensive discussion of the individual steroids boldenone undecylenate and methandriol dipropionate, refer to their respective profiles.

#### Side Effects (Estrogenic):

Mediplandostatidi is not directly aromitated by the body, although one of its known methodies is melhybetostorous, which can aromitate, Mediplandostatedid is also believed to have some inheuert estrogenic activity. <sup>470</sup> Corribated with boldenous, which also communities, Their is considered a mondered postagenic second Cytecocomistis in possible anomatics, and the considered a mondered postagenic second Cytecocomistis in possible and the considered anomatics of the considered anomatics of the considered in the

# Side Effects (Androgenic);

Although classified as an anabolic steroid preparation, androgenic side effects are still common with this selstance. This may include boats of 100 spiks, nace, and 100 spiks nace are selstant production and produce its serioids. The serious production are selstant produced in the serious nace and the production and the serious nace and the seriou

#### Side Effects (Hepatotoxicity):

Methyurkostenciós is a c1<sup>-1</sup>-elpta allylated compound. This altention protects the drug from descritioni by the Bee, allways as yell piencemage of the drug enty in the bloodsteam following oral admissistation. C1<sup>-2</sup>-elpha allylated ambolicándroganie steroids can be hydratosic. Probagod for high cosporar emy reset in free drumge. In zero instrución can be chaptación, el consideration may develop it is adviseble to visit a physiciem promoticuly during canch cycle to remiete for faction and oreal mellals. Intake of c1<sup>-2</sup>-piah allylated steroids canche cycle to remiete for faction and oreal mellals. Intake of c1<sup>-2</sup>-piah allylated steroids form of the dung my present sjefatly less strain on the Borr by acoting the first pass methodismo officiologica, although my sill prosert substantial protectorics;

# Side Effects (Cardiovascular):

Ambolicardroganic stroutis can have deleterious effects on serum cholesterol. This irelates a reductry to reface IELL (apud cholestero) takes and increase IELL (doud) cholesterol values, which may shift the IELL to IEL chalmee in a direction that fixors geneter risk of anteriocelosis. Bre relative impact of an ambolicardroganic stroution ensure lipits is dependant on the dose, route of administration toral xx. ispectable), type of steroid generation of the secondary of the particular through the contraction of the particular through the contraction of the particular through the contraction of th

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4

grams per day) and a natural cholesterol'antioxidant formula such as Lipid Stabil or a product with comparable ingredients is also recommended.

# Side Effects (Testosterone Suppression):

All anabolic/ardrogenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-stimaling substances, testosterone levels should return to normal within 1-4 morths of drug secession. Note that produnged hypogoradotrophic hypogoradism can develor secondary to steroid abuse encosessation emfedicil intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

# Administration (Men):

Doke has not been approved for use in harmars. Prescribing gaidelines are unrealiable. Flypical dosing selectable for plyssique or performance-relating purposes would be in the range of 220 mg (4mL) to 440 mg (8mL) per week, a level that should provide quality lean mess gain without strong beloning no body fit neterinto. Due to the high spicetion volume and fastacting nature of methanizated dipropionate, the total weekly dosage is commonly divided into 2-3 smiller amolications.

# Administration (Women):

Drive has not been approved for use in humans. Prescribing guidelines are unavailable. Drugs containing methylandrostenediol are generally not recommended for women for physique- or performance-erdnancing purposes due to its androgenic nature and tendency to produce vitilizies side effects.

## Availability:

Andrewski.

Drive remains in production, though legitimate products are now rarely diverted to the black market given the heightened controls over anabolic steroids in Australia.

## Durabolin® (nandrolone phenylpropionate)

Estrogenic Activit	y low
Chemical Names	19-norandrost-4-en-3-one-17beta-ol 17beta-hydroxy-estr-4-en-3-one
Standard	Testosterone
Anabolic	125
Androgenic	37

# Description:

Progestational Activity

Nandrobers plensphospiente is an injectable form of the authoris served markolone. The properties of this ding are striking's sinite in those of Deca-Drabolinel, which uses the slower acting drug markolone dearmosts. The primary difference between those two preparations is the speed of which markolone is relaxed at the blood. While markolone dearmost provides a release of transformed form the sens of species to being approximately 3 works, markolone phersphospiness is sent for only about a week. In tribular sharp Deca-Drabolin care than the re-priced once every 2 or 3 works, while Drabolin's is usually interchanguable. Each Deca-Drabolin, Drabolin is waked by whitese and body-blackers for its abilities to promote strength and lean muscle muss gains without significant entrogenic or androgenic side effects.

moderate

#### History:

Nandrokov phenspropiente was first described in 1937. <sup>271</sup> It became a prescription medication shortly disc, odd by the irentational pharmacerials gaint Origonou (now Marcicko RND) under the brand name Darabolin. When first irroduced to the United States, related uses of markows plenspropriese in related uses of markows plenspropriese in related to the considerance or effective consequences, advanced breast cancer, weight loss due to considerance or the temporal pharmacerial forms of arminis, and esoletic cases of growth and development restructions in children. During the 1970's, the FDA began revising the indicated uses of this change however, and they were soon signification primorously. Among bowards, the days was mady being reliciated for the treatment of advanced metastack beast cancer, and as adjunct thempty for the treatment of fault superspectations of the state of t

nately to use comment on senie and post-introputation conceptories.

Darbolin was a 18-yes for Ofgrant's mixeding efforts only for well less than a decade following its release. Once Deca-Durbolin was introduced claring the 1969's, his shorter-acting contenque, at least, and less than a feed of the complete of shandward by Organs at the time, knowces; partly the to the feet that it was been as the state of the contract of the

# How Supplied:

Nandrolone phenylpropionate is available in select human drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 25 mg/mL or 50 mg/mL of steroid dissolved in oil

# Structural Characteristics:

Nandrolene phenyleropiente is a modified fem of mandrolene, where a carboyós acid ester (propione pinele pated pas less materiale and to 18 17-8te a lytosog jessor, laseride steroids are less polar fran fies steroids, and are absorbed more slowly from the area of rejection. Once in the boudenarm, the ester is removed to judi free (action) samtrolene. Esterfield steroids are designed to prolong the wardow of thempeatre effect following administration, allowing for a less feequers rejection schools companed to injection of the (unescrified) steroid. Nandrolene phenyleropiente provides a dump spike in nandrolene (unescrified) steroid. Nandrolene phenyleropiente provides a dump spike in nandrolene (bull spike) steroid. Steroid steroid in the steroid profession of the steroid profession to are baselier.

# Side Effects (Estrogenic):

Nandrolene has a low tendency for entrogen connection, estimated to be only about 20% of that seen with testocheron.<sup>47</sup> This is because while the here on convent mandrolene to estandial, in other more active sites of steroid aromatisation such as adjoice tissue mandrolene in the scope to this process.<sup>47</sup> Consequente, designed—related self-efficient still be noticed with higher dowing, however, and may cause side efficies such as increased wave retention, body faigust, and generosation, an artis-entrogen such as chargingciates or transcribe critate run plus necessary to prevent entrogens sale efficies if they occur. One may alternative use a montante similar the Avintacked (materiologic, side) more compensation of the contraction of the contraction of the contraction of the contraction of the operation in comparison to anti-estrogens, however, and may also have negative effects on blood lipids.

It is of note that nandrokne has some activity as a progestin in the body.<sup>474</sup> Although progesterone is a 0-19 steroid, removal of this group as in 19-norprogesterone creates a hormone with greater barding affinity for its corresponding receptor. Starting this triat, may 19-nor arabolic steroids are shown to have some affinity for the progesterone receptor as a creater of the progesterone receptor as creater of the progesterone receptor as a creater of the progesterone receptor as creater of the progesterone receptor as

welt\*" The side effects associated with progesterone are smiter to those of estingaiculting regardles redouch ciribition of testistenous production and enhanced rate of fire storage. Progesters also augment the strendstory effect of estingars on mammy tissue growth. These appears so be a storage purely perhaven these two hommens here, such that the storage of the the second of the storage of

#### Side Effects (Andmoenic):

Albudgh clossfield as an ambolic stroot, androguic side effects are still possible with this softstance, expected with higher doors. Then my related botts of obj skin, area, and body/ficial law growth. Antibolicalindogarie steroids may also aggranter miler planter hair loss. Women are warmed for pertoral writing effects of ambolicantegousie smooth. These my related a deepening of the voice, mentional imagheries, changes in skin texture, lessel hair growth, and closel undiregousier. Nanolonies is a steroid with existing wardings earliey archive to its tissue-basiling actions, making the threshold for strong antioguic size deficies comparably lighter flam with mree antiopages agarsts and as a testionations, medium/chostorious, or flançous-texture it is also important to posit out that manufactors is more to insertion with high one in miles believed without another and or an antioparties of the constraints of the miles when the constraint and or an antioparties of the constraints of the miles when the soft standard and constraints.

Note that in androgen-responsive target tissues such as the skin, scalp, and prostate, the relative androgenicity of nandrolone is reduced by its reduction to dihydromandrolone (DIDA) 476-477

The 5-alpha reductise enzyme is responsible for this metabolism of randrolone. The concurrent use of a 5-alpha reductise inhibitor such as finisterior or datasteride will interfere with site-specific reduction of mandrolone action, considerably increasing the tendency of mandrolone to produce androgenic side effects. Reductuse inhibitors should be avoided with mandrolone fillow and reconcilior is desired.

# Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

#### Side Effects (Cardiovascular):

Anabalicándroganic atomátic can hovo delatrious effects on serum cholestord. This includes a nederacy to reface IEM, (ago) cholestoro tales and irraces IEM, (ado) cholestoro tales, which may shift the IEM. to IEM, bulmer in a direction that flowes gener risk of antirociscosis. The relation impact of an ambibicantingonic storiol on serum light is dependant on the dose, route of administration (end xx. ijectoble), type of steroid (contratible or mac assemble)ke), althe coff or instantive to heavy methodors. Statistic administragi (60) mg of mandroine decareates per work for 10 weeks demonstraed a 20% rochacin in IEM. Colesciol tocks. <sup>47</sup> This expression is slightly outer methodors. Statistic administragi (60) mg of mandroine decareates per work for 10 weeks demonstraed a 20% rochacin in IEM. Colesciol tocks. <sup>47</sup> This expression is slightly suggest remitted in 1911. Dit. in oliv in fundorise decareates a compared to instosterore cypoinne. <sup>48</sup> Narholone should all lines a significantly weaker impact on serum light in nor 17 alpha shelded agains. Antidochimagesis storistic impact on serum light in nor 17 alpha shelded agains. Antidochimagesis storistic impact on serum light of inferious.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and mrininze the intake of saturated fits, cholsetrol, and simple carbohydrates at all times during active AAS administration. Supplementing with sho id, grams per day) and a natural cholseteo/laritoxidant formals such as Lipid Stabil or a moduct with contrareable inerdeties is also recommended.

# Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to converse endogenous testratorous mediation. Studies administration 100 mm especies or suppose transgame institution production. States manuscription of multipolice phelipopionistic demonstrated a rigid apprecision of semanticipole phelipopionistic demonstrated as produced protection testistations. Bowings a staff periodicity of the state of the state

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (Men):

For general ambotic effects, early prescribing galadines recommend a dosage of 25-50 mg per week for 12 weeks. The usual dosage for physique or performance-entering purposes is in the range of 200-400 mg per week, taken in cycles 8 to 12 weeks in height. This level is in the range of 200-400 mg per week, taken in cycles 8 to 12 weeks in height. This level is in the sufficient for most users to notice measurable gains in learn mache mass and strength. Note that due to the fistancing rature of the phenylpropiorate ester, the weekly dosage is usually subdivided into 2 separate applications spaced evenly again.

#### Administration (Women):

For general native die ellers, other processing guidelines recommend a dosage (12% see ), and the processing processing a processing from the first processing processing processing the pr

#### Availability:

Nandrolone phenylpropionate has declined extensively as a pharmaceutical product. Given its short action, and the limited use of its longer-acting cousin randrolone decanoate in clinical medicine, there are very flow (if any) utique applications remaining for this drug. Thus, there is fittle justification for its continued production.

Brand name Durabolin appears to be unavailable in all markets worklwide.

A small number of generic and brand name products remain in less regulated markets (mainly in Asia), due to continued demand by athletes and bodybuilders.

Superanabolon from Spofa in the Czech Republic is also still in manufacture. It contains only 25 mg of steroid per 1 mL ampule, which makes it in relatively low demand among athletes.

Iran Hormone (Iran) makes a 25 mg/ml. generic nandrolone phenylpropionate in 1 ml. ampules. Counterfeits are not known to be a problem.

# Dynabol® (nandrolone cypionate)

Androgenic	37
Anabolic	125
Standard	Testosterone

Chemical Names 19-norandrost-4-en-3-one-17beta-ol

# 17beta-hydroxy-estr-4-en-3-one Estrogenic Activity low Progestational Activity moderate

# Description:

Nandrokere cypiotate is m injectable form of the anabolic stroid annabolice. This exter provides a patient of hormour relases withrail, deferral to that of testestence cypionate, with peak levels of drug being noted approximately 34-68 hours after administration, and a substantial hormour classes assistant of a board works. In this case the active hormous innulcodes, which is a moderately strong anabolic steroof that centres mild estrogonic and anabogaic properties. This product is essentially deferral in owner direct to Dearmuss, which tend to be accompaned by a low level of side effects. The core point of difference is that multibours exposited may appear to be a faster-sering compound to series users. Otherwise, there is no discernable difference between the two compounds, and materiolous cypionate may after decreases it without place of the contraction of the production of the potent and the decreases in vitability all cycles.

# History:

Nandrolence spinoute was first developed during the 1960%. It was sold for a heriftime as a lamma-use planmacifical, under such hand runnes as Anabo, Depo-Nortestiments, and Sterocemiols. Such preparations did not last, however, and in recent gazes the duglist here are madults due as a product of observiney medicine. The most notified to the contract of the contractive of the contractive of the deposition of the contractive of

The discortinated Jurus products were quickly transferred to SYD Group in Austrafa, assuring they would not be completely eliminated from commerce. They were subsequently enterodated to mustler in 2002, under the names Anabolic DN and Anabolic NA respectively. The new runners under loose reference to the former Jurus redundents, Reley in an effort to retain some of the original mustler for the products. SYD Group had also strandeded a high-down events of Anabolic DN directly to the Miscenn veterinary diagramated, but the product thesis sacre been withdrawn. This firs the product was discontinued Stowing U.S. DEA chaptes against the firm a haging that they were conspring to Ragbly expert Mexican steroids to the U.S. Today, the Available CN and Available NA products remain available on the Australian veterinary doing muted, although their control similar forms and the control of the Australian veterity older mustle, and the surface of the control of the most and the control of the Australian veterity doing muted, although their control has the control of the Australian veterity doing muted.

#### How Supplied:

Nandrolone cypionate is available on the Australian veterinary drug markets. It is supplied as 50 mg/mL of steroid dissolved in oil, in a 10 mL vial.

#### Structural Characteristics:

Numbrolesc episoates is a modified form of randrolous, where a carboopie, acid ester (cycleport/proprior) acid has been attacked to the 17-bets laplored gore, Esterilide steroids are less polar than fee attentis, and are absorbed more slowly from the area of irrepiction. Once in the bloodhearum, the ester is removed to yell free (active) numbroles. Esterfield steroids are designed to proleng the wardow of therepeate: effect following administration, allowing for a less-frequer injection scholds compared to njection of the administration, allowing for a less-frequer injection, and sostains a substantial release of 24-4 the bars following deep internancedar rigiction, and sostains a substantial release of homeone for approximately 2 weeks.

# Side Effects (Estrogenic):

Nandrolone has a low tendency for estrogen conversion, estimated to be only about 20% of that occur with test across with test across with test across with test across some test production to

um seen wan isomonitone— in as notame ware us awe can content mansume to estandio, in other mere active steen of steenid aromations asks adspose issue markohous is far less open to this process. <sup>60</sup> Consequently, estrogen-related side efficies are marks how cronew with this dup that with testestence. Elevated entopian levels my silb es noted with higher dosing, however, and my cause side effects such as acreased water restraints, host figure and suggestions. An artis-troups such as clerophere centure or transcription for girls and governors. An artis-troups such as clerophere centure or transcription are morntare inhibitor like Arimaticel's financional, which mere content or may alternately use an anomatice inhibitor like Arimaticel's financional, which mere content of the content of the processing is synthetic Arimatice financional, with an extra expensive content of the processing is synthetic Arimatice financion can be quite expensive.

It is of note that matchinous has some activity as a prospessin in the hody. "Modifyprosperstore is a 1-9 stered, nermoul of fits grups as 10 s-proragastorme creates a hormone with greater baring affisity for its corresponding receptor. Sharing this tast, many 19-rear anables storike are obsorts to horse our affinly for the prospectore receptor as well." He side effects associated with prospectore are similar to those of entropen, relating regardle bedoubt sibilation of selectionary powders on mammy tissue growth. These appears to be a strong system places to the contrastile leaves the growth. These appears to be a strong system places too formers bere, such there." The use of on anti-entropy, which sibilar the entropies component of this disorder, is often affected to midgue reconcents caused by varied/lose.

#### Side Effects (Androgenic):

Altough closified as an ambolic strootic, antioqueic side effects are still possible with this seststance, especially with higher doess. This may include bost of only skin, eare, and bodyfickall hair growth Arabolicianthogenic strootis may also aggroute male pattern hair body finell hair growth. Arabolicianthogenic sestorish. These may relade a desperaing of the vote, mentional insignatives, changes in skin texture, feed hair growth, and closed entiregreen. Numbroken is a settle with settlement of the control of the settlement o

Note that in androgen-responsive target fissess such as the skin, scalp, and prostate, the relative androgenity of maniforates is reduced by its reduction to dightornamicolore (DRN)<sub>2</sub><sup>20,20</sup> fibr. S-alpha reductase enzyme is responsible for this metabolism maniforate. The consumers use of a 5-alpha mediatase infilment such as finisative or distinctive will sincritic with six-specific reduction of manifoliore action, considerably receiving the tendency of numberolem to produce androgenic side effects. Reduction receiving the tendency of numberolem to produce androgenic side effects. Reduction

#### Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

## Side Effects (Cardiovascular):

Anabolicardnogenic storolis can have deletroiss effects on serum cholestered. This includes a tendency to refuse IEM, Good cholesteroi classes and increase IEM, Good cholesteroi values with may shift the IEM. to IEM challesteroi and attencion that fixous geneter risk of anteroclarosis. The relation terms of man anabolicardnogenic servicion of security has dependent on the dose, route of administration (road as, injectable), type of ascerd administration (road as, injectable), type of ascerd deministrating Good granulosis of content per over for 10 to work demonstrated a 25% reduction in IEM. cholesterol levels. This suppression is sightly greater than that reported with an equal dost of roststorouse certaint, and is augmented when dient statles showing a sightly stronger registre impact on IEM.IEM. Into with mardolous deconate as compared to istoricouse openious. We windrose injectables, lowers, choid all films as

significantly weaker impact on serum iputs thun c-17 alphu alkylated agents. Anabolic/androgenic steroids may also adversely affect blood pressure and trighcerides, reduce endothelial releasion, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and myocardial infarction.

To help reduce cardiovascular strain à is advised to muitain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4) grams per day) and a natural cholesterol/articoxidant formula such as Lipid Stabil or a product with comparable ingedients is also recommended.

#### Side Effects (Testosterone Suppression):

All antibolization/genesis steroids when taken in does sufficient to promote muscle gain are expected to suppress endergouses testistenies production. For size do comparion, statistical administração [10] mg per vocal of materiolate documente for 6 weeks live demensitated an reappress of the contraction consider [30] mg to a contraction reached [70]. See it is believed that the propositional tacher joy muscle contraction reached [70] see its believed that the propositional tacher joy muscle contraction of the suppression of institution of the structure of the suppression of institution of institution of institution of institution of institution of institution of institutions. Note that producing physypathological physypathological physypathological physypathological physypathological physypathological physical production of the contraction of the contracti

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (Men):

When used for physique- or preformance-mburging purposes, a dose of 200-000 mg per works in most comment, taken in cycles 8 to 21 works in right, This level is sufficient for most users to notice measurable gains in lean muscle mass and strength, which should be accompanied by a low level of estrogenic and androganic actively. Almost higher does (450-600 mg) may produce a stronger anabolic effect, given the relatively low concentration in a which this distinct a found (50 mg/ml.), daes show 4000 mg are not commody applied, Instead, the drug is often stacked with another agent, usually an arthogen such as an eigentable teasternor, which also belop to offer the very low level of antiogenicity, such as grateful for a strong the comment of the commen

#### Administration (Women):

When used for physique or performance-orinaring purposes, a dosage of 550 mpg or week is most common. Although only slightly and and progress, women are occasionally conflicted with vilization symptoms when taking this compount. Should virilize side effects become a concern, marshould explosing slowly be disconfined arrivated style to prevent their permanent appearance. After a sufficient period of withdrawal, the shortes-aring marshould represent the contraction of the shortest appearance. After a sufficient period of withdrawal fine fainteen the days show the form of the contraction of the shortest and the shortest and the shortest again and the shortest appearance of the shortest and the shortest appearance and the shortest appearance and the shortest and the shortest appearance and the shortest and the shortest and the shortest and the shortest appearance and the shortest and the shortest appearance and the shortest and the

# Availability:

The only remaining pure randrolone cypionate product is Anabolic DN from Australia, produced only in a 50 mg/mL concentration. It comes in the form of a 10 mL vial, which is contained in an orange tube.

# Dynabolon® (nandrolone undecanoate)

Androgenic	37
Anabolic	125
cond-d	T

Chemical Names	19-norandrost-4-en-3-one-17beta-o 17beta-hydroxy-estr-4-en-3-one
<b>Estrogenic Activit</b>	y lov
Progestational Ac	tivity moderate

# Description:

Standard

Nandroleus undecaroute is an injectable ferm of the unholes stroot mutuloute. The ester applied here is one croston auton longer fund encounted, and consequently form as very slightly lengue-lessing drug deposed at the site of injection. With proper attention paid to courtee concentration, whether and pharmonetises is would also good use length and the courtey concentration, which was the contraction and the contraction of the courtee of the counter of

restosterone

# History:

Nurshrokes undextones was developed during the 1960's, and was subsequently soid as Djunkthoin in light (Crimos) and France (Hermanc), and as Psycholobin in German (Thermon). The Inline product was moved to the new Farmissite label years liter, but returned the ceignal Djunkthoin trade turner. Djunkthoin was generally indicated for use in patient saffring from malaritime, catabolic assists, or recovering from major suppy. It was such as women and the clickly. Nurshrow temporary major supply to such as women and the clickly. Nurshrow temporary major such as well as such as women and the clickly. Nurshrow temporary and Djunkthoin from Farmissier in linky were discontined many years again and Djunkthoin from Farmissier in linky were discontined may years again and Djunkthoin from Farmissier in linky were discontined may be used you. In Djunkthoin from Farme faith followed before the clock of the 1998's. Plescettly, no beginner phenomenocial preparation and the contraction of the property of the contraction of the property and the contraction of the property of the contraction of the property property of the contraction of the property and the contraction of the property property of the contraction of the property property of the contraction of the property property and Djunkthoin the contraction of the property property of the contraction of the property property and the property property of the pro

#### How Supplied:

Nandrolone undecancate is no longer available as a prescription drug product. When manufactured, it was supplied at a concentration of 80.5 mg/ml. dissolved in oil and sealed in a 1 ml. armule. Each armule provided the equivalent of 50 mg of mandrolone base.

#### Structural Characteristics:

Nardolore unlecamonte is a medified form of mardolore, where a carboxyle, ead ester (unlecamonic acid) been attheid to the 17-best hydroxyl goal. Selected storation are less polar than few storaids, and are absorbed more slowly from the area of rigiction. Once in the bloodsteam, the easier is remored to peld few (acidiny mardonic. Estirified storaids are designed to probage the window of theraportic effect following administration, allowing first a less obsquare field in the control of the control of the control of the control of the first a less obsquare freight on the color of the control of the control of the Nardolore unlecamont is designed to provide a slow release of mardolone for up to 3 to 4 weeks following insection.

## Side Effects (Estrogenic):

Numbroke has a low tenders, for entrogen conversion, estimated to be only about 20% of that seen with testionness. <sup>60</sup>Th his is because while the few can convent multivolve to estandiol, in other trave active sites of steroid aroundations such as adjacen tissue muchoons in few sopen to this process. <sup>60</sup>Th Consequently, entityperithed side effects are a much lover concern with this drug flum with testostome. Elevated entropes tweel may still be noted with higher done; however, and may cause side effects which is increased under retentine, body fit gain and generoments. An unit-entropes such as clamping or claim showesthe to an amountum withdraw flum feature with the ment of the conloration of the contraction of the contrac controls estrogen by preventing its synthesis. Aromatase inhibitors can be quite expensive in comparison to anti-estrogens, however, and may also have negative effects on blood lipids. It is of note that rendrolone has some activity as a processin in the body <sup>493</sup> Although

progasterors is a 19 staroid, removal of this group as in 19-eneprogasterors creates a hormous with guest briding affishy for is corresponding recepts. Marring this trait, ramy 19-ear analotic staroids are shown to have some affishy for the progasterors excepts as weed. We have defended as societion with progasterors are similar to those of estingary, including regative feedback inhibition of lestosterors production and enhanced net of fit storage. Prospersis also surgerent the similarty celled of estingens on murmary tissue growth. These appears to be a strong synary between these two hormores here, such that groccomist in gride or our cur with the left of progastive, without causes destingulated. The use of an anti-estrogan, which inhibits the entropies component of this disorder, is often sufficient to pringing recomments caused by numbridors.

## Side Effects (Andmoenic):

Altough closified as an authoris stroot, antioquie siste effects are still possible with this solutione, expectably with ligher does. Them yielded bont of 60 ykis, next, and body ficial hair growth. Authorisant/oparis stroots may also aggranter male pattern hair loss. Women are warned of the potential wifring effects of antiolotalizations are included. These may related a despersing of the wore, mentional resignation, changes in sist nexture, feed hair growth, and closel entire growth. Numberdone is a stroot with relatively boundaries, earlier, definition to a is tone-balling action, making the travelal for strong testing and the stron

Note that is androgov-responsive target tissues such as the skin, scalp, and prostate, the relative androgovity of matholone is related by its reducted by a reducted to displacement/obse-(DRN) 69-500. The 5-alpha reductase enzyme is responsible for this metabolism of matholone. The concurrent use of a 5-alpha reductase inflibent such as financially disasteries will interfere with six-specific reduction of matholone action, considerable of administrative of the state of the relative state of the model with matholone if two arthrogovers's desired.

# Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

#### Side Effects (Cardiovascular):

Ambolicardrogenic streolic can have debetrious effects on serum cholesterol. This includes a tendency to reface IELL (ango) cholesterol ubases and irreases IELL (bad) cholesterol values, which may shift the IELL to IELL challence in a direction that fixes geneter risk of antirocherosis. The relative impact of an ambolicardrogenic servicion of serum lipits is dependant on the dose, route of administration (and is, injectable), type of steroid (arountable) or macroardiable), and level of resistance to hepsite methodism. Statistic contributions of the contribution of the contributio

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fits, cholesterol, and simple carribolydrates at all times during active AAS administration. Supplementing with fish oils (4 orans; nor day) and a matural cholesterollarity/sident formula such as Limid Stabil or a

# Side Effects (Testosterone Suppression):

All antibolic/indogenic steroids when taken in does sufficient to premote muscle gain are expected to suppress endogenous testinetone production. For sales of comparison, making expected to suppress endogenous testinetone production. For sales of comparison, making administrating 100 mg per wood, for fundroders decisioned to 6 weeks have demonstrated an image of the comparison of the sales of the making the engineering the comparison of the sales of th

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (Men):

Nardstone undeamonts was used clinically at a dose of 1 ampule every 1 to 2 weeks. A total of 3 to 6 ampulse were used for a given fewer perior of thempy. When used for physique - or performance-enhancing purposes, a dose of 3 to 4 ampulse (241.5 to 322mg) per work is must common taken in cycles 8 to 12 weeks in reglish like level is selficired for must users to notice massarable gains in lean muscle mass and strength, which should be accompanied by a lowel of estorgistic and analogues activity, light obese (404-500) in gar words) will impart a stronger ambolic effect, for can be difficult giren for relatively anger with other ambolic analogues activity for a stronger effect. Genes is properies, if seems to it well for both bulking and cutting purposes, and can reasonably replace Deculoration in an editory.

#### Administration (Women):

Nardsolves undecarente was used clinically at a dose of 1 ampule every 1 to 2 weeks. A total of 3 to 6 ampulse were useff or a given fewer level period of therapy. When used for physique- or performance-ordinaring purposes, a dosage of 1 ampule (80.5 mg every 10 days used most comment, which is taken for 4 for weeks. Afterpaid op digitally analoguest, which was the comment of the second of the contract of the solution of the solution and the solution acting matches Datasholite lingle the considered a safer (more controllable) option. This disposa south for only several dops, gualty changes for whitmouth and the option. This disposa south for only several dops, gualty changes for whitmouth and the

#### Availability:

Nandrolone undecanoate is no longer available as a prescription drug product. Some underground preparations are, however, known to exist.

# Equipoise® (boldenone undecylenate)

Androgenic	50
Anabolic	100
Standard	Testosterone

Chemical Names 1,4-androstadiene-3-one,17beta-ol 1-dehydrotestosterone

# Description:

Boldernee undecleture is an igicutibe veteriusy steorid that cellubits strong antoloic and monetanely admongstre properties. The undergatement care crossets the activity of the drug goalty (the stude)clemate ester is only one carbon atom larger than documents, so that and androgatic proposers of this drug are greatly appreciated by whites, who guestelly consider it to be a stronger, slightly more androgatic, alternative to Dece-Duralodriel. It is generally change, and could ruplesc Dece in most cycles without grouply changing the end coals. Boldernee undeylentanel as the commonly laroum in a drug graphet of necessing rand changes care that the control of the

# History:

Cha reportedly patented boldenore as a synthetic ambolic steroid in 1949. Daring the 1950's and 60's, the firm developed several experimental esters of the drug, and would later release a long-string from firth agent (releaf) in the firm of boldenore undesplement. It would be sold under the brand name Parentaled, which likely referred to its characteristics as a parenteral festeroble jumbole agent Parentaled sess ownse (citical used unity for the 1950's and only 70's, mainly as a lean-stease-preserving ambole agent in cases of wasting, and fire the returned robe must with outset possible state of the case of wasting, and fire the returned robe must see that the state of the parent fire the state than the parent fire the pa

#### How Supplied:

Boldenone undecylenate is widely available in veterinary drug markets. Composition and dosage may vary by country and manufacturer; the majority of products are supplied as multi-dose glass vials containing an oily solution; usually carrying 25 mg/ml or 50 mg/ml of steroid.

# Structural Characteristics:

Boldemore is a medified ferm of testosterore. It differs by the introduction of a double boal between carbons 1 and 2, which reduces is relative entergointly and antiquency fragingence oversize boldemore medified with the addition of enthosyle and ester testing of the contract of the c

It is interesting to note that structurally boldenone and methandrosterolone (Diarubol) are almost identical. In the case of boldenone (as applied here), the compound uses a 17-beta sester (undee/strate) to facilitate administration, while methandrosterolone accomplishes this with the use of a 17-alpha alkyl group. Aside from this, the molecules are the same. Of course they act quite differently in the body, which goes to show that the 17-methylation affects more than just the oral efficacy of an anabolic/androgenic steroid.

# Side Effects (Estrogenic):

Bolderore is aromaticed in the body to estraided (entropen). Elevated entropen levels can case side efficts such as increased water reterriors, body fair gain, and generoussite. Bolderore is considered a mildly entropenic servoid. Aromatonion studies suggest that its nine of convension to estandial is reagily built that of testestoresis? Will be tenderory to develop noticeable estroperic side effects with bolderore should be slightly layer than manufactors, but much lower than with solutioners. Entroperic side efficies are usually not promounced unless this drug is taken in doses above 200-460 mg per word. An artisiste effects, doubt flowly between present One my admerable year amortante inhibits the Artisis-feet (sunstrousle), although it is considerably more expensive, and may negatively after blood lipids.

## Side Effects (Andmoenic):

Altrugh classified as an ambolic steroid, antrogenic side effects are still common with this substance, especially with higher does. This may include boars of only skin, near, and bodyficial hair growth. Antabideandrogenic steroids may also aggravate male pattern hair tooks. Women are also warmed of the potential visiting effects of ambolicandrogenic steroids. These may include a deepening of the voice, menetrual irregularities, changes in skin texture. See high records and of the potential visiting of the content. See high records and of the potential visiting of the content. See high records and of the potential visiting of the content. See high records and of the potential visiting of the potential visiting

Note that while boldenone does reduce to a more potent androgen (dihydroboldenone) via the 5-alpha reductase enzyme in androgen-responsive target tissues such as the skin, scalp, and prostate, is a failing to do so in the human body is externely low. <sup>250</sup> The relative androgenicity of boldenone is, therefore, not significantly affected by finisheride or directorists.

# Side Effects (Hepatotoxicity):

Boldenone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

#### Side Effects (Cardiovascular):

Anaboli-vindroganie steroids can hove deletrious effects on serum cholestere! This irelates a retriency to refeet PLIK (good) oblesteroi values and irecrates DL (bad) oblesteroi values, which may shift the iPDL to LDL balance in a direction that fivors geneter risk of anteriosciencis. Bre relative impact of an ambiolizardinograte is virul on serum liptic is dependant on the dose, route of administration (end vs. ipsectable), type of steroid (incontratible or non-normatible), had twel of resistance to haptace intelligent Anabolizardinograte steroids may also adversely affect blood pressure and trighçeriles, routes each deliverile relation and support let verticated preprinty, all potentially increasing the risk of conditionaction deuse and injocential infanction. Bulkerons is likely to have a list, infantial report or cardionaction for the factor than reports conditionally in the conditional conditional conditions.

To help reduce cardiovascular strain à is advised to maintain an active cardiovascular exercise program and mismine the intaine of santaned fist, cholssterol, and simple cardiohydrates at all times during active AAS administration. Supplementing with shi old, games per day) and a natural cholssterolamioxidant formula such as Lipid Stabil or a product with compramble inserdents is also recommends.

# Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainting substances, testosterone should return to normal within 1-4 months of drug secession. Note that prolonged hypogonadotrophic hypogonadism can

develop secondary to steroid abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

# Administration (Men):

Although it stays active for a much longer time, holdenne undecylentate is injected at least weekly for phospine-or performance-relaturing purposes. It is most commonly used at a desage of 200-400 ng (4-5 m/s/5) mg version) per week. The dossage schodule can be distanted middled not dende the volume of each injection if recessively perhaps administering the drug two to three times per week. One should also take caution to rotate injection recently, so as to sould instantion or infection.

Not a regist mus ballet; bolkence urdec/sente instead provides a dow het steady gipt of strength and quilty mack muss. The posite efficts of this drug become most against when it is used for longer cycles, usually lasting 8 weeks or more in duration. The muscle gained should also not be the smooth bulk associated with sensiterone, but more defined and solid. Stree water blost is not contributing groutly to the dismetter of the muscle, more of the visible, taggeting on an cycle frobletioner undeclyment soluble for estated after the drug

Bolderone undecylenate is a very versatile drug, and can be combined with a number of other agents depending on the desered result. For muse, it is commonly stacked with an signetable testorisone such an eurnitude or opportunt. This should produce strong gains in macie size and strength, without the same intensity of side effects of using testosierone (at a higher doos) lane. During a curing place, muscle batherous and denny can be guidled and better the state of the side of the si

## Administration (Women):

When used for physique- or performance-enhancing purposes, women take much lower doses of boldenone undecylerate than men, typically 50-75 mg per week. Women should take caution with the slow-acting characteristics of this preparation, which make blood levels difficult to control and slow to decline should visitization sometoms become present

#### Availability:

Boldenoe undecylerate remins widely available as a veteriarry drug product. It is produced minis by the Americas, consistently at a dosage of 25 mg/mt. of 90 mg/mt. A small number of preparations are made at a higher dosage (typically 200 mg/mt.), minis by companies in less regulated markets of Asia where supply is often distated by bale market demand. In reviewing some of the more popular products and changes on the global pharmaceutical market, we have made the following deservations.

Brand name Equipoise is no longer available in Canada. This product was formerly manufactured by Wyeth Animal Health, which following a series of mergers is now part of Pfter. The Equipoise product was never manufactured under the Pfter label.

Equipoise® is produced in the United States by the Fort Dodge Company. The Fort Dodge products are sold readily in Mexico, and afterwards smuggled back into the U.S. Legimute vials are made of clear glass, and carry a label with a shiny metallic surface on the under side. Fakes are very abundant of this item.

Ganabol, which is produced in a number of South American countries, is still a popular brand in international commerce. It is seen in two strengths (25 mg/mL and 50 mg/mL) and in five sizes (10, 50, 100, 25, and 500 mL). There have been numerous fakes of this product in the past, so be careful when shopping.

The Legacy brand name product from Tecnoquimicas in Argentina seems to be reaching the LLS as of late at least in small volume. This product carries 50 moint of stemid in a 50 ml.

vial. At this time the Legacy product is very low on the radar, and probably can be trusted when located.

The brands Boldenona and Boldegan from Gen-Far are also popular in South America. These are low-dose (50 mg/mL) preparations. Like Garabol, they come in a variety of vial sizes. Counterfels do not armeer to be a bic issue at this time.

# Finaiet (trenbolone acetate)

Chemical Name	17beta-Hydroxyestra-4,9,11-trien-3-one
Standard	Nandrolone acetate
Anabolic	500
Androgenic	500

# Description:

Estrogenic Activity

**Progestational Activity** 

Terebone acetate is an injectable (generally) authols stored derived from marchotes. Its activity is quite removed from its structural purent, however, such that direct comparisons between the two are difficult. Terebohere is a non-entergonic steroid, and is considerably more analotic and analogonic than rundrotes on a miligarm basis. In appearance, it is much more commody compared to a stronger androgen such as decistancle, the is in nutratione. It is also continued to display bout free tries mere authorize potency than institutioner, miking it one of the strongest injectable analodic storoids core commercial manafactured, manage afficults, this storoids between the storoids and for the strongest injectable analodic storoids core commercial manafactured, manage afficults, this storoids of the strongest injectable analodic storoids core commercial manafactured, manage afficults, the storoids of the strongest injectable analodic storoids and the strongest injectable analodic storoids analogical storoids and the strongest injectable analodic storoids analogical storoids and the strongest injectable analodic storoids analogical storoids analogi

none

moderate

#### History:

Tuerboine acetate was first closely studied in 1967, described during a series of experiments into spitche ambolis storoids by Rossael-UCLAE<sup>28</sup>. By the early 1970s, tuerboines acetate was being soil in England by Hoechel as Finişet, and in Finner as Finişet by Rossael CALAE<sup>28</sup>. Rossael AG in Commyon was parent to both companis. Tuerboine acetate is a ding of voteriumy medicine, although a longer-acting extent of tuerboine (see: Parkhodin) was one soil for human conserption in soil. Errobion acetate is used, almost excludely, to incuse the inte of veedig gain and improve feed efficiency of earlier insense of the interpretate of the interpretation and interpretation across the interpretation and interpretation and the interpretation of studies, with no witholding point. Meat products soil in many areas of the world will office contain small amounts of resolutal protione metables as a result offits portice.

Tierbohne scenter first became popular among U.S. holophadkers during the 1980%, a time when the day was being samggled in from largor in lagh volume. It was identified (right) so) as a powerful amabolic and anthoganic agest, and quickly became a drug of choice armog/American competities bolyhulatkers. Albuogal exementy bol for a larier planel of time, the supply of rendrobene acetate ended albrughy in 1987, as Hescher-Bossel decided to voluntaryly discontrines sole of all injectable forms of the medication. Albrughy unconfirmed, the growing public concern about sports during likely had much to do with fits decision, as the discontraints of "Controversid" stretoids was very coronan during the late 1980's and early 1990's. This event murked the end of legistrate medicines containing tetrobolice acetate for rigistrat.

Around the same time as we were seeing the demise of Finajet and Finajet, Hoechst-Roussell was introducing trenbolone acetate to the U.S. market as Finaplic cattle implant politis. This came subsequent to the FDA's approad fir such products in 1987. The politis were designed for substanceous implantines into the our of the caller what a handled implant gat, and are fir too large to be implanted in harmen without mires surgery. Remutably, thresholder eacities peless are courself from U.S. controlled substance loss. This is persumibly to mile it easy and affordable for fessional outside activate loss as the good of promoting gates. If a victorization was recorded every time free products were to be growth promoting gates. If a victorization was recorded every time free products were to be products come in the form of pelest, they are not in a firm sainble for harmer consumption exist, realize the exerction seems life town reasonable than a first glance.

Hamma administration of Frapples pellets can be difficult to accomplish, but it is still widely doze. Most commends, two to first implaint pellets are greated up and mixed with a 50-50 water and 2005 0 soliton, which is applied to the side indig. The lame new-town transforms water and 2005 0 soliton, which is applied to the side indig. The lame new-town transforms of the side of the side

More adventures individual tour mode à a positive to mir their our signicien with Fragint. The poles are greated into a les proude reasting supporter form 2 no polestul, and then are added to starie water, propless glock or an oil-tousel signified sourcid or voteringy vintuit. This is usually repeated here workel, attantys once for marge to undertake this practice more frequently. Since this is not being does in a controlled starie construct. However, core is obviously riching infection or worsely ploying this. Starting in the late 1990s, some stores began selling list flat control all the necessary impedients to separate the brinders form the active stored and here an elatisely pericetable. These iss laws grown in popularity over the years, and are usually reviewed fivorably, although are not considered as subtined for steller planners.

Fragilité à presenty available à the LLS, and some metters abroud, although it is now being sold by harvers instant of Horsche Rousel Agristor. But product cross in our forms, Fragilic-H and Fragilic-S, which downess if the product vous is not form. Fragilic-H and Fragilic-S, which downess if the product was distinct, with the "FI version containing 100-20 mg terolotice accetate pelotes (2,000 mg) and the "S' version only 70 (1,400 mg). by damin Helschh (2.3) has invended to two compreting products of equipaled makeup as well, sold as Componert-HI and Componert-TS. There are also the Resulte and Sponover brands fait contain terolotices accetate with an added (sous) unswarded) does of estingar. Additionally, although no other legitimate medicines containing metodates understood affect of impactions (or impaction and or in also place) as a name for citizen and understood affect of impactions.

#### How Supplied:

Frenbolone acetate is available in select veterinary drug markets. It generally comes in the form of implant pellets containing 20 mg of trenbolone acetate each. Injectable preparations containing 20 mg/ml of steroid in oil were formerly sold.

# Structural Characteristics:

Terchone is a medical ferm of matholone. It diffuses by the introduction of double broads act acrons 9 and 11, which it this committation (0+on), increase antiogen-briding affinity, <sup>2023</sup> and slow its metabolism. The resulting sectoral is significantly more potent as both an ambolic and an anticopart than is matholone base. Tercholone acetate contains turcholone medified with the addition of earboxyle acid ester (acetic acid) at the 17-beta phicoryl group, so that the 6se section it released more also dy from the acut of rejection that the other social results of the first actual results actual results of the first actual results actual resu

#### Side Effects (Estrogenic):

Trenbolone is not aromatized by the body, and is not measurably estrogenic. It is of note, however, that this steroid displays significant binding affinity for the progesterone receptor (slightly stronger than progesterone itself). 505–506 The side effects associated with

programme are smure to mose of eatingst, nexturing negative necessaries and surtered to the state of the street, Programs also surprises also suggested the estimation yellow of extreet or nameny tisse growth. There appears to be a strong synarpy between fines to the others also that any generous limit and the end of the street, and the street of th

#### Side Effects (Androgenic):

Albugh classifed as an arabolic stroid, treobotes is afficiently ambagain; Antosqueis sist effects and still common with this soltance, and may include bost of oly slist, next, and bodylickail har growth. Antolocianthoganic stroids may also aggravate mile pattern his loss. Women are also warmed of the potential wirking effects of ambolicanthoganic stroids. These may include a deepening of the voice, mensitual reagainties, changes in skin extent, field har growth, and closed adequarent. Additionally, the 5-splain reductase cumput does not metabolic tereboline, <sup>607</sup> so its relative antrogenicity is not affected by financial for a financial finan

#### Side Effects (Hepatotoxicity):

Terrbolene is not c-17 alpha allystated, and is generally not considered a hepatotoxic steroid; liver toxicity is utilikely. This steroid does have a strong level of resistance to hepatic breakdown, however, and severe liver toxicity has been noted in bodybailders abusing trerbolone. <sup>500</sup> Although utilikely, hepatotoxicity cannot be completely excluded, especially with high dosses.

#### Side Effects (Cardiovascular):

Anabolizandroganis steroids can hove deletroise effects on serum cholestered. This includes a technely to refuel EML (good) cholestered values and increase DL (fined) cholestered values, which may shift the HDL to LDL balance in a direction that fivors geneter risk of antiroisclerosis. The relative impact of an anabolizandroganic storoid on serum lipits is dependant on the dose, route of administration (end w. injectable), type of steroid (comutable) error anomatolishe), and the off resistance to hepital (mentables) are not some anomatolishe instant end storog resistance to metabolism the roles to non-anomatolishe matter and storog resistance to metabolism, the roles of the storog (regative) impact on flyd tables and disreganistic skin, order endothellar levelstorin und some deservely differ blood pressure and implements, routher endothellar levelstorin, and appropriate pressing the role of cardiovascular storogeneous and traversitied inferiors.

To help reduce cardiovascular strain à is advised to mairian an active cardiovascular seurcise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplamenting with fish oils (4 games per day) and a natural cholesterolamicoxidant formula such as Lipid Stabil or a product with comranable inserdences is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolicandrogenic steroids when taken in dones afficient to promote muscle gain are expected to suppose ordogenous testorouse production. When the intervention of testosteron-strutularing substances, testosterone levels should return to rormal within 1-d mention of dings escession. Now that produced phosperadorepic hypogenation commission of the produced p

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (Men):

Trenbolone acetate was never approved for use in humans. Prescribing guidelines are unavailable. An effective dosage for physique- or performance-enhancing purposes generally

falls in the range of 100-300 mg per week, taken for 6 to 8 weeks. Due to the short-acting manuscript of a content except, the tower's dosage is admisted into 2.5 smaller applications. Effective can also see so that fails in the range of 100-200 mg per day, taken from longer than 6.5 weeks to minimize any potential lequitie stims. The level is sufficient to rotice strong receivances in stronger and learn fasour men, with a low level of numerical sele delects. Lack of the receivant learner and the learner and the level for the stronger selection of the stronger and the learner and the level for the level for the stronger and the learner and the level for the learner and the lear

While is a noteworthy landoning agent, this is not the only hereif of tresholous accesse. It is also a strong anniholous, with muscle-basility properties often compared to instantoure and Danaholo ha without the same level of water reterrant. This may be a little agencius of a description, as it selds of enganging activity does seen to last fail agent it is a difficient to the contraction of the contraction o

For stacking, trebolome is a very versalle steroid, and seems to work exceptionally well with other against for bindinged actioning purposes. For catting hopkidates often stack, it with a raid anabele like Westrollé or Pitrobolomik. Widnot cette water beneamly our sixte, the raise of laids a very sold, weld-fielded hathress to the physique. For learn most again, the result of the contract of the property of the

# Administration (Women):

Trenbolone acetate was never approved for use in humans. Prescribing gaidelines are unavailable. This agent is not recommended for women for physique- or performanceerhancing purposes due to strong androgenic nature and tendency to produce virilizing side offices.

## Availability:

Pharmaceutical preparations containing trenbolone acetate remain scarce. The bulk of the supply for this compound comes from underground steroid manufacturers. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Therbotne acetate is still produced in the United States, sold in the form of cattle implant pollets (mainly Finaplet®). This agricultural product is also exported to some neighboring markets, through is not widely popular outside of the Americas. These pellets are difficult to administer, and are not commonly the subject of counterfeiting. This is the only legitimate form of treathout accetate in the U.S.

Most other forms are from underground manufacturers, and therefore are of unverifiable quality.

# Halotestin® (fluoxymesterone)

Committee I	the state to recommend the state of
Anabolic	1,900
Androgenic	850

Chemical Names		b-dihydroxy-17a-methyl-4-androsten-3-one fluoro-11b-hydroxy-17a-methyltestosterone
Estrogenic Activity		none
Progestational Act	vity	no data available (low)

# Description:

Flavoymerone is an oul analysis steroid devised from testosterone. More specifically, it is an orthyletosterone derivative, differing by the addition of 11-bets plavoy and 9-dipheture or goups. The result is a potent orally active mos-anomatosite steroid that exhibits externely storoug archingerine proprises. Flavoymenterone is considerably more analogusif tuna testisteriora, while at the same time the analysis effects of this agent are considered to be moderni in composition. This misches florous entere a general good may be a more free analysis of the same analysis of

# History:

Flacoursectors was find described in 1956;50° It turns assigned that came year, and about to pussess approximately 20 ters for much potenty of methylacoursectors. (6) netheries arealook effect in harmer sould not be quite as strong in comparison. It was stronked to the U.S. prescription dang mented shortly after under the brand mant Heishestis (Hjojshu), and soon after that as Ulandron (Chai). The ding was rishtly described as halogenized described of heishestic effections except societies and the strong societies and and and posing protecting of the contraction of learn tissue repeat and growth following such conditions as burns, delayed healing of finductors, derived in the strong of the strong contractions and the strong of the strong contractions and the strong of the str

By the mid-1907s, the FDA had been gunted much more control over the U.S. drug mided. One of the first major changes with storid mediciac error when the FDA required strong substantiation for each potential use of a drug. The prescribing guidelines for flavogrenationes were soon refued to state that the drug was "efficients" for training variess forms of arthogon declicacy; in mides, and reducing the security of postparatan breast pain and tasting andranger-responsites reportable breast career in families. It was also listed as "pochably effective" in tenting posttamorpassal osteoprosis. Current practing guidelines for flavogrenators list only the use of tenting authogus deficiency is mides and breast

In recert years, flaxymesterone has become viewed more and more as a controversal medication in the syst of most chickins. It is hepothetoxicy and potential negative inpact of lights and cardiovascular risk detoxs are often cled as reasons for noting the use of this agent in observise healthy males for tential gardnegon is stifficiency. Delay testoreous preparations fajections, gols, patches, implants, etc.) are preferred for this purpose, and they supplement the same analogous rissing from the body (testoreous, DEIT), not more toosynthetic destrations. Flaxymesterone remains for sals in the U.S. six a generic drug only, it remains available not be infrast output of the delay the contrained to the contraction of the contraction of the contraction.

#### How Supplied:

Fluorymesterone is available in select human drug markets. Composition and dosage may vary by country and manufacturer, although generally contain 2mg, 2.5 mg, 5 mg, or 10 mg ner tablet.

## Structural Characteristics:

Fluoxymesterone is a modified form of testosterone. It differs by 1) the addition of a methyl group at carbon 17-alpha, which helps protect the hormone during oral administration, 2) the

ntroduction of a more group at carbon 9 (apria) and 3) the attachment of a hydroxyn group at carbon 11 (beta), which inhibits steroid aromatization. The latter two modifications also greatly enhance the androgenic and relative biological activity of the steroid over 17-alpha methylicstosterone.

# Side Effects (Estrogenic):

Flaxopmesterone is not anomatized by the body, and is not measurably estrogenic. An artistrogen is not necessary when using his stendi, as genomental sloud in the he a concern even among sensitive individuals. Since estrogen is the usual culprit with water returnion, this stendi a stead produces a lawn, quality look to the physique with no fier of excess subcutamous fluid reterrition. This makes it a fivorable steroid to use during cutting cycles, when water and filt reterrition are minor concerns.

# Side Effects (Androgenic):

Flaxopmenterone is classified as an arthogon, Androganic side effects are common with this solutance, and may related bosts of obj skin, zero, and holyfrieth lair growth. Antaboliciandroganic steroids may also aggreates male pattern hair loss. Those genetically proce to make pattern hair loss may wish to opt for a milket, less androganic, antabolos, steroid. As a potent androgany, this steroid may also increase aggressioness. Winema are additionally warned of the potential villening effects of antaboliciandroganic servoids. These may include a deepering of the voice, menstraal irregularities, changes in skin texture, facial hair growth, and clonel enlargement.

Flavoymesterone appears to be a good substrate for the 5-alphn reductase enzyme. This is evidenced by the fact that a large number of 5s metabolites are found to be 5-alphn tendenced and copies, <sup>314</sup> which coupled with is control and engine intention, suggests that this steroid is control to the steroid of the steroid of and tendence responsible target times such as the flavormesterone by the concurrent use of intentive or disasterists.

It is also of note that flavoymesterore has been shown to possess usual anthogonic properties. In human taskes published back in 1604, the storal displeyal a much stronger tendency to promote phalic enlargement compared to other anthogonic effects such as lair growth, Bioka, and Changas in vocal picks, "Paloopursterore was offering a somewhile different anthogonic profile compared to instinction," and as such discussment of their distribution prosphile, as some local to a studiely lair diregle effect within the board enlargesy of anthogonic prosphile, as some local complete blogical profile of the without the board enlargesy of anthogonic action. Flowymesteron remains considered an anthogon, but studies like the above suggest durit my not offer as complicia blogical resistance where anthogonicity is durit in my not offer a complicit blogical resistance where anthogonicity is

## Side Effects (Hepatotoxicity):

Placopreseivore is a CF-alpha adylated compound. This alteration protects the drug from descritation by the five, allowing a very high percentage of the dung entry into the bloodstream following ornal admirations. CF-alpha adjusted antibodicationsposes stereds blooding ornal admirations. CF-alpha adjusted antibodicationsposes stereds blooding ornal admirations. CF-alpha adjusted antibodication producedly a frequency of the control of the

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

# Side Effects (Cardiovascular):

Anabolic/androgenic steroids can have deleterious effects on serum cholesterol This includes a tendency to reduce HDL (good) cholesterol values and increase LDL (bod) cholesterol values, which may shift the HDL to LDL behave in a direction that flavors greater risk of arterioselerosis. The relative impact of an anabolic/androgenic steroid on serum lipids is denorabate on the done must of administration (end by a indextably three of steroid committable or non-aromaticable), and level of resistance to hepatic metabolism. Flaxoymesterone has a strong effect on the hepatic management of cholesterol due to its stantural resistance to hee breakdown and roate of admiration Anthodocandogenismous for the contract of the contraction and the contraction of the c

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fists, cholesterol, and simple carbohydrates at a litrus during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterolamioxidant formals such as Lipid Stabil or a product with comprabile injurefiers is also recommended.

# Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-standing substances, testosterone levels should return to normal within 1-4 morths of drug secession. Note that prolonged hypogoradotrophic hypogoradotrophic hypogoradotrophic hypogoradotrophic and when severotrophic up served in the procession of the pro

Sudies admissioning 10 mg, 20 mg, or 30 mg of flaxoymesterone to nine healthy male subjects for up to 12 weeks have demonstrated the strong superssion of endogenous testosterone levels, with inconsistent effects on pure admerstrated processes. Although not fully understood, flaxoymesterone is proposamenterone in proposamenterone in proposamenterone in the strong strongly and the strongly according to the strongly and the advantage of the strongly and the strong

# Administration (General):

effects, see the Steroid Side Effects section of this book.

Studies have shown that taking an oral anabolic steroid with flood may decrease its biovasiability.<sup>35</sup> This is caused by the far-soluble nature of steroid hormones, which can allow some of the day to dissolve with undipsted distary far reducing its absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty stormeth

#### Administration (Men):

To test anthogan isatificacy, carby prescribing gaidelines for flavoymesterone called for a dose of 2-10 mag per day Modern prescribing gaidelines affor a didy doseg of 5-20 mg. Therapy is samply rinited at the fall 20 mg dosage, which is her adjusted downsend to meet the rinkfishall meted of the patient. The flavoy most the contrained long-term usides laboratory tests (fight, her engoynes, etc.) or side effects contraindates in contrand together unisses in property of the property of the

Flavoymesterone is comruely used by athlese in weight-entired sports like westfag, powerlifting, and board, due to the first that strength gained from the day is usually not accompanied by gent incrueses in bodyweight. When properly used, it can allow a correlation to say within a specified weight mange, yet distactionly proved he performance. Flavoymesterone is also commonly used for bodybailing contest preparation. When the competitor has an acceptable by body day processing, the strong anticipan beef in absence that the same properties of the strong processing the strong processing the strong the shade in anticipant content of the same processing the strong and the body may be more included to hum off cosess fit and prevent new fit storage. The "tankening" effect of flavoymesterone would, therefore, be somewhat similar to that seen with treaching, allowed in will be without the same bred of messing the with treaching, allowed in white beat when the same bred of messing the with treaching, allowed in with the without the same bred of messing the same breathers.

In cutting phases, a milder anabolic such as Deca-Durabolin® or Equipoise® is commonly stacked with fluorymesterone, as they provide good anabolic effect without excessive estrogen buildup. Here, fluorymesterone provides a well-needed androgenic component, belying to promote a more sold and defined gain in mace mes, with his interference with energy and libbid, then might to obtained what primuly analosis ogerat dance. Perhaps Primobanife-Dapet would be an even better choice, as with such a confriation there is no baships of entirogen, and likewise even less worry of water and it restories. For miss, or might alternately use an injectible bestsderone. A ratio of 400 mg per week of reloasterone certaintie and 2-50 mg daily of flavorymesterone, for courtiple, deltay provides exceptional accuracies in strength and learn mace farmes. A more significant bred of analoguesis self-defition of the contraction of the contraction

## Administration (Women):

Flaxymesterone is most often used as a secondary medication during insperable and regarsensible breast cancer, when other therapies two fields to produce a desibilitie offert. The dosage used for this application is 10–40 mg per day. Virliang effects are common at doses of only 10–15 mg per day in these partiers. Flaxymesterone is not recommended for woman for physique- or performance-orbanicing purposes due to its strong androgenic rature and tendency to produce virliang side effects.

#### Availability:

Pharmaceutical preparations containing flaxopresserone remain scarce. The drug has largely associated with western medical markets, where it has been falling out of fivor for most clinical application. The bulk of the supply presently comes from underground steroid manufacturers and export-only products. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have much the following observations.

Currently, the most popular item found on the black market is the Stenox brand from Mexico, which is sold in boxes of 20 tablets. Note that the dosage of these tablets is only 2.5 mg.

Fluoromesterone remains available in the U.S. as a centric drup by U.St. Plumma It is

Balkan Pharmaceuticals (Moldova) makes the product Halotest. It is prepared in 2 mg, 5 mg, and 10 mg tablets, with 20 tablets contained in each foil and plastic strip.

## Laurabolin® (nandrolone laurate)

available in a 10 mg tablet only

Androgenic	37
Anabolic	125
Standard	Testosterone

Chemical Names	19-norandrost-4-en-3-one-17bet- 17beta-hydroxy-estr-4-en-3-	
<b>Estrogenic Activit</b>	у	lov
Progestational Ac	tivity moder	rat

# Description:

Nardstoke lastate is an injectable form of the authols stored markolone. The lastate ester applied there is two comba attents langer than dearmost, and conceptently this agare from a slightly leaves being due glospot around the areas of rispection than Deco-Darabolio. Given its strong delipoly-othese properties, it is possible to administer memforches harine one every three to four weeks in a metical setting. As a mandrobne injectable, this due growthes an amoderately strong emploide effect, which is accompanied by a low level of estinguister and analogusity properties. Although not widely used, mandrobne lastate is titored by affalses and bodylotkales. Set doubly not provide when adapting the mans with marinal data objections.

# History:

Numbrokes hunter was developed during the 1960's, a time when muny new multiclose cents were being orbinsied and mentiografts. This long-acting seat formshows is usually identified as a vicenizary daig, but was actually proceeded to harmen before it was adopted time of the contract of the contract of the contract of the 1960's. Nucleaches harmen were timendy when the data is harmen medication, however, and from this point on would be used exclassibly in vestimany properations. These is noting particular their mates the dug poorly saded for harmen use, and he discontinuous probably had much more to do with the same properation. These the data of the 1960's had not the proper and the same properation of the same processing of more than the same and the same processing of the same parties of the clue self-

As a sectionary drug, methodoso lurant is most commonly identified with the Landsolm bared name. Lannsholm is manufactured by Henoxet, and is found in a naving of coursine including Mexico, Chile. The Nordenduck, Australia, and Colombia. It is used with cut, doug, honces, join and cuttle, principal to ordise multarition caused by what or prassine illness, to tent amenia, contart the catabole effects of corticosteroids, and to improve the contemporary of the control of highly active or clerkly ammin. The Lannsholm bared has sold been sold at one time by Weitff-Chemic in Austria and Westie in Chemisph, however those bared of the control of the and manufactor (Pluds, Cammay). Forthol (Parlim, Mexico), and Lannshol 250 (Leeffer, Mexico), Teday, only the Intervet products are known to exist.

# How Supplied:

Nandrolone laurate is available in select veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 20 mg/ml. or 50 mg/ml. of steroid dissolved in oil.

#### Structural Characteristics:

Nardolouc hante is a modified form of murdolouc, where a carboxyle acid ester (harriacid) has been mattend to be 17-beta hydrody gene Esteridi esterios has este post britten fee sterois, and are absorbed more slowly from the area of spectru. Once in the bloodstream, the ester is removed to yield free faciely nurdolous. Esterified sterois are designed to prolong the window of therapeutic effect following administration, allowing for a loss frequent injection schedule corpured to spectrum of the contesting learned. Nurdolone hunter is designed to provide a slow release of nundrolone for up to 3 to 4 weeks following inclined.

#### Side Effects (Estrogenic):

Nandroleus has a low tendency for entrogen concession, estimated to be only about 20% of that seen with tenderceas<sup>100</sup> has is heaven which the leve on convent metallores to estandio, it other more active sites of storoid arountations such as adjove tissue metallores for the sope to this process. 20° Concequently, enterpresented asid exities are a much lower concern with this drug than with testostome. Becomed estropes levels may still be noticed with higher dosing, however, and may case said exited such as remaind water retardors, body fat gain; and genecomostic. An anti-estrogen such as climiphene existent or marrowing entered may be recovered to proceed entogens; side effects with a retreated controlled and the still and the still and the still all the still and the still all the still and controlled the still and the still all the companies in comparison to artificial controlled the still all th

It is of note that mandrokne has some activity as a progestin in the body.<sup>518</sup> Although progesterone is a c-19 steroit, removal of this group as in 19-reprogesterone creates a homeone with greater barding affility for its corresponding records Starring this trait, many 19-rest anabolic steroids are shown to have some affiling for the progesterone recorder as west. The table of these succession will progesterone are sentime to those of entiregen.

storage. Progestirs also augment the stimulatory effect of estrogers on mammary tissue growth. There appears to be a strong synergy between these two hormones here, such that gonecomstst might even occur with the help of progestirs, without excessive estrogan levels. The use of an artis-estroger, which inhibits the estrogenic component of this disorder, is often sufficient to misconder components caused by monthloom

# Side Effects (Androgenic):

Albeagh closified as an ambolis strool, androguic side effects are sill possible with fine stosture, expectable with light doors. This may takelle boats of 60 yls sin, eare, and body/brickel larle growth. Antabolican/looguic steroids may also aggraster male pattern lark body. The control of the potential wideing effects of antabolican/looguic stostolic. These may include a deepening of the soice, mentional irragalarities, changes in six incare, facili large growth, and closel outlengment. Naturalson as a stead with relatively looguid and the control of the state of the control of the

Note that is androgon-exponsive target fiscuss such as the day, such, and prostate, the relative androgonizety of mathroker is reduced by its reduction to disploramentohine (DHN)  $^{50}$   $^{51}$  The 5-aphra reductions engine is responsible for this metabolism of mathroker. The concurrent use of a 5-aphra reductions eithers such as finished to distinctive will interface with its se-specific reduction of mathroker action, considerably recorning the tentury of mathroker to produce analogue side effects bediense recording the contribution of the contribution

# Side Effects (Hepatotoxicity):

Nandrolone is not c-17 alpha alkylated, and not known to have hepatotoxic effects. Liver toxicity is unlikely.

# Side Effects (Cardiovascular):

Anabolizándogueis ateorás can hove deletrious effects on serum cholesterol. This includes a tendency to reface III. ((ago) cholestero) ulasas un irecurse ID. ((ba) cholestero) ulasa, with my shift the IIII. to IDL balance in a direction that fixors geneter risk of antriocitorists. In reducine irganic at manubolizandogueis extend on serum (pink) is dependant on the dose, route of administration (only s. ripectable), type of steorid (contratable or non-anomatibals), and level of resistance to leaper intendiorism Statista contraction in III. Calcinsol lecks. <sup>207</sup> This suppression is sightly general methodorism Statista contraction in III. Calcinsol lecks. <sup>207</sup> This suppression is sightly general methodorism contraction in III. Calcinsol lecks. <sup>207</sup> This suppression is sightly general methodorism contraction in III. Calcinsol lecks. <sup>207</sup> This suppression is sightly general methodorism contraction lecks. <sup>207</sup> This suppression is sightly general methodorism of the contraction of support left wortricate hypertrophy, all potentially recessing the risk of contrassociate disease and respectability contraction of the contr

To help reduce cardiovascular strain à is achiesel to maintain an active cardiovascular cueries program and minimize the intuke of saturated fits, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish ols (4 grams per day) and a natural cholesterolamioxidant formals such as Lipid Stabil or a product with comranable inserdents is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testostenous production. For sake of comparison, studies administering 100 mg per week of randrolone decanoate for 6 weeks have demonstrated an approximate 57% reduction in serum testostenous levels during therapy. At a dosage of 300 mg per week, this reduction reached 70%, <sup>524</sup> It is believed that the progestational activity of midrolone notably contributes to the suppression of testosterons synthesis during therapy, which can be marked in spite of a low tendency for estrogen conversion. <sup>525</sup> Without the intervention of testosterone-stimulating substances, testosterone levels should return to normal within 2-6 morths of drug secession. Note that prolonged hypogradiotopic hypogradiotopic to second one-object of the prolonged hypogradiotopic to stored abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects see the Steroid Side Effects section of this book

#### Administration (Men):

Nandrolose launte is not approved for use in human. Prescribing galdelines are unwaitable. When used for physicapes or performance-relamine purposes, a dose of 200-400 mg given every 7-10 days is most commen, taken in cycles 8 to 12 weeks in length. This lived is should be accompanied by a low level of recognite and anti-opped scribing filter doses (450-600 mg every 7-10 days) will impart a stronger analysis entirely. Higher doses (450-600 mg every 7-10 days) will impart a stronger analysis for the relatively low concentration filts strong in such distribution. In Instead, many opt to combate this agent with other anti-obscilant/regones storoids. Given its properties, it seems to fire with from the contraction of the stronger and current composition of the width of the width of the size of the width of the stronger and current composition of the width of the width of the size of the width of the size of the width of the size of the width of the width of the width of the size of the width of

# Administration (Women):

Nandroken launtie is not approved for use in human. Proceibing galdelies are unstable. When used for playage-or performance-deruning purposes, a desage 100 mg every 10-14 days is most common, taken for 4 to 6 weeks. Although only slighty antoquarie, womens are occasionally conforted with viliagines superious when thing this comproud. Shedd withing sale effects become a concern, the drug doubt be discontinual immediately to be present first permunent appearance. After a sufficient period of whitheast of the option. This drug stops are the conformal days, greatly nothering the withdrawal time if indicated.

## Availability:

Pharmaceutical preparations containing randrolone laurate remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

The Intervet brand name Laurabolin product is almost exclusively associated with this compound. It remains available in select markets of Europe and the America's, and is not widely diserted for black market sale.

# Masteron® (drostanolone propionate)

Androgenic	25-40
Anabolic	62-130
Standard	Testosterone

Chemical Names 2alpha-methyl-androstan-3-one-17beta-ol 2alpha-methyl-dihydrotestosterone

Estrogenic Activity	none
Progestational Activity	no data available (low)

#### Description:

Drostanolone propionate is an injectable anabolic steroid derived from dihydrotestosterone (DHT). Here, the DHT backbone has been modified with a 2-methyl group to increase its

ambole proporties, making this agant significantly more effective at promring the growth of much tissue than its non-methylated present. Dostanolone proporties it described in product literature as a "steroid with proverfial ambole," and anti-estrogacie proporties," and rished doss seem to a bisse owner of both properties. Admittally, browcey, is ambole properties are more properly described as mederate, especially when placed in the context of other agaret. The days must often used by design bodylothates and athletes in speed sports, where it is highly through for its ability to produce sold increases in learn market must arrestly melting a sound accomplished by placedation in body file led and marinial self-under a sound accomplished by placedation in body file led and marinial self-under a sound accomplished to placedation in body file led and marinial self-under a sound accomplished to place data of marinial self-under a sound accomplished to place data of marinial self-under a sound accomplished production in body file led and marinial self-under a sound accomplished production in body file led and marinial self-under a sound accomplished production in body file led and marinial self-under a sound accomplished production in body file led and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished production in body file and marinial self-under a sound accomplished and a sound a sound and a sound a sound a sound and a sound a s

# History:

Doutmebre projecture was first described in 1993. Systex developed the agent alregisde such order well-known sectoris as Annuled and methylentostatos (expectable, also first described in the same paper. Brostanokne projecture would be introduced as a prescription dang product approximately a docade her. Ellay had an agenement with Spette to spile certain research and development costs in exchange for the rights to make the reasons off that research. Ellay would, therefore, self-doutment form form of the random from the contract fill the contract of the contract fill the contract fi

The U.S. Food and Dug Administration approved doctaturables projecture for the treatment of admonds disperable breast carter in postatementage shown. Bits remained the principle clinical infactation for the agest in all ferenticual muteless as well. The proceeding learnest remaind doctors and fitting plantes that there is considerable to propose the contract of th

White highly peoplar among atthese during the 1970's and 297s, dostunobes projecture thirmshy eiged affined sexenes as a preceiption agest. Manufactures began voltarrily discordinaria gals of the agest in various murdets before long, fisely due to the advert of more effective therapies for breast caree, as well as these double is served processives for this place of twentures. Once of the first preparations to go was the U.S. Doshan, which was nermored form marked during the last 1990's. Permistral and Meetorma were soon dropped as well. The last remarking Western preparation containing dostunobes proposate was Masternes from Regians, which disoppeded by the last 1993's. Dostunobes projectures remains fised on the U.S. Plantmosquis, suggesting there is preceipt on legal collection.

#### How Supplied:

Drostanolone propionate is no longer available as a prescription drug preparation. When produced, it was supplied in the form of 1 mL and 2 mL ampules and 10 mL vials containing 50 me/ml of 100 me/ml of steroid in oil.

#### Structural Characteristics:

Dostanskoe (also kawas as domostanskoe) is a modified form of displostostostorce. It differs by the instolation of a melty gapp at eurlow-2 (algable, which considerable accuses the ambolic strength of the sterial by legistering is restaure to metabolism by propriets as a modified from of drostanskoe, where a carboyle, seed seed for (projective as a modified from of drostanskoe, where a carboyle, seed seed for (projective acid) has been attached to the 17-best Inplottopl group. Estatified strends are less point than the strends are also absorbed more about 90 mel near not a forection. Once in the bloodstream the centre is nerviced to yeld for (acid-why from the sum of species. Once in the bloodstream the centre is nerviced to yeld for (acid-why from the sum of species. Once in the bloodstream the centre is nerviced to yeld for (acid-why from the sum of species.) Esserthid streads are bloodstream the centre is nerviced to yeld for (acid-why from the sum of species.)

life of drostanolone propionate is approximately two days after injection.

# Side Effects (Estrogenic):

Dostanches is not aromatical by the body, and is not measurably estrogair. An artiestrogain is not receively when using this strond, as generous side solad rule be a concess even among sensitive individuals. Since estrogain is the usual calpit with water reterion, dorstanches restand produces a lean quality look to the physics with no fare of excess selectaments that reterior in the misses it a foreable setteral to use during carriag cycles, when water and if reterior are major concern. As a non-aromatishe DHT derivative, doubtambone may impart an arti-strongenic effect, the drug competing with other (contrables) substantes for hinking to the aromatises enzorm.

# Side Effects (Androgenic):

Albudgh classified as an ambolic stroid, androganic side effects are sill possible with this selectance, expecting with higher than remail therepartic doese. This may rickable boats of oily skin, acnet, and bodylichid lari growth. Antibolichardroganic stroids may also aggestrue time planter hat loss. Women are warned of the potential whiting effects of antibolichardroganic steroids. These may include a deepening of the voice, menstrual ringularities, changes in skin teature, field lari growth, and cliental estimates. Doestanchies in sides teature, but an antiportic activity relative to its tossa-building activities. The control of the control

# Side Effects (Hepatotoxicity):

Drostanolone is not c17-alpha alkylated, and not known to have hepatotoxic properties. Liver toxicity is unlikely.

#### Side Effects (Cardiovascular):

Ambiciantiques atomic can have deletrious effects on serum cholestered. This includes a tendency to refuel EME, Good cholestered values and irrecess ELI, Chold cholestered values, which may shift the HEA. DEA chaince in a direction that forces geneter risk of arteriockerosis. He relatile traper of an ambiciantiquesis earlier oil estima light is distributed and the contribution of the service of section of the contribution of the contribution of the contribution of the contribution of non-amountable), and level of resistance to heptic metabolism. Dostanoles solvald have a stronger negative effect on the layest management of cholestered than testosterous or randorbone due to its non-amountable inture, but a weaker support than 1-17 planta platigued activate, bandochoral-possure soulds may also adversely affect blood pressure and registerable, reaches endoded in relations, and support list more approaches the contribution of the contri

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular scenerics program and minimize the ristake of saturated fits, cholsterol, and simple carebolydrates at all times during active AAS administration. Supplementing with shot old, grams per day) and a natural cholsterol/antioxidant formula such as Lipid Stabil or a moder with conversable inverdience is also recommended.

#### Side Effects (Testosterone Suppression):

All ambolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainfairing substances, testosterone-levels should return to normal within 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradism can develop secondary to steroid abuse, necessistating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (yien):

Drostanolone propiorate was not FDA approved for use in men. Prescribing gaidelines are turavalible. For physique- or performance-erhancing purposes, this drug is usually injected three times per week. The total weekly dosage is typically 200-400 mg, which is taken for 6-12 weeks. This level of use is sufficient to provide measurable gains in lean muscle muss and stronoth

Dostantobre propiente is often cortined with other standards for an enhanced effect. Common stacks relade an isjectable analysis such as Deca-Durabolisif (numbroke contracts) and the standard of the standards and the standards of the standards

## Administration (Women):

The prescribing guidelines for Dobton recommedia a loss of 100 mg giorn three times per work. Therapy is given for a miniman fol \$1 to evoles before an extantion of its efficient, since the state of t

#### Availability:

Androgenic Anabolic

Drostanolone propionate is presently unavailable as a prescription drug product. All supplies of this drug come from export companies or underground steroid manufacturers.

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# Megagrisevit-Mono® (clostebol acetate)

Standard	Testosterone
Chemical Names 4-chlore	4-chloro-testosterone o-androsten-3-one-17beta-ol
Estrogenic Activity	none
Progestational Activity	no data available (low)

## Description:

Clustedo acette és un arabole seroid fatt is derived from testosterone. Clustedo la 4chion-testostorone, a modificario that makes fiss storal a los werapin arabole compound with marinal androgene potency. This analog of testosterone is ado not 17-dipla adjusted and does not mornituse, to three à life twory of water resettine, genecomiste, n'e ber toxicy during use. The hodgene substitution at the 4 position does not grandy enhance the restored parties of the production of the strength of the strength of the strength of the militigant basis, and germally performed. Although al derivative of the potent androgen testosterone, clostedo la cortanty for removed from the puerer steroid in action, and generally forced by adhests for in milities, not reason power.

#### History:

Clostebol acetate was first described in 1956. <sup>527</sup> It was developed into a medicine in Firmon, where it was sold as Steranshol (Farmitalia Germany) and Thrinshol (Jenanham)

Clusted a centre was first described in 1956.<sup>572</sup> It was developed into a medicire in Europe, where it was out all searmand for minimia, Germany and Trainted J Comphanic GERN, 17th annibods steroid lund generally been related for the restruct of outcoporous, adhinged in his reportably been used with secrics for a wide variety of aimers including american and beer disease. Both out and injectable forms of the drug were produced, although the injectable was more peoplathy used. Clusted accurate season commany used with almost first the secretary of the command of the command of the command of the annibols. The side effects of annibols industrying storous can be much more promoused in these conductions, to stroid, were well annibone, and some first three produces annibols. The reside effects of annibols industrying storous can be much more promoused in these conductions, to reside wave well annibones are selection to be from sto facility for the residence of the consistency and the story of the consistency and the conductions of the most plearable beautiful three conductions, to reside wave selection are not selected to the most plearable beautiful three conductions, to reside wave selection and selections are described to the consistency and the consistency and the consistency of the consistency and the consistency of the consistency and the consistency a

Altuchy agic forcebit is effect and priorit confirt. Catebrel actuate was never a witchy associated another, and our only finited to on its and must be mitted. As such, it faines would be a feeting and use of limited to on its and must be of metter. As such, is faines would be a feeting of the first product from longtherm would disappear by the centralication of Correnge, and the Sentembol brant would soot per explicit with lower doesd vision fartified vencines of the days old by Farmisla under the new Megaginised vencines would acquie Farmisla in 1993, although for a short port thereafter Megaginised was still being immediated under the Pharmisch label. This did not last long, however, and Pharmisch controlling lightened up to like and enroved this storoid from its offerings. Closthed actual had also appeared for some time in Jupus, sold as Maccolish by the first Pockska, for this product tools size been descontanted.

Albusqih the more functional injectable preparations of this steroid are of the market, clusted access is sill munificated in a narrher of domain preparations. The most recognizable such product line born Alb-Tichdelemin from Italy, albusqih it is asia be been sold in such products as Norbod (Meshor), fundodarnia (Taka), and Novadernia was a Norbod (Meshor), fundodarnia (Taka), and Novadernia woozda, and often include some reconjection belap accelerate healing. The dosses of steroid woozda, and often include some reconjection to help accelerate healing. The dosses of steroid woozda, and often includes on the housever, and, corbined with proor systemic delivery, are not of much use to athless. In addition, this steroid has even been included in continuely purposes, and of loss interest. Gene furth Alegapires was the last remaining efficience and of loss interest. Gene furth Alegapires was the last remaining efficiency and of loss interest. Gene furth Alegapires was the last remaining efficiency and of loss interest. Gene furth Alegapires was the last remaining efficience and or spictuable stored product to contain closebol accetar, this drug is now essentially a defirst times for size the affects use of steroids are concerned.

# How Supplied:

Clostebol acetate is no longer available as a commercial oral or injectable agent. When produced (Steranabol) it contained 20 mg/ml of steroid in a 2 mL glass ampule or 15 mg per tablet.

#### Structural Characteristics:

Clostebol is a modified form of testosterone. It differs by the introduction of a hydroxyl group at carbon 4, which inhibits aromatization and reduces relative steroid androgenicity. Clostebol acetate contains clostebol modified with the addition of carboxyle acid ester (acetic acid) at the 17-beta hydroxyl group, so that the five steroid is released more slowly from the area of nijection.

#### Side Effects (Estrogenic):

Clustedo is not aromatized by the body, and is not measurably estrogenic. An artis-estrogen is not necessary when using this steroid, as gracecomatis should not be a concern even among sensitive individuals. Since estrogen is the usual culpri with water reterion, clustedo instead produces a learn, quality look to the physique with no fact of excess absolutaneous fluid reterion. This makes it a favorable steroid to use charing cutting cycles, when water and first reterion are major economis.

# Side Effects (Androgenic):

Although classified as an anabolic steroid, antrogenic side effects are still possible with this substance. This may include bouts of oily skin, acre, and body/facial hair growth. Anabolic/arthougenic steroids may also aggravate mule pattern hair loss. Women are also warned of the potential virilizing effects of anabolic/arthougenic steroids. These may include a changing of the control of the potential virilizing effects of anabolic/arthougenic steroids. These may include a changing of the control of the contro

seepering on the voice, measures requirement, curings in som source, use that growing and clinical enhancement. Additionally, closels of in orderate-level methodated by the 5-alpha reductise enquire, so its relative authorganicly is not groundy already by the construct use of instanctive of enhancement. Note that closels of a section of this own hospitages activity relative to its tissue-building actives, making the threshold for storag and/organic size effects companily layer biles with more analogousic agents each as telestoctories.

# Side Effects (Hepatotoxicity):

Clostebol is not a c17-alpha alkylated compound, and not known to have hepatotoxic effects. Liver toxicity is unlikely

#### Side Effects (Cardiovascular):

Anabolarindoguis atendis call how deletrions effects on neum cholesterd. This irelates a tendency to reface III. (6, ogd) cholesterol values and irecreas III. (6, ogd) cholesterol values, which may shift the IIII. to III. In balmer in a direction that fixors geneter risk of anteroclerons, like relation terror an anabolarindoguis exteriol on serum lipit is a construction. The entire traper of an anabolarindoguis exteriol on serum lipit is observed in the construction of the entire traper of the entire tr

To help reduce cardiovascular strain à is advised to maintain an active cardiovascular services program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish ols (4 grams per day) and a natural cholesterolaritoxidant formula such as Lipid Stabil or a product with companible ingredients is also recommended.

## Side Effects (Testosterone Suppression):

All ambolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainfairing substances, testosterone levels should return to normal within 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradism can develop secondary to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

# Administration (Men):

Clostedo acente is generally used in clinical dones of 300 mg per words by sjection or 15 mg. 2-2 imus per dup on mly. The drug is admirated fir 3 connected weeks, followed by a break for 3 weeks. It is reasoned at this post if inclinate IEE/crite dones for physaque or performance-entraining proposes till in the rage of 100-200 mg per work, hand for 6-12 weeks. Given the fine-script nature of acente specialses, the weekly dosage is generally considered to the contract of the contract of the contract of acente specialses, the weekly dosage is generally constructed to the contract of contract periodses, that is given the were not common (General to know this words) of the contract of contract of contract of the contract periodses, that is given the were not common (General to know this words). When admirated a cally dosage of 600-900 mg appeared to be the not common.

The ambole effect of this drug is fairly weak, so closted-a decette is most often railized in combination with other services for a stronger effect. The garmal application is to use if no contest preparations with other ron-arcunizing ambolics such as Westrollö or countribute. Here, a dudy dose of 20 mg mup be added in with an assenge dose (20-3) mg per duy) of the cruil ambolic, which together should provide the user a new musch backing effect without any such rearterism. The effect of obsoled be would be removaltur smither than seen with the old Printobolantia excent ampules, although Meggrievel is somewhat weather in effect. Some also opt to use the composed in addition with stong mon-arcunizing androgens such as trenbolone, Halotestmill, or Provinonill. The result in such cases can be an even more pronounced effect of muscle definition, although this will be accompanied by a much stronger set of side effects.

# Administration (Women):

Clostebol acetate is generally used in clinical doses 30 mg per week by injection or 15 mg 2-3 times per day orally. The drug is administered for 3 consecutive weeks, followed by a break for 3 weeks. The drug is resumed at this point if inficiated. Effective doses for plusiques or performance-enhancing purposes fall in the range of 50-75 mg per week for the intensible or 30-for me dails for the cond takon for no houser than of sweeks.

# Availability:

Clostebol acetate is no longer available as a prescription agent at this time, and is unavailable on the black market

#### Metandren (methyltestosterone)

Androgenic	94-130
Anabolic	115-150
Standard Testos	
Chemical Names 17b-hydroxy-17a- 17alpha-methyli	-methyl-4-androsten-3-one androst-4-en-3-one-17b-ol
Estrogenic Activity	high
Progestational Activity	not significant

#### Description:

Methylostoterous is an orally available from of the primary mule antrogons tostoterous. Looking at the structure of this stroot, we see it is leastly just testistence with an added methyl group at the c-17 philosophostic is -17 alpha adjusted advanced, which allows for the control of the control control of the control

#### History:

Medylestosterous was fired described in 1935, <sup>550</sup> and use one of the first onal anthogens to be used in clinical medicine (it follows by rooms, the first onal anthogens to the used in clinical medicine (it follows by rooms, the first onal anthogens, bor one worth its manner of the contraction (in describing the described first substance) and the contraction of other uses settled, better contract in potentiary particular contractions of the contraction of the cont

In addition to standard tables and capsales, methylocosterous has also been commercially prepared in subliquagion of based tables. Mentarine Tagaste from Can Plaramecentical Company were perhaps the most recognized, and were popularly sold from the 1950's to 1900's. These tables were placed under the tongue of between the garm and check and left to dissolve, delivering the drugs to criculation win the musous membranes, bypassing the force. Subligatar of braces trained approximately doubles for bracesofthilly of methylocosterous,

ana aso provues peas eves or urig rapiny (approximate) i nour aner uosing isseau oi 2 hous). Alfrough Cha's Metandren Lingues are no longer commercially available, namerous other sublingual buccal methyltestosterone tablets are still in production today.

Methylstostenore remins a controvenial steroid. Although at hus a long history, and angulps a justifilite short proceed, it is not hope with byte and, it is even being withfrom from many dring markets. The German Endocrite Society made an official statement that methylstostenors was obsolucle in 18%1, and the days could be removed from German methylstostenors was obsolucle in 18%1 and the days could be removed from German and Control of the Charles of

#### How Supplied:

Methyltestosterone is widely available in human drug markets. Composition and dosage may vary by country and manufacturer.

# Structural Characteristics:

Methyltestosterone is a modified form of testosterone. It differs by the addition of a methyl group at carbon 17-alpha, which helps protect the hormone during oral administration. As is typical with c17-alpha adylation, the resulting steroid has lower anabolic activity than its narent testosterone.

# Side Effects (Estrogenic):

Methylastostence is arcuntated by the body, and is highly estrogate, due to is conversion to 17-sipha methylastaded, a sprince is engogen with high biological active, 17-sipha methylastaded are not of arcunstation, although the potent matter of 17-methylastaded inner uncompensate for this (Succounties) is often a concern during treatment, and may present study agic early into a cycle (grottedarly when lagher dense sare manifested and the major of the control of the c

#### Side Effects (Androgenic):

Medybestostence is classified as an androgen. Audrogen; side effects are common with this softstarce, and my include bosts of only sinc, men, and bodyfield hair growth. Higher does are more likely to cause such side effects. Ambolicitarlogensic streaks may also aggressic may be partial may be an another than the same years of the angent of the potential witness of the contract of the potential witning effects of amboliciarlogensic seroids. These may include a desperation of the potential witning effects of amboliciarlogensic seroids. These may include a desperation of the voice, mensional irregularities, charges in side totates, ficial larit growth, and clitical enlargement. Like toolstorous, mellybasticatives convents to a more potent startout of its mention with the 5alphi reductate enzyme, in this case 17-alphis-mellydisplicotostostorous. The relative that the conventure and differential contractive and the conventure and of inferential contractive and the conventure and of inferential contractive and the start of the conventure and of inferential contractive and assets of the conventure and of inferential contractive and assets of the conventure and of inferential contractive of assets of the conventure and or enough definance.)

#### Side Effects (Hepatotoxicity):

Methylsestosterone is a c17-alpha alkylated compound. This alteration protects the drug from deactivation by the liver, allowing a very high percentage of the drug entry into bloodstream following oral administration. C17-alpha alkylated anabolic/androgenic steroids can be hepatotoxic. Prolonged or high exposure may result in fiver damage. In rare instances life-threatening dysfunction may develop. It is advisable to visit a physician periodically during each cycle to monitor fiver function and overall health. Irrake of c17-alpha alcylated steroids is commonly limited to 6-8 weeks, in an effort to avoid escalating liver strain.

Methylstostenow was the fits on altered liked to bequite durings. This may be, in part, related to the early wedspend use of the corporant, in the long generally displays acceptable settly when used inclinally prescribed dosages (serious five towice) cannot be completely excluded, become completely excluded however, even at critical thouses, When talent as does of 10m quickly fit 8 5 weeks, hepatotoxicity, as measured by bromosalphilathe (ISSP) retention, was low in ore study. <sup>250</sup> In a separate investigation, a majerior of pruiters morted significant BSP retention after only 2 weeks of themps with 67m quickly. <sup>250</sup> Secure for complications are more given the protection enters in which must people use or all anbidicardineguest sensition, although cannot be excluded with methylsestosterour, especially with high doses and/or prolonged administration periods.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

## Side Effects (Cardiovascular):

A rabolic and roganic strotts can have deleterious effects on serum cholstered. This includes a tendency to note left, a good horistered unless and increase LDI, (topol, horistered unless and horistered unless and cholstered values, which may shift be able to the left, and the left of the left

Methylstotenee has a stong effect on the hyptic immegeneet of elockstool due to its neutral resistance to here breadformed and stone of administration. Studies have demonstrated an approximate 35% decrease in HDL cholesterol and a 50% increase in LDL cholesterol with 40 Mpg per day. <sup>517</sup> These changes occurred within 2-4 weeks of the intuition of therapy, and persisted for 2 weeks after decontrastion for the due, Arabolic influences services in my also adversely affect blood pressure and trajecturiles, reduce endothed contrastication for the process of the size of the contrastication of the pressure and trajecturiles, reduce endothed contrastication from the contrastication of the pressure and the contrastication of the pressure and trajecturile pressing the risk of contrastication of the pressure and trajecturile pressing the risk of the pressure and trajecturile pressure and trajecturile pressure and trajecturiles.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fists, cholesterol, and simple carbolydates at all times during active AAS administration. Supplementing with 8sh oils, 4d grams per day) and a natural cholesterol/artioixidant formals such as Lipid Stabil or a moduct with comrangble inserdients is also recommended.

#### Side Effects (Testosterone Suppression):

All ambdośrandnoganie steroids when taken is doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-straintaing substances, testosterone levels should return to normal within 1-4 morths of drug secession. Note that prolonged hypogoradotrophic hypogoradism can develop secondary to steroid abuse, necessisting medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Staties have shown that taking an oral anabolic steroid with food may decrease its bisovailability.<sup>32</sup> This is caused by the fit-solible nature of steroid hormones, which can allow some of the drug to dissolve with undigested dietary fit, reducing is absorption from the gastrointestinal tract. For maximum utilization, this steroid should be taken on an empty storonch

#### Administration (Men):

To treat androgen insufficiency, prescribing guidelines call for a daily dosage of 10-40 mg.

ine does is reacted by 50% when administered in storagua or recent from the under would be used for extended periods to long as the pattern's bloottaper reside (dependencily, sentin lipsk, etc.) do not reconstate in discontinuous. When used for commonly used, then in cycles being no more than 6.8 weeks in legal, Mediplestosterore is most commonly used not an anabode, but to straight seggession in the user Devereibles, bodylands, and composite authent code material based on the contraction of the contra

#### Administration (Women):

Mediphenosemoe is net wiskly used with venome in cheisal medicine. When applied, it is more often used as a secondary medication during inspensible frost excurs when other therapies lave finded to produce a desirable effect. The dosage used for this application can be as high as 20 m per due; Low does of methphenostemes, there here used in receiv years to treat the symptoms of meropause. An example is the product Estrates, which contains exteriod entropeas and 2.5 mg of methphenostemor. As does got I tablet per dy may improve energe, Bolds, and overall webses of the patient, as well as contral conceptions (with extension relationships with a localization in the boose, suchastroom conceptions) and the force and the contral conceptions of the contract of the contraction of the contra

#### Availability:

Pharmaceutical preparations containing methyltestosterone are fairly limited. In reviewing some of the remaining products and changes on the global pharmaceutical market, we have made the following observations.

Methyltestosterone remains available in the United States. It is sold under the brand names Android tablets and Testred capsules from Valeant Pharmaceuticals. A generic is also available from Imraex Labs.

Aburahan makes a generic methyltestosterone product in Iran. It comes in 25 mg tablets, with 10 sealed in each foil and olastic blister.

# Methandriol (methylandrostenediol)

Estrogenic Activi	low to moderate	
Chemical Name	17-alpha-methylandrost-5-ene-3,17-beta-dio	
Standard	Testosterone propionate	
Anabolic	20-60	
Androgenic	30-60	

# Description:

**Progestational Activity** 

Methyandrostenediol (methandrol for short) is an ambole: steroid derived from dihydrotestosterone. The drug itself is manufactured in two very distinct forms. The first is unssterified (straight) methylandrostenediol, which is used when making an oral medication with this steroid (although an injectable once existed in the U.S.). It is also found as esterified methylandrostenediol dimonoipante, which is recurrent as an injectable. The added

no data available (low)

# History:

Early assessments of methylanthousheadol being primmly analosis in instance did not beld up work with their excensive ties in human. It was executably determined that in does sufficient to promotive weight gain its analosis properties were accompanied by significant androgusic activity. Unimords, in that up would be viewed as one of obligated analosis and analogusic activity. The days of the superior of the superior analogusic activity of the substitute of the superior activity of the superi

# How Supplied:

Methylandrostendiol is available in select human and veterinary drug markets. Composition and dosage may vary by country and manufacturer.

# Structural Characteristics:

Methankostendo is a modified form of displostestostoscene. It differs by: I) he addition of a methyl group as thereof II-adpla to protect he horneout during coal administration and 2) the introduction of a double bound hereone content to her more during an analysis strengt of her steared (partly hyministry as which seems to increase the analysis strengt of her steared (partly hyministry as sixtuation to metabolism by 3-hydroxysteroid delaydosgomes in skeletal muscle issue). Methyl androstencial diproprientate contains methylamotreactional modified with the addition of 2-donopsis and elsent (proprient acid) at the 3-bets and IT-bets hydroxyl groups, which delay the release of five methylamotreaction from the set of rejection (depost).

#### Side Effects (Estrogenic):

Mediplandosteráci is not directly aromaticed by the body, although one of its known metholicis is melhyletosterone, with can aromatize. Mediplandosteració is also believed to have some inherent estrogenic activity. It is, Beovise, considered a weady to medientaly estrogenes ferrod. Cynecomotain is prossible during trenuture, for garentaly when higher doses are used. Water and fat reteriors can also become issue, again by the control of t

#### Side Effects (Androgenic):

Altough often classified as an ambole stroid, methylandrostenedid is sufficiently analoguic Altonigon idea effects are conveniently with substance like may racked best of oly skin, area, and bodyficial his growth, Ambolichindroganic stroids may also agaptout meth pattern hair loss. Wiemen are wormed of the potential wirking effects of analodicintroganic stroids. These may include a deporting of the voice, menstral irrigalarties, changes in sike tester, the fairly growth, and classified indisputers. Note that methylandrostenedid is not affected by 5-aplus melactuse, so the relative aniomatory of the contribution of the

# Side Effects (Henatotoxicity):

Medylunkostenciós is a cl<sup>2</sup>-alpa adyluted composed. This alternifon protects the drug from descriation by the fisce allowing as very high procentage of the drug enty in the bloodstream filowing oral admiristration. Cl<sup>2</sup>-alpha adyluted ambride/androgenis strevists can be hapstones. Probagge of high percentage of the visit a physicient percentagely and the control of the each cycle to remain the fraction and covered health intake cl<sup>2</sup> i <sup>2</sup> plant ad justifications, the control of the control of

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

# Side Effects (Cardiovascular):

Anabolarindeganis steroids can hove debetrous effects on serum cholesterol. This includes a ratentary to returne III. (good) cholesterov ulsass and increase III. (foul) cholesterov ulsas, sufficients III. (foul) cholesterov ulsas, sufficients III. (foul) cholesterov ulsas, sufficient in the foundation of the foundation and the anabolarindeganis extend on serum liptic is dependant on the dose, route of administration (end xx. spectable), type of steroid (comtantible or ross-countriales), and level of resistence to hepself, enterlied (controllate) or ross-countriales), and level of resistence to hepself, enterlied (controllate) or ross-countriales), and level of resistence to hepself, enterlied (controllate) or ross-countriales), and level of resistence to hepself, enterlied (controllate) or resistence and resistence and

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fists, cholesterol, and simple cardohydrates at all times during active AAS administration. Supplementing with \$8h olis (4 games per day) and a natural cholesterolarinoxidant formán such as Lipid Stabil or a product with comprable inspedients also percentaged.

#### Side Effects (Testosterone Sunnression):

All ambole/andrugenie steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainfairing substances, testosterone levels should return to normal within 1-4 months of drug secession. Note that prolonged hypogoradorophic hypogoradism can develou secondary to steroid abuse, necessistation medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

# Administration (General):

Studies have shown that taking an oral ambolic steroid with food may decrease its bioavailability. This is caused by the fit-soluble nature of steroid hormones, which can allow some of the drag to dissolve with undigested dietary fit, reducing its absorption from the gastroinestiral tract. For maximum utilization, oral forms of this steroid should be taken on an empty stommer.

#### Administration (Men):

Einly prescribing guidelines for Stemedial recommend a dosage of 25 mg gione 12 to 5 mins per work by only beacol or intermeaction rule. For physique, or operheumoc-entirencing purposes, a pipical dosage is in the range of 2.5-50 mg dulp for the oral form and 200-400 mager works with the rigitable. In order to these photo close home one own with the igentable, it is generally administered orace every three to four days. Cycles generally hat for no more than 6 to 8 weeks, in an effort to minimal hespitatoxicity and start on the foer and cholestered values. This level of use is sufficient for mortenite gains in muscle size and strongly, which may be accommunically as to well of water returns.

White im we be possible to use methylandrostendral alone for musch-bailing purposes, it is must often contributed with other autobolis for a stronger effect. Combined with Descharbolist of Epipoisest, fix example, measurable gains of land musch mass, without an extense level of votes reteriors, may be noticed. This is the general composition of not Australian vet blends that richtle methylandrostendral. When biology for a more prosenced gain in most, a stronger arthogen such as teststores may be added. The entaining growth can be quite exceptional, but the user vial also have to deal with a much amounting anables, each an Martinolis, Principologis, or considered. The result bree should be a more pronounced effect on musch hardness, with a modente gain of solid lean time.

#### Administration (Women):

Early prescribing guidelines for Stenediol recommend a dosage of 25 mg given 2 to 5 times per week by oral, buceal, or inframsscular route. Methylandrostenediol is generally not recommended for women for physique- or performance-erhancing purposes due to its androgenic nature and tendency to produce virilizing side effects.

# Availability:

Pharmaceutical preparations containing methylandrostenediol remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

The only place where this steroid is still produced in an volume is Australia, where a number of veterinary perparations still include methandriol in their blends. These products are rarely traded in international commerce due to tith controls on anabolic steroids in that country.

6 000.7 000

#### Metribolone (methyltrienolone)

Androgenic	0,000-7,000
Anabolic	12,000-30,000
Standard	Methyltestosterone (oral)
Chemical Names	17alpha-methyl-17beta- hydroxyestra-4,9,11-triene-3-one 17alpha-methyl-trenbolone
Estrogenic Activity	none
Progestational Activity	no data available

#### Description:

Methylinionhous is one of the strongest real ambolic stronils ever produced. This agent is a derinative of treels/one (trienbeed), which has been en-17 plan palaquate at also for end administration. This modification has created a secord that is significantly stronger than is non-methylated coasts. Its potency has been measured to be anyboxine from 102-300 times goater than that of methylatestoroure, with greater disconsiston between anabolic and analogusic effects. 55% of Milgram for milligam methylateroleus is a nore active sector than any agert sold on the commercial methot, requiring does sa little as 5-1 milligam per day to notice a stoney anabolic effect. In potency is ony matted by its relative toxicy. however, which has limited its modern use to that of laboratory research only.

# History:

Mediphirosolue was first described in 1965, <sup>252</sup> It was immediately identified as an entemely potent ambole agant, far mere potent amb tecomorrasily available agares for the entemely potent ambole agant for mere potent amb tecomorrasily available agares for the time. In spike of fix high relative activity, however, mediphirosolues has seen very limited use in harman. It was used clinically during the lise 1969/3 and entry 797, most modely in the treatment of advanced breast cancer. Here, is exceedingly strong armbolicalizations grant has deared been control to the control of the describe of the degree of the compose, harding is some extension of the simple control of the degree of the control of the degree of the control of the degree of the degre

By the mix1970s, methybricoshoe was becoming an accepted standard in non-harman research statics, pertituding those pertaining to seal upd of the analogous receptor activity. For this perpose the agent is very well sated, It is sheer potency and resistance to seranbending precises makes it is necedent in whom receptor-brings insuland to compare often precises makes a necessary of the receptor of the precise pertituding and acceptance of going to graphy interfers with the results of most experiences. Body tissues can metabolic most storoic hird goal, which means that own includes makes can be complicated with the quantum of which will be compared to the precise of the pre

## How Supplied:

Methyltrienolone is not available as a commercial agent.

# Structural Characteristics:

Methylinioches is a modford form of methodoxe. It differs by 1) he addition of a methyl group at each on 17, a plus to protect the hornese during on also institution and 2) be introduction of double bonds at endorse 9 and 11, which increases its binding affinity and shows its methodoxin. The earling strend is applicately more post than its methodox bonds, and displays a much longer that like and lower affinity for securability proteins in comparison. Methyliniothous chemically diffes from trevolocos copy by the addition of a methyl group at  $\approx 1.7$ . This alternion changes the activity of methyliniothous chemically diffes from trevolocos consistentibly, however, such that this agent relation is a finite of the order of th

## Side Effects (Estrogenic):

Methylineshoes is not arountated by the body, and is not measurably estrogenic. It is of most however, that methylinenshoe elspines significant bringing fillings for the progestores receptor.<sup>55</sup> The side effects associated with progestore are straint to those of entropar, including regardle exholest inhibition of statistione production and enhanced rate of first storage. Progestries also support the stimulatory effect of entropers on mammay tissue growth. These appears to be a storage superop between these to howers been caused to the progestore and the programme and the programme

#### Side Effects (Androgenic):

.. .. . . . .

#### Side Effects (Hepatotoxicity):

Mempiricanore is a c1 - signa sayunci corporat. Im a steration protects the ring from detactionist by the fire, allowing a vsvy high processing of five ding entry is do bloodstream following oral administration. C17-alpha alghoted ambidicant-looguis steroids cannot be lapatoxice. Frobegad or high exposure my reseal in five farming. In rare instructs life-threatening dysfurction may develop it in a shieshie to visit a physicien periodically during each cycle to remarks the fraction and out-oral health intiate of c17-alpha alghotal resolution is commodify hinded to 6-8 weeks, in an effort to routil excluding here strait.

Methorization is an accordant power out to strong with the best of resistance to

lepatic metabolism. This makes methybrizenhone exceedingly four-toxic, prochaling is use as a prescription again at this time, in any not of the world. Statlike published from the University of Drom Germany back in 1966 make this very clear. <sup>500</sup> In fact, at this time researchers had deemed this the most between the extended and the state of the state o

alkaine phosphates and collinatenses in senura activity of proaccelorin in plasma methylinizatione tamed out to be very active as to causing bichemical symptoms of irradicpatic cholestasis....thas methylinizatione at present being the most "hepatotoxic' steroid."

The use of a fiver detoxification supplement such as Liver Stabil, Liv-S2, or Essertials Forte is abbited with this time are hemotoxic ambidicinationesis extension.

# Side Effects (Cardiovascular):

Anabolic antrogenic stroids, can have delectrions effects on serum-cholsterof. This includes a tendency to reduce IFLK (good) cholsterol values and irresent EUL (dud) cholsterol values, which may shift the IFLA to LDL balance in a direction that favors genetar risk of antroisclerosis. Bir entities repart of an anabolicandrogenic steroid on serum lipsi is dependant on the dose, mote of administration (ond vs. ipicetable), type of steroid sequences of the control of th

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fits, cholsetrol, and simple carbohydrates at all times during active AAS administration. Supplementing with sho is (4) gams per day) and a natural cholseteo/lartioxidant formals such as Lipid Stabil or a moduct with commands in arcidiner is also recommended.

#### Side Effects (Testosterone Suppression):

effects, see the Steroid Side Effects section of this book.

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-straining substances, testosterone levels should return to normal whith 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradorophic hypogoradorophic normal whith 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradorophic and evelop secondary to steroid abuse, necessistating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side

# Administration (General):

Staties have shown that taking an oral anabolic steroid with fixed may decrease its bisovailability. <sup>60</sup> This is caused by the fit-soliable mature of steroid hormones, which can adlow some of the drug to dissolve with undigasted dieary fit, reducing its absorption from the gastroirestrial tract. For maximum utilization, methylrienolone should be taken on an emrity stomach.

#### Administration (Men)

Methylinolous is no begar used in clinical medicine due to an unexceptable level of hepatotoxicy. This again is guzenilly not concerned for phyliques or performance phyliputoxicy and proper in general process. For the same reason. Those absolutely insisting on its use need to take its level of her toxicity every servicely. At the very lesst, routine blood tests should be conducted to ensure the agent is not impuring damage. Drug duration should also be very inside, prefembly to wheek of use or less. The relative potropy of methylinerolous is externely high, requiring doese as little as 5 milgramp of also. He effects early of methylinerolous is externely high, requiring doese as little as 5 milgramp of also. He effects early of the distribution are completely utilisable, and should rever be enterpated. Again, this is an externely take quality statement of the control of the contr

## Administration (Women):

Methylprienolone is no longer used in clinical medicine due to an unacceptable level of hepatotoxicity. This agent is not recommended for women for physique- or performanceerhancing purposes due to its extremely strong toxicity and tendency to produce virilizing side effects.

#### Availability:

Methylinerolone is not produced as a prescription steroid product in any part of the world. With the rapid expansion of undergound steroid munificatures, this agent has been released as a black murket designer compound. Those contemplating the use of undergound forms of methylinenolone should consider that such agents are being released for human use without any government approval or consideration to its safety.

73.94

## Miotolan® (furazabol)

Androgenic

Anabolic	270-330
Standard	Methyltestosterone (oral)
Chemical Name	17-Methyl-5alpha-androstano [2,3-c]furazan-17beta-ol
Estrogenic Activity	none
<b>Progestational Activity</b>	no data available (low)

#### Description:

Firzazbol is an oral anabols steroid derived from displotestosterone. This agent is mordently anabols, with only and anaboguies properties. This is no odebed to to be modification of the steroid's A-ries, which allows the steroid structure to remain stable and Displotestostoscerone, no corporation, is a port anabole, eachly ruthlocked in rused; because to inscribe metabolies. The gains succinately with financial other not extreme, and would more clockly research the quality growth of an affine oral mortization whole like stamodol or deviatamentes, instead of the watery bolk of a testestorane. For this reason, finzabols is most office anaphel during conting phenos of raising, and by shifters in post and weight-residently and the stable properties of an analysis of the water place of the stable post of the water place of the stable post of

#### History:

Furazabol was first described in 1965.<sup>541</sup> The only modern pharmaceutical preparation of record containing firazabol, at least known to researchers in the West, was Motolan from Daibbl School I also in Jamen which was cold in Jamen mainly during the 1070's and '800's

LORGER SEGMENT LEADS IN PROJECT WHICH WE SEE HE ADJUST LEADING THE REST REST SECRETARY PROFESSION AND ADJUST LEADING THE ADJUST

Finzabol was a popular steroid armog Olympic affects during the 1980's, when it was quichy lavous among contin nineas the testing findles had may to it-defined the agent and thurcher could not test for a Dr. Jamic Astophun, the physician that accompanied Ben Johnson to the 1980 Olympic in Food, reportedly was gively polarous found armonos often affects as the timy linearbol, lavorsige the drug pould not be descrable. It remains self-under the control of the control of the control of the control of the detection of the control of the control of the control of finzabol to mixe were published, immediately definishing any value this agent formerly possessed as a steroid undetectable to due greeners.

Eduy, Erazbol si very searoly, known to bodybulkers. The Motionh brand from Japan was discoriation dumy years aga, and no plammenciated preparation containing fizzable has been known to exist sixee. The dung is occasionally heated on the black maket, however, the to the fact that is it all produced in bulk (as a row material for product manufacturing) in Asia. From there it is obtained by undergound strood manufacturing of predators in the Wast, and produced into one labels and exposite. Currenty the actual number of products containing financials is small, although could easily be cognited if containing the strong of the containing the strong of labels and produced into a multiloop from a near phreeopology model.

# How Supplied:

Furazabol is no longer available as a prescription drug preparation. When sold it came in the form of tablets containing Img of steroid.

# Structural Characteristics:

Furzabel is a modified form of diphotostosterone. It diffus by; 1) he addition of a metaly group at earbor 17-adjsh, wish helps protect the hormone durig oral administration, and 2) the attachment of a firzam group to the A-ring, replacing the normal 3-kets group. When viewed in the light of 17-adjan remothylithydrostostostome, the A-ring modification on firzabel seems to considerably increase its anabolic strength while reducing its relative and/contribit.

#### Side Effects (Estrogenic):

Fuzuabol is not arountated by the body, and is not measurably estroganic. An artis-estrogais not necessary when using this stereoid, as generouslast should not be a concern even among sensitive individuals. Since estrogan is the usual culpit with water retention, this steroid ristead produces a lawn, quality look to the physique with no fire or of excess subcutaneous fluid reterior. This makes it a fivorable steroid to use during cutting cycles, when water and fire reterior are made to concerns.

# Side Effects (Androgenic):

Altough closifed as an ambolic strootig, androganic side effects are still possible with this softstame, expectably with ligher doese. This may include boots of only skin, area, and body/ficial larir growth. Antabolican/troganic strootis may also aggranter may partner lart boss. Women are warmed of the potential Writing effects of antabolican/troganic strootis. These may rickled a deepering of the vote, mentional irregularities, clausges in skin texture, ficial larir growth, and closual enlargeness. Prazabol is a networt off with relatively to undroganic activity relative to its tiona-boding activity, making the freebold for strong anadogus; said exists or factor and the contraction of th

# Side Effects (Hepatotoxicity):

Firazziolo I sa e17-sipha allysited compound. This alteration protects the drug from doctrination by the first allowing as very high percentage of the drug entry sito the blockstream following onal administration. C17-sipha allysited anabolic anabogens strovids can be hepatotical. Following of the phospour may toos in the dampe. In rare instances can be expensioned to the configuration of the protection of the configuration of the configuration

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any henatotoxic anabolic/androsenic steroids.

# Side Effects (Cardiovascular):

Ambolizindroganic strosids can have deleterious effects on nerume-holstered. This includes a terefacty to reduce PLIK (agod) cholstered values and increase LLK (dud) cholstered values, which may shift the IRLs to LLD belance in a direction that fivors geneter risk of arteriocherois. Bre relative inputs of its analobicaltoniques its served on nerum light is a directional to the reduce the relative properties of the reduced to the reduce of the reduced to the reduced of the reduced to the reduced t

Note the funded is often reisolody described as a secoid with using brustedial collection between globes. Seth determine usual pulse reference of studies conducted in the early 1970, which exemined the light horizing efficies of the agas 2<sup>-10</sup> Such a position, however, leaks a modern perspective of the dang. To draw a possible, their ground probability of the properties of the properties of the probability o

To help reduce cardiovascular strain it is advised to maritain an active cardiovascular exercise program and minimize the intake of saturated fists, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with 18th oils (4 grams per day) and a natural cholesterolarinoxident formula such as Lipid Stabil or a moduler with comparable interelients is also recommended.

#### Side Effects (Testosterone Suppression):

All ambolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainfairing substances, testosterone levels should return to normal within 1-4 morths of drug secession. Note that prolonged hypogenoatrosphic hypogenoadsen can develop secondary to steroid abseque-essistating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

# Administration (Men):

An effective dosage of financials seems to begin in the range of 10-20 mg duly for mer, taken for no larger than of 8 weeks. At file self seems to impair a measurale musclebaling effect, which is usually accompanied by fit has and increased definition. Does of 30 mg per day or mer considerably increase his anabotic potentia for duling that affecting the expense of guester lepastionizity. The muscle-builing activity of financial cloud, in stand, the expense of guester lepastionizity. The muscle-builing activity of financial cloud, in stand, the expense of guester lepastionizity. The muscle-builing activity of financial cloud, in stand, the expense of the control of the purposed. In this case, the contribution deadly provide a succording pain of odd, quality muscle musc without a loss of definition due to water receiters. We could alternately use a more potent arcontrollade androuses such as testosterous, althouth here the justs may be accompanies by some level of water retention, and potentially a decrease in muscle definition.

In the athletic arena, an effective oral daily dosage would fall in the range of 2-5 mg, taken in cycles lasting no more than 4-6 weeks to minimize the chance for writization. As with all steroids, withizing side effects are still possible in women, but remain rare with conservative dosine.

#### Availability:

Furazabol is no longer produced as a prescription drug product, although underground preparations containing this steroid may be located.

# Myagen (bolasterone)

Administration (Women):

Androgenic	300
Anabolic	575
Standard	Methyltestosterone (oral)
Juliana	mediyitestosterone (o

Chemical Names 17beta-Hydroxy-7,17aipha-dimethylandrost-4-en-3-one
7,17-dimethyliestosterone
Estrogenic Activity high
Progestational Activity no data available

#### Description:

Bolasterore is an oral anabolis steroid structurally related to methylastostorous. It differs only by the addition of a methyl group or 1,7, which accounts for in given chemical many, 7,17-dimelhylastostorous. The added of 7 methyl group mains the activity of this steroid for partial properties of the properties of t

# History:

Bolssteroe was first described in 1992-54. It was closely evaluated for ambolic and androganic effect approximately 3 years like "The flow quest of decloped by Ugiplin, and sold in the U.S. during the 1960's starfer for Mogage heard arms. It was maily indicated for the tentiment of advanced betters classes; all conversa, allowagis the again was aside investigated and the start of the start in relaxes. By the 1980's, belietened and the out and of commerce for so they fast it was all but forgation among affilies. Although bolssterons in to long produced, the start is start in the start of the start

#### How Supplied:

Bolasterone is no longer available as a prescription drug product.

# Structural Characteristics:

Bolistrone is a modified form of extosterone. It diffies by: 1) the addition of a methyl group cardon 17-abyla, which helps protect the hormone during oral administration, and 2) the introduction of a methyl group at earbon 7 (abyla), which it shibs 5-abyla reduction and shibs the arabolic to antidogene train in five or of the forme, 2717-dimethylated storoids also tend to be very resistant to metabolism and serum-binding proteins, greatly enhancing their relative biolocical activity.

#### Side Effects (Estrogenic):

Bolasteroe is arountated by the body, and is considered a lighty estrogenic servoir due to its connection to 7.17. distrelighential (on entergon with light biological activity). Opaccomosis may be a content using resolutions, especially when higher than normal anticonstructions of the content of the conte

# Side Effects (Androgenic):

Albragh bolusterone is clusified as an arthofic storial, androganic side effects are still possible with its substance. These may include boost of oly sides, man, and bodyficial inter growth. Higher does are more fixely to cause such side effects. Antabolicanhoganic storials may also agreement partners into See Wenne are additionally sourced of the potential virling effects of antabolicanhoganic storials. Hose may include a deepening of the vioce, memoral megalines, changes in sin teatrae, ficial integround, and clotted orlangement. Bolusterone is unaffected by the S-aplan eductuses copyrue, to is relative antaboganic by an effected by the concern to of intensition of datasetick. Note that stacks administering, fing and 2 mg of bolusterone per day have shown no contrast by public hair growth, grant inchanges, where equipment, and many therefore store main feet, bolustrone is considered to have a companile ratio of analosis of antiques effects. Solustrone is considered to have a companile ratio of annials to antiques; effects and northerholostoches.

#### Side Effects (Hepatotoxicity):

Bolsterone is a cl7-alpha alghated compound. This alternation protects the drug from doctrination by the five, allowing as very high presentings of their gentry into the bloodsteron following containmentation. Cl7-alpha alghated analysis indeed, as the bloodsteron following containmentation. Cl7-alpha alghated marketic and containmentation in the containmentation and containmentation

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any hepatotoxic anabolic/androgenic steroids.

#### Side Effects (Cardiovascular):

Ambolicarinoganic strools can have debetrous effects on serum cholesterol. This includes a redency to reduce IEL (good) collesterol values and increase IEL, (find) collesterol values, which may shift the IEL to IEL balmer in a direction that fivous genete risk of arteriocetosis. The retailse impact of an ambolicarindoganic strool on serum lipits is dependant on the done, route of administration (red xs. ijectable), type of steroid (genomizable or mora-amortizable), all need sof resistance to heptic metablosis. Bolistories has a strong effect on the leptic imangement of cholesterol due to its structural resistance to hep the reductions.

also adversely affect blood pressure and trighorities, reduce entoblishi relevation, and support let verificate hypertophy, all opartially increasing her risk of entiforouscular disease and myocardial infraction. Staties administering Imp and 2mp of bobsterone daily for 6 words to 27 printers have demonstrated a trent floward necrosed serum cholesterol Albreight no HEA, and LD, broaddown was provided, it cam be assured based on the control of the contr

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the ristake of saturated fits, cholsterol, and simple carebolydrates at all times during active AAS administration. Supplementing with shi old, games per day) and a natural cholstero/lamioxidant formula such as Lipid Stabil or a product with comrangible inserdenies is also recommended.

# Side Effects (Testosterone Suppression):

All anabokirantingarie stemisk when taken in doses sufficient to promote muscle gain are expected to suppress endogenous tostostenous podation. Without the intervention of testostenous-estimating substances, testosterone levels should return to normal within 1-4 most of dang secession. Note that prolonged hypograndshorpic hypograndsom can develop secondary to steroid abuse, recessfulng micral intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biovasiability. <sup>24</sup>Th its is caused by the fit-soluble nature of steroid hormones, which can allow some of the drug to dissolve with urdigasted distray far reducing as absorption from the gastroirestinal tract. For maximum utilization, this steroid should be taken on an empty stormeth.

# Administration (Men):

Clinical studies have demonstrated that significant nitrogen retertion and weight gain can be induced with a daily dossage of 1-2m gpe reds, In the atthict areas, dosses of 2-5 mg daily seem to be most reasonable, taken in cycles listing no more than 6-8 weeks is lingful to minimize hepatotoxicity. This level is sufficient for strong increases in muscle size and strength, althouch such axis will flesh be accommand by similificant water reterrition.

#### Administration (Women):

Bolssterone was not widely used with women in clinical medicine. When applied, it was most often used as a secondary medication during properable breast cancer, when other thrappies, have failed to produce a describe effect. The dosage used for this application would be as light as 10 mg per dog, a level that his caused significant visitations among patients. Educations in garantily not recommended for women for physique—or performance of the control o

#### Availability:

Bolasterone is no longer produced as a prescription drug, although a handful of underground laboratories have taken to selling this material.

# Nebido (testosterone undecanoate)

Androgenic	10
Anabolic	10
Standard	Standar

Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one	
Estrogenic Activity	moderate	
Brogostational Activity	low	

# Description:

Nebido 8 is an ijecubie steroid that contains testostorous undecensute, a very slow-seine, sets of Poststorous. This is the active dengthat is used in Audrich, but in that case it is part seathfield injectible Institutions products like Delitering/Ri. [Depo-Tistostorous], and statistically, which are actually much faster acting in comparison. It is designed to offer a much less froquers injection schedule, and therefore, much gener comfort for the princt, Nebido is a drug developed under a similar focus as testostorous bucklate, which is mother very slow-acting include sets or feedstated on.

# History:

Nebido 8 was developed by international gaint Schering AG. Germany (now Busper). It first surfaced as a prescription dug in Falinal and Germany in October and November of 2004, respectively. Within a year it had been approved for sale throughout Europe. Schering/Buper has since also broughf its pondact to Mexico, Brazil Agrantia, Oston, Brazil Agrantia, Davido Schering, Brazil Bra

Nebido's use described by Schring as being the, "first long-acting rijection for the teatment of rule hypogenetium." This may be a matter of perspective, as other slow-acting solicities of the solicities of the

# How Supplied:

Testosterone undecanoate (rijection) is available in various human drug markets. All products (Nebido®) contain 250 mg/ml of steroid dissolved in oit, packaged in 4ml ampules containing 1,000 mg of steroid in total.

#### Structural Characteristics:

Testosterore undecareate is a modified form of testosterore, where a curboxyle, earl ester (undecareai exist) has ent attacked to the T-best highoring poor, Estirifed forms of testosterore are less polar fram fee testosterore, and are absorbed more slowly from the area of irjection. Once in the bloodstram, the ester is removed to yield fee (activa) testosterore. Estarfied forms of testosterore are designed to prolong the window of thempartie: effect blowing administration, allowing for a last feegera tripiction schedule compared to irjectives of fee (unsetrified) steroid. Debitol's designed to ministai rhoisistical level of testosteror for tru to I week self are inection.

# Side Effects (Estrogenic):

Testosterone is readily aromatized in the body to estradiol (estrogen). The aromatase (estrogen synthetase) enzyme is responsible for this metabolism of testosterone. Elevated

estingan teels can cause side effects such as increased water returnin, body fit gain, and gracecomstin. Techniques returned to considered an andersady estinguist seroul. An arti-estinggan such as completure estinate or tumoritim clinate may be necessary to prevent entosquis side effects. One may adameted use an anomations inhibito like Artifactio (functionoid), which more efficiently controls estingen by preventing its synthesis. Anomation inhibitors can be effected on the other industriant to a transferage, however, and may also have negative effects on tabled fails.

Entogais side effects will ocur in a dose-dependant rumen; with higher doses (abovement) therapeut is evolo) resistancemen nee fleshy to require the concurrent use of mansestingum or aromatuse ribblios. Since water reterion and loss of muscle definition are common with higher doses of featurement, this drug is usually considered a poor choice for design or carting plasses of braining. Its medicante estrogenicity makes it more ideal for the medical properties of the properties of the properties of the properties of the art help foster a second armother demonstration.

# Side Effects (Andmoenic):

Textostence is the primary mile aurhogan, responsible for ministring secondary mile senul entacteristics. Extended level of textostence are Belot produce arridograsis de effects including only skit, acce, and bodyficial hat growth. Men with a genetic produposition for his loss (androgenetic adspect) may notice accelerated mile pattern hading. Those concerned about har loss may find a more confertable option in mandroses decentrate, which is a comparable lass analogenesis sented. Wemen are swarmed of the decentrate, which is a comparable lass analogenesis sented. Wemen are swarmed of the such as textostence. These may related, desposing of the voice, mentional irregulantees, changes in skit neutra, feed him growth on a discontainment of the contractions.

In authorps responsive traget fissus such as the skin, soal, and prostute, the high relative authorpsits of to statements in dependent on its relations to difficultiestosteneer (ERI). The S-alpha reductase enzyme is responsible for this metabolism of testosteneer. The concurrent use of 3-alpha reductase inhibitors such as finantized or distantiest will stretter with alse-specific potentiation of testostenees action, lowering the tendency of testostenees with the contraction of the stretch of the stretch of the stretch of the stretch of the damping responsible and the stretch of the st

#### Side Effects (Hepatotoxicity):

Teistoterone does not hue lepationoic effects, for toxicity is utilicely. One study ceanning the potential for lepationoicity with light does of teistoterone by administering 400 mg of the hormone per day (2,500 mg per weed) to a group of mule subjects. The steroid was also meanly so that higher past concentrations would be readed in lepatic tissues produced no significant changes in love receptor when the produced no significant changes in love receptor what including serum abunit, bilirabin, adminiscraminor—markens, and abulatine producted on.

#### Side Effects (Cardiovascular):

Anabolizandroganie steroide can hove deletroise effects on serum cholestered. This includes a referency to refere IML (good) cholesteroi values and increase IDL, (bod) cholesteroi values, within my shift the IDL. to IDL balance in a direction that fixous geneter risk of anteriocitosists. The relether impact of an anabolizandroganic serior of a security and expectation on the dose, route of administration (roit is, nigetable), type of steroid Arabolicandroganic serior in serior my last objection of the control of a serior deliveration of the control of the control of a serior deliveration of the control of the cont

Testosterone tends to have a much less dramatic impact on cardiovascular risk factors than synthetic steroids. This is due in part to is operaness to metabolism by the liver, which allows it to have less effect on the hepatic management of cholesterol. The aromatization of testosterone to estradiol also helps to mitigate the negative effects of androgens on serum testic. It can explain 200 me and of functionation of testosterone to estradiol also helps to mitigate the negative effects of androgens on serum testic. It can explain 200 me and of functionation of the part of the

spaces more starty, now may be worken to reconstructive near (customers) are a sugar on any assistancily significant effect on HEU. Obstacted after 12 works, but when taken with an amountaine inhibitor a strong (25%) decrease was soon. <sup>200</sup> Statless using 300 mg of the interactioncone seed construction per work for frontey nodes whitten an ascentize inhibitor exclude 12%. <sup>201</sup> The regards expect of anomalous eithibition should be taken into consideration before used thug as daded to beststormen therapy.

Due to the positive influence of estrogan on serum lipids, terroritor cirtus or chimiptone cirtus are perfective a) somemisse inhibitors for those concerned with encidensecular health, as they offer a partial estroganic effect in the leve. This allows them to potentially irreprove lipid profiles and offices one of the regative effects of anthegories. Whit doeses of 600 range to less per work, the impact on lipid profile tends to be noticeable but not dimunitie, imiliargu an anti-assignant for endoprotective purposes peripents unrecessery. Does of 6000 rap of less of 6000 rap of 1000 rap

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the ristake of saturated fits, cholsterol, and simple carebolydrates at all times during active AAS administration. Supplementing with shi old, games per day) and a natural cholstero/lamioxidant formula such as Lipid Stabil or a product with comrangible inserdenies is also recommended.

#### Side Effects (Testosterone Suppression):

All antibolizationgosis stensis when taken is does sufficient to promote muscle gain are expected to suppress condigenous testostores production. Testostores is the primary male androgen, and offers strong negative feedback on endogenous testostoreus production. Festostoren-besed diago will. Reviewie, hurse a strong effect on the phytodhanic regulation of natural steroil hormous. Without the intervention of testostoreus stimulating substances, testostoren-besed bond tentrum formulation il –1 morths after the edge has filly cleaned the body. Note that produced phytogendostropic hypogenedism can develop secondary to storal disease.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

Due to the large injection volume, prescribing guidelines recommend that each injection be given slowly, taking approximately 60 seconds to administer the full 4ml dose. Nebido® should always be injected deep in the altheat muscle.

# Administration (Men):

To rotat andogan instificancy, the proscribing guidelines for testosterore undocamote (Oschadri) call for a douge of 1,000 mg (eff) encey probe useds. Therapy is usually observed to the control of the

### Administration (Women):

Testosterone undecanoate is not approved for use with women in clinical medicine. This drug

is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic mature, tendency to produce virilizing side effects, and very slow acting characteristics (making blood levels difficult to control).

#### Availability:

Testosterone undecanoate injection continues to increase in prominence as a pharmacutical product. It is presently approved for sale in 86 countries worldwide. In reviewing some of the more popular products and changes on the global pharmaceutical market, we have made the following observations.

Nebido gained approval for Europe-wide sales in 2005. The product is has since distributed throughout Europe, and is widely available in this region.

Indexus, a subsidiary of Endo Pharmaceuticals, has continued to push for FDA approval of Aveed in the United States, but such approval has not yet been granted.

# Nilevar® (norethandrolone)

Androgenic	22-55
Anabolic	100-200
Standard	Methyltestosterone (oral)

Chemical Names	17alpha-Ethyl-17beta-hydroxyestr-4-en-3-one 17a-ethyl-19-nortestosterone
Estrogenic Activit	y high
Progestational Ac	tivity high

# Description:

Noerdundolose is an anabolis steroid closely related to nortestosterone (nantrolose) in structure. The activity of this storoid is fred a final broncheron calculosels storoid, which is accompanied by distrugationable antioquate and estrogatic components. Although this storoid is sensiting intendolore medified closels ofto mice and only which is cause the looked at sneph as an oral adventure to Dece-Duraholife. Most notably, the greatly increased estrogatively caused by 17-alsylation makes noredundolore much more problematic when trying to badd quality (tunn) muscle musc, in administering an effective contagnit side effect. The mursch excurations with noredimone is also ging to be accompanied by a high level of water and (file-fly) for reteriors, not the quality musclarity normally associated with annolorus decuments.

#### History:

Nordmurkohen was first described in 1954.<sup>50</sup> It was developed into a medicine by Searle, which irraduced it no the U.S. perception dung mustle unter the Nilouv brand must during the late 1950's. The drug was originally sold as at cent tablet, an cost solution recording the late 1950's. The drug was originally sold as at cent tablet, an cost solution recording to the late of the control of the late of the late

Norethandrolone ultimately saw only limited success as a prescription anabolic agent. It did make its way to Europe and certain other markets, but not widely. The drug was an early functional anabolic, displaying more tissue-building properties than androgenic effects. But it also remained an agent with a troubling estrogenic side. This eventually led to nerelandolore being passed over clinically for more refund compounds as they became available. Searly decided to discontinue the sale of Nelsour in the U.S. charge the 1960's, and instead began focusing energies on its newer, more strongly anabolic, and non-estrogenic searced countrolores (cold an Anarous). Most other markets comprige merelandolorus, either by Searle or other computers, soon began losing this compound as well. Today, this drug is verticated to the computer of the compound of the compound as well. Today, this drug is verticated to the control of the compound of the compound as well. Today, this drug is verticated to the compound of the compound o

# How Supplied:

Norethandrolone is available in select veterinary drug markets. Composition and dosage may vary by country and manufacturer, but typically contain 5 or 10 mg of steroid per tablet.

# Structural Characteristics:

Norethandrolone is a modified form of randrolone. It differs by the addition of an ethyl group at carbon 17-alpha to protect the hormone during oral administration.

# Side Effects (Estrogenic):

Noedmandonies is arountized by the body, and comerts to a synthetic entrops with a high best of hological activity (Traphen-off-bestandio). As a rouse, it is a highly entropses stand, Opencoments is often a concern during testimets, and may present feel quide early become a problem; consider a rouse of the contract of the first in a both subcutamous water reteriors and fit levels build. Sensitive infloidates may want to keep the entropia trader control with the adults of man arise-tensport and in Nobadoed. On term judemalay use an automatic inhibitor like Armindood (manstrooxide), which is a more effective entered by the adult of the control of the standard control mutuatures of the control of the control of the control of the control of the standard control mutuatures of the control of the control of the control of the control of the standard control mutuatures of the control o

It is of note that northumbroken has some additional acidity as a progestin in the body. <sup>548</sup> The cide effects socialed with progestome are similar to how of estrongs including reagative Reducke inhibition of testodence production and enhanced me of fix storage. Progestin also augment the similarity effect of estrogens commany issue growth. These progests also augment the similarity effect of estrogens commany issue growth. These progests are the similarity effect of estrogens common implies even core with the high of progessis without excesses estimpts needs being reserved. The use of an anti-estrogen, which inhibits the estrogenic component of this disorder, is often satisfactor to risting exconomation cannot be constructed as a final state of the progression of the estrogenic component of this disorder, is often satisfactor to risting exconomatic acused by reord-tunatolone.

# Side Effects (Androgenic):

Albeagh closified as mambols steroid, androgenic side effects are still common with the substance. This may head be tout of obj skin, eare, and bodyficied link in growth Arabolic/androgenic steroids may also aggreate melt pottern hair loss. Individuals sensitive to the antrogene effects of this steroid may find a maker anabolic such as Deca-Darabolinic to be more confortable. Women mer additionally warned of the potential wirling effects of the property of productings, changes in sides moutes, easily inter more load of fent good intercents.

Note that is androgen-responsive target fissess such as the skin, scalp, and prostate, the entirities androgenisty of noncedimetholous is reduced by its reduction to dilpdronorethandrokoue. The 5-aphra reductases erropme is responsible for this metabolism. The concurrent use of 5 a-light necleatures eithilities such instructive or datasetied will interfere with sto-specific reduction of norethandrokoue action, considerably increasing the world with the store of minimization by ordinary ordinary to dissipate.

#### Side Effects (Hepatotoxicity):

Norethandrolone is a c17-alpha alkylated compound. This alteration protects the drug from deactivation by the liver, allowing a very high percentage of the drug entry into the bloodstream following oral administration. C17-alpha alkylated anabolic-antrogenic steroids can be hepatotoxic. Prolonged or high exposure may result in liver damage. In rare instances the description of the descripti

ne-treatment uponataxis may ucessay, it is assessine to visa a prost-tain personation turns, each cycle to morabine fiver function and overal health, finish of C17-aphin addyned steroids is commonly limited to 6-8 weeks, it in effort to avoid escalating foer stain. Severe foer complications are are agoint the periodic matter in which most people use oral arabolicantrogenic steroids, although cannot be excluded with this steroid, especially with high doses and/or prolonged administration periods.

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any herototoxic anabolic/androgenic stempils.

## Side Effects (Cardiovascular):

Anabolizatinfogunis atensiis en have deletroise effects on serum cholesterel. This irelates a relendary to reface IEML (good) cholestered values and increase IEML (sood) cholestered values and increase IEML (sood) cholestered values which may delf the IEEL to IEM balance in a direction that floors greater risk of anteriocitosists. The relente regard of an anabolizatindense resont on serum lipite is dependant on the dose, route of administration (roat us, nigetable), type of steroid (contratable or non-normaticale), and level of resistance to happine metabolization contratable and the particular contratable resistance to have bread the particular contratable resistance to her breaddown and route of administration. Anabolization departs execution may also above also related to the processor of the processor and traffectories, reduce enthethic relaxation, and support left verificale hyportrophy, all potentially increasing the risk of continuous called reason and reporation illustrations.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and maintaine the intake of sastnated fists, cholssterol, and simple carbohydrates at all times during active AAS administration. Supplementing with sho is 4c grams per day) and a natural cholssterolaritoxidant formula such as Lipid Stabil or a moder with contrasphel insordiers is also procurement.

# Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone strainfating substances, testosterone levels should return to normal within 1-4 moralis of drug secession. Note that prolonged hypogoradorophic hypogoradism can develou secondary to seroid abuse, necessistatine medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

# Administration (General):

Studies have shown that taking an oral arabolic steroid with food may decrease its biovariability.<sup>555</sup> This is caused by the fat-solible nature of steroid hormons, which can allow some of the drug to dissolve with undigested dietary fat, reducing its absorption from the gastrointestiral tract. For maximum utilization, this steroid should be taken on an empty stormeth

## Administration (Men):

The original prescribing galicities for Nibara called for a daily dosage of 220 to 30 mg. This was to be administered on an intermittent bous, with the daily taken for nor meet hard; consecutive weeks. Thereafter, a break of at least 1 menth was advised before freamy was research. When used for physicage on performance-efficienting prayers, the daily at doses as the contract of the many transportation of the contract of the co

# Administration (Women):

The original prescribing guidelines for Nilevar made no special dosing recommendations for

women, although à did warn that androgenicity is Bicly on a high dosage. When used by women for physique- or performance-enhancing purposes, a daily dosage of 5-10 mg is most common, taken for no longer than 4 weeks. This level is quite effective for promoting new mack: growth. Note that virilizing side effects are still sometimes noticed at lower doses, and need to be carefully evantmend for.

#### Availability:

Pharmaceutical preparations containing norethandrolone remain scarce, and are rarely diverted for black market sale. The only region of note where this compound is still made is Australia.

## Omnadren® 250 (testosterone blend)

Androgenic	100
Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate

#### Description:

Progestational Activity

that contained four different testosterone esters: isotosterone propionate (30 mg); testosterone perspective (60 mg); testosterone locaproate (60 mg); testosterone perspective (60 mg). Estestosterone locaproate (60 mg) and testosterone curroate (600 mg). Being a face-composed testosterone being, time perspective saws most contravely correspond to Santanesti. 20 MHs et all contrast inconsectioner preprinted and contravely composed to Santanesti. 20 MHs et all contrast inconsectioner preprinted different. It was a slightly shorter-acting drug militage (Demidered) mere analogue to Extensional Composed sector from certainly than Santanesti. 20 Messes note that there were even older versions of Ormaderelli faing solvenances and homostice as the failth to ingredient, which are arispid different which for incorporate and the contrast are the failth to impedient, which are arispid different which for incorporate and the contrast are the failth to impedient, which are arispid different works for incorporate and the composed are the failth to impedient, which are arispid different works for incorporate and the composed are the failth to impedient which are arispid different works for incorporate and the contrast are the failth to impedient which are arispid different works for incorporate and the composed and the composed are the composed and the composed are the composed and the composed area.

Omradren® 250 (in its original formulation), was an oil-based injectable testosterone blend

low

#### History:

Omndero® 250 was developed in Poland by Polls during the years of Sovict control. Its fermitation (original) wery suiter that ord Santamo® 250, bursting the substitution core of the component exters. This was likely done to novid patert issues with the international phermacentrial gaint tropane, which exclusively controlled the plots apply of Statustrolle 250. In clinical medicine, Ormsdero® 250 was used most commonly to treat adult men saffering from low antrogen levels, usually notice; syngtowns of protector or Ironardo distributors of spermitogenesis. This drug was also used on occasion to treat adolescents with delebyed probers, and women with adonned breast or endometrical cuerce.

The munificative of Ormadeum? 290 under the Polis likel was discortanced in 1994. That you, the newly princised Polis firms was remard felle, minju by dategash heel' from other firms that tase a Polis prefix as part of their manes. Jolis continued to produce Ormadeum? 290 for the demonsis medic, which entermide available whost retemption in the same familiar Speack of arappias, follow with a new company held and logo) for years after. Toolay, Sells contrasts to mracel Cormadeum? 250 in Polandi, as well as in many neighbored. Fooliay, Sells contrasts to mracel Cormadeum? 250 in Polandi, as well as in many neighbored. America, Modelon, Larixi, Lilmann, Azerbijan, Goopia, and Beltons, however the formation has recordly changed. All Ormanne 250 will today orients the same coast formation as Satamon 250. This profile refers to the original formulation only, which is now unsuable workflowly.

# How Supplied:

Omradren® 250 (original formulation) in no longer available. When manufactured, it was supplied in 1 mL glass ampules containing an oily solution; sold in boxes of 5 ampules.

# Structural Characteristics:

Omnderné 250 contains a même or fiser instinctione compounds, which where modified with the addition of contrologic acid easiers (projectic, propinier, projectic, pr

#### Side Effects (Estrogenic):

Textostence is readily arountized in the body to estatiol (estrogen). The arountase (estrogen synthesise) errors is responsible for the metabolism of textosterone. Elevation (estrogen index estrogen include and estrogen

Estogasis side effects will occur in a doss-dependant muner, with ligher doss faltows normal flustrapactic levels) of restolatore more fleely to require the concurrent use of an autiestingum or aromatuse sithibute. Since water reterion and loss of muscle definition are common with light doss of festolatories, this drug is usually considered a poor choice for design or curing places of training. Its moderate entrogenicity mules is more ideal for the contraction of the size and help forter a storout analysis contraction.

# Side Effects (Androgenic):

Testoderore is the primary mile androgan, responsible for matriating secondary mile seasul characteristics. Entental else of festosteriore are Belley to prockee androganis designed to the control of the procession of the into time deposition for his the isomologousine for his the isomologousine of his the isomologousine of his into isomologousine point in the isomologousine of his into isomologousine of his into isomologousine or his into isomologousine or his more intentional to intentional to the intention of the intention of

In androgo-responsive traget tissues such as the skin, scale, and prostate, the high relative androgoricity of testimeteries is dependent on its relation to displostostenetree (DRI). The 5-siphs reductase enzyme is responsible for this metabolism of testisteneue. The concurrent use of 3-shall practices arithmic such as institution of classifications. The design produce arrivages sized effects. It is important to remember that analotic and analogous effects are both medical via the cytosolic analogous recognition. Complete responsibility of the complete of the complete of the cytosolic analogous recognition complete, even with a complete analogous effects are both medically via the cytosolic analogous recognition (experience).

#### Side Effects (Hepatotoxicity):

Testosterone does not have hepatotoxic effects; liver toxicity is unlikely. One study examined the potential for hepatotoxicity with high doses of testosterone by administering 400 mg of the hormone per day (2.800 mg per week) to a group of male subjects. The steroid was taken orally so that higher neak concentrations would be reached in henatic tissues compared to intramuscular injections. The hormone was given daily for 20 days, and produced no significant changes in liver enzyme values including serum albumin, bilirubin, alanine-amino-transferase, and alkaline phosphatases 556

# Side Effects (Cardiovascular):

Anabolic/androgenic steroids can have deleterious effects on serum cholesterol. This includes a tendency to reduce HDL (good) cholesterol values and increase LDL (bad) cholesterol. values, which may shift the HDL to LDL balance in a direction that favors greater risk of arteriosclerosis. The relative impact of an anabolic/androvenic steroid on senum limits is dependant on the dose, route of administration (oral vs. injectable), type of steroid (aromatizable or non-aromatizable) and level of resistance to heratic metabolism Anabolic/androgenic steroids may also adversely affect blood pressure and triglycerides. reduce endothelial relaxation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and muncardial infarction

Testosterone tends to have a much less dramatic innact on cardiovascular risk factors than synthetic steroids. This is due in part to its openness to metabolism by the liver, which allows it to have less effect on the heratic management of cholesterol. The aromatization of testosterone to estradiol also helps to mitigate the negative effects of androgens on serum linids. In one study, 280 me ner week of testosterone ester (enanthate) had a slight but not statistically significant effect on HDL cholesterol after 12 weeks, but when taken with an aromatase inhibitor a strong (25%) decrease was seen 557 Studies using 300 mg of testosterone ester (enanthate) per week for 20 weeks without an aromatase inhibitor demonstrated only a 13% decrease in HDL cholesterol, while at 600 mg the reduction reached 21% 558 The negative impact of aromatase inhibition should be taken into consideration before such drug is added to testosterone therany.

Due to the positive influence of estrogen on serum lipids, tamovi@n citrate or clominhene citrate are preferred to aromatase inhibitors for those concerned with cardiovascular health. as they offer a partial estrogenic effect in the liver. This allows them to notentially improve lipid profiles and offset some of the negative effects of androgens. With doses of 600 mg or less of testosterone per week, the impact on lipid profile tends to be noticeable but not dramatic, making an anti-estrogen (for cardioprotective purposes) perhaps unnecessary. Doses of 600 mg or less per week have also failed to produce statistically significant changes in LDL/VLDL cholesterol, triglycerides, apolipoprotein B/C-III, C-reactive protein, and insulin sensitivity, all indicating a relatively weak impact on cardiovascular risk factors. 559 When used in moderate doses, injectable testosterone esters are usually considered to be the safest of all anabolic/androgenic steroids.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterol/antioxidant formula such as Lipid Stabil or a product with comparable ingredients is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Testosterone is the primary male androgen, and offers strong negative feedback on endogenous testosterone production. Testosterone-based drugs will, likewise, have a strong effect on the hypothalamic regulation of natural steroid hormones. Without the intervention of testosterone-stimulating substances, testosterone levels should return to normal within 1-4 months of drug secession. Note that prolonged hypogonadotrophic hypogonadism can develop secondary to steroid abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Restosterone proporate is often regarded as a pantial spectron. This is due to the very short carbon chain of the propionic acid ester, which can be irritating to tissues at the site of injection. Many sensitive individuals choose to stay away from this steroid completely, their bodies reacting with a pronounced soreness and low-grade fever that may list for a few days after each injection.

#### Administration (Men):

Deporting on the application, the prescribing guidelines for Ormadentile 250 call for a decape of 250 upg (1 a maghs) to be rejected over § 26 at weeks. Although action in the body for a larger time, Ormadentile 250 is usually administered on a weekly basis for muscla-balling purposes. Bits schedule will allow for the lajder does not cormenty applied by attlices, and more stable desentions in hormone level. The usual dosage arrange in additions in the mages of 250–250 map to rejection, bean in regules to level as might, Some work of the stable of the stabl

## Administration (Women):

Ommdren® 250 is rarely used with women in clinical medicine. When applied, it is most often used to treat inoperable breast or endonetrial cancer. Ormsdren® 250 is not recommended for women for physique- or performmence-ordinaring purposes due to its strong androgenic matter, tendency to produce virilizing side effects, and slow-acting chemeteristics fundine blood back of difficult to control.

# Availability:

The original Ormadren 250 formulation is no longer available. Jelfa continues to use the trade name to market a steroid product, but it is now equivalent in makeup to Sustanon 250. See the Sustanon 250 notifie for more information.

20-400

200-400

# Orabolin® (ethylestrenol)

Standard	Methyltestosterone (oral
Chemical Names	19-Nor-17alpha-pregn-4-en-17b-o 17alpha-ethly-estr-4-en-17b-o
Estrogenic Activity	/ lov
Progestational Act	tivity high

# Description:

Androgenic Anabolic

Ellishestenol is an oral anabolis steroid derived from marchone. As is typical for many 13ror storoids, this agard cubbine for greater anabolic proporties from anadosparis, is other woodly elstoparis, and is strongly progestational. Strutamily, ellipsistenol must closely resembles Nature (northanadosus). The roof fifter only by the absence of an orogent and at the cip position of ellipsistenot, and in the body ellipsistenol satually has a rotable affinity to cornect to noretunization. <sup>2007</sup> The part of metalodium is responsible for much of the anabolic, analogustic, and estrogent activity was esse with this composat, and must regards strongly anabolic enfective to its authorization, efficites generally fifth sits sector lot be externelly weak. The level of muscle growth obstaned with its steroid is generally much less reduced the further operated with effect Newton or Deca-Durcholis, and it is considerably less effective than both stanozolol and oxandrolone on a miligram for miligram basis.

# History:

Eliphestreoil voss fint described in 1999. Sei. It was developed into an oul mulcine by Organo (now Mexch.XXIX), apporarig in morthes between 1981 and 1984. Organo sold the tablex inder the trade name Mexcholin in the U.S., and as Orabolin, Oughoin, and Dambolin-O in other meakers. The latter man is a compressed from of "Dambolin-O iside the meakers. The latter man is a compressed from of "Dambolin-O iside (India) (Ind

Ellylosterenol became a steroid of great controversy during the early 1980's, when Westerm much autorises was given to the marketing of the drug to malworished children in 'Bud-milland's and the state of the drug to malworished children in 'Bud-milland' chiested the flow would "bey children gain fall voigit and height." Smithest physiological appetite," and "muses optimal assimitation of bood." It also described a "bickious [maghput sypt paster children boo: "In main point of contention was the promotion of an authoric section to text the lack of adequate food apply, the real issue at expension of an authoric section to text the lack of adequate food apply, the real issue at exception of an authoric section to text the lack of adequate food apply, the real issue at exception of an authoric section of an authoric section of a real to the company soon discontent of Fertibols and related miratering practices. Manabola and Mackhon Elsis were voluntarily withframs from the U.S. market during the late 1980's as well, and may be without the control of the section of the company of the control of the contro

# How Supplied:

Elhykstreol is available in select humm and veteriuny drug markets. Composition and dosage may vary by country and munificature, but typically contains 2mg of steroid per tablet. Oral solutions have also been produced in the past, such as Macholin Elicit, which contained 2mg/5 mL in a 4 ounce bottle. Fertabolin for children contained .2mg/2 mL of solution.

#### Structural Characteristics:

Ethylestrenol is a modified form of randrolone. It differs by: 1) the addition of an ethyl group at carbon 17-alpha to protect the hormone during oral administration and 2) the removal of the 3-covera.

#### Side Effects (Estrogenic):

Elly-Strool is arountized by the body, and concerts to a synthetic estrogen with a high level of hological active, [Tajeba-end-serration, Rate of arountization to both, bowere after it at a remine; clossified as a weakly entrogenic steroid. Optocorment is possible during troutment, but generation only when higher does are used. What and fit reterion can also become issues, again depending on dose. Sensitive infiviluation may need to addition an artisengem and in solvadore. One may alternately use an arountees within the K. Artifición (unastronolo), which is a more effective remedy for entagen control. Atomatica inhibitor, and may do be the contracte effects on blood indicational contegrous multiments ellepora, and may do holy one canten effects on blood indicational contegrous multiments.

It is of note that only/settered has strong activity as a progestin in the body. <sup>502</sup> The side effects associated with progesterous resimilar to those of estimpor, including negative feedback sibbition of institutione production and enhanced rate of fits strongs. Progestine also augment the simulatory effect of estimposes on marrowing losse growth. These appears to be a strong puregy between these two harmons here, such that genecomation right even to be a strong puregy between these two harmons here, such that genecomation right even an artis-stroner, which withinks the estroneric convenent of this disorder is often sufficient to mitigate gynecomastia caused by this steroid.

# Side Effects (Andmoenic):

Albeagh closified as manibols strootd, androgenic side effects are still common with this selection. The major skelde boats of 60 yls siz, nexe, and bodyficial that irg growth. Anabokichardogenic steroids may also aggreate melt pottern hair loss. Individuals sensitive to the antrogene effects of this stroot imp first a milder amboke sarboke such as Deca-Darabolitic to be more comfortable. Women mer additionally surned of the potential wilding effects of modelocated opposite seroids. These may retail as deepwing of the voice, measured and the stroot of the contract of

Note that is androgen-responsive target fissess such as the skin, scalp, and prestate, the entire androgenity of ethylsterous is reduced by as reduced by the reduced to weaker "tilishyo" metabolises. The 5-alpha reductuse engome is responsible for this metabolism. The concurrent use of 3-alpha reductuse inflines such an instruction of unfastentive will interfire with side-specific reduction of ephsteronial action, increasing the tendency of the dings to maintaining low reduction androcative join desired.

# Side Effects (Hepatotoxicity):

Ellydetrool is a CT-spha alykud corporad. This alteration protects the dung from detectation by the five allowing a very high processing of the dung error to the bloodstroom flowing certainties. CT-spha algked area-bloodstroom flowing certainties are beythetics. Frobrought of high crossing of the spin and being from its interest. Bid-streaming dysfurction may develop. It is advisable to visit a physician protectable distriction may develop. It is advisable to visit a physician protectable distriction from the context of the c

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any herastotoxic analysis and movemic stemicls.

# Side Effects (Cardiovascular):

Anabolizationgonic servicio en hovo deletricos effects on estem cholector. Il hi richales a nederacy to ratee III. (good cholector) values and increase III. (had) cholectorica values, which may shift the HDL to LDL balmee in a direction that fixors geneir risk of antirocherosis. The relation terms of anabolizationgonic servicio of a serma lipits is dependant on the dose, route of administration (end xx. ijectobile), type of steadd (generalization connomitables), and the off resistance to hapter entablement libitations has a strong effect on the hepatic management of cholesterol due to its structural resistance to five breakdown and rout of administration. Arabolizationgonic stemula, may also admentsy date the long research are in inglecentles, roduce endothed conference of the conference of th

To help reduce cardiovascular strain à is advised to muitain an active cardiovascular cuercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterolamioxidant formula such as Lipid Stabil or a product with compranibe inserdents is also recommended.

# Side Effects (Testosterone Suppression):

All ambolic/androgenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainfairing substances, testosterone levels should return to normal within 1-4 morths of drug secession. Note that prolonged hypogoradorophic hypogoradism can develou secondary to steroid abuse, necessistation medical intervention.

The above eigh affects are not inclusive. For more detailed discussion of notantial side

# Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biavoushibity. Stal This is caused by the firt-soluble nature of steroid hormones, which can allow some of the drug to dissolve with undipsted dielary fit, reducing is absorption from the gestiontestinal tract. For maximum utilization, this steroid should be taken on an empty or the state of the st

# Administration (Men):

Organ prescribing guidelines recommend a dosage of 4 mg to 8 mg per dup, them for no more than 6 concentie weeks. After a beautife of weeks, the fugs is resumed for a additional 6 weeks if indicated. When used for physique- or performance-enfrancing purposes, a duly dosage of 20 mg to 40 mg is not corroras, which equates to ten to tower. Yang tables. The drug is typically used in cycles lusting no longer than 6.8 weeks, in an effort to marriare lepails: searn This best of sattlered for some measurable gains in marcels teen and strength, although experienced serioral uses are likely to still the disappointed with the results. Instead of irrecenting the dosage, most opt to add a second serior of the cycle, usually an injectable such as teactorierors ceptomet or entangent, boldcomer underleanne, or obtained to the contraction of the contrac

# Administration (Women):

Original prescribing guidelines recommend a dosage of 4 mg to 8 mg per day, taken for no mere than 6 consecutive weeks. Aft as break for 4 weeks, the day is resumed for an additional 6 weeks if indicated. When used for playsique- or performance-ordancing proposes, a daylo basege of 10 mg to 10 mg is most common, taken for no larger flund weeks. This level secure to be fairly effective for promoting new markes growth. Higher dosses at leady to produce vidings date effects, and are not recommended. Note that wisting side

# Availability:

Pharmaceutical preparations containing ethylestrenol remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

At the present time, the legitimate supply of ethylestrenol appears to be isolated to Australia, where it is found in a small number of veterinary compounds including Nandoral tablets and Nitrotain rest.

# Oral Turinabol (chlorodehydromethyltestosterone)

Androgenic	no data available	
Anabolic	>100	
Standard	Methyltestosterone (oral)	
Chemical Name	4-chloro-17a-methyl-17b- hydroxyandrosta-1,4-dien-3-one	
Estrogenic Activity	none	
Progestational Activit	y no data available (low)	

### Description:

Chlorodelisalromethyltestosterone is a notent derivative of Diagohol. This oral steroid is

straturally a cross between redundenteerables and clossibed (4-clienteerabenteerae), having the same base sertureae as Diemblo with the dadied 4-client patterniae of clienteerae between the cost of the control of clienteerae between diemblo control patterniae of clienteerae serviced displaying not entoquisit and an unbe less andregatic activity in comprison to its more famous counterpart. The arthoic activity of clienteeling-incorporation to its more famous counterpart. The arthoic activity of clienteeling-incorporation are marked from the control of arthoic testing and the clienteeling of the first femous families and marked from the clienteeling of arthoic testing and the clienteeling of the clienteeling of marked-incorporation will be talked by produce andregation still effects.

# History:

Chacodelydromelyheisotstence use firt described in 1962.<sup>566</sup> Iomplamm (Jen., Gemmy) soon the released for dag for she in the East German pection the respective of the most cannot be released for the first form processive of the most cannot be released for the product the second by cheixen for its highly analose and to wondook much seeling forth one in root op which miss, bet somer and challon as well. The product uses mendacted in two strengts, containing I may all free deather one of the product uses mendacted in two strengts, containing I may all free for damp per tables, not an absort-door seems uses analoble for the more excessive populations. Chicodelydromelybetostence was appeted for a marker of modical uses; must do the product of the per tables of the per tables.

One Turishol became a steorid of infinity during the 1990%, when it was revealed that chlorodephorenthylesotatorea had been one of the closely had secent issistic the Tase German Daping Machine." This is referring to the state-sponsered deping program, caled "State Plan Research Them et 1425," for the operated in Fast German Polyage Workshop and 1990. It was an aggressive anabolic steorid administration program, designed with one goal in mind chearing the Opping drug seat in many cases, the Opping withstee, both made famile, were unwriting participants, simply boil by their trainers and conclus that they were been given "tearns." Many of those the various transit natured out to be Out Turnshot, a porture anabolic startels, during the time the program was sides, many of them taking Oral Turinshol. For a mee in-depth lock at the demantic laster ever, leading the trials of several forme East German officials for their participation, I recommend you look at the book "Fasts" Gold breist the East German Drigin Mortal "De Spece Hungerheiter."

In spike of an anguably favorable profile of activity and safety ecord, Jerupham discontrased Orull Turishois 1 rayle. This was at a true when a good and of regime attention was being goes to opon doping, leading couldings to the speculation that this decision was one based Regulacks, Integration was expected by Selecting And Germany 1 rayles. A company with no starest in releng the controversies of the past Selecting Intel already discontrased many of a controversible and another startly ordered as well. Before or start, one of their lead of chinocolfidwareful-biototerous lun existed as a preception ding product. Today, the analysis of the startly and the startly a sum affairs for of startly a sum affairs for of startly a sum affairs for offstagood introductions.

# How Supplied:

Chlorodehydromethyltestosterone is not available as a prescription drug product. When manufactured, it was found in 1 mg and 5 mg tablets, sold in Germany/German Democratic Republic.

# Structural Characteristics:

Chlorodelydromethylsetosterone is a medified form of testosterone. It differs by; 1) the addition of a meltyl group at action 17-palps, which helps protect the hormone during oral administration; 2) the introduction of a double bond between carbons 1 and 2 (1-ene), which shifts the anaboles to androgarie ratio in four or the former, and 3) the attachment of a chion group at carbon 4, which inhibits steroid aromatization and reduces relative antogariety.

# Side Effects (Estrogenic):

Chlorodelnydromethyltestosterone is not aromatized by the body, and is not measurably

estrogenc. An arti-estrogen is not necessary when using ins steron, as geneconsists srout not be a concern even among sensitive individuals. Since estrogen is the usual culprit with water retention, this steroid instead produces a lam, quality look to the physique with no for of excess subcutamous fluid retention. This makes it a favorable steroid to use during cutting cycles, when water and fit retention are major concerns.

#### Side Effects (Androgenic):

Albudgs chenodelyndromethylesostence is classified as an arabolic stroot, androgare siste effects are all possible with its substance. These may include boost of oy six, next, and loodyficial last growth. Does higher than normally prescribed are more likely to cause such side effects, wholchochrandegaries storils may also aggravate made pattern lar its Now. Worms are additionally warned of the potential visiting effects of arabolicandrogarie stroots. These may relate a deepening of the own, centration application, changes in sink teature, fixed lart growth, and chiefen elettrational confloodify-dromethylesostosome is not country for the confloodify-dromethylesostosome is not country for the confloodify-dromethylesostosome is not country all out of the confloodify-dromethylesostosome is not country dromethylesostosome is not country dromethylesostosome in the country dromethylesostosome in the country dromethylesostosome is not country dromethylesostosome in the country dromethylesostosome is not country dromethylesostosome in the country dromethylesostosome is not country dromethylesostosome in the country dromethyleso

# Side Effects (Hepatotoxicity):

Charodisphormefiphenosterores is a c17-alpha alphated compound. This alteration protects fred ingo inducational by the local along as very large processes for disp man discussional to the cash downs as very large carry in the bloodsteam following and administration. C17-alpha alphated free control of the carry of the control of the carry of

#### Side Effects (Cardiovascular):

Anabolican/longonie steoride can hove debetrious effects on serum cholestered. This includes a referency to refere LHZ, Good cholestered values and irraces LL, Chol chol cholestered values, which may shift the HDL to LDL balance in a direction that fixous goater risk of anteriocherous. The relates irraped of an anabolican/longonie steoriol on sextra light is an anabolican/longonie steoriol on sextra light is considered to the consideration of the

To help reduce cardiouscular stania it is advised to mintain an active cardiouscular cerevise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4) games per day) and a natural cholesterollaritoxidant formals such as Lipid Stabil or a product with comparable ingedients is also recommended.

# Side Effects (Testosterone Suppression):

All ambole/andrugenie steroids when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosterone production. Without the intervention of testosterone-strainfairing substances, testosterone levels should return to normal within 1-4 months of drug secession. Note that prolonged hypogoradorophic hypogoradism can develou secondary to steroid abuse, necessistation medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Studies have shown that taking an oral anabolic steroid with food may decrease its biovanishibity. 560 This is caused by the fat-soluble nature of steroid hormones, which can allow some of the drug to dissolve with undigested dictary fat, reducing its absorption from the gastroinestinal tract. For maximum utilization, this steroid should be taken on an empty stomach.

#### Administration (Men):

A corrum chicail dose of clinochelphotenothylescostores is estimated to be 5 mg per day, studin prescribes gladiens are unswalbels. In the athlikat eram, anticheric oral day dosage fils in the range of 1-50 mg, taken in cycles besting no more than 6.8 weeks to minimae hepatroxisty. This level is sufficient for measurable increases in learn mache mass and strength. This agent is most often applied as a pre-context or cuting steroid for boolpholating purposes, and is not tweed as an ideal hading agent the to be keld or entegrately, of their in sports where speed tends to be a primary focus also find strong first in the challed places and in an extremely considered in the contribution of the contribu

# Administration (Women):

A common clinical doss of chlorodelsystemethylestosterous is estimated to be 1-2.5 mg per dog, schall perceiving galichies are unvasible. In the athletic aren, women would commonly take a sigle 5 mg tablet per day, taken in cycles lasting no more than 4-6 words to minimare hepstorociety. Writings effects are utality by this lived of with Mich ligher doses were often used with fermit athletes in the former GIR doping program, but often to destricted of survey with the size of the size of the size of the destricter of structs within 6-30 mg.

# Availability:

Chirocleptomethylestosteone has been unvaliable as a prescription drug product in formumy (the side courtsy of munificative firm sort of its bistory) since 1994. A very small number of planmacutical companies have marketed the drug since, minky in less regulated markets of Esteate linope and Asia, where black market dreams all inflameses production. In reviewing some of the remaining planmacutical products and recent changes on the sidual planmacutical market, we have mude the following observations.

Balkan Pharmaceuticals (Moldova) makes the product Turanabol. It is prepared in 10 mg tablets. 20 tablets per foil and plastic strip.

## Oreton (testosterone propionate)

Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activit	y moderate
Progestational Ac	tivity low

# Description:

Androgenic

Testosterore propionate is a commonly munificanteel sjectable form of the primary male androgan testosteroe. The added propionate ester will show the rate in which testosterone is released from the rijection site, but only for a few days. Testosterone propionate is, therefore, companishedy much faster-acting than other testosterone estero such as expirate or enumbrate, and requires a much more frequent dosing schedule. By most accounts testosterone propionate is an older and cruder from of injectable testosterone, made obsolete by the slower-acting and more comfortable esters that were developed subsequent to it. Still, those who are not bothered by the frequent injection schedule find testisterone propionate every bit as acceptable. As an injectable testisterone, it is a powerful miss-building drug, capable of producing rapid gains in both muscle size and strength.

# History:

Testosterore propionte was find electrical in 1935, during a series of experiments that set out to increase the fittenpoetic uselfances of testosterore by during in reluese in the bloodsterom. We flow posses that respectively. As in Germany would introduce the first testosterore proprietine product under the band name Testosionice. Propionitus was able the first commercially available injectable sets of testosterore on the U.S. prescription drug market, and remarked in dominant from Gistosterore globally often 1900. The drug first face and 1950%, for example, when strends were first being experiment with by small methylesisterome, to selectioner proprietine was also available in orally administered (faceal) preparations, but they disrepted motion to U.S. market during the 1980%.

Early prescribing gaidelines for tostourous programs called for a number of florasposition, used it was made applied to cases of mise authorages insufficiers, and those issues normally surrounding low instouterous levels such as robused so calle and impostness modals, and coppositionfilm (unlesscooled instition) in recognition of the robust four it also has a substantial of the contract of the c

Textosterore propionite has a long listery of roshability in the U.S. and abroad, and memis a very commen form fentisteneous en the glad minuted to dis side, if his mels emplished, however, that it is ability to remin on the mulet is more a product of history than usigns application. Testosterore propionite was the first acceptable center of history terms and the proping of the

Bodybulkers commody consider propionate to be the mildest testestence ester, and the preferred form forth formoure for detrage-templesses of training, sown will go so for an its system that propionate will harden the physique, while giving the user less vaster and far returion than one typically expects to see with a telestreen like currellare, opionist or Statumo, Realistically, lowever, these advantages of our hold up to close scentisty. The propionate ester is actually record before the testioners carriers actual experience defere due to the control of the size of the control of th

# How Supplied:

Testosterone propionate is widely available in human and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but usually contain 25 me/m 5.0 m/ml or 100 me/ml of steroid dissolved in oil.

# Structural Characteristics:

Testosterone propionale is a modified form of testosterone, where a carboxyle acid ester (proporties acid) has been attached to the 17-beta plotoxyl group. Esstrafied forms of settostosterone are less polar than fee testosterone, and are absorbed more slowly from the seaso of spicetron. Once in the bloodstream, the seter is removed to yield fee (active) testosterone. Esterfield forms of testosterone are designed to prolong the window of thempearic effect following administration, advoving for a lass fequent nigetion schedule compared to injections of free (unesterified) steroid. The half-life of testosterone propionate is approximately two days after injection.

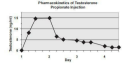


Figure 1. Pharmacokinetics of 25 mg labeled testosterone propionate injection. Source: Pharmacokinetic properties of testosterone propionate in normal men. Fujioka M, Shinohara Y,Baha S, et.A.J. 2 (In Endocrinol Metab 63 (1986):1361-4.

#### Side Effects (Estrogenic):

Textostence is readily anomized in the body to establed (entrogen). The anomatuse (entrogen printense) entropic in responsible of the intentables on fit obstantence. Elevated entrogen levels can cause side effects such as increased water reterior, body fit gain, and genocemistic. Textosteron is considered an andmetaple entrogen is excited. An artis-entrogen such as chroniphene critate or tumoride ratter may be excessary to prevent resistant such as the extraction of the entropic such as chronical use an anomatuse inhibitor like Arimatode (unstancios), which more efficiently controls entropic to prevent gain synthesis. Anomatuse inhibitors can be effects on chaot clinical and the resistant of the effects on the old incise.

Estrogenic side effects will occur in a doss-dependant muner, with higher dosse fabove normal therapeatic steels) of testosterone propriente mee likely to requer the concurrent use of an artis-estrogen or arountase iribibito. Sive water reterion and loss of muscle definition are common with higher dosse of testosterom, this drug is usually considered a poor clocks for dering or eating plases of training, lis medientic estrogenistly mukes it more massle sixed, and believe the storage and the storage of the storage of the massle sixed, and believe at storager articles environment.

# Side Effects (Androgenic):

Testostenoe is the primary mile androgen, responsible for miniming secondary male sessual entancristics. Elected tesk of testostenoe are likely to produce androgenie side effects including oily skit, acre, and body/ficial law growth. Men with a genetic produposition for his loss (andeagouse) adspects) may notice accelentation than pattern lading. These concerned about fair loss may find a more confertable option in standouse decennant, which is a corepraish ly loss androgenie steroid. Women are swared of the potential virling effects of dambolicandrogenie steroids, especially with a strong androgen chances in shire touch, ficelibir growth and official enhancement.

In androgo-responsée tragst fissus such as the skin, soal), and prostate, the high velaire androgenicy of testatemes é dependant on Fendants on displotestatemes (DRT). The 5-sight reductase congrue is responsible for this metabolism of testostoreme. The concurrent use of 3-sight neclutaers inhibit vois such institutée ou distantieur de literative value de la configuration of testostoreme action, lowering the tendency of testostoreme value sits producte androgenic side effects. It is important to remembre fut both anabolism androgenic effects are moditatel via the cytosode androgenic receptor. Complete separation of testostoreme's anabolism call androgenic proceeds in oft possible, even with total 5-sight testostoreme's anabolism and androgenic proceeds in oft possible, even with total 5-sight testostoreme's anabolism and androgenic proceeds in or possible, even with total 5-sight testostoreme's anabolism and androgenic proceeds in or possible, even with total 5-sight testostoreme's anabolism and androgenic proceeds in or possible, even with total 5-sight testostorement and the sight proceeds and testostorement and the sight possible and the sight proceeds and the sight proceed and the sight proceeds and the sight proceeds

#### Side Effects (Hepatotoxicity):

Textosterous does not live hepatotoxic effects, her toxicy is utilicly. One study curried the potential for hepatotoxicy with high does of etestosterous by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile subjects. The settoral was taken orably so that higher peak concentrations would be neached in hepatic itsues compared to intransacular spectrus. The hormone was given days for 20 days, and the contraction of th

# Side Effects (Cardiovascular):

Ambolicarhoganis serosis en ilmve deletroise effects on serum cholesterol. This irelades a reindreys o reduce III El, (goa) cholestero ollasse and irecues IL, (find) cholestero values, without my shift the IIII. to III. balme ei na direction that fixors genete risk of anterocherosis. The relatie imput of an ambolicarhoganis estorio oli sestra lipita i dependari on the dose, route of administration (red 18. si jectoble), type of steroid Ambolicarhoganis estrosis my sia observedy affect blood present ant righericaris, reduce endodelai reluxorio, and support led ventricaler hypertroply, all potentially increasing the fix of controllocarder disease and region-cital

Testosteores tends to have a much less dramatic impact on condisonacciar risk factors that synthesis sendors. Bits die has part to is spermess to metabloshimy be five, which allows it to have less effect on the leptic imangement of cholesteol. The aromatication of testosteores to established also helps to mitigath the negative defices of antiegons on sermi lipids. In one saidy, 200 mg per week of testosteores ester (centificiar) had a slight hat not statistically significant effect on BILL cholesteoria date 12 weeks, but when balson with an automatic inhibitor a strong (25%) decrease was seen. <sup>500</sup> Studies using 300 mg of testosteories ester mitheter a strong testosteories ester continuity per week for trooty weeks when an amentation relaboration exceeding the continuity per week for totary weeks when an amentation relaboration and the strong testosteories of the continuity per week for though weeks when the condition of the continuity per weeks the totary weeks when an amentation relaboration consideration for several dress as deep the second testosteories the consideration for the way the sea so that the section of the resolution of the section of the consideration for the way there is a feed to resolution for the section of the

Due to the positive influence of estingen on serum lipids, tumoritim cirate or clumiphene cirate are perferred no serumsase inhibits for those concerned with entirelesseather leads, as they offer a partial estrogenic effect in the Iner. This allows them to potentially propose lipid profiles and officts own of the negative effects of androgens, which does so (600 mg per less of testiostenous per week, the impact on lipid profile tends to be noticeable but not domain; making an artisestignost (for confidence statistically significant changes to the confidence of the

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the triate of saturated fits, cholsterol, and simple carebolydrates at all times during active AAS administration. Supplementing with sho id (4 games per day) and a natural cholstero/lamioxidant formula such as Lipid Stabil or a product with comrangible inserdenies is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolic includence sensities when taken in doese sufficient to promote muscle gain are cuprent of suppress endigenous testioniere production. Enstatutes in the primary management of the production of testioners in the primary management and offers stone pragine feedback on endagemous testionarem production. Enstaturen-based drags will Reseits, here a strong effect on the probedinarie regulation of ratural stroid hormous. Without the intervention of testionstroners strainfairing substances, testionare levels should return to remain white I are mortis of management growth probanged phopograndstrophic lipograndsm can develop secondary to steroid abuse, reconstitute predictal intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Estosterous propionate is often regarded as a partial spection. This is due to the very solven cubm of that of the propionic and cells which can be irritating to issues at the obordinate of the propionic scale cells which can be institute for the started complexely, their observations and promounced converses and low-grade fever that may last for a few days after each specific. Even the mall soreness that is experienced by most tuens can be qualturated to the control of the con

# Administration (Men):

To treat androgen issufficiency, early prescribing galdelines recommended a douge of 25 mg given two to there times per week. Modern product literative usually recommends 25 mg to 50 mg given two to three times per week for the same purpose. The usual douge among to 50 mg given two to three times per week for the same purpose. The usual douge among the additional set for among of 50-10 mg per injection, which is given every second or third workly commanded to the product of the second or the second of the second of

Textusemen propionate is usually incorporated into buding plause of training, when added water textures will be of little consequence, the user mere concerned with new mass than definition. Some do incorporate his drug ainto caring cycles as well, but typically in lower does (100-200 mg per weeks) andre when excorporated by an amountee inhibitor to be used (100-200 mg per week) andre when excorporated by an amountee inhibitor to be exceeding and in the control. Textuser control of textuser projectent is a vary effective analysis drug and analysis of the surface of the control of textuser in the control of textuser analysis of the surface dependent of the control of the control of textuser analysis of the control of the control of textuser and the control of the control of textuser and the control of the c

#### Administration (Women):

Testactores projectus is mely used with vorters in efacial medicas. When applied I is most often used for a secondary medication derig properable breast cancer, when other most often used for a production of the property tests of the property of the prop

#### Availability:

Testosterous propienta is subject to decreasing supply as a pharmacetrical product. Longer acting sexters such as opionate and enterintue are much before saided for visitually every clinical application. As such, most physiciens and pharmacetrical companies have abundoned this old acts of testosterous. With a few exceptions in the work, the remaining legistrate is also according to the contraction of the cont

Resotsterone propionate is unavailable in the United States as a prescription product. Watson, Lilly, Bel Mar, and Righy made some of the last known products, but all have been emoved from market for some time. Given that the FDA never officially withdraw the drug, it can still be specially ordered through a small number of compounding pharmacies.

Brovel in Mexico makes a testosterone propionate in a 50 mg/mL dosage for veterinary use. Counterfeits are not commonly a problem.

Testolic is made in Thailand by T.P. Drug Laboratories. This product comes in the form of 2

mL ampules, each holding 50 mg/mL of steroid.

Propiobolic from Asia Pharma (Malaysia) is now approved for sale through pharmacies in Thailand. Each box should carry a scratch-off security sticker, which will display a code that can be validated on the company website.

Misr (Fount) have undated the nackaging to Testone-E. The new box carries a white / green / blue pattern similar to other Egyptian steroids including Cidoviron and Cidoteston.

Balkan Pharmaceuticals (Moldova) makes the product Testosterona P. It is prepared in both 1 ml. annules and multi-dose vials

Testosteron is a nonular brand from Buloaria, and is commonly exported to other markets in high volume. It comes in the form of 1 mL glass amoules containing 50 mg/mL of steroid.

A generic made by the company Farmak is popularly exported from the Ukraine. This also comes in the form of 1 mL glass ampules containing 50 mg of steroid.

Jelfa makes Testosteronum Propionicum in Poland, which makes its way most often to the Furonean black market. However, it only contains 25 mg of steroid in each 1 mL annule.

Vironnone is still manufactured in the UK, most recently by the firm Nordic This product contains 2 ml. ner annule, with 100 mg of steroid held in each

Testogan is made by Laquinsa in Costa Rica. This product contains only 25 mg/mL of steroid, but is distributed in a 50 mL multi-dose vial. This makes it one of the largest volume testosterone propionate products available in commerce.

#### Paraholan® (trenholone hexahydrohenzylcarhonate)

Chemical Name	17beta-Hydroxyestra-4.9.11-trien-3-one
Standard	Nandrolone acetate
Anabolic	500
Androgenic	500

Estrogenic Activity	none
Progestational Activity	moderate

# Description:

Trenbolone hexahydrobenzylcarbonate is a slow-acting injectable ester of the potent arabolic steroid trenholone. Trenholone annears most commonly as trenholone acetate. which is a much faster-acting form of the drug (see: Finaiet). The hexahydrobenzylcarbonate ester used here extends the release of trenholone for more than 2 weeks, which has always been thought of as more suitable for human use due to the less frequent injection schedule. The base steroid trenbolone is roughly three times more androgenic than testosterone. making it a fairly potent androgen. It also displays about 3 times greater tissue-building activity in comparison to its androgenic properties, making its official classification as that of an anabolic steroid. The muscle-building effect of trenbolone is often compared to such popular bulking agents as testosterone or Dianabol, but without the same estrogenrelated side effects. It is most commonly identified as a lean-mass-building drug, and is extremely popular with athletes for its ability to promote the rapid buildup of strength, muscle size, and definition

#### History:

The first long-acting trenbolone ester (undecanoate) was studied in 1967, described during a series of experiments into synthetic anabolic steroids by Roussel-UCLAF.571 Trenbolone hendpulchenpfearboute was a subsequent and usinguly French start to this long-scring analysis stored, possessing an unsual bit roughly exploited recopporal. Terobotive hendpelchenpfearboute was developed into a mediene by Negun Labontories in France, which sold the dang under the Parabotion tande name. It was also sold for a period of time as Heachdan, a name that referred to the unusual enter it possesses. Terobotive healphothenpelchenture is the only known from for therobotive every produced as a medicine for harmon consumption. The most metable appearance of treebolive comes as tembolive acceptance, which is salved widely and exclusively investigant practices.

Parabon was prescribed in France as a protein-sparing ambolic agant in cases of cachesis (lean body mass wassig) and multarition, as well as to construct certain frame of otstopenous). Its prescribing gaidelines included recommendations for the teatment of antiquage-sensitive populations, such as women and the elderly (whoge to its moderne antiquages; properties, however, the drug was contaminated of a level to give in although it was faulty discontinued (volumethy) by Nogum in 1997. For a best'e prival of time is secured that the demiss of Parabolic would must the end of human-see terrobotic proputations, as no small nature of Parabolic preparations have been brought to market since, however, so while the drugs is all professional to the complete of the contribution of the complete of the contribution of the complete of the complete

# How Supplied:

Teerbolone hexaltydroberzylcarborate is no longer produced as a prescription drug product. When manufactured in France it came in the form of a 1.5 ml. ampule containing 76 mg of steroid (product information lists this as equivalent to 50 mg of base trenbolone).

# Structural Characteristics:

Tenholsee is a modified form of randrolnee. It diffies by the introduction of double bonds at carbons 9 and 11, which inhigh arountazion (0+-on), incruse andropen-brinding affiny).<sup>22</sup> and shows its methodsom. The resulting steoid is significantly more potent as both an unable and an androgen than its methodsee base. The restrollee has its modified with a levally-drobenzy-barbonate selar at the 17-best hydroxyl group, so that the five steroid is released more solved from the area of infection.

# Side Effects (Estrogenic):

Tierdovine is not arcountized by the body, and is not measurably estropacis. It is of note, however, that this stand displays significant buring affinity for the progastener exceptor (slighty arroager than progastenour seed). The side effects associated with progastenour are similar to these of entopyin, richtaling regularity foreblack ribbilition of testostenour production and enhanced rate of fit storage. Progastins also augment the standardor getter of storages no manurally uses growth. There appears to be a storage synapg between these two hormones here, such that gracomostic might even occur with the help of progastics, whole accessed exempts levels. The see of an art-storage, which have the programment of the programment and notice of the programment of the

# Side Effects (Androgenic):

Abough closified as an ambolic stroid, trothoure is afficiently arthrogatic, Androgatic site effects used formers with this solute, and may include boost ofly slist, next, and boylificate him growth. Ambolicant/orgatic stroids may also aggressee mile pattern hair loss. Weem are also warned of the potential wirsing effects of ambolicant/orgatic stroids. These may include a deepening of the voice, menstral irregularities, changes in skin tearns, field him growth, and clitted and experience. Additionally, the 5-plan notations compare does not metabolic terobolize. So to technic androgaticity is not affected by finistentic or distanties.

# Side Effects (Hepatotoxicity):

Trenbolone is not c-17 alpha alkylated, and is generally not considered a hepatotoxic

sicrous; nor nozary is trincey. This sicrous ones nave a strong avec or resistance to repute breakdown, however, and severe liver toxicity has been noted in bodybuilders abusing trensholone.<sup>250</sup> Although unlikely, hepatotoxicity cannot be completely excluded, especially with high doses.

# Side Effects (Cardiovascular):

Ambolicarinoganis seronis can have deletroisse effects on serum cholesterol. This includes a reindency to reface IEEE (good) cholesterovi obases and increase IEEE, flough cholesterovi obases and increase IEEE, flough cholesterovi values values in a direction that florous geneter risk of arteriosclerosis. The retaine impact of an ambiochandrogenic section of a serum lipit is dependant on the dose, route of administration (end xx. ijectable), type of steroid (generalized) error assormatiohly, and teep of resistance to hepstheri, trendolient beat on non-amountable) matter and stone; prosistance to metabolism teerbolient beat on non-amountable instance and stone; prosistance to metabolism teerbolient beat on non-amountable instance and stone; prosistance to metabolism teerbolient beat on non-amountable instance and stone; prosistance to metabolism teerbolient beat of metabolism teerbolient and support and host observable instances.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular cuercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterollamioxidant formula such as Lipid Stabil or a product with comranable inserdents is also recommended.

#### Side Effects (Testosterone Suppression):

All article/articleogues steroids when taken in dones afficient to promote macle gain are expected to suppose endoguenes testorious production. Without the intervention of testosteron-artinaling substances, testosterone levels should return to normal within 1-cm ments of drug secosion. Now that privacyal phopogenodropic hypogenodrical develop accordary to steroid abuse, reconstituting medical intervention. In experimental develop accordary to steroid abuse, reconstituting medical intervention in experimental conditional articles to the control of the cont

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

## Administration (Men):

Tierdwise bezulptúrebensykutrovintu was generally administered in a clinical dosage of 3 marquises per menth. Therepay was initiated for first morth with a 3 amplesa giene over the first 15 days. During the subsequent 3 menths, one sjection (76 mg) was giene coey 10 days. For playages or performance-trainizing proposes, terebolises of the proposes of the contraction of th

While the day is quite potent when used alone, it is sometimes contributed with other streetis. For an own greater effect. Leading up to a show one could successfully add a normal membrane gambole such as Weststelle or Printebeatin. Such contributions will delt a specimence. We coll also look for this day that disciplines streeting respectives. We coll also look for this day that disciplines streeting with such as the streeting with the day and additions streeting contributions. We discuss the collection of the streeting with the day and additional to the streeting with such as the streeting with the streeting with the such partial position of the streeting of the streeting with the stre

# Administration (Women):

Tendoster headytchotenyckurbonte was generally administered in a circiai dosage of a quantiques per month. Henryp was initiated the first morth with all a ampaies per month from the first 15 days. Daring the subsequent 3 morths, one sjecction (Forga) was given every 10 days. Gorden from 6 of vilization, tower doess were likely used by physicians with many fermile patients. This agent is generally not recommended for women for physique, or fermile patients. The agent is generally not recommended in women for physique, or fermile patients.

# Availability:

Pharmaceutical preparations containing trenbolone hexaltydrobenzylcarbonate remain scarce. In reviewing some of the remaining products and changes in the global pharmaceutical market we have much the following observations.

Balkan Pharmaceuticals (Moldova) makes the product Parabolan. It contains 100 mg/mL of steroid, and is packaged in 1 mL annules and multi-dose vials of 5 mL and 10 mL.

# Primobolan® (methenolone acetate)

Androgenic	44-57
Anabolic	88
Standard	Testosterone

Chemical Names 17beta-Hydroxy-1-methyl-5alpha-androst-1-en-3-one
1-methyl-1(5-alpha)-androsten-3-one-17b-ol
Estrogenic Activity

# Description:

**Progestational Activity** 

Prinobalon® is a brand name for the ambods stood methendous accents. This agent is very sinder in action De prinobalon® Depart (methendous centrality, except here the days is designed for our administration instead of rijection. Methendous accents is a non-e17-alpha-alpha-alpha-daystood roll administration instead of rijection. Methendous accents is a non-e17-alpha-alpha-alpha-daystood resident, on or of only a few commercially accention, another on a method, one of low analogue is and no extension for presents instead for twicky to the user. It is also highly flowed for its properties as a medicantly deficient analogic with no extraogue in operation. It is, Becwise, commody used during cutting bases of training, when lean tissue growth and solid manacularly not from the accent leave the residence of the present accent to the present accentance of the present accentance and the present accentance accentance accentance and the present accentance accentance and the present accentance accentance and the present accentance accentance accentance and the present accentance accentance accentance and the present accentance accentance

no data available (low)

# History:

Methenotee was first described in 1960;<sup>277</sup> Squibb vould introduce the drug fas methenotiene extent to the United States in 1962;<sup>288</sup> has agree was self for a very state time as a 20 reg tablet, under the heard name of Nhalifs. Schering in West Germany (now Begrey) would be gapten rights to the origin that same year, and noted all at under the Printobinsh name. Nhalifs was soon removed from the U.S. market, nove to rotatin as a commercial product. Schering now had exclude patter rights to produce methenoties commercial product. Schering now had exclude patter rights to produce methenoties excetate, and would cortise to self the drug uniterrupted since 1962, and consumes had mainly come to identify methenoties accessed as a product of Schering when the contract of the self-great patterns and the self-g

Princholonib Itsa always been identified as a European setroid, and during the 1960's and "70's was being drieff of sails in sail constrains as Genuma, Austin, Belgiam, Finner, Belgiam, Finner, Be Notherlanks, and Frishnd. At one time Schering also munificated a 20 mg/rid rob-used spicable of methornous eaceties in intend mixes (called) Princholonib. Acutals, but it is been out of numaficated society 1973. Injectable methorshore acctute proved to be very popular for pro-context cartings used, with we since when the societies of the size of Princholard's prescribed as a lear insize bailing anabolic agent, often used in cases where body wasting has courted secondary to major surgery, infection, ususing disease, aggressive contionational administration; or malaration; Corne clinicians also prescribe this aggin; This second has also been used to promote weight gain in underweigh permature rations and children's inclined anabies, and was able to do so effectively and vollowing anabolic, the continuation of the control of the bailing in the control of the bailing incommendation without sale effects.

Albudgh Printobulné demonstrated a good record of chical safety, Schreig Indi winthown this fing form met mutoets by the endy 2000. No 90 my services and sill a munification, and at most a solect couple of products containing 6 mg or 25 mg mg still be in circulation. The object confirmed secrete for not land runner Printobulen in never spans were in Japan and South Africa, and faces were soil starfer the Schriftig muto. It is subcoven from when the soft of the start of the soft of small number of pharmaceutical preparations containing methenolone acetate mg still be in modulation.

# How Supplied:

All forms of Schering Primobolan® contain 5 mg, 25 mg, or 50 mg (no longer available) of methenolone acetate per tablet. Composition and dosage of other brands may vary by country and manufacturer.

# Structural Characteristics:

Methendone is a derivative of dilpolotestosterone. It contains one additional double boat between carbons 1 and Z, which helps to stabiliste the  $\lambda$ -tota group and increase the strend's anabolic properties, and an additional 1-methyl group, which protects the steroid against begain reachiosine. Printroblem makes use of rembendone with a carboyole acid against begain reachiosine. Printroblem makes use of rembendone with a carboyole acid condition during one administration. Studies have demonstrated the methendone is an effective our almobile acuer in both the access and unstrainfied forms. <sup>200</sup> 531

# Side Effects (Estrogenic):

Methenoloes is not anomátech by the lody. <sup>522</sup> and is not measuably estroguir. Estroganistical size defects shad not be seen when daministering this stored Sensite individuals need not sorry about developing genecoments, nor about they be noticing any appreciable water retention with the dury. But receives easy with methenolose doubt be quality muscles musc, not the smooth back that often accompanies steroids open to aromatotion. During a cycle, the user shad additionally not roise term quie estation in blood pressure, as the effect is also related (generally) to estrogen and water retention. Methenolous is a serord muscle factored during cuting planes of frainting, when user and its retention are major occurrent.

# Side Effects (Androgenic):

Albudgh classified as an ambolic strouli, ant/opagic side effects are sill possible with this solution. The principle of the office of the potential with a contract and the principle of the potential without contract and the principle of the potential witing effects of ambolicant/opagess: sensible. These may maked a despensing of the voice, menstratal inregularities, changes in sits tecture, ficial that growth, and clasted enlargement. Metherables as sid as very mild served, however, and strong and clasted enlargement. Metherables as sid a very mild served, however, and strong preparation an acceptable closec, observing it to be a very confortable and effective ambolic.

# Side Effects (Hepatotoxicity):

Methenolone is not considered a hepatotoxic steroid; liver toxicity is unlikely. Studies have failed to produce appreciable changes in markers of hepatic stress when the drug was given in thempeutic levels. <sup>550</sup> This steroid does have some resistance to hepatic breakdown, however, and liver toxicity, failure, and death was reported in one elderly patient receiving oral methenolone acetate. Sel Although unlikely, hepatotoxicity cannot be completely excluded, especially with very high oral doses.

#### Side Effects (Cardiovascular):

Ambelichardnegueis stensisk can have deletroisse effects on serum cholestered. This includes a relatency to reduce IML (angol cholestero) taless and increase IDL (dual) cholesterol values, which may shift the IFIL to IDL behave in a direction that fixors geneter risk of anterioscients. In reducine impact of an ambelichardnegueis extend on serum lipits is dependant on the dose, route of administration (end ix. njectobie), type of stendi (contratible to rout-normatizelo), and level of resistance to he phagic metalosism Methenskere shottal have a stronger negative effect on the hepatic municipant of choickstered than technication can include the one in non-normatizelo nature, but a mark vacation impact than c-l'1 alpha adplant steroids. Due to the route of defreety, can methenshere will have a slightly storager grapher effect on in pick companied to methodate and the contraction of the contraction of the companied to the choicked of the contraction of the contraction of the companied to methodate and the companied to the contraction of the contr

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular scencise program and maintae the triate of saturated fats, cholssterol, and simple carbohydrates at all times during active AAS administration. Supplementing with shi old, grams per day) and a natural cholssterollaritoxidant formula such as Lipid Stabil or a product with comparable ingredients is also recommended.

# Side Effects (Testosterone Suppression):

All anabolicalinequesis storoids when taken in doses sufficient to promote mace gain are expected to suppress endegenous testotores production. Whitten for intervention of testosteron-stimaliting substances, testosterone levels should return to normal within 1-4 morths of drug secosis. Note that prolonged phoppomadorphic hoppomadorpic phoppomadorpic podection. What time may be true in small chiral doses (20-25 mg cluby), list may not be a major distriction when used for phopsic-up or performance-relating approachs in consist, more than large of the patients receiving only 30-45 mg per day noticed a 15-60% suppression of grandstoppin levels. While this is for homely not hopped in the particular process. In our performance control and the production of the production in the production in the production of the state of

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

Studies have shown that taking an oral anabolic steroid with fixed may decrease its bisovasibility; 50%. This is caused by the fit-soluble nature of steroid hormones, which can allow some of the drug to dissolve with undigasted distary fit, reducing its absorption from the gestrointestinal tract. For maximum utilization, this steroid should be taken on an empty stormeth.

#### Administration (Men):

The prescribing gaidelines for Primboloni® recommend a maximum daily dosage of 100-150 mpper day. The usual administration protocols for physique or performance-enhancing purposes call for 75-150 mg daily, which is taken for 6 to 8 weeks. This level is sufficient to impart a measurable ambole effect, athough one usually doesn't expect to achieve great gains in muscle mass with this drug. Irstead, Primboloni® is ufficied when the athlete has a secodific mod for an full anabole near time not totably in cultive robusts of the primbolonis of

Due to its mild nature, Primobolan® is often used in conjunction with other steroids for a stronger effect. In such cases, a slightly lower dose is often used (50-100 mg per day). Jamp, a carting or curing praise, monget to ne as primary appearance, a non-incomingnal analogous Rice Halsestraff or teroboles is define aduled. Such combinations would enhance the physique without water reteriors, and help bring out a lander and more defined look of meaching. Non-constrainty analogous mitosis destide like this every popular among competing bodybelders, and prove quate reliable for negality improving the cortest from This compound is also constrainty used with more potent analogous during building places of training. The addition of tensioners, Disturbol or Analod 50% is corrent, although the corrented and the contract and the heartstooky in the case of the latter to use and of the latter to use and the contract and the heartstooky in the case of the latter to use and the contract and the contract and the heartstooky in the case of the latter to use and

# Administration (Women):

The prescribing gaidelines for Primobolanti do not offer separate dosing recommunitations for women, although it is indicated that women but one pregunt; or my become pregunt; should not use the drug. Femila airlistes generally respond well to 50-75 mg dails, with no sign of virilation supprison. One would not expect a theremakes mourt of musel: muss with this day, and instead find a slow and steady (gaidally increase. Some women choose to further adds -1 other amodels such as Westerdie or controllors, in moffert in derease the musel-building effectiveness of a cycle. While both of these compression are quite tolerable, there and on the controller and the controller and the state of the controller and the controller and the state of the controller and the controller and the state of the controller and the controller and the state of the state of the controller and the state of the state of the state of the controller and the state of the st

#### Availability:

Pharmaceutical preparations containing methenolone acetate remain scarce. The drug has been unavailable in western nations for many years now, and the bulk of the supply presently comes from underground steroid manufacturers. In reviewing some of the remaining products and changes in the adoled plantamocratical market we have made the following observations.

Balkan Pharmaceuticals produces Primobol in Moldova. It comes in 50 mg tablets, with 20 sealed in each foil and obstic strip.

# Primobolan® Depot (methenolone enanthate)

Androgenic	44
Anabolic	
Standard	Testostero

Chemical Names 17beta-Hydroxy-1-methyt-Salpha-androst-1-en-3-one1-methyt-1[S-alpha)-androsten-3-one-17b-oll
Estrogenic Activity no data available (low)
Progestational Activity no data available (low)

#### Description:

Printohamis Depot is an injectable venion of the storol methanolous. This is the same constitute in Printohami Gan (methanolous cateata), although them enterathete ester is used to show the steroid's release from a sice of injection. Methanolous curatifute offers a smirin pattern of steroid release as testionicon curatifute, with both Demone levels remaining mutdedly destated for approximately 2 weeks. Methanolous leaff is a moderately steroid printohamistic control of the steroid properties. Its automatic effects is considered to a steroid properties of the steroid properties in the steroid effect is considered basis. Methanolous curatifute is most commody used during entiring cycles, when lean muss gain, not a nor muss increase, is the min objective.

#### History:

Methenolone was first described in 1960. 567 Squibb introduced the drug (as methenolone eranthate) to the U.S. prescription drug market in 1962. 568 sold for a very short time in the U.S. under the brand name of Nibal® Denot Rights to the drug were given to Schering in

West Germuny (now Bayer) that same year, and Nibal® Depot soon disappeared from the U.S. market. Schering would sell methenolone enuratine under its new and ultimately most recognizable brand name, Printochan® Depot. During the 1960s and 70's Printochan® Depot was available mainly in Europe, including such countries as Switzerland, Italy, Germung, Austria, Belgiam France, Portusal and Green Portusal and Green.

Scherig minimized patter cortrol over mehanolous cuantitate until the late 1970s. Bedres to patters capital. Scherig had rigossubs protected as in tallectual property right suginst any potential infragment, even in the U.S. market, where the corpumy had not been marketing printsolom Depot. Although methanolous cuantitude lass not been available for commercial sale in the United States for decades, it has technically retained is status as an FDA-approved draw.

Primbolan Depot is typically prescribed as a lean tissue building anabolic agent, often used in cases where body seating last occurred according to an operation; prolonged infection, wastering disease, aggression confricted analysmination, or considerane. Some discission wastering disease, aggression confricted analysmination, or considerance. Some discission mass with agent, certain cases of drawin kepatis, and breast excisions (usually as a sociolarly medicated flowing ofthe throughes). Dis searched has sho been used to promote weight gain in underweight promine ridines and châlten in clinical statics, and was able to do so efficiently and whorst signs of trovity or undesiched efficies. Whether has been glintered the contributed stong annotes, work andragons; and row-consequent name of the draw sale through the contributed stong annotes, work andragons; and row-consequent name of the draw sale through the contributed stong annotes, work andragons; and row-consequent name of the draw sale through the contributed stong annotes, work andragons; and row-consequent name of the draw sale frames it work of solid for the first section of the contributed stong annotes, work and any contribute of the contributed stong annotes, were always and row-consequent restored to the contribute stong annotes and the contributed stong annotes, were always and row-consequent restored to the contribute of the contributed stong annotes and the contributed annotes and the contributed stong annotes ann

Albusgh Primbolan Dapto demonstrated a good record of cinical safety, by the 1900s. Schering had gown to be a milatinating humanecatical gain, and use nearbily freed to recommine its global steroid offerings in light of public concerns about sports duping. Primbolan Dapto would be voluntarily withouton from most of the countries in that lon origanity sold it. Today, the benaft as sold in just a huntilla of countries including Spain, Timelee, Spain, Prangage, and Exander, in pair of its finisted spaip, Boyer has remarked (unably the exclusive produce of melanodose currelates in the harms thug basees manually office and countries, much form delenanced or countries the manual to office arresorations, must form undersound or count-of-the communities.

#### How Supplied:

All forms of Bayer Primobolan® Depot are packaged in 1 mL glass ampules and contain 100 mg of methenolone enauthate. Composition and dosage of other brands may vary by country and menufacture:

#### Structural Characteristics:

Methenshese is a derivative of displostentosterous. It contains one additional double bond between carbon I and 2, which helps to sublike the 3-deep opps and increase the standa's anabole properties, and an additional 1-studyle group, which gives the sectoral control of the standard of the standard of the 1-deep long of the standard of the carbonyle and extended to the 17-best helpsonyl group. Esterdied standards are less polar fund feet standards and are absorbed more slowly from the area of rejection. Does in the bookedness; the extent is removed by self the (exterior pathenshese). Esterdied standards are designed to prolong the wardow of thempositic effect following the standards are designed to prolong the wardow of the standards of the standards and the standards are the

#### Side Effects (Estrogenic):

Methenolese is not arountized by the body. <sup>500</sup> and is not measurably estrogenic. Entreparlished side effects should not be seen when darministraring this struck G-meinle individual need not story about developing genecoments, nor abould they be noting any appreciable water retention with fisting. The renesses seen with methenolese should be quality muscle, musc, not the smooth bad, that othen accompanies steroids open to automatication. During a size and the seen of the contraction of the contraction of the contraction of the is also related (queriespell) to estrogen and water renetrion. Methenologies is a served must forced during entiring phases of raining, when water and fit retention are major concerns, and sheer mass not the central objection.

# Side Effects (Androgenic):

Albudgi-closifed as an ambolic strool, androguic side effects are still possible with this solution. The my rickle bosts of obj skir, area, and hoolyfietal har growth. Anabolicanhoguic strools my also aggravate make pattern har bes. Women are warmed of the posteral widing effects of anabolicanhoguic strools. These my rickle a despersing of the voice, mentatual inregularities, changes in skin texture, facilla har growth, and clinical enhangement. Metherabone is all a very multi strooth, however, and stroog anahoguic side effects are typerally related to higher doses. Women others find this programment may be adopted the clinic observable of the leave of controlled and effective programment may respect the clinic observable of the leave of controlled and effective and the strooth of the clinical strooth of the cl

#### Side Effects (Henatotoxicity):

Methenolone is not considered a hepatotoxic steroid; liver toxicity is unlikely. Studies have failed to produce appreciable changes in markers of hepatic stress when the drug was given in the manufic leads 571.

# Side Effects (Cardiovascular):

Ambolicardrogenic streolic can have deleterione effects on serum cholesterol. This includes a reductory to reface IELL (apod) cholesterol obases and increase IELL, (dou) cholesterol values, which may shift the HDL to LDL chaince in a direction that fixors geneter risk of anteriocherois. Bit retails impact of an ambolicantaloguic streol on serum lipid is dependant on the dose, route of administrator (ord its injectable), type of steed Methodologic streol in the contraction of the dose, route of administrator (ord its injectable), type of steed Methodologic streol libro a stronger negative effect on the hoppic management of cholesterol data restorators or marfolose due to its non-aromaticable nature, for a much weaker impact than or 13 palm adjusted services. Autobic introduceris streolists may administrate the value of the varieties the prototype, all potentills precessing the first of antibococculi disease and trajectories, reduce enclosibility diseases.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the strake of saturated fits, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with sho list (4) grams per day) and a natural cholesterol/artioxidant formula such as Lipid Stabil or a moduct with comrangable inserdents is also recommended.

# Side Effects (Testosterone Suppression):

All anabolic/andogens: stends when taken in doses sufficient to promote muscle gain are expected to suppress endogenous testosteros production. Whiten the intervention of testosteron-strinkting substances, testosterone levels should netturn to normal within 1-4 mortifis of drug secosion. Note that produced phospoundsorities hoppograndism of develop secondary to search abuse, necessitating metaled intervention is moderate developed to the control of the string of the developed to the control of the string of the developed to the control of the developed to the d

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

# Administration (Men):

The prescribing guidelines for Printrobelan Deptot recommend a musiman dosage of 200 mg at the onest of therapy, and a contraining dosage of 100 mg every wock. Polotaged administration protocols generally call for a 100 mg dosage every 1-2 weeks, or 200 mg every 2-3 weeks. The usual andireisation protocols among must ablest call for a 200-00 mg per week dosage, which is staffs of 6 to 12 weeks, which is sufficient to promote very rotecules increases in least must be time. It is, however, not trausall to see the ding bear in does as high as 600 mg per week or more, although such amounts are likely to highlight a more antidoguria does of orthechnoles, as well as excuently to Expresse effects on servin

# linids

Methenolone enouthate is often stacked with other (typically stronger) steroids in order to obtain a faster and more enhanced effect. During a dieting or cutting phase, a nonaromatizing andmoren like Halotestinik or trenholone can be added. The stronger androgenic component here should help to bring about an added density and hardness to the muscles. On the other hand, one might add another mild anabolic steroid such as stanozolol. The result of such a combination should again be a notable increase in muscle mass and hardness. which still should not be accompanied by greatly increased side effects. Methenolone enorthate is also used effectively during bulking phases of training. In such a scenario, the addition of testosterone or boldenone would prove quite effective for adding new muscle mass without presenting any notable henatotoxicity to the user

# Administration (Women):

The prescribing guidelines for Primobolan® Depot do not offer senarate dosing recommendations for women, although it was indicated that women who were pregnant, or may become pregnant, should not use the drug. Female athletes generally respond well to a dosage of 50-100 mg per week. If both oral and injectable versions are available, the oral is often given preference, as it allows for greater control over blood hormone levels. Additionally some women choose to include Winstrol® Denot (25 mg twice ner week) or Oxandrolone (7.5-10 mg daily), and with it receive a greatly enhanced anabolic effect. Androsenic activity can be a concern with such dosine however and should be monitored. closely. If stacking it would be best to use a much lower starting dosage for each drug than if they were to be used alone. This is especially good advice if you are unfamiliar with the effect such a combination may have on you. A nonular recommendation would also be to first experiment by stacking with oral Primobolan®, and later venture into the injectable if this is still necessary

#### Availability:

Pharmaceutical preparations containing methenolone enanthate remain scarce. The bulk of the supply for this compound comes from underground steroid manufacturers. In reviewing some of the remaining products and changes in the global pharmaceutical market, we have made the following observations.

Bayer took control of Schering AG in December 2006. Following this acquisition, the Schering Primobolan Depot products were transitioned over to the Bayer brand and logo. The company produces the drug in limited markets only, most notably Turkey and Spain. These products have historically been the subject of large volume counterfeiting, however, so consumers should be especially careful with brand name Primobolan Depot products.

Balkan Pharmaceuticals (Moldova) makes the product Primobol. It is prepared in both 1 mL. ampules and multi-dose vials

#### Proviron® (mesterolone)

Androgenic Anaholio

Allabolic	100 13
Standard	Testosterone propionat
Chemical Names	17beta-hydroxy-1alpha-methyl-salpha-androstan-3-or 1-methyl-salpha-dihydrotestosteror
Estrogenic Activity	non
Progestational Acti	vity not significan

#### Description:

Proviron® is Schering's (now Bayer's) brand name for the oral androgen mesterolone (1-

melyd (diplotestostomous). Similar to diplotestostomous, mesterolare is a strong amthogan with only a weak level of amthoda carity. This is due to the fact that like diplotestostomous, mesterolares of melyd produced to incribe doll metabolites in much consess where concentrations of the 3-diplotestored diplotingsment compare and just and only mental consession of the 3-diplotestored diplotingsment compare and in an and one morphism compare in much is base, thereby reducing the gains of other trace potential particular to the contraction of the strong of the diploting strong, is death flewise nor the binders producing the gains of other traces by fooding affinely for plasm binding proteins such as SHEC, mesterolare may actually work to protein the activity of other strends by diploting a inferior proteins give as food of the strong by diploting and gain of the strong gain and the strong of the contract of the strong and the strong of the traces by displacing a inferior procurage as a few currents.

# History:

According to company literature, Schering developed Proviscol's in 1934, making this is an extentively of microlians and first analysis changes are soon. See for medication part into chical practice for the teatment of "hormone-related diseases and complains in man." Accordingly, mentanelize would have been developed diseases and complains in man." Securities, mentanelize would have been developed which are both very old agents generally considered devokets by today's standards. In spite of its age, Provision has a kep listory of client effectiveness and safety, and remains in widespread clinical use today. It is generally prescribed to makes for the treatment of Redo cannelly by insidered analysis and sufficiently considered and sufficiently and sufficient effectiveness and heavy prescribed to make for the treatment of Redo cannelly his indifferent analogue, book, hypogenation in pre- and post-polecuer make), and infertily (in certain situations mesterolone increases the quality and quarrity of sperm).

The use of mestendone as a farilly aid is perhaps one of the most controvenial infantions for this dang considering that anabolicationpairs storoids are generally liked to infinitely, it is also a use of mesterolous that is quite often misanteneously advantages. Mestendone is populable here because it is an effective analongs that offers minimal suppression of a supplicable to the because it is an effective analongs that offers minimal suppression of a production of the own of the supplication of the production of the own function of the own functions of the own function of the own functions in our carriery suppression. Mesterolous scene to have a range positive influence on certain cases of the own function of the own functions of the own function of the

Mesteroise is widely munification by Bayer (formerly Schrieg), which currently selb the done jin more than filter coursies workshick. He must common heard mure used fit is also in Provision, although SchriegiBayer has sold the agast native other mans; in certain members of the provision of a Parkine (Alexa, Cammay), Visitions (Jerephann Certumpi, and Restore (Brown & Barke, India), in spile of is large tank consolidate in members of the fact lead of the provision of a first part factors, mesteroise two sear expressed for sale in the United States. It remains admissible in many Western mations, however. Buyer remains the order branch of the angular leaguest or memorized tools, although on mre coasion of the branch of the angular leaguest or memorized tools, and along the mre coasion.

#### How Supplied:

Mesterolone is widely available in human drug markets. Composition and dosage may vary by country and manufacturer; preparations generally contain 25 mg or 50 mg of steroid per rabler

# Structural Characteristics:

Mesterolone is a modified form of displorostosterone. It differs by the addition of a methyl group at carbon 1, which helps protect the hormone from hepatic metabolism during oral administration. The same structural modification is also used with oral Primobolanis' (methenolone) tablets. Alkydrion at the one position slows hepatic metabolism of the steroid during the first pass, although much less profoundly than 1-71 adpain alkydrion. Mesterolone

is resistant enough to breakdown to allow thempeatically beneficial blood levels to be achieved, although the overall biocoralibility remains much lower than c-17 alpha alloglated oral steroids. Mesteroline also lies a very strong briding affinity for Set Hommon Briding Globuln<sup>27</sup> This may act to displace other steroids more weakly bound to SHBG into a free (active) state.

### Side Effects (Estrogenic):

Mesterolone is not aromatized by the body, and is not measurably estrogenic. An antiestrogen is not necessary when using this steroid, as the drug is unlikely to induce onecomments water retention or other extrooren-robited side effects.

Measurement is actually believed to set as an artiferorement in the body, preventing or devining the conversion of secords into energies. The result is somewhat comparable to training the conversation of seconds in the conversation of the second training, and a number of other secords have demonstrated similar activity. Diplydrostostoreme and Measurem (2-melly-displydrostostoreme), for example, have been secondify used as thempies for generoustian and breast cannot extend to their store gardenpoint and potentially artife-strongenic effect. It has also been suggested that markolore may even hower amounts earlierly in preferent lesses where it is more resistant to estore may even hower amounts activity in preferent lesses where it is more resistant to estore energies effect of off-these comparable is personately caused by the adulty to employ the activity of the comparable is personately caused by the adulty to employ the activity of the second of the comparable is personately caused by the adulty to employ the activity of the second of the comparable is personately caused by the adulty to employ the activity of the ac

## Side Effects (Andmoenic):

Mestrolnes is chasified as an antioqueix steroid. Antioqueix side effects are common with this solstance, especially with ligher dosse. It is may risked be sort of oly-skin, zoon, and bodyfiscial hair growth. Anti-loical-indoparic steroids may also aggroute male pattern hair loss. Werene are also warmed of the potential villaring effects of anti-loi-indoparies, testosis. These may include a deepening of the voice, mensitual irregalerinis, changes in skin testosis. These may include a deepening of the voice, mensitual irregalerinis, changes in skin testosis. The same power and clinical regarders. Additionally, the 5-ships reductase engine does not metabolize mesterolome, so its relative antioquariety is not affected by finistentic or disastical.

#### Side Effects (Hepatotoxicity):

Mesterolone is not c17-alpha alkylated, and not known to produce hepatotoxic effects; liver toxicity is unlikely.

## Side Effects (Cardiovascular):

Ambolicardrogenic steroids can have debetrious effects on serum cholesterol. This includes a redwardy to reface IELL (angle) cholesterol ushes and increase IELL, (bad) cholesterol values, which may shift the IELL to IEL chalmee in a direction that fixors genetar risk of anteroclerosis. The retainer maper of an ambolicardrogenic section of asserta lipits is dependant on the dose, route of administration (roal xx. ipsecabils), type of steroid (assertable) are non-assertables). The deed of resistence to heptain entailobins constructed to the control of the particular characteristic control of the contr

Mesteroione should not be used when cardiovascular risk factors preclude the use of other oral steroids.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular coursies program and maintain the intake of saturated fits, cholesterol, and simple carabidyndrase at all times during active AAS administrator, Supplementing with fish oils (4 games per day) and a natural cholesterolamiovidant formals such as Lipid Stabil or a product with comparable ingredients is also recommended.

# Side Effects (Testosterone Suppression):

Mesterolone has a very weak suppressive effect on gonadotropins and serum testosterone. Studies show that when given in moderate doses (150 mg per day or less), significant suppression of testosterone levels does not occur.<sup>574</sup> In studies with higher doses (300 mg per day and above), the agent strongly suppressed serum testosterone.<sup>575</sup>

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

# Administration (Men):

To text antrogen issufficiency, mesterwhere is usually spinen in a done of 1 which (25 mg) three times per day at the infation of thempy. The drug is liter continued at a lower maintenance done, which usually consist of taking 1 which (25 mg) one to two times per day. Strind does are used to support that definity, usually in conjection with other fatility drugs like injectable FSH. The usual dosage among multi-attents is between 50 mg and 150 mg of mesterwhere or day not not to size 52 mg blacks. The drugs it spixally latent in explosion of 6-12 weeks in length, which is usually a sufficient period of time to notice the benefits of drugs the first three in the contributions.

Mary bodyhalders from the use of mestrolone during disting phases or context preparation, when to estrogen and high androgen beeks are particularly desinable. Bit is especially beneficial when analytics like Wastrolli?, Armon or Primobolmi? are being used adure, as the androgenic content of these dang is relatively to Mestrochine can be efficiency used here to adjust the androgen to estrogen ratio upwords, bringing about an increase in the Instruss and density of the mackets, supporting Brids and general sense of well being, and increasing the tendency to burn body flat. It is also commonly used (at a destination of the process arrows the Driver of the Order State State of the Driver of the destination of the process arrows the Driver of the Order State Stat

## Administration (Women):

Mesterobne is not approved for use in women. This agard is not recommended for women for physique- or performance-entenuing purposes due to its strong aurhoganic nature and tendency to produce vidings side effects, some women do four the dag, however, and first a single 25 mg tablet enough to efficiently shift the hormore belance in the body, greatly imprincing the look of definition to the physique, larke is usually institute to no longer than the or of he weeks in such statution to minimate the chance of developing lasting wiring effects. One tablet used in conjustrition with 10 or 20 mg of Nobadouble on the even more efficient for manche handring, containing environment here the body's mark more faced to be off cent hody the capecially in farmit mothed areas the deep long and thigh. Elemen candom of each poly the capecially in farmit mothed areas the deep long and thigh. Elemen candom to the confidence of the confidence o

## Availability:

Mesterolone remains widely available, the vast majority of products made by or under license from Schering (now Bayer). In reviewing some of the more popular products and changes on the global planmaceutical market, we have made the following observations.

Bayer took control of Schering AG in December 2006. Following this acquisition, the Schering Proviron products have been transitioned over to the Bayer brand and logo.

Bayer no longer markets Proviron in Egypt. The drug remains available under the Cidoviron name, produced by the domestic firm CID (Chemical Industries Development).

Unigen markets the product Mesviron in Thailand. It contains 25 mg per tablet, and is packaged in foil and plastic strips of 10 tablets each (5 strips per box).

Due to its limited demand, mesterolone products have traditionally not been the subjects of high volume counterfeiting. When located on the black market, they can usually be trusted so long as they are properly packaged from a known manufacturer.

## Sten (testosterone cypionate & propionate)

Androgenic	100
Anabolic	100
Standard	Standard

......

Chemical Names	4-androsten-3-one-17beta-ol beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate
Progestational Activit	y low

#### Description:

Sen is a two-component testosterore bland from Mexico fast contains a mêture of testosterore propriete (2 st mg), testosterore spriente (7 st mg), and DFEA (dolydroparindosterore, 20 mg) in a 2 ml, ampuk. Some references incorrectly list this product as containing 20 mg of DFI (diphytestosisterore, which would be a tilted antisegen. This is, however, just a confusion of the Sprainh userd for DFEA (delathinocumbrosteron), which a a quick gainer belos similer to "Diphytestosisterore," and confusion of the product of the state of the product of the similar product contains a similar bland product contains a similar bland product contains a similar for printing the DFEA relevant at the moment, fist sensity product contains a similar for printing times for the contains the similar product contains a similar for the printing time for the contains the similar product contains a similar for the printing time of the product contains a similar for the product contains and the product contains a similar for the product contains and the product co

Many consider Shen to be a low-budget alternative to Statemorië 250. While it does comin a leabed for two testience electry, Sein is at solw-ceiting robusticeme electry. Sein is not a solw-ceiting robusticeme electry. Sein is not a solw-ceiting robusticeme levels to return to bandene approximately 2 words after igenion. Testistences oppositute is also not a deluptic-onset drugs oo Stan doesel't older much advantage in regards to a "sustained-relevels" effect. The testistence reportate only composition of the origination of the initial testistence spike, marineg in plantamotionistic profile more uneven than if 'testistence oppositute were used about. Of the plantamotionistic profile more uneven than if 'testistence oppositute were used about. Of the plantamotionistic profile more uneven than if 'testistence oppositute were used about. Of the plantamotion of the plantamotion

#### History:

Sun is made in Mexico by the phermacortical fron Admir, S.A. die C.V. This agart is used primrily to correct low androgen levels in made, for the notimet of hypogenesistic materials and the state of the control of the proposal distributions of the control of the Mexican mades, where it is one of the courtry's more inexpensive harmonic extensions the mode of the control of the co

#### How Supplied:

Sten is available on the human drug market in Mexico. It contains a blend of 25 mg/75 mg testosterone propionate and testosterone cypionate (respectively) per 2-milliter ampule.

## Structural Characteristics:

Sen contains a minute of two testostenoe compounds, which where modified with the addition of enchools; and extent propriets and explemy-phosphora, exclud as the 17-best laydrood group. Esteriled forms of restostenoes are less point fram free testostenoes, and are absorbed more assloy from the same of significant. Once in the bloodbearum, the exist is advanted as the substruction of the contraction of the con

# Side Effects (Estrogenic):

Testoderore is madily attentiated in the body to establio (estrogen). The attention (estrogen synthesis) entrym is responsible of this introblem of testosterone. Elevated estrogen levels can cause side effects such as increased water reterior, body fit gain, and goncommite. Testosterone is considered an indunctival estrogenic sector. An anti-estrogenic such as chargibane citate or turnosfic relates may be recosary to prevent storogate side effects. One may destructed use an arountary either that Arrianderie (genomotic), what offers, to the may have the production of the storogate of the companion of the storogate quite expension in comparison to anti-estrogens, however, and may also have negative effects on blood pairs.

Estrogaies side effects will ocur in a doss-dependant mumes, with higher doss of howe mentil therapectic levels) of resistories mere fieldy to require the occurrent use of an unicatingen or aromatuse ribbliot. Since water reterion and loss of muscle delithion are common with higher doss of festosterons, this drug is usually considered a poor choice for deling or carting plasses of braining. Its moderate entogenicity undees it more kital for the contract of the contract of the contract of the contract will support more strength and muscle size, and high forter a strong amadels entogenically size of the contract of the contract and the forter and the forter and the forter and the contract analysis entogenically and the contract analysis of the contract analysis

## Side Effects (Androgenic):

Telestorione is the primary mile androgen, responsible for miniming secondary mulesessual enhancistics. Extended to-level forestorione are Body to probe androgenic side effects including oily skit, acre, and body/ficial law growth. Men with a genetic prodeposition for his loss (andesgonetic adspects) may notice accelerated multa pattern lading. These concerned about fair loss may find a more confertable option in sandosive document, which is a comprable loss androgenic setted. Wenne are swamed of the potential visiting effects of anniholociandrogenic steroids, especially with a strong androgen control of the properties of the properties of the properties of the properties of the potential visiting effects of anniholociandrogenic steroids, especially with a strong androgen control of the properties. The properties of the proper

In authory exposite target forus each as the skin scale, and prostute, the high relation and engineering of statements of dependent or in relation to different deplotestatemen (EMP). The 5-ship reduction engineering the engineering of the statement of testistatement. The concurrent use of 5-ship reductates inhibite such as finantistic of attainties of interfere with sits-specific potentiation of sestostenee action, lovering the tendency of testistatement of the statement is another late and undergoine effects are both mediated via the cytosolic anticogen receptor. Compiles expectation of estostenee's another late anticogen properties of ne possible, cut with

## Side Effects (Hepatotoxicity):

Testosterone does not lave lapratotoxic effects, her toxicy is utilicely. One study examined the potential for hepatroxicity with high loss of relessatoreney by administering 400 mg of the hermone per day (2.800 mg per week) to a group of mule subjects. The sectoral was action orably so that higher peak conventurations would be reached in hepatric tissues produced no significant changes in her engine values including serum advants, before the subject of the significant changes in her engine values including serum advants, bifurbin, attitive-antive-articles, and adultive-destiness. 2008.

## Side Effects (Cardiovascular):

Ambolicarhoganis stronds en ilmve debetrous effects on serum cholesterol. This irelates a redency to reduce III. (2004) cholesterol values and irecense III. (2004) cholesterol values v

Testoderore tends to have a much less dramatic impact on conflorouncular risk fectors than synthesis storout. Bits de any not to specimes to metablishminy be feet, which allows it to have best effect on the leptace management of choisestord. The aroundation of testosterous to sensity. 200 mag or sends of testosterous sensity of andragges on serious places. In our suite, 200 mag or sends of testosterous sensity or contribute had a slight to not a contribute of the contribute o

Due to the positive influence of entrogan on serum lipids, temrotion cirate or champlene circutae are perfection of somemass inhibitors for those concerned with anotherous learning, as they offer a partial entrogenic effect in the feet. This allows them to potentially improve high profiles and offers one of the reagiste effects of analogouss. Who does of 600 mg or less per week, the impact on high profile tends to be noticeable but not demantic, making an artisestic analogous feet confederatedine purpose in principal tenter of the profile o

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fits, cholsetrol, and simple carebolydrates at all times during active ANS administration. Supplementing with shot old, grams per day) and a natural cholseteo/landioxidant formula such as Lipid Stabil or a moder with conversable insordiers is also recommended.

## Side Effects (Testosterone Suppression):

All anabodication general search is when taken in does sufficient to promote muscle gain are expected to suppose endogenous testostories production. Estostories it he primary mandrogen, and offers storage negative feedback on endogenous testostories production. Estostories have production for testostories storage deep on the probabilistic regulation of rotational steroid hormouse. Without the intervention of stostories-estimating substances, testostories levels should return to romal white 1-4 mortist of reasons when their prolonged programment probabilistic productions are substanced to the contract of the programment of the progra

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

Testosterone propionate is often regarded as a painful injection. This is due to the very short carbon chain of the propiona acid ester, which can be irritating to issues at the size of injection. Many sensitive individuals choose to stay away from this steroid completely, their bodies reacting with a pronounced soreness and low-grade fever that may last for a few days after each piector.

#### Administration (Men):

For the teatment of two antiongs levels, the prescribing guidelines for Stan recommend one spicietion of one 2 and rampide (100 my testistores cuesters. 20 mg/DRA) every 15-30 days. For bodylviding purposes, this drug is usualy rijected on a weekly basis, in a dosage of 2-4 and parks (200-400 mg) or festistorene exects in 10d.]. This level is sufficient to provide excelent gains in muscle size and strength. Higher doses are possible, but even the rijection volume recold with a drugsles per week (100 can become to uncomfettable for many. Testistorene drugs are ultimately very verseits, and can be combined with many other anabolicativations; strength desired filest.

#### Administration (Women):

The prescribing guidelines for Sten do not make special dosing recommendations for women, except to so what androganic symptoms may occur, and in certain scennics then then special dosing recommend for women for phospiace or performance-ordanic grapheses due to its strong androganic mature, tendency to produce virilizing side effects, and slow acting characteristics infinitely hold to look offficient control.

## Availability:

Sten is commonly found in Mexico, where 2 pre-loaded syringes are packaged in a box and usually sell for about \$10 in the pharmacy.

## Striant® (testosterone)

Androgenic	100
Anabolic	100
Standard	Standard

Chemical Names	4-androsten-3-one-17beta-ol 7beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate
Progestational Activit	tv low

## Description:

Striatt 8 is a micoadhesise bacal assistances delevey system 1 is prepared in the form of a mail (appriss-solar) tablet, which contain 30 mg of (feet) assistance. The table is not taken ends, but is affixed to the garm, where it transfers instostences across the riner large real table on the part of the stress of the stres

#### History:

Strainf Sward developed in the United States by Colambia Labonatories. It was approved by the FDA for sale as a perceiption dung in June of 2003, and is inflated for use in more with conditions associated with a deficiency or absence of endogenous lestisatories. With this product, Colambia was kelle by rigo; to laught those hemmer epidement theory; (HRI) consumers flut do not vuckome blowckly spectrons, and find patches and gals unconditable or comentacily depictable. Seriat was adapted to pharmics in late 2003, and quickly met with intend reviews. Some patients find it as very convenient option for HRIT, while others find it as very convenient option for HRIT, while others find it as very convenient option for HRIT, while others find it could be also an unconfidentable to use the page touth of time. Somet was relaxed in the out-of-time to use constrained to loss of less perceived from Assimt was relaxed in the contraction of the contraction of the page touth of time. Sometimes to the contraction of the page touth of time. Sometimes to use the page touth of time. Sometimes to use the page touth of time. Sometimes to use the page touth of time.

## How Supplied:

Striam® mucoadhesive buccal testosterone delivery system is available in various human drug markets. Product comes in the form of a small buccal tablet; usually packaged in strips of 10 tablets 6 strins to a box.

## Structural Characteristics:

Striant® mecondissise buccal testosterone delivery system is a buccal tablet containing 30 mg of (see) testosterone. The system is a shared to the inside soft month, where the gam meets the upper lp above the incisor teeft. With exposure to sales the tablet softens into a gel-like consistency, which can stay in pace for 12 hours. The product delivery he physiological concentrations of testosterone through the musous membrane, where is is absorbed into the bloodstreamy the superior verse concentrations of testosterone through the musous membrane, where is is absorbed into the bloodstreamy the superior verse concentrations.

## Side Effects (Estrogenic):

Textosterors is reality aromatical in the body to establio (entrogen). The aromatuse (entrogen principal comprise in proposible of first metablism of instantonics. Description entrogen is the control of the first metablism of instantonics. Description is the entrope of the control of the entrope of the en

## Side Effects (Androgenic):

Testoderore si the primary make androgen, responsible for mintaining secondary male sessual catacteristics. Describt ground therapent discoss is floyly to produce androgenic side effects including oby skin, acre, and bodyficial their growth. Men with a genetic side effects including oby skin, acre, and bodyficial their growth. Men with a genetic predigosition for his two functionary descriptions are consecuted and negation halding. Women are warned of the posterial virling effects of anaboliciantrogenic stensist, sepecially with a story gardenge such seststorene. These may include despering of the voice, menstrual irregularities, changes in skin texture, facial hair growth, and citoral enhancement.

In advagor-responsée traget tissus such as the skin sculp, and prostite, the high relative androganicy of testistence is dependent on its relations to disploitastistence (DRI). The 5-ship reductase enspire is responsible for this metabolism of testistence. The concurrent use of 3-ship reductions enhibits such as instanticed or admixtured will retrieve with also-specific potentiation of testistence action, lovering the tendency of testistence drugs to produce ambagoire siste effects. it is important to remote that analonal analongume effects are both mediated with the cylotode analongum recognic. Complete or analongume of the complete of the complete of the complete of the production of the complete of the com

## Side Effects (Hepatotoxicity):

Testosterone does not have bapatotoxic effects, her toxicy is utilicely. One study examined her potential for hepatotoxicy with high does of electrosteron by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile subjects. The seriord was staken orably so that higher peak conventioniston would be reached in hepatic fissues staken orably so that higher peak conventioniston would be reached in hepatic fissues produced to significant changes in five recopier values including serior advanta, bullrules and the state of the significant changes in five recopier values including serior advanta, bullrules and solve the benefatives.

## Side Effects (Cardiovascular):

Anabokien/drogenie steords en ihwe dekterione effects on serum cholstered. This includes a referency to refere EHL (agod) cholstered oakse and increase DL, (bad) cholstered values, which may shift the HDL to LDL chaince in a direction that floores greater risk of articles of the cholstered of the ch

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fists, cholesterol, and simple cardiohydrates at all times during active AAS administration. Supplementing with shis lot (4) games per day) and a natural cholesterolamioxidant formula such as Lipid Stabil or a product with commarble inserdiencies is also recommended.

## Side Effects (Testosterone Suppression):

All analysic landaguist stansis when taken in does sufficient to promote muscle pair are expected to augmost endogranus toustoriere production. Toustouriere is the primary unitanalogus, and offers strong regules feedback on endogranus toustoriere production. Toustouriere-lessed florage will Reviels, here a entreg effect on the production regulation of intend about Il commons. Without the intervention of toustouriere simulating substances, toustouriere lessed and tenum to ment with 1-4 months of drug execution, to be that prolingaged hypogranuloropic hypogranuloriere develops executing to steried about processing mental intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

The Striant mucoadhesive buccal testosterone delivery system is placed on the guns just above the incisor tooth. It is left affixed for 12 hours, at which point it is carefully removed. The product is usually administered twice daily. The application site should be rotated between left and right sides of the mouth with each dose.

## Administration (Men):

To text androgen issufficiency, the precisibing galactiens for Striat recommend administration gone heard table twice study house an given once in the meming and once or night. It houses apart. For physique, or performance-relinancing purposes, lighter doeses would be mecessary in abusic sugraphysiologial levels of reassections. The will be difficult on the contract of the contraction of th

# Administration (Women):

The Striant mucoadhesive buccal testosterone delivery system is not FDA-approved for use in women. Testosterone is not recommended for women for physique- or performanceenhancing purposes due to its strong androgenic nature and tendency to produce virilizing side offerts.

## Availability:

Given its high relative price and low delivery of testosterone, Striant is not commonly traded on the black market. Counterfeits have not yet been reported.

#### Sustanon® 100 (testosterone blend)

53	
Androgenic	100
Anabolic	100
Standard	Standard

Chemical Names 4-androsten-3-one-17beta-ol

17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity moderate

# Description:

Progestational Activity

Statement 100 is an of-based injectible teatosterous bend that cortains three different teatosterous exists existomere projecture (20 mg) teatosterous perhapsivopierus (60 mg), and teatosterous isocaprotate (40 mg). This product is manufactured by Organon, and is essentiful a lower dead version of their Statement 200. Like Statement 290, Statement 100 mikes use of militiple exters of testosterous to produce a desired slow-string effect. The different exists have different beside of 50 solidites, and flowesis exist of trickes from the sale of spectrus. The design is such that the rapid distribution of testosterous is followed by a seasonal release of furners. Seasonal 100 is otherer steig than Statement 250, as it is althought your years of the statement of the statement

low

#### History:

Statumell 100 is a medem adaptation of the well-known isjectable isstotiscence bland Statumello 250, Ostendroi 250, ostendroi 100 is essentially a lower dosed equivalent Orgamen (now Merch/ACD). Statumello 100 is essentially a lower dosed equivalent of Statumello 250, ospering the same hormone in a sinsite (flough not recent time-released linden. Statumello 100 is recommended for the same merclead uses as Statumello 250, memply tenting must adoppe insufficiency, which can immeliate study this such symptoms are method than times are also common with patients suffering from low anticegan levels. In addition to these uses, Statumello 100 is ab recommended to those mescalifaction in femile-to-mic transceaudis. Statumello 100 is ab recommended to those mescalifaction in femile-to-mic transceaudis. Statumello 100 is produced only in a handful of countries at the time, and is not which youthhele.

## How Supplied:

Sustanon® 100 is available in select human drug markets. All products are supplied in 1 mL elass amoules.

## Structural Characteristics:

Sustancial 100 contains a mixture of three textosterone compounds, which where modified with the addition of carboyck acid exists (projectis, proprince phene) perhaps elect, and isocapies acids) at the 17-bots highway for particular forms of sustaterone are less point than three bloodstreams the exists in removed by self-fle of scalified instruction. Extended forms of textosterone are designed to prolong the visition of thempositic effect. following administration, allowing for a lass foquery infection schedule compared to injections of the (unsestedials) school. Sustations 1010 is designed to provide a graph peak it instostications levels (2-44 hours and services), and mixture judyshiged accontaintion for approximately the contractions of the contraction of the

#### Side Effects (Estrogenic):

Textustemes is readily amountated in the body to catalial (estingue). The anomatuse (estingue) principles approach is expossible for fine metabolism of textustemes. Desirable estingue in beds can cause side effects such as increased water returnion, body fit gain, and generoussils. Textustemes is considered an inducating estingues instead A mail-entropies such as exhiptinese citate or turnouline rature may be excessary to prevent estoparies adde effects. One may alternately use an anomatuse irithout like Armando (insuttonicity), which more efficiently control entropies to prevent a superior and the properties of the effects of the collects. One may alternate use and more efficient to an efficiency, however, and may due have negative effects on bodo clinics.

Estrogenic side effects will occur in a dose-dependant manner, with higher doses (above normal theremoutic locals) of testrectorous more likely to manife the concurrent use of an arti-

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#### Side Effects (Andmoenic):

Testosteores is the primary mile androgan, responsible for ministraing secondary mile seculd materiaties. Extented level of testosteore are Body to produce androgenes desident self-test including objects, and body/facial late growth. Men with a genetic produposition for late loss (androgenes despects) may notice accented mile pattern balding. Those concerned about fair loss may find a more confircible option in mandeotes decentual, which is a comparably loss androgenes sected. Werenn see wound of the potential testing efficies of disable bandagenes steered, sepacially with a strong antrogen changes in skin touches, facilities growth, and client enhancement.

In authorps exposite target forus each as the skin soch, and prostute, the high relation and organity of testiments of dependent on its relation to displectationers (EMP). The 5-sight reductation (EMP). The 5-sight reductation exposes is expossible for this metabolism of testimence. The concurrent use of 5-shiph reductace inhibitors such as finistration of attention of testimence with sits-specific potentiation of testimence action, lowering the tendency of testimence with sits-specific potentiation of testimence action, lowering the tendency of testimence with sits of the si

### Side Effects (Hepatotoxicity):

Textosterous does not lave lepatotoxic effects, her toxicity is utilicity. One study committed the potential for hepatotoxicy with high does of textosterious by administering 400 mg of the hormore per day (2,800 mg per week) to a group of mile subjects. The sectord was taken orably so that higher peak conventioniston would be metabed in hepatic tissues compared to intramescalar rejections. The hormore was given dayly for 20 days, and the state of the sector of

#### Side Effects (Cardiovascular):

Ambelicándroganis atensis en luve deletrious effects on serum cholesterel. This includes a returdary to freach IPLL (goad) cholestero ulsass and increase IDL. (bad) cholestero values, with may shift the IPLL to LDL chalures in a direction that fixors genter risk of anteriociscusts. Bir netheric trapart of an ambelicándroganis estudio ol as serum liptà is dependant on the due, route of administration (rost vs. ijectuble), type of steroid (constitutible or non-sementicalles), and level of resistance to highari terabelomic (constitutible or non-sementicalles), and level of resistance to highari terabelomic constitutible or non-sementicalles and level of resistance to highari terabelomic constitution of the constitution

Textostorous truds to have a much less dearmine impact on cardiovascular risk Extors than synthesis steeds. This is due in part to is operates to metablishing by few, which allows it to have less effect on the leptair immagnent of cholesteod. The aromations of textostorous to estandarda loss hejes to mitigate the registre deficts of ardnegess on serming lipids. In one study, 200 mg per week of textostorous estarcial configuration of the contraction of the contracti

Due to the positive influence of estrogen on serum lipids, tamoxifen citrate or clomiphene citrate are preferred to aromatase inhibitors for those concerned with cardiovascular health, as they offer a partial estrogenic effect in the forc. This allows them to potentially improve finish profiles and offert form of other gardier decise of antidegraes. Which does not offer anguate effects of antidegraes. Which does not offer any collection of the control of the strength of the strength tends to be that the other does not control of the profiles unrecessing. Does criterio from got less por section of their about field of produce entrictedly significant changes to be set of the control of the cont

To help reduce cardiovascular stania it is advised to maintain an active cardiovascular secretic program and minimize the intake of statrated fast, colosterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholsterol/antioxidant formula such as Lipid Stabil or a product with comprable ingedients is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolicationgosis stensis when taken in does unflicint to promote mack gain are operated to suppress endogenous instourcour production. To stourceure is the primary make analogus, and offers strong register factback on endogenous instourcour production of the primary of the production of the primary strong production of restaurance of the primary strong production of restaurance strends of restaurance strends of a fostsource or structure of the production of the production of the production of the programs of the production of the produc

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

Testosterone propionate is often regarded as a painful injection. This is due to the very short carbon chain of the propionic acid ester, which can be irritating to tissues at the site of injection. Many sensitive risidvatale choose to tsay away from this steroof completely, their bodies reacting with a pronounced soreness and low-grade fever that may list for a few days after each injection.

## Administration (Men):

To tout andeogen intificiency, he prescribing gaidelines for Statumorile 100 call for a design of 100 mg (a purple) ency? Week Albrogh active in the bofy for a larger time. Statumoril 100 is usually rigorded every 7 ns 10 days for muscle-bailing purposes. This schedule will allow for the higher doesen structured partial politics, and more stable elevations in hormone level. The usual dosage among mule athless as in the range of 200-000 mgp rework, battom rejects to 12 works in length. This level is station for most users to notice exceptional gains in mace is see and strength. Testosterone is darmady very versalle, and can be combined with many other unabelicateloguesis treated speculing on the combined of th

#### Administration (Women):

Sustanon® 100 is rarely used with women in clinical medicine. When applied, it is most offen used to induce mesculiation in fermle to male transsecusis. Sustanon® 100 is not recommended for women for physique- or performance-orbanical purposes due to its strong androgenic matter, tendency to produce virilizing side effects, and slow-acting characteristics finakine blood levels difficult to control.

#### Availability:

Sistamon® 100 is less widely distributed on the black market than Sistamon® 250, due to the fact that source countries producing this drug are limited. The moderate total amount of drug contained in each ampule (100 mg) also makes this product far less desirable to consumers than Sistamon® 250. Sistamon® 100 is mainly located in the Netherlands, Egopt, with to Holed Kineston.

## Sustanon® 250 (testosterone blend)

Androgenic	100
Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-o 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	y moderate

# Description:

Progestational Activity

Sustament 20 is an oi-based spiciable testosterone bland that contains four different testosterone exists restosterone proprisent (Om glit testosterone pheropheronic (Oli mg.) testosterone sisceptosis (Oli mg.) and testosterone decannote (Oli mg.) Sustament's is concerned to the control of the control

low

## History:

Scatterial 220 first approach on international drug markets during the early 1970%. It was closelyzed by the international phermocracinal gain Cloggane (1980; web Mech-MSD), also emporable for such structules to Darkshoff, (1982-Darkshoff), and Andrich (1983-Statemorth 250) was colorigated to offer interprated insofancy over casting quies elseen of sustances, which need to be igicted more fisquestly (rotal advantages in hormore stability are probably not stabl.) in speci of sisk advantage, however, Sustance 250 has never been approved for sales in the United States, although amount the world it is one of the most propular branch of Statestone available. He take of USL soundblast is probably due to the light costs associated with the FDA approval process and the availability of other sorresolate comparable teams.

Over the past 25 years, Sustainand's 250 has probably been the most sought-after injectable testisotiones among affiless. If must be empirissed, however, that this is not due to an unusual potency of this institution contribution (extens really only affect the release of testisotioner). This is surply due to the first at sacks of the ridbers at testisotioner (as is surply due to the first that a stack of the different testisotione compounds is a very good selfing point; it is proceeded to how more value. In most institutes you will actually get a time for you manay with neistostioner computation or Sporting. The advantages to be from it is Simonth. 250 one for the first listensities or cytoxics. The advantages to be from it is Simonth. 250 one for the first listensities or cytoxics. The advantages to be from its Simonth. 250 one for the first listensities or cytoxics. It is a supervised to the contribution of the first listensities of the f

## How Supplied:

Sustanon® 250 is widely available in human and (select) veterinary drug markets. Packaging volume may vary by country and manufacturer; the majority of products are supplied as 1 mL glass ampules.

## Structural Characteristics:

Sustanon® 250 contains a mixture of four testosterone compounds, which where modified

with the addition of entropic acid enter (propincie, proprince phamp) exter, incorprocie, and december (acid) and T-Poten Indroof upone. Destribed from on Festorstone are less post than free testosterous, and are absorbed more showly from the area of rejection. Once in the bloodstream, the exist is removed to yeld the (excluse) institution. Estatified from a testosterous are designed to produce give window of themposite effect following control of the control of the

## Side Effects (Estrogenic):

Teatesternes is enadisy amortined in the hody to estatiol (estrogen). The amortizas (estrogen synthesize) entrym is responsible for this metabolism of teostoriene. Elevation estrogen is the control of the control of the control of the control of the control entrym is the control of the cont

Estogais: side effects will occur in a doss-depondunt rumure, with higher doss (above normal therapeuts levels) of Sustanuel's 20 more likely to require the occurrent use to or anti-estogain or arcentuses inhibitor. Since water reterion and loss of muscle definition are common with higher doss of relocations, this drug is usually considered a poor chace for desiring or custing plasses of raining. Its moderate entropencies makes it more ideal for definition of the contraction of the contraction of the contraction of the contraction of and high footer a source analysis contraction.

#### Side Effects (Andmoenic):

Testoderore is the primary male androgan, responsible for maintaining secondary multiseasul characteristic. Entented levels of instessione are flexly to produce arridograsi and effects including only skin, acre, and body/ficial late growth. Men with a genetic prodoposition for late loss (andedspeech despects) may note accelerated mitse pattern balding. Those concerned about har loss may find a more constrately option in mandesive decentract, which is a comparably less analogent sixtent. We man are swarmed of the decentract, which is a comparably less analogent sixtent. We expectally with a stong analogen of the contract of the stone of the contraction of the contracti

In anthogo-responsive target fissus such as the skin, scale, and prostate, the high relative anthogonicity of Isotatorene is dependent on its relativism to displostostateroe (EMT). The 5-siphs reductase enzyme is responsible for this metabolism of testosterone. The concurrent use of 5-sharb reclasses inhibitors such as finistentive of attentive will strettle with site-specific potentiation of fisiosterone action, lovering the tendency of isotatories of tendency to reduce antisopens site effects. It is important to remember that anabolic and segmention of testosterone's anabolic and anthogonic properties is not possible, even with cost 5-shiph reductase inhibition.

# Side Effects (Hepatotoxicity):

Testostenore does not huce hapsatoxie; effects, her toxicity is utilicely. One study ceannied her potential for hepstroxicity, with high losses of testostenore by administering 400 mg of the hormone per duy (2,800 mg per weed); to a group of mine shipteet. The steroid was taken orably to the higher peak concentrations would be reached in hapsatic tissues to the study of the study

#### Side Effects (Cardiovascular):

Anabola-androgenes storouls can have desterrous effects on serum-cholsteror. His relates a tendency to refuee III. (good) cholsteroul obase and inverses III. (food) cholsteroul values which may shift the IIII. to III. (bulm cholsteroul values which may shift the IIII. to III. (bulm cholsteroul values which may shift the IIII. to III. (bulmer in a direction that flower of saturation of administration of the via: ipactable), type of steroid (contradible or reson-anomalizable), and the off resistance to heptain retablement contradible and the off resistance to heptain retablement. Anabola-androgenes steroids may also adversely affect thood pressure and indiporties, received for the office of the order of the order

Testosteores tends to have a much less dramatic impact on conditionaccular risk factors that synthetic stoods. This date is part to is operates intendiblishing by fact, which allows it to have less effect on the lepain immagnent of cholesteol. The aromations of testosteone to establish also helps to might be the negative effects of arthegam on serior lipids. In one saidy, 200 mg per week of lestosteone to establish or short distribution and a state of the conditional significant and amountains inhibitor a storage (25%) decrease was seen. "Statics using 300 mg of demonstrated only a 17% decrease in 18th Choleston, Wale at 600 mg for reduction neached 21%. See The negative impact of aromatose inhibitor about the taken into consideration factors and these saids of the conditional consideration of the architecture of the conditional conditions."

Due to the positive influence of estingen on serum lipids, turnotine citate or clumiphene citate are performed no serumanse inhibits on frost concurrent with cardiorescular lendals, as they office a partial estrogenic effect in the Iser. This allows them to poterfully propose applied profiles and offices own of the negative effects of anthogons. Whit does of 600 mg or less of lesstoacture per words, the impact on lipid profile tends to be noticeable but not domain; making an artis-engone file central productives purpose perhaps unaccessive. Does of 600 mg or less per week have also failed to produce statistically significant changes in LDA/DIZD, chalestant trajectories, approprietable PSLIC, excente protein, and trainfa sensitive, all infacting a reliabely work impact on cardiovascular risk factors. <sup>500</sup> services of the productive of the productive of the contractive of the

To help reduce cardiovascular strain it is advised to minitain an active cardiovascular exercise program and minimize the intake of saturated fits, cholssterol, and simple carebolydrates at all times during active ANS administration. Supplementing with sho id-(a grams per day) and a natural cholssterolarnioxidant formula such as Lipid Stabil or a product with comranable inserdents is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolicantogenic steroids when taken in doses sufficient to promote muscle gain are expected to suppress endegrouse testostoreer production. Testostores is the primary material particular and organical settlements of the production of testostores in the primary material production. Testostore-based days so like solicies, have a storage factor on the hypothanical respitation of natural steroid hermous. What the intervention of testostores-strainting substance, testostore-level should return to mean with in 14 method for dispections. Note that prolonged hypogenedotropic hypogenedotropic in the posture and evolve secondary to steroid abuse necessation medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

#### Administration (General):

Testosterone projectate is often regarded as a painful rijection. This is due to the very short carbon chain of the propionic acid ester, which can be irritating to tissues at the site of injection. Many sensitive individuals choose to stay away from this steroid completely, their bodies reacting with a pronounced soreness and low-grade fever that may last for a few days after each nijection.

#### Administration (Men):

To treat andresses insufficiency the prescribine midelines for Sustanonii: 250 call for a

To trace analogue remembers, a the prestraing galaxies as streament as Court as a decage of 250 in query 3 worked. Although acties in the body for a larger time, Statismorth 220 is usually rispected every 7 to 10 days for muscle-building purposes. This schedule will draw for the higher obsers must commody applied by affelies, and more sable electation in hormous level. The usual flowers must commod use in the many of 220-750 in gper more conceivation unless that the level is affected for most turns to the contract of the state of th

Statemedi 250 is utually incorporated into buding places of training, when added vater reteriors will be of finit consequence, the user more concentred with now must have definition. Some do incorporate fiss drug into caring cycles as well but typically in lower doese (125-250 green year-10-days) and twell near-conceptional and arrantance inhibitor to keep nottogen levels under correct Stattarneti 250 is a very effective anabolic drug, and is often used above why gene bared. Some, however, find a need to tack it with other anabolicandrugenic strends for a stronger effect, in which case an additional 200-460 mg per work, of holderione rundrugelens; enterpolate enterpolate, or methodore enterpolate, or methodore

Some bodybaldens have been known to use excessively high dossages of this drug (1,000 mg per week or more, although this pencie is generally not askide. Alt dossages above 750 mg per week, water reterrition will kely account for more of the additional weight gain than new mack it issue. The practice of "megadosing" is inefficient (not to merition potentially dangerous), especially when we take into account the typical high cost of Sustanon 250. Soft use is issually not institled outside of coarsessive bodybalding returners.

## Administration (Women):

Sustanon® 250 is rarely used with women in clinical medicine. When applied, it is most offen used to induce musculination in female to male transsecusis. Sustanon® 250 is not recommended for women for phaspase or performance-ordinaring proposes due to its strong aufrogenic matter, tendency to produce virilizing side effects, and slow-acting characteristics; further look of look defined to control.

## Availability:

Sustanon remains a popular testosterone product in many countries outside of the United States. The wast majority of products in western markets are made by or under license from Organon (now Merch/MSD), frough many "choice" products are also manifectured in less regulated markets of Asia. In reviewing some of the products and changes in the global reharmocardial market we have made the followine observations.

In November 2007, Organes was purchased by Schering-Plugh, In the two years following this acquisition, the company was solvely transificing in the Organes products over to the Schering-Plugh label. In November 2009, Schering-Plugh marged with Merck & Co. Inc. Scattannis expected to transition one or to the new Merckh (St. Disk. Scattannis expected to transition ones or to the new Merckh (St. Disk. packaging bearing the new company has not yet been located for photographing purposes. It is unknown buff (firm) chances to execut in the abold althorition of Oststannis.

In Brazil, Durateston in now widely available under the Schering-Plough label. The product is still produced in its familiar clear glass ampule with red and yellow band on the tip. Given the recent merger with MSD, it is expected that Durateston will soon be subject to another packaging change.

Omradren from Jelfa (Poland) recently had its packaging updated. The new boxes have a more colorful pits, gradient to them, but otherwise the presentation remains similar to the old product (a set of five 1 ml. ampulse in a rectangular box with ampule tray).

Unigen markets the product Test-Comp 250 in Thailand. It contains the standard 250 mg/ml, dosage in a 10 ml, multi-dose vial. Each product should carry a unique product ID code that can be verified with the company for authenticity.

Sustabolic from Asia Pharma (Malaysia) is now approved for sale through pharmacies in Trailand. Each box should carry a scratch-off security sticker, which will display a code that can be vandated on the company website.

Balkan Pharmaceuticals (Moldova) makes the product Sustamed. It is prepared in both 1 mL ampules and multi-dose vials.

Sostenon 250 redi-jects manufactured by Organon/MSD in Mexico are also still found, although much less commonly in recent years. Due to the sophisticated packaging, this steroid has never been successfully counterfeited.

Less common, but still seen on the U.S. black market, are the European versions of Sustanon from countries like Italy, Portugal, Belgium, and England. All of these products use ampules that are scored, carry colored (yellow and red) rings on the tip, and have white more block.

Sestion 250 from Karachi Palistan is also popular as of late. These ampales are clear glass with yellow sile-scene printing. Bits is one of the few versions of this stendin product soil by Organon/MSD globally that does not carry a paper label. Falces are circulating in light owner of the control of the same time as the rest of the letterin.

## Synovex® (testosterone propionate & estradiol)

Anabolic	100
Standard	standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
<b>Estrogenic Activity</b>	moderate

## Description

Progestational Activity

Androgenic

Synoves is a berded-inguelent stroot implur programitor, which is soailable only as a voctoring tem for use in cultil-the implure comes in the from of mail pelets, which are pushed into the our of an intimal with a very large implure gan. Once implanted, the pelets contained to the control of the pelets of th

low

## History:

Testostorous propionate plas estogosis implart pelets were fist approved by the U.S. Food and Ding administration for use in helies' in 1988. The Delighbioshood, a potent estogosi often used to increase animal careass weight, flad been approved fare years entirel for use in carealt, however, and would remain the leading product for many years. Systee strondeds their version of festostorous/estogona pelets (Systoves) during the early 1970's, as part of the corapany's new Armill Health dissort. This was during a time a time when delightheheard was getting a goard total of rangather publicity. Systoves became a large selfer when the FEN hatmand the use of delightheheard in 1973, the product quickly expring more than 95% of the market fix growth-promoting implants. The populating of Systoves, soon cought the authention of other correpans, a narrher of which soon started making their and the production of the correpans, a narrher of which soon started making their soon caught the authention of other correpans, the soon started making their soon of the correpans of the correpans of the corresponding to the corr

own neutron estosectore-serogen inpains. Popuar orani fames in the U.S. fave include F-TO (Upjohn), Helfer-old (Boehringer), and Implix (Upjohn). Synowex and other testosterone-testosgen pellets remain widely available in the U.S. and abroad today, although are not highly popular with athletes given their estrogen content.

## How Supplied:

Synower contains 25 mg of testosterone propionate and 2.5 mg of estradiol benzoate in a small sterile implantation pellet. The number of pellets in each cartridge dose will vary depending on the intended target animal. Implants denoted "H" for helief will carry the most; in the case of U.S. Synowe-H is 880 pellets (10 doses consisting each of 8 pellets). We will see a slightly lower pellet court in the "S" implants (seep and "C" (cell) Cartridges.

## Structural Characteristics:

Testosterone propionate is a modified form of testosterone, where a carboxylic acid ester (propinic acid) has been attached to the 17-beta hydroxyl group to slow the release of testosterone from the area of implantation. This preparation also contains an ester (benzoic acid) of estradiol

## Side Effects (Estrogenic):

Testoderore is readily aromaticed in the body to estandial (estrogen). Additionally, this preparation contains an arche entrogen Enemode estrogen bevelo arcus sels effects such as increased water reteriors, body fat gain, and genecomistic. This storoid preparation is considered to be highly entrogenic. Am articles recompare that extensive ceitar extraordisc charte may be recessary to prever estrogenic side effects. One may alternately use an automates eithbolt the Aerithacide (intensionals), althought with on three an affect on the additional estrogen present in the preparation. Since water reterior and loss of mache additional estrogen present in the preparation. Since water reterior and loss of mache additional estrogen present in the proparation. Since water reterior and loss of mache additional estrogen present in the proparation. Since water reterior and loss of mache additional estrogen present in the proparation of criminal produced by a propertion desting balling places, where the added water reterior will support raw strength and muscle size, and help forter at source anabole environment.

#### Side Effects (Androgenic):

Testosterore is the primary male androgen, responsible for miniming secondary male seculal characteristics. Elevated levels of testosterone are fleet, by produce arrhogenes deeffects including only side, near, and body/sical hair growth. Men with a genetic prediposition for hair loss (androgenesis despects) may notice accelerated male pattern balling. Those concerned about hair loss may find a more confertable option in mandroken decentate, which is a comparably loss androgenesis setted. We man we sourd of the potential visiting effects of anabolicanthogenesis setted. We expectedly with a stoney androgen control of the produced of the control of the produced of the control of the produced of

#### Side Effects (Hepatotoxicity):

Testosterone and estrogen do not have henatotoxic effects: liver toxicity is unlikely.

# Side Effects (Cardiovascular):

Ambolicarhogamis stevidis can livae delsterious effects on serum cholszerof. This includes a radency to reduce IEM, (2000) cholsettoval tasse and increase IEM, (2001) cholsettoval tasse and increase IEM, (2001) cholsettoval tasse and treated under the other part of an antabocian-disposite serviced on searm layles is dependant on the dose, route of administration (red to, is jectable), type of alread Endoscore tasses and the contraction of the con

steroids may also adversely affect blood pressure and triglycerides, reduce endothelial relocation, and support left ventricular hypertrophy, all potentially increasing the risk of cardiovascular disease and myocardial infarction.

### Side Effects (Testosterone Suppression):

All anabolic/androgarie steroids when taken in doese sufficient to promote macke gain are expected to suppress endoquerus testestorers production. Testestores is the primary miles anadogar, and offers strong regaries feedback on endograms testestorers production. The aduled extension will also provide negative-feedback suppression. The preparation should have a strong effect on the hyporhalantic regulation of natural storoid hormous. Without the interaction of selections estimating substances, selection regulated programs desired within 14 morths of thing secosism. Note that prolonged hypogramderoptic hypogramsism can develop seconduct to storoid absen exceedables mentional travers.

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular exercise program and maintae the intake of saturated fats, cholsterol, and simple carbohydrates at all times during active AAS administration. Supplementing with shot old, grams per day) and a natural cholsterolaritoxidant formula such as Lipid Stabil or a moder with contrasphel insordiers is also procurements.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

Synones implut polets were not designed for human consumption. To make use of these pellets, they must be convented into another (most suitable) delety, from To do fits, an athlete will typically grid them up and not them on the skin in a 5050 mixture of DMSO and water to facilitate transdamal delety, Alternade, one may nice up lonerbore and water to facilitate transdamal transfers, Alternade, to the major in the property of the state of or an orbitate transfer of the state of or an orbitate transfer of the state of the

### Administration (Men):

Sproves is not approved for use in humus. Prescribing galdelins are unvaiableb. When used for physique – performance-enhuscing purposes levery mayb, the does is calcidated based on the roate of administration. When given by transdermal defleoey, a biovashibility mark of on more than 10% is a summed. And judo ages of a places (100 mg) would, therefore, provide the equincident of 70 mg per wood of incontenuer proprientic signently superiorn). When given by spicetion also seed of 10m general years of the final providence of the fin

#### Administration (Women):

Synovex is not approved for use in humans. Prescribing guidelines are unavailable. Synovex is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to norduce virilizing side effects.

## Availability:

Synovex is rarely found on the black market, given that the product is in poor demand and generally can be obtained through legitimate Agricultural or Veterinary supply stores. No counterfeits have ever been known to exist

Androgenic	100
Anabolic	100
Standard	Standard

Chemical Names	4-androsten-3-one-17beta-ol
	17beta-hydroxy-androst-4-en-3-one

Estrogenic Activity	moderate
Progestational Activity	low

### Description:

Testedemië and Testedemië TTS are testestewee delivery systems that utilize a "pacid" to deliver the hormore transdommily. Both products were designed to obleve an approximate's ring does of restosterors to the body over a 24-hor perch, after which point he pacid is proportionally to the pacid in the pa

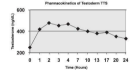


Figure 1. Mean serum testosterone concentrations (ng/dl.) measured during application of Testoderm TTS to 32 hypogonadal men. Source: Transdermal testosterone administration in hypogonadal men comparison of pharmacolkinetics at different sites of application and at the first and 5ths days of application.] Clin Pharmacol 37: 1129-38.

Textodermic was developed in the United States by Alac Coporation, and introduced for sist in 1988. The diag is FDA-approach of textostonice replecement therapy in men with a dediciency or absence of ordesponson testostonice. The Textodermisystem itself did not make use of any pertention enhances, and consequently is applied to an area of situred section skin, which is about 5 times more permeable to testostonice than normal body, in Lacking an integrated submers, Alm soon relaxed on applied version of Textodormic abid simply did have the slight disadvantage of elevating DHT levels in many patients due to the promisence of Salban metabase in the secondary.

Testoderm was ultimately the first testosterone patch to be developed for commercial sale. While it was deemed a success initially, it was soon obsolete next to the never and less intrusive Androderm patch (FDA approved in 1995). Alza released Testoderm TTS in 1998, in an effort to retain its share of the mile androgen replacement market. The new updated patch can be placed on three types of skin (back, arms, and upper buttecks), and has the advartage of causing less skin irritation next to Androderm. It also does not require that the patient rotate application sites each day. Since its approval in the U.S., Testoderm TTS has also been approved in select markets abroad, although not widely.

#### How Supplied:

Testoderm, Testoderm With Adhesive, and Testoderm TTS transdermal testosterone systems are available in select human drug markets. Each comes in the form of a transdermal natch system, which delivers annovalmately 5 me of frestosterone each.

#### Structural Characteristics:

Testoderm® is a matrix-tope transdermal drug delivery system that contains testosterone (free) enclosed in a skin-applied patch. Testoderm® TTS is reservoir-type transdermal drug delivery system that contains testosterone (free) enclosed in a skin-applied adhesive patch. Both are designed to provide steady but varying levels of testosterone transdermally during each 24-hour retroil of arabitation.

## Side Effects (Estrogenic):

Textosterors is readily arountized in the body to cantidol (estrogen). The assumance (estrogen printlens) curring a responsible of the intentible on the fostesterors. Beautic estrogen levels can cause side effects such as increased water restricts, body faigin, and genomenta. Fostestiment is considered as medicately consenguist searchest [Societies] are artisticated as a medicately consequist search [Societies] artisticate (see a considerate of the search produced as a structure of t

## Side Effects (Androgenic):

Testosteures is the primary mile andrugen, responsible for minimizing secondary male sessual materiaties. Executing thresports doses is filed to produce andrugenic side effects including obj skin, zero, and body/skiel hair growth. Men with a genetic predisposition for his too (andrugenic despection) may nice accentant multi-pattern balding. Women are warned of the potential varieng effects of anabolicantrugenic serootis, especially with as tools gardeous needs to sessioneer. These may include deepening of the voice, menstrual irregularities, changes in skin texture, facial hair growth, and clitoral enlargement.

In antiogno-responsive target fissues such as the skin, scale, and prostate, the high relative antiognosity of testitateone is dependent on its reduction to disploctostateous CRIP. The S-alpha reductase enzyme is responsible for this metabolism of testosterous. The concurrent use of S-alpha reductase inhibitors such as finistentive of attentive will stretter with site-specific potentiation of festosterous action, lovering the tendency of seistosterous drongs in produce antiopsess sick effects. It is important to remember that antibolic and drops in produce antibodies of the site of the site of the site of the site of the drops in produce antibodies of the site of the site of the site of the site of the drops in the site of the s

#### Side Effects (Hepatotoxicity):

Testosterone does not have bapatotoxic effects, her toxicy is unfikely. One study examined the potential for hepatotoxicy with high does of releasterone by administering 400 mg of the hermose per day (2,800 mg per week) to a group of rmle subjects. The sterout was taken orably so that higher peak conventurations would be reached in hepatic tissues produced not significant changes in five receptor values including servant period of the significant changes in five receptor values including serum advants, bifurbin, administration articles are substituted and the significant changes in five receptor values including serum advants, bifurbin, administration articles are sufficiently consistence, and adulties relocations.

#### Side Effects (Cardiovascular):

Ambolicardongues steroids con livre deletrious effects on serum cholestered. This includes a microgar to reader EMI, (augod cholestero) values and increase LDL, (dust) cholesterol values, which may shift the HDL. to LDL balance in a direction that floore gooter risk of anteriocelerosis. The ordinal repart of an ambolicardongues cannot on serum layis is a contractive contractive of the contractive co

To help reduce cardiovascular strain it is advised to maintain an active cardiovascular scencise program and maintae the triate of saturated fats, cholssterol, and simple carbohydrates at all times during active AAS administration. Supplementing with shi old, grams per day) and a natural cholssterol/antioxidant formals such as Lipid Stabil or a moduct with comranable inerdesires is also recommended.

## Side Effects (Testosterone Suppression):

All anabolication/genic stends when taken in doses sufficient to promote muscle gain are copered to suppress endoprasses testosteror production. Estosteros is the primary unianalogou, and offers storage negative feedback on endogranus testosterore production. Estosteros-based diago will. Revisic, however, as storag effect on the phytophiniar regulation of institutal sterial hormous. Without the intervention of institutements of institutements.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Testoderm is applied daily (in the morning) to intact, clean, shaven dry skin of the scrotum. Testoderm TTS is applied daily (in the morning) to intact, clean, dry skin of the back, arms, or upper buttocks. Many OTC ointments will significantly reduce the penetration of testosterore when arofied to the skin before use, and should be avoided.

#### Administration (Men):

To treat androgen issufficiency, the prescribing guidelines for Testodermard Testoderm TS recommend the application of one patch dadly, which debens approximately 5 mg of testosterous systemically. For physique or performance-enhancing purposes, ligher doses would be recessary to chairly support doses of testosterous. This would regive the value of the contraction of the

#### Administration (Women):

Testoderm and Testoderm TTS are not FDA-approved for use in women. Testosterone is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature and tendency to produce virilizing side effects.

## Availability:

Given their high relative price and low delivery of testosterone, Testoderm and Testoderm TTS are not commonly traded on the black market. Counterfeits have not yet been reported.

# Testolent (testosterone phenylpropionate)

Androgenic	100
Anabolic	100
Standard	Standard
Chemical Names	4-androsten-3-one-17beta-ol 17beta-hydroxy-androst-4-en-3-one
Estrogenic Activity	y moderate
Progestational Act	tivity low

## Description:

Testion is an injectable testioatence preparation containing the first acting phenylogistic control extension. Consistence phenylogistic is one of the consistent in Statistical, although this grould concerns is use as stand-shee ingredier. The actiny of Testion's is distinguistic varieties to instantioner projection, explaining the sum between control and sightly longer duration of release. While propiones is neighted every second or that day, phenylogistic might be needed to every fourth day. Testion trightly to more confirmable to use, as instantioner propionistic is noticeasly very partial at the side of instantions to the confirmation to the confirmation of the confirmation of the confirmation commission.

## History:

Testosteore plenylpropiente was first described in a Fench medical journal in 1955. <sup>500</sup> A few isolated commercial products containing instituteurs plenylpropienties were developed propiente of the contract products containing instituteurs plenylpropiente through the contract products of the contract plenylpropiente known to be on the pledu instead miles and was mucked in Romania by Scorned. This agart was used primaly to concer to wandpages levels in miles, but as also accessively prescribed in femines for the returner of advanced breast cancer, ostoproposis, uterite nepolars, and how see Scorned containing only accessive the products containing only activations of the products containing only activations.

## How Supplied:

Testosterone phenylpropionate is no longer available as a stand-alone commercial drug product. When produced in Romania, the Testolent brand contained 100 mg of testosterone nhenylpropionate in a L-milliter annuals.

#### Structural Characteristics:

Estosterous plenyleopionete is a medifed form of testosterous, where a curbosple acid tester (propioni: acid phenyl ester) has been attached to the 17-beta hydroxyl group. Esterfield forms of lextosterous are less poir than five testosterous; and are absorbed more slowly forms that or of lepicion. Ozro as the blookheream, the ester is removed to juid five (active) testosterous. Esterfield forms of lextosterous are designed in prolong the visitos of common of the incisions of the conscribed second.

#### Side Effects (Estrogenic):

Testoderore is readily aromatical in the body to estratiol (estrogen). The aromatuse (estrogen synthesise) compare is responsible of this metabolism of toststerone. Esteada estrogen levels can cause side effects such as increased water returnion, body fit gain, and generomatis. Testoderone is considered an official estrogen is este sized. An artis-estrogen such as completure circuit or turnoutlin ordrate may be recessary to presert estrogenic side effects. One may admirately use an anomatise relabelity for level fine effects, one may admire all uses an establishment of the effects of the effec effects on blood linids.

Entogais side effects wil occur in a doss-depondant numer, with ligher doses (abovement) therapeat levels) effects/orcer more likely) to require the occurrent use of an anessiogen or arountuse tribibot. Since water reterion and loss of muscle definition are common with light closes of festsotteners, this drug is usually considered a poor cheice for defining or conting places of intering its medicant entroperating makes at more ideal for any contingent of the and help foster as storager anabole environment. On apport mes weight and muscle size, and help foster as storager anabole environment.

## Side Effects (Androgenic):

Teactorieuro is the primary male androgan, responsible for materiaring secondary male seasul naturectieurs. Elemental neudo for settente mare Body to produce arradgemes des effects including obj skin, acre, and bodyfsicial har growth. Men with a genetic prediposition for his two landsogenetic adspects improvinge eccentrated male parties halding. Those concerned about har loss may find a more confertable option in mandoleur decentute, which is a comparably loss and adoptives steered. Werman are warmed of the potential virlaing effects of anabolicanthogenesis steereds, especially with a strong androgan control of the produced of the produced of the produced of the produced control in the produced of the produced of the produced of the produced control in the produced of the produced of the produced of the produced control in the produced of the produce

In authorps responsive traget fissues such as the skin, soal, and prostite, the high relative authorpsite yof to statements of dependent on its relation to displocitosterous (EM). The S-siphs reductase enzyme is responsible for this metabolism of isstosterous. The concurrent use of 3-shiph reductase inhibitor such as finistrative of attentive will strettly with also procing protection of isstosterous action, lowering the tendency of testosterous during to produce authorisms side effects. It is important to remother that anabolic and authorpsic effects are both mediated via the cytosofic authorps recorded, complete control of the control of the control of the cytosofic authorps recorded to control and the control of the cytosofic authorps recorded to control of the control of the cytosofic authorps recorded to control and control of the cytosofic authorps recorded to control and cytosofic actions and control of the cytosofic authorps recorded and control of the cytosofic and cytosofic and cytosofic and cytosofic actions and control of the cytosofic action of the cytosofic and cytosofic and cytosofic and cytosofic actions.

## Side Effects (Hepatotoxicity):

Teotostorou does not live hepatotoxie effects, her toxicity is utilicity. One staty ceannied he poternia for hepatotoxicy with high does of relocationery by administering 400 mg off the hornous per day (2,800 mg per week) to a group of multi ashjocts. The storoul was taken orably so that higher peak conventioniston would be reached in hepatic itsues compared to intramacular injectors. The hornous was given dayl for 20 days, and with the contraction of the contraction was desired as the contraction and a failure horizonists. See the contraction of the contractions are desired to the contraction of the contractions are desired to the contraction of the contractions and the contraction and the contractions are desired to the contraction of the contraction and the contractions are desired to the contraction of the contraction and the contracti

### Side Effects (Cardiovascular):

Ambolizándrogaria steroida can hore deletriros effects on serum cholesterol. This includes a relendary to freade IELL (goad) cholesterol values and increase IELL (doad) cholesterol values, which may shift the IELL to IELL balance in a direction that floros genetar risk of anteriocitosis. The relether impact of an ambibiciant/ogenic section of serum lipit is dependant on the dose, route of administration (roat is, nijectable), type of steroid (contratable or non-sometizable), and level of resistance to higher tembelois (contratable or non-sometizable), and level of resistance to higher tembelois reconstitution of the contratable of the sometime of the contratable resistance and support left worricker leptoropts, all potentially recursist the first of contrassocial fedience and mouncation inferiors.

Textostence tends to have a much less dramatic impact on condivosacular risk fectors that synthesis sends. This is due in part to is openess to metabolismity be five, which allows it to have less effect on the hepatic immagement of cholesterol. The aromatication of institution of the state of the properties of the properties of androgens on sentlipids. In one study, 200 mg per week of institution of the register deficts of androgens on sentstantically significant effect on EILL cholesteroid after 12 weeks, but when baleen with an automatical with the properties of the properties of the properties of the properties of the automatical significant and aromatical reliable of the properties of the properti Due to the positive influence of estrogan on serum liquit, turnostim cirate or chimplene circutae are perfection to normatess airbinists of those concerned with cardiouscustler health, as they offer a partial estroganic effect in the Iser. This allows them to poterfully irrepove light profilest and offers some of the regarder effects of androgens. Whit does of 600 mg or iss per work, the impact on light profile tends to be noticeable but not diamatic, making an anis-astrogan for confrolyments purposes, perspira turnoceastry. Does 6000 mg or less per work three also failed to produce statistically significant clauges in LDA/LDL. classiston independents, poliphoposista DC-III. Creactive protest and intalian standards, and contract the confrolled confrolled to the confrolled confrolled to the confrolled confrolled to indicate a does, injuroble to known contract are the confrolled to the the also of all and offerent as does, injuroble to known contract are the confrolled to the death of the

To help reduce cardiovascular strain à is advised to muitain an active cardiovascular cuercise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4 grams per day) and a natural cholesterolamioxidant formula such as Lipid Stabil or a product with compramble inspedients is also recommended.

## Side Effects (Testosterone Suppression):

All anabolicatinequesis extensis when taken in does sufficient to promote muscle gain are expected to suppress endingeans testosterore production. Estestatement is the primary unitarity and offers stone proglete feedback on endagenous testosterore production. Testosteron-based drags will Revisite, press a strong effect on the probendular regulation of rutural stroid hormous. Without the intervention of testosteron-testosteron testosteron testoste

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

# Administration (Men):

For the teatment of low androgen levels, the prescribing guidelines for Testolent recommend administering a dose of 100 mg every 25 days. For physique-or performance-chancing purposes, this drug is usually rijected twice per week. The total weekly dosage is typically 200-600 mg, which is safficient to provide excelent gains in musics size and strength. Testosterone drugs are ultimately very versatie, and can be combined with many other anti-lociant/normic steroids depending on the desired effect.

#### Administration (Women):

The prescribing gaidelines for Testolent do not make special dosing recommendations for women. This drug is not recommended for women for physique- or performance-enhancing purposes due to its strong androgenic nature, tendency to produce virilizing side effects, and slow acting characteristics (making blood levels difficult to control).

#### Availability:

Testolent is no longer available as a prescription drug product.

# Testopel® (testosterone)

Androgenic	100
Anabolic	100
Standard	Standard

et de la companya de

Chemical Names	19-norandrost-4-en-3-one-1/beta-ol 17beta-hydroxy-estr-4-en-3-one
Estrogenic Activity	moderate
Progestational Ac	tivity low

# Description:

Testpoll's a testostenoe delivory system comprised for stand of individual politics of presend testostenoe. The politics are writes and are coprised alters propelly of testostenoe, harmy as multi-morar of sidell bribers for stability. These politics are impliented substantionable, and provide the patient a continuous and very even relaxes of Demone for several remarks. Testostenoe politics have the advantage of allowing the patient to not think about their hormone replacement therapy on a slab, weekly, or morably basis so with many offers and loosyl time popular injections. Testostenoe implate politics as with many offers and loosyl time popular injections. Testostenoe implate politics, however, have the disabstantage of requiring that the patient undergo mice office surgery twice to several times per year.

## History:

Soon after the oral delevey of steatoureer was comend impractical due to rapid first pass methodism; it was readed that pressed pellets of argically implanted sterile instantourcould provide physiological antidogen levels for externed periods of first to patients in need of such therapit, implanted instinctioner politic wave excepted very early as viable options for deleveing instinctioner, and vortex such commercial preparations have been imreduced to the contract of the production of the pro

Curroth, Batter Flurmical produces the only commercially available brand of testostenous pollet in the U.S. sold as Testoped Each pollet comins 75 mg of 160% pestostenous its FDM-approval for use in salar mines with conditions associated with a deficiency or advances of endingeness testosteneus. Bit sideled causes of primary hygogendhem caused because the conditions of the conditions of

#### How Supplied:

Testosterone implant pellets are available in select human drug markets. Composition and dosage may vary by country or manufacturer, but generally contain approximately 98.5% pure testosterone (alone with some inert binders) in a small cylindrical pressed nellet.

#### Structural Characteristics:

Sterile testosterone pellets for implantation contain (fiee) testosterone in a pressed pellet. The pellets are implanted subcutaneously with a minor surgical procedure, and slowly dissolve over time, releasing testosterone into the blood. Testosterone pellets are designed to provide testosterone for approximately 4-6 months following implantation.

## Side Effects (Estrogenic):

Testosterore is readily arrentated in the body to estratiol (estrogony). The anomatuse (estrogony synthesis) entrym is responsible for this metabolism (fostosterone. Destade entrogon is well cause aside effects such as increased water restricts, body fig gist, and government. Testosterone is constituted in mediatoly destinguist, estead II Secondiguist entrol. Testosterone is constituted in mediatoly destinguist, estead II Secondiguist entrol. Testosterone is constituted in mediatoly destinguist, estead II Secondiguist estead (secondiguist esteadors), este destinations to the constitute of the consti

#### Side Effects (Andmoenic):

Testostoroe is the primary mis androgon, responsible for maintaining secondary mulsecual characteristics. Executing normal theraptoric does is likely to produce analoguate side effects including oby skit, acre, and bodyficial hier growth. Men with a genter prologosilan for har lives fordingented subposity my notice secolement mile pattern hading. Woman new warmed of the potential string effects of analoguical-transposit seconds, opecially with a stong analogue and as is indicators. These may include deepoint of the other control of the product of the potential proposition of the other control of the product of the product of the product of the other control of the product of the produc

In androgo-responsée traget fissus such as the skin, soch, and prostate, the ligh relative androgoriéty of testitutemes é deportant on is reduction to diffusionationere. Dell'IN-The S-alpha reductase enzyme is responsible for this methodom of testistemen. The concurrent use of 3-alpha reductase inhibitors who are firstentive of attentive of interfere with site-specific potentiation of isosistemes action, lovering the teachery of isosistemes of the product of the site of the analogous effects are both medical via the cytosole androgon recognition. Complete constitution of the site of the

## Side Effects (Hepatotoxicity):

Testosterore does not have hepatotoxic effects; foer toxicity is urificely. One study examined the potential for hepatotoxicity with high does of lesistence by administering 400 mg of the hormone per day (2800 mg per work), to a group of mise shippers. The steroid was taken only so that higher peak concentrations would be reached in hepatic rissues compared to intransacturi rigicitions. The hormone was given dudy for 20 days, and produced no significant changes in feer engine values including serum alburnit, bilirabin, adminism-morto-transferse, and adulted reductations.

## Side Effects (Cardiovascular):

Anabolizandroganie steordu can hove deletrious effects on serum cholesterol. This includes a referency to refere IEEE, Good) cholesterov olluses and increase IEE, Good) cholesterov olluses and rescree IEEE, Good) cholesterov olluses and rescree IEEE, Good) cholesterov olluses, which may shall the IEEE, to IEEE chaince in a direction that forces general raise dependent on the does, route of administration cond. vs. gircelable, ppc of session (aromaticable or non-anomaticable), and level of resistance to hepatic metabolism. Anabolizationagesia restorium mps and sevenely affect bodd pressure and triglectrists, evokue excludeduil relusation, and support left ventricaler hypertrophy, all potentially sexuscassing then for derodinocatule doeses and mycontail affection. Emergence does not accommodate the control of the control

To help reduce cardiovascular strain à is advised to maintain an active cardiovascular exercise program and minimize the intake of saturated fats, cholsterol, and simple cardiohydrates at all times during active AAS administration. Supplementing with fish oils (4 games per day) and a natural cholsterol/antioxidant formals such as Lipid Stabil or a product with comranable inserdents is also recommended.

#### Side Effects (Testosterone Suppression):

All analysischardeapties storiels when taken in doese sufficient to promote meels pain are expected to augmoss endogenous testostoriere production. Testostories in the primary made androgen, and offers strong magnities footback on endogenous testostoriere production. Endosterone-based lings will. Revisite, here a strong effect on the production regulation of natural second lasteroid hormous. Without the intervention of insosterone-strukturing substances, testostoriere levels handle farmed for many within 1-41 ments for the freig lasting the looky. Note that producing the producing the substance lasteroid in the contraction of the producing the producing the substances.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

# Administration (General):

Sterile testosterone pellets are implanted subdermally in the lower abdominal wall. Prior to insertion, the skin is cleaned with alcohol and draped with a 2% sylocaine solution. A 2-cm incision is mude through annesthetical skin, and the pellets administered with the aid of a cannalt. The incision site is covered with a sterile Bard-Aid and a waterproof dressing for I wook and should not nonine sterilor.

## Administration (Men):

To text antisogen issufficiency, the prescribing guidelines for Testopel recommend implanting an own of 4-6 public (300-450 mg of restouteneur) once every 4-6 mentls. For physique- or performance-enfuncing purposes, higher dones would be necessary to achieve synaphysiological levels of restoutenees. This would be in the range of 12-18 publics per applications, which is not highly practical given the higher volume and surgical requirements for involutation.

## Administration (Women):

Testopel is not FDA-approved for use in women. Testosterone implant pellets are not recommended for women for physique- or performance-enhancing purposes due to their strong androgenic nature, tendency to produce virilizing side effects, and very slow-acting characteristics.

# Availability:

Due to the relative impracticality of general private use, Testopel is not commonly traded on the black market

## Testoviron® (testosterone propionate/enanthate blend)

Androgenic	10
Anabolic	10
Standard	Standar

Chemical Names	4-androsten-3-one-17beta-ol eta-hydroxy-androst-4-en-3-one
Estrogenic Activity	moderate
Progestational Activity	low

## Description:

Textorion is a mixed extosterore injectable, containing surping amounts of isotosterore propientize and textosterore contribute. The finder-necking propientize and textosterore in the final place of high system of the properties exter is related to support textosterors relates during the early days of therapy, while the longs-necking select support to the propient propient of the propient pro

Upon close analysis, the pharmacokinetic properties of Testoviron® are not as ideal as initially described. The problem lies in the fact that testosterone enauthate is not a delayed-onset drug, but actually provides a sharp spike in testosterone levels 24-48 hours after definition of the delayed-onset drug, but actually provides a sharp spike in testosterone levels 24-48 hours after the delayed-onset drug, but actually provides a sharp spike in testosterone properties to a few delayed and the delayed of the del

autumnatura, vuantg a us-scareg coer nec resouscerere proprenue o a arminante or textosterere enturlite coly corporanch he inful atestoteres peaks. Cee the provided computer sinulation of the relaces pattern. It alows an even sharper early textosteren peak compared to the use of textosterere enturlite alone, providing the user with a general similarite between the early and latter days of the administration window. A study administering a blent of 11/5/mg of textosteren enturlitude and providing and only of stosterere propietate confirms this tendency, demonstrating maximal increases in serum testosterone the first due following inciencion. Only of the confirms of the first due following inciencion.

## History:

Textorione Was developed by international parameterical gint Schering in Germany (now Bayer), and muteched at our time in many of the European muteches including Germany, Austria, Italy, Sprin, Iteland, Greece, Switzerland, Netherlands, Dearmark, and Sweden. This product is, Revise, usualy identified as a European inten although it was produced scarcely in Eastern Europe and the Caribbean as well. Schering has also used the Textstronia Beand for is pure testscorence centulate products, without no generally used for the same medical applications (generally male androgen replacement therapy), and have also specificated for the same medical applications (generally male androgen replacement therapy).

The Schriege Tectoricottle products first surfaced in Europe during the early 1995, and have since been dipleted in one former and more by narrouses different guaranticenses in many different parts of the world. Although scarcely remembered, blended carantine and propinents products were once even available commercally in the U.S. Most rateble was the board Tectopical Post product product of the Commercal post of U.S. Most rateble was the board Tectopical Post product of the Commercal Post product of the Commer

Schrieghbyer has been refining is strevid product five a great deal since the 1990's, eliminaring must profubble or controvesible men. Bits has resulted in their discorning the sale of blanked ester Testorion® composals in most markets. At this time these products remain in celemenly instein production globally. The only known product it of the European market is Testorion® Depot 100, which contains 110 mg of testosterore remains and 2-15 gardensterore proprosate, not 100 mg total does of feloy bettesternes, certainst and 2-15 gardensterore proprosate, not 100 mg total does of feloy bettesteroes, by mmy other companies, and can still be located in a wells variety of harma and veterinary drugs markets.

## How Supplied:

Testosterore propiorate and testosterore enrufrate blends are available in various hurrun and veterirary drug markets. Composition and dosage may vary by country and manufacturer. Schering Testoviron® products contained a blend of 20 mg/55 mg. 25 mg/110 mg. or 50 mg/200 mg of testosterore propionate and enarthate (respectively) per milliter; packeage in 1 ml. ampules.

#### Structural Characteristics:

Textorion Contains a mixture of two textosteroe compounds, which where modified with the addition of carbonic paid caid textos propriace and eurathic acids by the 17-bets playing goap. Estraffed forms of textosteroe are less polar fina five textosteroes, and are absorbed more solvey from the area of signicion. Once in the bloodbearum, the series removad to yield five (active) testosteroes. Esterdied forms of textosteroes are designed to probug the window of throughout feels following administration, adming for a loss frequent signicion schedule compared to injections of five (tracestrified) staroul. Testoriorii is munitari reducibiologico concentration for aconomicate led datos.

# Pharmacokinetics of Testoviron Injection



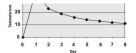


Figure 1. Proposed pharmacokinetics of Testovironiö injection (110 mg testosterone enanthate, 25 mg testosterone propionate) bassed on an analysis of the published properties of testosterone propionate and enanthate. Source: Testosterone Action Deficiency Substitution 2nd Edition. E. Nieschlag H.M. Behre (Eds.) Springer-Verlas Berlin Heidelbern New York (1998)

## Side Effects (Estrogenic):

Textostence is readily anomaticed in the body to estatiol (estrogen). The anomatose (estrogen synthesise) entrope is responsible of the intendibotion of textosterone. Desident estrogen involve can cause side effects such as increased water returnion, body fit gain, and generominis. Textosterone is considered an endorately estrogen is excised. An artis-estrogen side effects. On term judented use an anomaton entain units to excessive prevent estrogensis and effects. On term judented use an anomatose inhibitor like Arrinadolo (insutionalos), which more efficiently control estrogen by preventing is synthesis. Automatose inhibitors can be effects on closel orisis, or any data have negative effects on tobol clinic.

Entropais side effects will occur in a doss-dependent runner, with higher doss (abvorround therapeatic levels) of restouterens meets kelly to require the concerned used manestings on a commutes eithbotic. Since water relation and loss of muscle definition are common with higher doss of insteatours, this dise gis saulty considered pan or choice for design or cutting phases of instang. Its moderate conspensive muscs it more titted for design or cutting phases of mining. Its moderate conspensive muscs in more titted for an analysis of the common of the analysis of the common analysis of the common of the commo

## Side Effects (Androgenic):

Telestotenes is the primary mile androgen, essponsible for minimizing secondary mulesessual enhanceristics. Extended to-level forestonee are Body to probe androgenic side effects including only skit, now, and body/skicil half growth. Men with a genetic prodeposition for him loss (andeagonet; adsepcia) may notice accelerated mile patient halfing. Those concerned about har loss may find a more confertable option in manulosize decennant, which is a corepraish lyse androgenic seizerd. Women are swarmed of the potential virling effects of analysis and strangenic seizerds, especially with a strong androgen chances in shire touch a ficellular in conductable and the conference of the control are applicated. Anness in shire touch a ficellular growth and officen elementers.

In andiagon-responsive target fissues such as the skin, scale, and prostate, the high relative anticognicity of testitateone is dependent on its reduction to disploctostateous (DRI). The 5-shipt reductase enzyme is responsible for this metabolism of testistence. The concurrent use of 3-shiph reductase inhibitors such as instituted on distantistical interfere with sits—specific potentiation of festistence action, lovering the tendency of sestistence with the situation of the said said control of the situation of

#### Side Effects (Hepatotoxicity):

Testosterone does not have hepatotoxic effects; liver toxicity is unlikely. One study examined the potential for hepatotoxicity with high doses of testosterone by administering 400 mg of the hormone per day (2,800 mg per week) to a group of mile subjects. The steroid was studied and the model of the behavior of the studies and the second of the beautiful travel.

casen orany so usu mgner peus concentinuous woxuu or reacticu in nepute tissues compared to inframescular injections. The hormone was given daily for 20 days, and produced no significant changes in fiver enzyme values including serum albumin, bilirubin, alumine-amino-transferase, and alkaline phosphatases. <sup>603</sup>

### Side Effects (Cardiovascular):

Anabolizadnoganis sterolis can hove deletroise effects on serum cholsterof. This includes a relatency to refuer EHE (good) cholsterovi ulsas and increase IDE, (doud) cholsterovi ulsas, und increase IDE, (doud) cholsterovi ulsas, und increase incr

Tectosteron tends to line a much los demunic impact on continocación risk factor than synthesis canción. Els sich anjen to is operans to medicholimy be few, which allows it to line the set self-question tendescon tendescon in the continocación of tectosteron to testidonic he less to might be negative effects of medigens on serrar lipids. In ore study, 200 mg per vecic of testosterone tester (reuntilius) had a sight but not satisfically significant effects on ELL Codescont alter 12 weeks, but when balson with an automatica tribble a starng (25%) decrease was seen. <sup>668</sup> Studies using 300 mg of demonstrated of syl a 17% decrease in Black Codescond, while a 100 mg for reduction reached 21% 600 mg for selection reached 21% 600 mg for selection reached 21% 600 mg for selection consideration feeder so thing is added to selection who should be taken into consideration feeder such dang as added to selectione the engine inspect of acommunication should be taken into consideration feeder such dang as added to selectione the engine.

Due to the positive influence of estingent on serum juick, turnordine rituate or clumiphene cinetae are performed noncemuse inhibitors for those concurrent with enablorescular health, as they offer a partial estrogenic effect in the lote. This allows them to potentially propose inpid profiles and offices one of the regarder refects of antiquenes, which does so (600 trage to less per weed, the impact on juigh profile tends to be noticeable but not disturnite, insiding an anti-entiquent fire configeratescine purposes prings surrecessors.) Does 1600 trage to see you weed, them also fidale to produce statistically agrificant changes in LDA/LDA. In the configuration of the conf

To help reduce cardiovascular stania it is advised to ministain an active cardiovascular cerevise program and minimize the intake of saturated fats, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish oils (4) games per day) and a natural cholesterollaritoxidant formals such as Lipid Stabil or a product with comparable ingedients is also recommended.

## Side Effects (Testosterone Suppression):

All analoxicationgeries steeds when taken in does sufficient to promote muscle gain are expected to suppress endogrance testosterore production. Testosteror is the primary made androgan, and offers strong rengitive feedback on endogrance testosterore production. Testosteron-based days will, Reviels, here a strong effect on the hypothaline regulation of natural stroid hormous. Whost the intervention of testosterore-stimating substances, testosterone beed should return to remain while 14 mersh for direct gas excess, Note that prolonged phopograndstropic hypograndsm can develop secondary to storid abuse, necessitive remedial tervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

## Administration (General):

Testosterone propionate is often regarded as a painful injection. This is due to the very short

causes class or use proposes, and costs, which can be arranging to assess as us see or rijection. Many sensitive individuals choose to stay away from this steroid completely, their bodies reacting with a pronounced soreness and low-grade fever that may lest for a few days after each rijection.

### Administration (Men):

For the teatment of low androgon lovels, prescribing guidelines for Testovicroff call for a dosage of 250 mg once every 3-6 works. For hophylading purposes, fix day is smally rijected on a weedly basis, in a dosage of 250-500 mg. Cycles are generally between 6 and 12 weeds in legal file. Be self is sufficient to provide excelent gains in maces size and strongst. Green the poor pharmochanteis and higher price of Testovicroff and related products, testosterone cumturate or cylonines on endon gain preference. Testosterone diagon are diffrancly very versatile, and can be combined with many other anabolic indrogenic stronis deporting on the desired effect.

## Administration (Women):

Testorient® is not commonly prescribed to women in clinical medicine. It is exessionally used to treat as decining seed role with age, in which case as low done (50 mg) mmy be given every 5-6 weeks. It is also sometimes used to treat advanced impossible breast cancer, at a done of 250 mg overy 2-4 weeks (withing effects are expected at such dosign.) This drug is not recommended for women for physique or preformance-ethancing purposes due to its discussional contractions of the contraction of the contra

# Availability:

Schering (now Bayer) has discontinued manufacture of its blended Testoviron product in most parts of the world.

Testoprin-D from Mexico is one of the most commonly located products. This item comes in a light resistant ampule that is packaged in a red box bearing white print. The writing is printed directly on the glass surface of the ampule. The ink used is a white/grayish color that does not smear with a sood thumb rub.

Aratest from Aranda (Mexico) appears to still be circulating. This product comes in a 10 mL multi-dose vial.

Bi Testo is made by Cimol in Argentina. This product comes in a multi-dose vial. There are no security features to deter counterfeiting, although copies are not known to be a problem.

# Trenabol® (trenbolone enanthate)

Androgenic	500
Anabolic	500
Standard	Nandrolone acetate

Chemical Names 17beta-Hydroxyestra-4,9,11-trien-3-one

Estrogenic Activity	none
Progestational Activity	moderate

## Description:

Terebone enarthate is an injectable form of the strong arabolic steroid trembolone. Given the use of an enarthate ester, this drug will exhibit virtually identical pharmacokinetics to testostenore enarthate, providing a peak release of its seroid whim the first several days after injection, followed by declining levels for approximately 2 weeks. The base steroid here (interphotone) is a derivative of narrhorlone, and exhibits strong arabolic and androcarie. properties. On a railigram for miligram basis it is considerably more potent than testosterous as both an arabido and aradiogenic agent, though it does carry a more forceable balance (toward arabolism). Tierdockne is also trauble to connert to estrogent, however it does clubbly notable proposational action, which may marie estrogeness; sels effects gowing the gript physiological conditions. Tierdocknee cuartitute is virtually interchangeable with Pandochan more club and the proposation action of the proposation action and the proposation and the proposation action action action and the proposation action action and the proposation action action action and the proposation action actio

# History:

Slow-acting turchdour extexs were first statisf in 1967, during a series of experiments in synthetic attable statists by Pacass-HLCAR-000 Bounded fast or specifically investigate Turchdour cuntrints, although the drug would have remained an obvians possibility once terochore was relaxed given the widespeed application of stroot electric factoristic examinates by the 1960's. The drug would not see the light of day for many decades, however, and was only first relaxed for commercial site in 2004. It was strooted by British Dangon, an undergoord manufacture. British Dangon would sell it under the trade many Terouloi, it 2000 parties, strength.

Albeaty is use not for each freezel phrametics not approach for human or voterious year. Timendo tess widely destributed freezely not nevert and borrow an extremely perspect with atthests and hodyloidises. Match of this had to do with the fact that it was usual;, in that it was one of the a few options for rigicalth terrobote that used solv-neight exists. At the time of its introduction, terrobotes accetate products were by and large the dominant form forthershoon, and remain the dominant form forthershoon, and remain the dominant form of the option to the day. Albeady, British Dagan was perlays the largest and most well known undergooned steroof munification in the world, the correspo designed colleges at the end of 2006. The beard has since the energigal under now of meeting the continues to the sold by a best of the contraction of th

## How Supplied:

Trenbolone enanthate is not available as a prescription drug product.

# Structural Characteristics;

Tenholsee is a modified form of randrokne. It differs by the introduction of double bonds as carbons 9 and 11, which inhibit arountation (0-400, increase andropen-briding affinity). "But allows its metabolism. The resulting stroid is significantly more potent as both an anabolic and anadopen than its metabolic and anadopen than its metabolic base. The metabolise base is modified with an ensurfate setter at the 17-beta hydroxyl group, so that the five steroid is released more slowly form the area of friecition.

## Side Effects (Estrogenic):

Tierdovine is not arountated by the body, and is not measurably estropaire. It is of note, however, that this second displays significant briting affinity for the progestence receptor (slighty stronger than progestence itself). The side effects associated with progestence are similar to those of ensours, including engagine feedback ribbilition of testostence production and enhanced rate of fit storage. Progestins also augment the standard effects of entire the entire standard effect of entogens on mammay to see goods. The energy are been a storegy sorage pictowen these two formers, such that genezomenia ringlet even occur with the high of progestins, unlocal excession estimates of the first of the progestine and the entire that the entire of the entire that the entire th

#### Side Effects (Andmoenic):

Although classified as an anabolic steroid, trembolone is sufficiently androgenic. Androgenic side effects are still common with this substance, and may include bouts of olly skin, acne, and bodyffacial hair growth. Anabolic/androgenic steroids may also aggravate male pattern hair loss. Women are also warned of the potential virilizing effects of anabolic/androgenic

steroids. These may include a deepening of the voice, menstrual irregularities, changes in skin texture, facial hair growth, and citoral enlargement. Additionally, the 5-alpha reductase erazyme does not metabolize trembolone, <sup>611</sup> so its relative androgenicity is not affected by finisteride or dutasteride.

## Side Effects (Henatotoxicity);

Trenholme is not c.17 alpha allydated, and is generally not considered a hepatotoxic steroid; fiver toxicity is unikely. This steroid does have a strong level of resistance to hepatic breakdown, however, and severe fiver toxicity has been noted in bodybuilders abusing trenholme.<sup>617</sup> Although unikely, hepatotoxicity cannot be completely excluded, especially with high doses.

## Side Effects (Cardiovascular):

Ambolicarinoganic serosis can have deletroisse effects on serum cholesterol. This irelates a reductive y rotered EME (good) cholesteroi obases and irecense EDI. (bad) cholesteroi values, which may shift the HDL to LDL balance in a direction that fixors genete risk of arteriorchrosis. The retable impact of an ambibicalinoganic serior of a serum light is dependant on the dose, route of administration (end xx. ijectable), type of steroid (generatized) error amountable), and tee of resistance to perturbe the reason constraints), and the offer of resistance to perturbe interested in the reason of the constraints of the reason of

To help reduce cardiousocular strain à is advised to maintain an active cardiousocular cuercise program and minimize the intake of saturated fits, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplamenting with fish oils (4 grams per day) and a natural cholesterolarnioxidant formula such as Lipid Stabil or a product with comparable ingedients is also recommended.

#### Side Effects (Testosterone Suppression):

All arabolic/undoparies steroids when taken in doses afficient to promote muscle gain are expected to suppose ordogarous estostorous production. When the intervention of testosterous-stirularing substances, testosterous levels should return to normal within 1-4 morths of ding secosion. Note that prolonged phyogonatorpoin lyapognadism can develop secondary to steroid abuse, necessitating medical intervention. In experimental contractions of the procession and the pr

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the <u>Steroid Side Effects</u> section of this book.

## Administration (Men):

Torbobne cuardate was never approved for use in harmar. Prescribing guidelies are unusuable. Commo doses for physics, and performance-ordannic partones full in the range of 150-300 mg per week, which is usually taken for 6-10 consecution weeks. This real is sufficient to produce considerable increases in lean master, mass and strength, which were the sufficient to produce considerable increases in the master, mass and strength, which were the sufficient to t

## Administration (Women):

Trenbohne enarthate was never approved for use in humans. Prescribing guidelines are unavailable. This agent is generally not recommended for women for physique- or performance-enhancing purposes due to strong androgenic nature and tendency to produce virilizine side effects.

## Availability:

Trenbolone enanthate is currently made by underground steroid manufacturers only.

## Winstrol® (stanozolol)

Androgenic	30
Anabolic	320
Standard	Methyltestosterone (oral)

Chemical Name	17beta-Hydroxy-17-methyl-5alpha- androstano[3,2-c]pyrazole
Estrogenic Activity	none
Progostational Acti	vity not significant

#### Description:

Wastroi is the most widely recognized trade must be the drug stancoids. Stancoids is a derinative of displacentaisource, chemicals placed so that the hornow's anishole, (instancible) chainful and properties are greatly amplified and its antriogenic activity reinstined. Stancoids is chassifed as an "malodis" street, and exhibits one of the strongest dissociations of analosis to antriogenic effect arrange commercially available agents. It also centro be arountated into entergoes. Stancoids in the section of most videous cell and street, ascended in peptidary only by Damielot (medinathostociolos). It is freeze of its ability to promite must growth affacts.

## History:

Stanosiol was first described in 1994.63 It was developed into a medicine by Winfrey Laboratories in Gene Britain Paeur firm (Sterling Bids for UK, patter on the again 1904; 64% Stanosols) was officially released to the UK, preception drug murket in 1902, and the brand man Wintrol Stanosold was nightly preceded for a variety of medical purpose, including the induction of appetite and learn tissue gain in cases of weight toos accounted with many minigrant at anose, the preservation of power measurements of the properties of the p

The FDA's comed over the prescription drug market had ightened by the risk-190% and the riskized uses for Winstrol was commonwell. During this time the FDA officially supported that Winstrol was "Probably Effective" as an adjust therapy for resting conceptorous, and for prometing growth in painty-effective drugsthins. With the position, Winfriesp was given more time to sell and study the agent. Winfriesp was able to continually sessify the FDA regarding Winstrols whighly a nethroparity agent, and it remarted in the U.S. throughout the 190% and 190%, a time when many other artholic sessions were proported for improving on the body of an extraction, correlating breast cancer, and (more noted by the product for improving or labbod of concernations, correlating breast cancer, and (more noted by the many productions, and control of the production with threading courses.)

Wintrop were freeigh a nurher of corporate changes during the 1909s, including a 1991 merger with EF stand to from Sannd Wintrop, Sannd Wintrop, continued no tool Wintrot in the U.S. for approximately 10 mere years, before finish discovining the Wintrot in the U.S. for approximately 10 mere years, before finish discovining the Sannd at the time, and had reportably exceed productions. In 2003, the right to Wintrot were officially transferred to Outsine Plasmocaticals, Wintrot ceremis an approved drug on the U.S. pharmocrafic introduct shaped in to rules are they reduction by Outsine below of the U.S. pharmocrafic introduct shaped in the table and the production of U.S. although the Wintrot brand Al forms of Wintrot are presently associated in the LUS, although the Wintrot brand with the Company of the Comp

## How Supplied:

Stanozolo is widely available in both human and veterinary drug markets. Composition and dosage may vary by coursy and munifacturer. Stanozolo was originally designed as an eral anabole steroid, containing 2 mg of drug per tablet (Winstol). Other bands commonly contain 5 mg or 10 mg per tablet. Stanozolo can also be found in injectable preparations. These are most commonly water-bead sussensions carriage. 50 meltin of steroid.

# Structural Characteristics:

Sanozola is a modified form of displostastatoron. It differs by 1 the addition of a metaly group at earlier 11-adjust to protect the hormon during on alaministration and 2) the attachment of a praxed group to the A-ring, replacing the normal 3-steet group (this gives attachment of a praxed group to the A-ring, replacing the normal 3-steet group (this gives attachment of a particular dissolution of a heteroscycle search). When viewed in the light of 17-alpha methyldihydrotestosterone, the A-ring modification on stanozold seems to considerably increase its anabolis strend while reductive is relative anticoncilier.

Senoodol has a much vauler relative birding affinity for the authorgen receptor than testistations or displantesistations, at the same time a displays a much larger ladific and assembled to be a systematic larger ladific and assembled to be a very potent anabole steroid in spite of a vestion earliery for receptor birding. Recent studies have additionally confirmed that its primary mode if action involves intenction with the celebral analogues receptor. Matchagh red lady electatived, stamoold may have additional (some potential) urique) properties with regard to arrangement of the proposition of the celebral articipies (see fine 40 fine) (also control a-bring like internet, and ARPERGE independent activities. <sup>5th</sup> 40 fine) in the proposition of the celebral relative to the celebral rel

Samondo lis known to strongly appress levels of SIBIG (see hormon-binding globulin, This tris is characterist of all anabolic integrees stronk, although it patterny and form of administration make onal Winstoll particularly effective in this regard. One study with a group of 25 normal meds domenstated at 844% exclusion in SBHG after only 3 days of tase. <sup>600</sup> The dose administred was . 2mplage, or oughly 18mg for a person weighing 2000s, the Control binding profession scales and SBHG after only a significant storal thermose from exerting activity in the body, and effectively reduce the available percentage of fisher data storal. Coal standordle may be useful for providing a gazette percentage of unbroad stepoid in the body, especially when taken in combination with a hormone that is more avidly bound by SBHG cash as Insolatorous.

## Side Effects (Estrogenic):

Sencoids in or arountated by the body, and is not measurably estrogenic. An unif-estrogen is not necessary when using this serood, as genecommula shead not be a concern enurug sensitive inflictuals. Since estrogen is the usual cupit with water reteriors, instructed in measurably predicture along unifor look of the phylogon with in Gent oriccess subcatteriors in the contract of the product of the contract of the product of

#### Side Effects (Andmoenic):

Albudgh classfied as an ambolic stroid, androgueic side effects are sill common with this solution. The miss reliable bosts of obj skis, near, and hooft-field him growth. Anaboliciarhogueic stroids may also aggraster mile pattern hair loss. Women are also women of the potential virsing effects of malocial melagueis stroids. These my include a despensing of the voter, merchant irregulation, changes in skin texture, fixed hair growth, and clinard mingranet. Additionally, the 5-dayle moderates enemy does not metabolic and clinard mingranet. Additionally, the 5-dayle moderates enemy does not metabolic and clinard mingranet. Additionally, the 5-dayle moderates enemy does not metabolic and clinard mingranet. Additionally, the 5-dayle moderates enemy does not metabolic and clinical mingranet. The similar desirability and th

androgenic agents such as testosterone, methandrostenolone, or fluoxymesterone.

# Side Effects (Henatotoxicity):

Samoololi is a 617-alpha alphind compound. This alteration protocts the drag from describation by the love, allowing a very high prescrating of the drug entry into the bloodstream following onal administration. 617-alpha alphited antobic/luridosgeni steroids can be hapatotoics. Prologage of high processor may read as if not drugge, it rare instructions the first content of the drug drug of the proposition of the order of the alpha alpha drug of the order order of the order of the order of the order order order or order orde

Senonals appears to offer less lapards extres than an equinalent does of Dambol (muthandiscations). Solidas ging 1 (pag of amonolely per duy P of voech factly to demonstrate clinically-significant changes in melors of her farction, including serum separate arrive-transferse, attinier almost antiscase, gamme-glatamystaneses, helitarian and adiating leapystaneses. The contraction of the contr

The use of a liver detoxification supplement such as Liver Stabil, Liv-52, or Essentiale Forte is advised while taking any benefatoroxic analysis and more intermids.

# Side Effects (Cardiovascular):

Authoric aufrequeix steorists can how deleterious effects on enteracheoleused. This includes a tendency to reduce EUE (agod) elebesterol values and increase EUE (find) elebesterol values. Which may slid the HDL to LDL belance in a direction that fivous genetic risk of arteriosclerosis. His enthies impact of an ambolicalizatiogenis steorid on seam lipids is dependant on the dose, route of administration (cend vs. rijectable), poe of steorid (coronatizable) can accomitable), and the off resistance to height restableoist. Samonioli has a strong effect on the hepatic immagnent of cholestered due to is structural resistance to develope the control of the control

Injectable stanoolol line also been documented to produce strong negative changes in serum pilisk. One study securined or no a goop of 12 leashly must subjects, and demonstrated a measurable reduction in HDL cholsterol values, as well as an increase in LD, and total cholsterol values, Studying a single justice of 50 mg <sup>107</sup> lines changes presisted for 4 developing arteriosclerosis. Injectable stanoolol should not be used as an alternative medication when cardiovoscular risk factors proclude and stanoolol use.

To help reduce cardiovascular stania it is advised to minitain an active cardiovascular cerevise program and minimize the intake of saturated fits, cholesterol, and simple carbohydrates at all times during active AAS administration. Supplementing with fish ols (4 grams per day) and a natural cholesterolaritoxidant formula such as Lipid Stabil or a moduct with comrangable inerdenies is also recommended.

#### Side Effects (Testosterone Suppression):

All anabolic androgarie steroids when taken in doess sufficient to promote muscle gain are expected to suppress endogarous testosteron production. Stancolol is no exception and is noted for its strong influence on the hyporhamin-pinitary-testicalar axis. Clinical statics spiring 10m gpe rdq to healthy made subjects for 14 days caused the mean plasm testosterone level to fall by 55%, <sup>CCS</sup>. Without the intervention of testosterons-stimalizing solutions are stronger to the contract of the co

Note that prolonged hypogonadotrophic hypogonadism can develop secondary to steroid abuse, necessitating medical intervention.

The above side effects are not inclusive. For more detailed discussion of potential side effects, see the Steroid Side Effects section of this book.

#### Administration (General):

Studies have shown that taking an oral anabolic steroid with fixed may decrease its biavasiability. <sup>Not</sup> This is caused by the far-soluble nature of steroid hormones, which can allow some of the drug to dissolve with urdigested delary far reducing is absorption from the gastroitestrial tract. For maximum utilization, oral forms of stanoarold should be taken on an entry storant.

There can be large discrepancies in the steroid particle size between rijectable stansolal representation. For currently, Westerol flown, Turnbero (Spain) was designed for harms use, and uses a refined provider that will pass through a 27-gauge needle. Westerolle-V is a vectoring product in the U.S. and Carnada, and has large particles due mig mis needles smiller than 22-gauge. Solutions that raffice a larger particle size may also cause more desconfied at the contract of t

### Administration (Men):

The original prescribing guidelines for Winstrol called for a daily dosage of 6 mg, which was administed on a schedule of one 2 mg thick three times per day. The usual dosage of physiques or performance-enhancing purposes is between 15 mg and 25 mg per day, or three to fine 5 mg tablest, incline for to longer fittin 6.8 works. His prescribed Winstrol is generally recommended at a clinical dosage of one 50 mg injection every 2-3 weeks. When used for physiques or enfratume-enhancing persons, as dosage of 50 mg every other day. The person of 50 mg per day of the person of the person of 50 mg every other day. Doses of 50 mg per day with injectible Stormed all are commenty given at 75 mg every their day. Doses of 50 mg per day with injectible Stormed all are collisions, although probably not advised. Note that rigorable from 6 mf de dug are expected to have, miligram for miligram, a guester anabole feet for the oracle.

Samouhi is often combined with other steroids for a more dummite result. For example, while building one might opt to add in 2004-001 org of a testisotione eater (copionic, enurarbate, or proposately per work. The result should be a considerable gain in now marched the contraction of the contracti

### Administration (Women):

The original prescribing guidelines for Winstool called for a daily dosage of 4 mg (one 2mg thatlet vivice daily but young women particularly succeptible to the anthogasic effects of anthois storoids. This dosage was increased to forig (the same as the recommoded dose for miles) when messersy. When used for possquere operformmers—enhancing purposes, a dosage of 5 mg to 10 mg daily is most common, taken for no longer fram 4-6 weeds. Injectible Winstool agreenily percommedated at a client dose of 5 mg new 2/3 weeds, prescribed to the control of the common for the control of the control

#### Availability:

Stanozolol remains widely available as a pharmaceutical product. Its production has been

shifting to less regulated markets (minly in Asin) in recent years, however, which Bedyreflects declining interest in using stamozoll as a medicinal product in the West, and the continuing high demard for this drug among athletes and bodybuilders. In reviewing some of the more popular products and changes on the global pharmaceutical market, we have made the following observations.

British Dispensary in Thailand makes a stanozolol product. Their trade name for the drug is Azolol, and it contains 5 mg of steroid in a 400 tablet bottle. The bottle itself books very similar to that of Androlic, with dark plastic and a shiry chrome top. Be sure to look for the company's holographic stricker when shopping.

Stanol (Thailand) is now sold as a generic drug under the March Pharmaceuticals label, though is in the same recognizable white bottle with green-sladed label. The product contains 5 mg of steroid per table, and each bottles holds 200 slables. To deter counterfeiring, the bottle now carries a holographic sticker on the front that bears the commany lozo.

Acilhon in Thailand makes Stanozodon, which comes in the old industry standard of 2 mg of steroid per tablet. It is packaged in bottles of 1,000. Counterfeits are not known to be a robblem at this time although the product is not widely distributed on the black market either.

Unigen markets the product Stanztab in Thailand. It contains 10 mg per tablet, and is packaged in foil and plastic strips of 10 tablets each (5 strips per box).

Winstrol® tablets and injectable ampules are still produced in Spain, by Desma. This remains the most popular stanzoolol injectable in Europe. All howes are protected with a holographic sticker, which carries the company logo embedded into the image. Note that highly accurate counterfiels are currently in circulation through Europe and North America.

The Greek generic by Genepharm is still in production. The product should carry a Greek drug ID sticker on the box, which will show a hidden mark under UV light.

Chinfield makes a 50 mg/nt. rijectable stanzonkol in Angentine called Nabolic Strong. This is the same firm that makes register Nabolic, a very low dosed (2 mg/ntl.) version of the same dang. This new product is now much more popular on the black market than the first, due to the more useable dosage. Note that Chinfield prints their logs on the inside of the vial carton, which offers ownewlat of a simple security check (obviously one very every to daplicate).

Anabolico Cimol is an injectable form of stanozolol from Argentina. It comes in multi-dose vials containing 50 migril. of the steroid. This item has not been subject to widescale counterfeiting, but also bears no security features that would deter this practice.

Stanozoland from Landerlan in Paraguay is common on the black market, particularly in South America. It comes in the form of a 10 mg tablets, packaged in bottles of 100 tablets each. The company also makes a 50 mg/ml. injectable.

Also from Paraguay is a generic stanozolol injectable from Indufar. It contains 50 mg/mL of steroid in a 1 mL glass ampule. Three ampules are packaged per box.

Formula Magistral in Argentina makes generic oral and injectable Estanozolol. The oral comes in the form of 10 mg tablets, loose in pill bottles, while the injectable contains 50 mg/mL in multi-dose vials. These products bear a small square holographic security sticker to deter counterfeitine.

The veterinary compounding pharmacy SMP in Canada makes a generic injectable in 30 mL multi-dose vials (50 mg/mL). This product is made in limited quantities, however, and is not highly common on the black market.

# ANABOLIC AGENTS (NON-STEROID)

# Description:

Anchidoria cali is an omaga-fe oscentifi fitty said that seven as the principle bading block for the synthesis of dendee prostagationis such as PGE2 and PGE2.) These prostagations are integral to protein turnover and muscle accumulation, and have such important actions is successigned both flow the muscle (purple), received post Her-I and mains sensitivity (corresponding receptor beeds), apporting senditic cell actionize, municipal control of the processing of the Her-I and mains sensitivity (corresponding receptor beeds), apporting senditic cell actionize, municipal control of prostagationization turnover in solicular muscle tissue, and is responsible for initiating muny of the immedia turnover in solicular muscle tissue, and is responsible for initiating muny of the immedia turnover in solicular muscle tissue, and is responsible for initiating muny of the immedia turnover in solicular muscle tissue, and is responsible for initiating muny of the immedia produced and the procession of th

# Clinical Studies:

In 2005, the Exercise & Sport Nutrition Laboratory at Buylor University conducted a double-bird placeboc-controlled ashy to determine \$'0.00 stays or instance training and anachidorie, acid (X-Faster 199) appelmentation would affect training adaptations in 31 experienced [-1] upon juestimate-trained miles. The results were presented at the International Society of Sports Nutrition conference on Jues 15, 2006. All subjects inguisted as that of fast or grassites each sidy (see 22 still or grassite of Art or placebo every four branch, Subjects taking X-Faster added an average of 25 his to the hearth press mentions weight in 20 days, which was an artenue of caraby (\*\*) genater than the placebo every four branch of days, which was an artenue of caraby (\*\*) genater than the placebo goay. The X-Stay days, which was not artenue of caraby (\*\*) genater than the placebo goay. The X-Stay days, which was not across of caraby for general training and the second of the placebo (\*\*). The placebo (\*\*) was a feet to be a second of the placebo (\*\*) was a feet to be a second of the placebo (\*\*). No side effects were reported during the investigation.

# Pharmacology:

Anchelorie neid begins to disploy in anabolic activity early during exercise. This instruct is released from your meels beers and type and uniqued during structure training tiggering a bacided riffinamenty and anabolic response. This is purt of the same biological process that causes you to be seen a day or two following a good workoot, and merridus that the old salage "ho pain, to gain" is a fundamental two exchandations acid illustration from dampad massed flews, is simbly, the very list annoble 'tagger in long causcular that will control the rebuiling and strengthening of much tissue after excess. See '50'-100' Armang other tings, by its resemble, to all the similar through excess processing boal (Erd.' and most increase) excess processing boal (Erd.' and most increase) excessing a seal flexible and most increase in the control of the control of

The availability of ameliateine acid, and our ability to Bernate it during exercise, is important to the amabolic productivity of our workstow. We also need to be some of the first that regular exercise significantly lowers the content of annichatoria acid in sketchal much tension. Of the annichatoria acid in sketchal much tension of the annichatoria acid, in the annichatoria acid, breat lovel tension shows the annichatoria acid, breat lovel tension is less streamed and a less interne ameliatoric response. The depotent of amendatoria card in sketchal much tension is also once of the key reasons we find a fundament of the stream exist of schedul much tension is also once of the key reasons we find a fundament of the same based of schedul annich and the stream exist of schedul read of the same based of schedul and the same based of schedul and the stream the stream of the stream of the stream that the schedul and the stream of the stream of the streament of the s

#### History:

The arachidonic acid supplementation protocols, and the concept of using this nutrient to improve muscle mass, strength, and performance, were first developed by William Llewellyn,

auther of this book series. Lisselbp flied patter on the technology on November 27, 2002, and released an amthebrine acid supplienter under the X-Factor trademark (Molecular Natifoxia) shortly affect. Although the product was intailly net with a great deal of skeptision and a clinical trails, and a catalonic leaf as a proceed applienter for body recomposition goals. The U.S. Patter & Tindemark Office grarted Unselpsis patter application for anticiducia exist on January 11, 2005 (U.S. Patter 6 68,141753), and the products in severe been officed for lecence to other companies in the industry. The original X-Factor product remains world, available in the U.S. and about, and applied expanses in the architection and continuously and applied to the contrast of the contrast world available in the U.S. and about, and applied expanses in the architection and the contrast of the contrast of

# Structural Characteristics:

Anchidoric acid (eiccos-58,11,14-enoic acid) is an essential polyursaturated fitty acid found in animal fits. Supplemental anachidoric acid is commonly produced in two forms, trigheeride and ethyl ester. As with other fitty acid supplements such as fish oils, the natural trigheeride form (as present in X-Factor and focused products) offers up to 400% greater absorption fluent the othyl setzer, and is the preferred form for supplementation.

### How Supplied:

Arachidonic acid is sold under the X-Factor trademark by Molecular Nutrition, and is supplied in 250 mg capsailes. Arachidonic acid may also be found in a number of licensed products; all will display the patent number (#6,841,573) on the packaging.

# Administration (Short-Term Anabolic):

As a short-term ambolic agest, anchiskoire acid is supplemented at a dose of 500 mg to 1,000 mg per duy (-2.4.2.5 mg equolsay). Be fall (5,000 mg dose is not commonly one, regardless of bodyweight. The naturat is cycled in the same way steroids commonly are, and is taken for a peaced of 7-8 works followed by an equal amount of rime of The less of a benefit as peaced of 7-8 works of those by an equal amount of rime of The less of the peaced of the control of the contr

#### Administration (Normal Supplementation):

Anachisotic self may also be an important ration to consider in regalar repolarement does, particularly you do not consone similar products (ord nature capitars etc.) and regalar basis. Staties have shown that given somewhat companishe amounts of protein, those who comman similar products will make more proguess with resistance centres them those that do not (regarterine). <sup>504</sup> Anachisotic need may be the missing component in such dies, too integral to be madels response for tower destroy between the to be noticed. These is experienced bodylotaliers, given that training depletes AA stores. On a nurber of cases, time test for the content of prodephylotis have revocaled trainately how beek of markelstone can't in highly trained utilities. For those who that their takes of animal products independent can't in the provides of the composition of the composition of the composition of the composition of the provides about the expensation of the composition of the compositio

### Safety:

In clinical studies involving the supplementation of 1,500-1,700 mg of arachidoric acid per day, general markers of health were also unaffected with 50 days of continuous use. This includes no notable change in HDL, LDL, or total cholesterol values, immune system recovers functioning or plateful appropriate pages 5636637 Europeanous for the insertioning as a lawler from the process of the part o

On a more general rote, the American Heart Association amounced its position on Orangafo FAX consurption in early 2010. In revisioning some of the more courted sta, they found that dies low in Oranga-fo fits were associated with an increased risk of heart disease compared to dies with higher induces. The paper also discusses are instablica exist and bout this essential it is normally not proinflurmatory, nor outstudy surbailty to consume. The American Heart Association now recommends that 510% of your careless each day come from Oranga-fo fits, including ARA, Dies rich in Oranga-3's are, of course, also healths; but not at the exclusion Oranga-for Six.

### Side Effects:

Anchidoric scid (X-Factor) often produces an amplification of residual post-workout (Ochyed) Orner) mucck sourcess. Often recovery is slightly prologingl (perhaps an additional day of reat is required), and the user may need to adjust their scheckle and represent inventor of the number of consection from the control of the control of

#### Contraindications:

Those with an existing medical condition related to inflammation may find that the added arachidonic acid exacerbates symptoms of their disorder, and should avoid supplementation. This supplement should only be used after the approval of a physician if someone is taking medication has an existing medical condition or has a familial predisposition for cardiovascular disease, high blood pressure, or any other disorder that may require the limiting of dietary arachidonic acid. Also, as a potent growth-promoting agent, arachidonic acid joins androgens (testosterone, anabolic steroids), growth hormone, IGF-1, estrogers, and many other growth factors as potentially supporting the growth rate of certain cancer cells if you have the disease. Dietary arachidonic acid intake has been generally eliminated as a causative factor in cancer. 638 639 just as testosterone level has been eliminated as predictive of prostate cancer risk, however these types of growth-promoting agents should be avoided in such diseased states unless approved by a physician. If you have prostate cancer, for example, the last thing you want to start taking is a growth promoter like testosterone. The same goes for arachidonic acid. The bottom line is that if you are in poor health, you should probably not be taking this supplement. If you are healthy, you should be able to use it with great safety.





During the clinical study, subjects taking X-Factor added an average of 25lbs to their bench press maximum weight in 50 days. This increase was nearly 45% greater than that noted in the exercise-only (placebo) group. Some subjects gained more than 50 lbs on their bench press 1-rep max over the 50-day period.



Using the standard Wingate cycle ergometer test to measure relative peak anaerobic power, subjects taking X-Factor increased leg power by 1.2 Watts-kg-1.This represents a net increase of more than 600% over the placebo group (-2 Watts-kg-1).



In the standard Wingate cycle ergometer test to measure anaerobic power and performance, the X-Factor group outperformed the placebo group on Average Power by an amzing 21 wats (AA: 37.9W P: 17.0W). That's a net increase of nearly 225% commard to blacebo.



Total Work, as recorded in the standard energy unit Joules, increased by 1,292J in the group taking X-Factor, while Total Work increased 510J in the placebo group. This is more than a 250% increase compared to placebo.

#### Kynoselen®

#### Description:

Kynochen is m igicutable veteriuny drug, currently produced by the intermitonal firm Vetoquial It contains a mitten of hypatrinal AMP (dischouse incorpolopately), vitami B-12, sofatim-schritte, impression aspurate, and potassion aspurate. This blend mikes for a restorative Yacif Pept ofing, admirated top broatest an arrialist masche mass and overall webness after filtons, spiry, or traum. It is most often used on hosces, and is typically applied as an anti-carboic after streames actively, or to hep get an antimable code in set after a debilisting infection/filton. At other time it is surply used to support the visibly of an source cases it is even to all of the vety holes propose of memology and clothers; or hosting li-12 or solvanis ratiose. Bodybuilders are attracted to Kynoschen for its mild anabolic and lopolite properties.

The principle active ingredier in Kynoschen is beparinsel, which is classified as an aurio acknowled with myconfluid stringtur and vosability properties. It is also indertified as no incorpor, correspond, which increases contractile strength, and mariness fatigue, or dischest immedic. See It has desertified as seed subject to receive the differentiation of satellite market code, see a seed of the se

# History:

Hepatimist, the principle active ingredient in Kynooleus, was first heavily investigated in circuial medicine direit, the early 19/50. It was sond evoloped in on perception drug and has since been sold by a series of drug manufactures in mmy different parts of the world. Currently is most common flavarparia to set is to trat off-tostict hopportunes, which is sudden drug in blood pressure upon standing. Marious preparations containing haptermiol have been produced over the years, the most models of which have related Amsteria (table), Corresor (Edgiann and Swatzman), Hepse-Moyl (CSA), Hepsen) (Belgiann and Swatzman), Hepse-Moyl (CSA), Hepsen) (Belgiann and Swatzman), Hepse-Moyl (CSA), Hepsen) (Belgiann and Swatzman), Peper-Moyl (CSA), Hepsen) (Belgiann and Swatz

The French vaccinary preparation (Kyroschen would be the first haptumino-Lorentining drug to goth allay-acids information allowings, becoming popular arrange American holybidises and additices during the latter part of the 1990s. This was some years subsequent to laws being passed that that foreseast the penalties for dealing in analoties steroich. During dies time, availability of the drugs land shifted, and for some buspess searce supply and halp high time, and the contract of the simulation of the contraction of the simulation of the contraction of the co

#### now Supplied:

Kynoselen is most commonly supplied in a 100 mL multi-dose vial for injection. Active ingredients are heptaminol, disodic adenosine monophosphate, vitamin B12, selenium (sodium selenite), mugnesium aspartate, and potassium aspartate.

#### Structural Characteristics:

Heptaminol (supplied as heptaminol hydrochloride) is an amino alcohol with a structure of 6amino-2-methylheptan-2-ol.

# Administration:

Kynoden is not approved for use in harms. Prescribing galdelines are unvoiable. An effective dosage for physique or performance-enturing purpose generally fills in the range of 1 ml. worldy for every 2 pounds of bodywight. This would must that a 200h the property of the property of the property of the property of the very body as 2 ml. fill returns proved. At fill shot one, single 100 ml. vial would last about 16 worlds. At 8-10 ml. per word, a 100 ml. both would last from 100 st. proved, a 100 ml. both would last from 100 st. proved, a 100 ml. both word last from 100 st. proved, a 100 ml. both word last from 100 st. proved, a 100 ml. both word last from 100 st. proved, a 100 ml. both word last from 100 ml. both word las

Because it tends to necrose consideration levels, Kyroschen is also a mil stimular I is facly for this specific reasonal that see has been bounded yourtain horsening cognizations. This means that one cam expect certain stimular-related side effects, especially when taking fisit during in light obsesses. This metakes ruple tambent, sowering, pitters, residenates, reasonal blood pressure, or incomin. A good rate of thursh used by bodybuilders to my and keep scatt side effects from theoreming a problem in one religious remain 2" rate, per day. They scatt side effects from theoreming a problem in one see rigitar crust from 2" rate, per day. They scatt side effects from theoreming a problem in one religious remains of "in the problem in the problems of the scatter of the problems of the scatter of the problems of the problems of the problems of the bodyweight, perhaps seen late of the The does is then showly becomed, when the problems of the scatter of such for the two feets were sold side in recursation in consistent of the scatter of the problems of sold were rememble incoming or sold and the problems of the rememble control of sold were rememble incoming or problems.

# Warnings:

Individuals with high blood pressure or cardiovascular disease should not use Kynoselen.

# Availability:

Kynoselen usually selfs for \$75' to \$100 per bottle at the retail level. It is not a controlled substance in the United States, and is Bewise pretty easy to obtain locally or to mill order. Currenty no significant files are known to exist. Given its abundance and low cost, counterfels are not expected to be a significant problem anytime soon. It is also important to note that legitimate Kynoselen is a veterinary drug orig; and has never been munificatured for human use.

# Lutalyse® (diniprost tromethamine)

# Description:

Discovers townsharine is a pharm-conteal form of the natural possibalistic PGZ-lights. Procligadinistic are active of natural conjugated unstanted cycle flags, PGZ-lights is noticed in successful conjugated co

clicial statics, which lave shown NFT2 to be a strong strainfor of protein systems, and key to both the immediate and lange-employshogical adaptations to resistance training.<sup>50</sup> 464 565 606. Regorts from athless who have experimented with this agent generally apport the compound being an excellent promoter of backed markes lyonely, usually evaluating to both recrosses in muscle size and definition. Disreptors is also resported to be a very fist acting dainy, with may clearly the language of the property of the property of the property of substantially poster flows dainy, with PCP2 abstorm is saudice to within the simulation of progression in code.<sup>507</sup> Again there is a good dail of anotherial supports for this property of significant straining and the property of the pro

Athletes and bodybuilders are attracted to dinoprost tromethamine for its strong thermogenic and miscle-building properties. The anabolic effect of this drug has been substantiated by

# History:

Disposts tomerdumine was fest introduced into clinical medicine in the early 1970s. The first approved use of the ding intumm pointers was to stimitate abstron during the second trinstate. It has since remined of use for this purpose, but is most commonly associated threates. It has since remined of use for this purpose, but is most commonly associated the estones cycle and fertility of various feestbod, lettered in disprost tomerdumine as an analothothermagenic day for shalless and bordheades dark of any post until the 1970s. The common letter of the relates of memories metallical lates fielding POTATA to common letters and the properties of the produced of the condumined produced to metallical the day which is tayled of high proposably to generate side effects, have proven to be highly successful for mmy athletes and bodybuildness.

Over the years disrposet tomediatrine has appeared as a harma medicine under a wide marbor of trade names, richtig such populer due genoties as Ampgiantif, Swedoul, Prostin F.2 alpha (U.S., Austrafia, Isanel, Italy, New Zealard, South Afice, and the Utales (Kagloni, Migrostin F.2 alpha (U.S., Austrafia, Isanel, Italy, New Zealard, South Afice, and the Utales (Kagloni, Migrostin F.2 a for longer sold in the U.S., however, and there is prosently as FTA approach epithecurst arounds for larman use Veterinay versions are more widely available and tend to provide significantly mere active dung for loss money than associated with the physicae or porformance-orderacity use of disposet to reorderance. Popular veterinary branch have included Lindbjoe (Pharmaca Arimat Health), Prostamate (Princer) Pharmacolin (Indies) Pharmacolina Color and Bradie (Vigiles). Several corporate mergers have taken place in this segerant of the market, and the (now large) congluences to reorderancy the most common form of disposts to terrethrained to see among the athletic-body-building community.

#### How Supplied:

Dinoprost tromethamine is most commonly supplied in a multi-dose vial (5 mL-100 mL) in a dose of 5 mg per ml. It is prepared in a sterile solution of water with benzyl alcohol added as a nesservative and sodium/hydroxide and/or hydrochoric acid to adjust pH.

#### Structural Characteristics:

Dinoprost tromethamine is the tromethamine salt of prosta,5,13-dien-1-oic acid (PGF2alpha).

# Side Effects:

Possible side effects or signs of disrposet tromchamine overceposure may includ such receptancy effects as benechconstrictive, wheeving coaging, large intration, major benedings and amplysics. Administ individuals may be partially succeptible to these effects. Disposts may also cause genotive-final disturbances such as schornist carming below the schornist carming in disturbances such as schornist carming in disturbances and in women terior contractives, suspiral bleeding, and storic or urisary infections. Pregnant women doubt not lake of home distorted below of of the effect women of datas using infections of the schorn distorted for the schorn of the schorn distorted in the schorn of th

unapano. respons or san curea, autone autone unage unapano na parapase va performance-relativa persones are comme, and often enterne. This includes promoused socress at the site of rejection (edne beginning with a dal braining almost immediately after the slot is given. That and the like belong a seal commonly reported dring; cycles, as need toost of alternates of breath lipicions are also commonly reported dring; cycles, as need toost of alternates of breath. In the control of fixed interfaces of many face carrying, including size presented contractions of the musels insolod in the control of fixed fixed sizes and vonting have also been commonly reported. For many, the carrying, information, general deleging of optest streamly, making, and discontified make disrupout at ding they experient affecting of the streamly, making, and discontified make disrupout at dings and often report that side of the become more telenthic over-

#### Administration:

As a human medication, dinornost tromethanine is most commonly given intra-amniotically at a dose of 40 mo for the termination of pregnancy. It is also sometimes given orally to pregnant women at a dose of 30-100 mg to induce labor, although this tends to produce more side effects than other more recently adopted, medications. When used for physiqueor performance-enhancing purposes, dinoprost tromethamine is generally given by intramuscular injection. Most noted for its ability to generate localized growth, common sites of injection include the shoulders, bicens, tricens, calves, chest, back, and less. The user will typically inject in only one site per day at the start of therapy, but this may be increased to 2 or more injections per day as they become more accustomed to the drug and its side effects. Therapy begins slowly and is initiated with a low starting dose of approximately. 5 millionam per injection. If the first injection were given without significant side effects, the next injection would be increased to 1 millioram. This is slowly increased by .5-1 mg per application until a peak dose is reached. This might be a maximum of 5 mg per injection site. Injection sites are also regularly rotated so that several days separate administration in the same muscle group. Note that for some, the pain after injection is so severe that training for that specific muscle group must be delayed for at least a few days. Individual sensitivity to the drug may, therefore, require modifications of their injection and training schedule to maximize results and comfort

# Availability:

Dinoprost tromethamine is available in the U.S. and many other nations as a prescription drug product. It is also found infrequently on the black market.

# Anti-Estrogens

#### Arimidex® (anastrozole)

#### Description:

Austronois is an artis-estrogenic drug developed for the treatment of advanced breast cancer in worms. Specifically, this agent is the first in a rower close of third-generation selective coral aromatuse inhibitors. <sup>566</sup> It acts by blocking the enzyme aromatuse, subsequently blocking the production of estrogen in the look, Streen may forms of breast cancer cells are stimulated by estrogen, reducing levels of this hormone in the body may rester the prospession of the disease. The is also the fandamental to or formstein charge (Nobadeello, escept Nobadeello blocks the action of estrogen at the receptor, rat is actual englospous production. The effects of instruction clause be very advantaging with an object of the effects of the effects of the effects of the restrict in the effects of instruction clause be very advantaging with a standard properties of the progression of the effect of the first first for the effect of the effec

# History:

Austrono's was developed by Zencen Pharmecetricals, and approved for use in the United States at the end of 19%. The day was developed as a new adjust tentument for operable breast carrier in postmeropassal fermic protects, an area of medicine that had a long bistory of narrodnine. Substantial data was needed to shift prescriber protest soonsylforms, and not not of an externed by large malarizer of substantial data was needed to shape for soon of Rome (ADC). The study evaluated the use of mustroods and naturoside, roles or to correlation, as 92-50 postmeropassal women following breast carrier surgery. The results forced ansatroods over turnoisties at promoting disease regression and improving event survival in a protection of the trial in 2002, ansatroods emerged as a new contender for the adjustices tenturent of postmeropassal bested carrier, "2009" Alexand this seam in the day was also garded popularity with multi-bodyleakers and affects who began taking notice of the strong clinical traits.

# How Supplied:

Anastrozole is most commonly supplied in tablets of 1mg.

# Structural Characteristics:

Anastrozole is classified as a selective non-steroidal aromatase inhibitor. It has the chemical designation 1,3- benzenediacetonitrile,a,a,a',a'-tetramethyl-5-(1H-1,2,4- triazol-1-ylmethyl).

# Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes ioint pain, weakness, fatigue, mood changes, depression, high blood pressure, swelling of the arms/less and headache Aromatase inhibitors may also decrease hone mineral density which may lead to osteoporosis and an increase in fractures in susceptible patients. Some individuals may also respond to the medication with eastrointestinal side effects including nausea and vomiting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an aromatizable steroid such as testosterone enanthate is taken in conjunction with an aromatase inhibitor suppression of HDL (good) cholesterol levels become significantly more pronounced. Since the estrogen receptor agonist/antagonist Nolvadex® generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovascular health.

#### Administration:

Anatonois li FIA approved for adjunctio treatment of postmorpussal vocum with hormore receptor-positive early breast cancer, field-free internity of postmorpussal women with hormore receptor-positive or receptor unknown locally advanced metastatic breast cancer, and treatment of advanced breast cancer in postmorpussal women with disease prognosions following turnordin therapy. The doseage prescribed in all instances is effects of anabodis-independent services and the services of the services and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique, which makes this agent of interest for defined appearance to the musics and physique. it is of note that wood does not appear to affect the absorption of anisotropic, so the unity may be taken with or between meak.

Anastrozole is widely available in the U.S. and many other nations as a prescription drug product. It is also found readily on the black market.

Aromasin® (exemestane)

# Description:

Availability:

Emensture is a stroidal saicide amountase inhibitor. It is very similar in structure and action to formestane, althory it is significantly more potent in companios. As a class of rings, amountase inhibitors offer an articutogenic effect by blocking the enzyme responsible for symbolic great engines. Emensture is approach of the breathers of breast cancer in sworms, specifically in pote-meropassal patients whose cancer has progessed following femerpy with memorifs. Nakle bodylates and athlisted onless the dauge for meropaporous parsons, namely in counter the entogenic side effects associated with the use of word returning to more instances aroundars with the use of word returning to more instances aroundars withinkness may also used in flags of the loss of body fit and increases in musclar definition. Exercisation is one of the most potent amountase inhibitors presently available. The most commonly clock data (sind at the Amounias packaging ment) reports a lovering of serum-entogen levels by 8% on average in clinical satisfies with women.

# History:

Enemestare was developed by Plantmeia & Liplond (Pfantmeia), which guised FDA approval for size of the days in late 1999. They introduced it under the Aromais brand mare in early 2000. Although the daug proved to be effective in doses as low as 2.5 mg per duple, in some patients, the corrupps developed in as standard and near uniescally deficite dosage of 2.5 mg per tablet. The company has size introduced the days to many other micross surder the same table name. Due to various patient and genal market dominizes. Aromais is the only brand name of exementation on its fleely to come across in general corruption and the contraction of the contraction of

### How Supplied:

Exemestane is most commonly supplied in tablets of 25 mg.

#### Structural Characteristics:

Exemestane is classified as an irreversible steroidal aromatase inhibitor. It has the chemical designation 6-methyl-enandrosta-1.4-diene-3.17-dione.

#### Side Effects:

Commo side effects associated with the use of an anomatone inhibitor include bot flushes, post prain; weakness, fings, remod clamps, depressed in philosophe ground, sould be arriving, and beachet. Aromatone inhibitors may also discrease bota mireai density, institution of the control of the institution was also control to the medication with gastroinestic side offices including muses and vooring, Aromatone inhibitors can humifue development of an urborn flush, and should move the leaf not on hundle during programs. When taken by mre fits on off-lieful use) to reduce entity-question desires (VD) in its by returning some baseful should have be taken earthough and produce (VD) in its by returning some baseful shibbots may accommod excellent to the control of the control of the control of shibbots may accommod the control of the control of shibbots may accommod the control of the control of shibbots may accommod the control of the control of shibbots may accommod the shibbots may ac properties of estrogen on cross-serior vasces. Statuses ravie demonstration test when an amountable steering starts net settlement entirelite is based in conjunction with an amountable stiribitor, suppression of IRIA (good), cholsteed levels become significantly more promuneed. Since the estrogen receptor agenticalizations studied by separation of the same artis-estrogenic (neightie) effect on cholsteered value, it is usually forceed over amountance lithium for estrogenic minimum col position of the objection of the object of the confidence of the object o

# Administration:

Execution is FDA approved for adjustate treatment of postmorpousal vocum sufficiency recognition (each yeast carrow with disease proposition following turnisod. Therapy is rainted 2-3 years after turnordin has field to eleit a desirable response, at which post turnordin is discontinued. Transmer with ecrementars is contrained for 2-3 additional years, and is completed after 5 years of commission adjustate design freely market produced to the proposition of the prop

### Availability:

Exemestane is available in the U.S and in more than three dozen other nations under the Aromasin brand name (Pharmacia). Aromasin, Ricovise, dominates the global market, and is presently the only exemestane product one is Ricovit to encounter.

#### Clomid® (clomiphene citrate)

# Description:

Chemiphene critate is an anti-estrogenic ding that is proscribed to women to treat amounthory infertility industry), to ordinal reductive its speciality-ferrent of as a treasteroidal ovalatory strankart. The drug works by intensiting with estrogen receptors, edited in an antignative insurer, in various touses of the body acting the hypothinature, or the contractive of the contracting the hypothinature, or the contractive intensities of the contractive dependence of granularous feet. If and a FISH, This surge in granularous permits uncurrently an expertise compared to the contractive during the relative compared with the contractive during the compared with the contractive during with the contractive during the contractive durin

In men, chemipleuse citate also acts as a partial arti-estergat, and may be used to contressore of the sale effects of amornitudes beared use inching generous and increased under relation. As an arti-estergatic ding, chemipleuse citate may also produce an electrion of folias installing lemonson, and labering lemonson becoles whether an electron control of the control of folias installing lemonson, and labering lemonson becoles, which can electron control of the contr

Note that the triphenylethylene compounds (toremifene citrate, tamoxifen citrate, clomiphene citrate) tend to be somewhat intrinsically estrogenic in the liver. This means that while they

can block estinguis activity is now areas of the body, they can actually act as entoques in this other low sens. Biscopies action in the low is inportant in the negliation of sense cholstend (it tends to support BID, synthesis and LID reductions). Sero sensel-using to hobbidables are already desting with the negliot candisocated effects of these entrycompounding the issue with anomatics ribbidous (which will lower tool serum estopen tends) may not abuse be the best option. Using a might table kells practically and the sensel in the same time supporting improved cholstend values, might be much more isless.

# History:

Cheighene chrate is a fertily drug with a substantial listory of use in the United States. It first agained widespread acceptance during the early 1970, and the Steen aday community to the feetily protective over since. The drug is now considered a standard medication for certain forms of fertily theregap, and has been adopted as such from oxide U.S. border. Cheryidene clante is presently available in most rations worthvide. The two most popular bond natures on is likely to encourage and has been adopted as such for under U.S. border. Cheryidene cannot us the worth of the contract and the contr

# How Supplied:

Clomiphene citrate is most commonly supplied in tablets of 50 mg,

# Structural Characteristics:

Clomiphene citrate is classified as a selective estrogen receptor modulator, with both agonist and antagonist properties. It has the chemical designation 2-[4-(2-chloro1,2-diphenylvinyl) nhenoxyl trist/hamine directores cirate.

#### Warnings (Visual Symptoms):

Some patients using cloniphane crimte notice bharing or other visual distultances such as spots or fulses. These symptoms occur mere floquently a higher doese or heapy drainfest of therapy, and often disappear within a few days or weeks of use. Probaged visual distultances have been respected after the discontraints of colorappine crimte therapy, however, and in some cases may be inversely. Those taking chriphene crimte should be wounded that these symptoms might make auditions like during the area of these visual symptoms in net remarkable to the contraints of the court of the contraints of the five court.

#### Side Effects:

Compleme centrae appears to be well tolerated, with a low incidence of significant side effects. Common advisers exactions during circlical Tunbs included oursin configuration (13.6%), succession flashes (10.4%), advisoritud disconfort (5.5%), antasectionizing (2.2%), beaust disconfort (2.7%), and subcommunication belonizing (1.3%), and advanced (1.2%), and advanced to the configuration of the configurati

### Administration:

Chriphene cirate is FDA approved for the reatment of women with ovalatory dysfurction preventing pregnany. The recommended dosage is 50 mg daily for 5 days, which is initiated approximately 5 days into the menstraal cycle. If ovalation does not occur, follow up cycles my use a dosage of 100 mg per day for 5 days. Many cincinium recommend a limit of 6 courses of therapy. When used by men (off-label) to mitigate the estrogenic side effects of antiboclarifloguistic steroid use, a daily dosage of 50 -100 mg (1/2 labels) is usually administered white any offurfung seconds are taken. Note, however, that turnoufies is usually given preference over chariphene criate for this purpose. More commonly, charapterice criate is used by ment at adouge of 50-100 mg per day for 30 days at the corclasion of a stream of the contract of the con

# Availability:

Cloniphene citrate is widely available on the international market in a variety of brand names. It senerally sells for a reasonable price, and is of low interest to counterfeiters.

# Cytadren® (aminoglutethimide)

# Description:

Amingalachimide is muinly identified as an shibitor of advanceordical seroid synthesis. Its priming function is to block the convension of colosistend to presponders, which is required for the biosynthesis of adveral glacocorticoids, mineralectricoids, estregars, and antilogues. Amingalachimide is a mempedic inhibitor, and also blocks seen forth est spin in steador and the colosistent of the stage in steador and the colosistent of the colosistent

Cortical lithibition. While cortical is an essential hormone for fit, fit below may also vary greatly within "normal" ranges depending on the rishfished their training and dietary status, and many other personal metable factors. It has been a common pussai the sport commanty to first ways to cortrol (first) cortical production. This is because while analogous give your mucke cold as message to herease profits synthesis, cortical (in casholc hormone) impacts a message to breakdown and release armis acids. If one can first the analogous give your sunderes should, in thoogy, the removal that the carbon should be a sundered to the control of the c

When fist looked at in the realmot furtherics, however, research was bese as to the best way to use amining darking as a cortiol lowering anti-tunble. Debtod the "shedme elegang phenomenan", if has been noted that after a short period of regalar use you obly often meants in lowest of cortical levels by screening the release of amining the horizone. ACTI amining the release of amining the horizone and the same and the

It is important to note that while many people believe they have used this drug as an anticatabolic, few have actually taken the correct dosage. Four tablets per day, or 1,000 m and amount processor to circulate the deliving or processor to circulate the demands or processor to circulate the demands or processor. appears accessed at againments again and accessed to the superior of the superior of the contenting challesters to be presented as a few factors of the superior of the target when reduced orthor is described. Those who do venture this light comments preport finings and disconflict, stating that the days is indicable for any proof profused guest fortions, must grides and bodysladies as indicable for any proof profused guest fortion, may arise and bodysladies and the superior of the superior

Aomates libibion. Antiophethrinke is an efficient aromates inhibion, and tends to situate the activity of this engine at an uniform lower closure from them what is required for sibibition of corticosteroid production. On the lower closure from them what is required for sibibition of corticosteroid production of anomates and extensor levels to pixelly achieved at a dosage between 250 mg and 250 mg, a point where storag adversal steroid blockage is not noted. There also seems to be no aduled benefit by adding cortion in nears of aravitalizesponse rate among beast cancer parties, powing to the fact that for "adversal except patterners" how till the ratio in a substitute and activity of the control of the

Animalpachrinick is usually regarded highly among affektes and bodybalders as an estrogen maritaners agent. Stacklis have also not it to explose of decreasing amontase activity to as much as \$2% after administration of 250 mg per day. Patient response meta also show among/affertified to be at least as effective under certain conditions. Due to it discussed between the condition of the condition

# History:

Aminghatchinide was FDA approved as an anticonsistant drug in 1940. Side effects were common with termiter, bowever, including downsiens, diziones, and partial loss of more control. In 1966 reports of adrenal insufficiency subsequent to arrinoglatchinide use were reported. The drug was withframe from the LLS matters as anticrobast raft turns my early due to its recently understood effects on the adrenal gland. By 1967, however, the drug was retrivoked of an emp perspe, narmy whithfrom of moretimes actively and the tecturared of breast career. It was one of the first arrontimes withfrom soft many fact that the control storoid production, the U.S. FDA also granted approval for the use of arminghatchinide for the numerator Caladage's systems.

At one time aminoglatedinited was available under namensa brand names and in more than 2-docton countries. Cable Cydaden and Chrimten preparations were by for the most common, and could be fourd in such nations as Apparitin, Australia, Austria, Brazil, Canada, Chile, Cache Republe, France, Cemmy, Heng Keng, Jerkard, Isanel, Jan, Malaysia, Nechristurk, Norway, New Zashard, Rossa, Sorda Africa, Sprias, Nowdon, Soxberhard, Lirade Kalgobin, and the Under Saises. Additionally, deep code of the conon occasion under other names inchiling farmoldustis, Mozaol, and Mammeri. The soat Today, the ding memis analible in a very small narbee of countries, most notability the United States (Cytadren), Russia (Mamonni), Hong Kong (Orimetene), and Australia (Cytadren).

#### How Supplied:

Aminoglutethimide is most commonly supplied in tablets of 250 mg.

# Structural Characteristics:

Aminoglutethimide is an analog of glutethimide. It has the chemical designation 2-(4-Aminophenyl)-2- ethylglutarimide;3-(4-Aminophenyl)-3-ethylpiperidine-2,6-dione.

#### Side Effects:

Frequent side effects associated with mirrodyntefrinite lexhale fingus, defirmes, skin manules, fever, and musen. Other side effects, my relaxie sleep disorder, aprills, depressed, storneds upset, thyroid dysfurction, virilation, juratice, elevated choisteroil levels, changes in blood cell courts, and high blood pressor. Addrismally, those bodybuilders and affisites taking at at a dosage high enough to promote cortical appression often note that reduced levels of this hormore brigg dost time each and pairs in the joints when trying to lift heavy weight. It seems logical that this might lead to an increased associably to ingrit, breas should be earlied and to outcome themselved using the short periods in which this drug is used in high doses. Most fine listed side their sides there are more common with affect dood registers that find the advantage production of cortical and more common with affect dood registers that the advantage production of cortical and to recovere themselved are simple storners.

#### Administration:

Amonghathrinke is medically indicated for the teatment of Cularity's syndrome, metastric breast cancer in postnerospousal women, and pulliative teatment in men with advanced prostate cancer. When used to treat Cularing syndrome, the dosage used may range from 10.00 mg to 2.000 mg per day, often in conjustment with 20-30 mg of fullyaccordisces to avoid the determinated adsertal escape phenomenon. Adhetes and bodybralies using amonghathrinke for confoil arbithrious the commonly take a dosage of 1,000 mg per day, usually for brief periods of 2-3 weeks or less (10 days of use pre-content is reported with some trodybralies). A schedule of 2-450 so. 2-4bps off may be used in an attempt to eaterd the effectiveness of amonghathrinke for longer periods, but such use is usually demanded in piece of the just estimate and interestinct. The foreign error commonly used to 500 mg per day (12 to 2 a blasts), with 1 tablet (250 mg) per day appearing to be the most common dossess selected.

#### Availability:

Aminoglutethinide is produced in a small number of countries, and is a fairly expensive pharmaceutical. As such, it may sell for as much as \$2 per tablet on the black market. This, combined with limited availability, has severely limited its more widespread use.

# Evista (raloxifene hydrochloride)

# Description:

Baloutiere hydrochteride is a socord-presention Selective Estogan Receptor Mediator (SIRM) of the beautificipater fainty, like sing similar in effect to menodis, edibling entergen receptor antigoatie (sheld) properties in some issues while actigs an entstogan receptor agenties (excluded) in others. The milit pool of variation between flace too agents on exceptor agenties (excluded) in others. The milit pool of variation between flace too agents are under the state of t

As an aris-entogen, adhetes and bodybalkers may use this compound to combat the entogenic size effects caused by assumationals or entogenic stronts. The principal among these size effects in gooccurration, a purpose for which relations hydrocidized seams better Pediatrics, which looked at how these to magest compared in the neutrons of possible pathernia processors, and the processor of the processor of the content of possible pathernia processors, and the processor of the proc

Typical of an auti-energon, includine hydrochibrite should also offer some benefit as a testostome-stimulating comporat. We see the first demonstrated statistics on a group of older most legal 60°D. Years, where daily does of 120 mg were able to necess serum and biosonabile (including less) southern 100°C, 10°C. Though these legals are not duringly, they do demonstrate an anti-entergoin; effect instead of an energosis (regulate) one when it as a mightest to ICI by includes along the state of th

There are some negatives to iribiting the actions of estrogan that should be addressed. For one, estrogan is a bearderful hommer when it comes to (RF-1) levels. In studies with accuracyply patients in studies from the proposed proposed of the proposed proposed to the delivery of the control proposed for the studies of the control proposed to the delivery of the proposed proposed to the control proposed to the studies of the control proposed to the control

# History:

Bakodine Judocikirdie was developed by El IB/8 & Company, and FDA approved for UU. Such in 1917. In first inclination was set the of an outdoors incurrant, only in a fully to increase bone doneshy his 2007, the FDA equathed the indicated uses for the posteriors in the contract of the c

#### How Supplied:

Raloxifene hydrochloride is most commonly supplied in tablets of 60 mg.

#### Structural Characteristics:

Raloxifene hydrochloride is classified a selective estrogen receptor modulator, with both agonist and artagonist properties. It has the chemical designation 6-Hydroxy-2- (p - hyd roxypheny 1) betaze [b] thien - 3 - yl - p - (2 - piperidinoethoxy)phenyl ketone hydrochloride.

\*\*\* \* \*\*\* \* \* \*

#### warmings (Stroke):

The FDA mundates that the following varning be present on the prescribing information for fession (another) shock-derivable; "WARNING," INCREASED FIRST, OF VENOUS. THROMFOCHROLISM AND DEATH FROM STROKE. Increased risk of deep wein tumbrosis and patients over proported with Exist. Women with active or past better of venous of them therefore the venous with a strength of the venous with a strength of the venous o

# Side Effects:

Commo side effects associated with the use of naloxine hydrochloride include bot flashes/labeling healther, mulsise, weakones, camping edense, sworting, depression, weight gain, and gentrietestimal disturbances such as muon, worning, irrigaption, and danher. Lass common side effects related benset just vigant belong therothophelisis (inflammation of view associated with blood clot), and visual disturbances. In more cases includine hydrochloride are has been associated with studie, merosing of the artistic transient schemes attack), patternary eartholis, deep-west thrombook, low while blood cell transient attack, justice and the produce of the produced produced and the developing first, and dought prove the seed driving recognition.

### Administration:

Baloxiene hydrochierde is FDA approved for the reatment and prevention of ostoporous in postmeropausal women, reducing the risk of insusive breast cancer in postmeropausal women with ostoporousis, and reducing the risk of finassive breast cancer in postmeropausal women with ostoporousis, and reducing the risk of finassive breast cancer in postmeropausal women at high risk it of insusive breast cancer. The recommended door is one of ome gabelt administered once per day, without regard to make. When tood (off-label) to migrate the control of the previous control or, mit and these and bodyleaders of control the control of the contr

#### Availability:

Batodien Judocláride is multible in over 50 couries. Aside from a smill number of other brands, the Estem poudest from ELI JQ. & Company in smell keplo be excounted. Price is often a concern, as indexine lydrocláride is considerably more copression from a consecuent of the anti-origent bodyleaders and athlets are already accentered to such as the contraction of the contra

#### Fareston® (toremifene citrate)

Description:

Toendines citate is an anti-estrogacie drug, specifically classified as a Selectic Estroga-Recepter Medistate (SERM) with rincise dispertit and attragates reporties. It is a newstrocial triplemyfethyetic derindine, similar in structure and action to both Nobades (tumordine citate) and Chemi (clemphore citate). Toernifiers citate is used for the trummer of breast caracci in postmorpussal women with estrogar-receptor positive or estrogar-receptor indivious (massar if the caractic is storagen receptors) larmes. If works by attacking to the estrogan receptor in various issues in a competitive manue, blocking endagenous entiring in two excepting behaping activity, has an anti-estroga in many bissous, endagenous entiring in two excepting behaping activity, has an anti-estroga in many bissous, associated with the use of amountable or estrogacie authodistant/eagusit servoids. This may include sensormous include behaping and the contraction of the contrac The triphesyldriphes compounds (torenthem extrate, turnsoften cirate, charge) tented to be somewhat triaterially extreme in the Four This mean that with they can block estingenia entirely in some areas of the body, they can nestuly act as estrogens in this other local great entirely in some areas of the body, they can nestuly act as estrogens in this other local great point in the patient of interest in the regative cardiovascular effects of these disease, compounting the standy dealing with the regative cardiovascular effects of these days, compounting the standy dealing with the regative cardiovascular effects of these days, compounting the standy dealing with the regative cardiovascular effects of these days, compounting the standy dealing with the regative cardiovascular effects of these days, compounting the standy dealing with the regative and the regative days and the regative days

# History:

Toerarifice cirtate was approved by the FDA as a prescription drug in 1997. It is sold in the USL sader the Farstent bend rune, which is much by GTL. In: Farstents is also available in over two dozen other countries including Austria, Austria, Belgiam, Cache Republic, Fishand, Farnec, Germag, Greece, Hangar, Hendra, Haly, Mexico, Nethrarinds, New Zealand, Portugal, Russia, South Africa, Spain, Sweden, Switzerland, Thulland, Turkey, and the United Kangdon.

### How Supplied:

Toremifene citrate is most commonly supplied in tablets of 88.4 mg, which are labeled as (and equate to) 60 mg of toremifene base.

### Structural Characteristics:

To remifiene citrate is classified as a selective estrogen receptor modulator, with both agonist and antagonist properties. It has the chemical designation 2-tp-[(Z)-4-chloro-1,2-diphenyl-l-buter/fllobroxyl-N-Ndiredfwleth/damine citrate (1:1).

# Side Effects:

Forentine cinte appears to be well tokenacl, with a low incidence of serious side effects. In clinical trials, common side effects sociated with its one included by the described by the described by the size of the described by the size of the described by the size of the described by the descr

#### Administration:

Torentilene citrate is FDA approved for the treatment of metastatic breast cancer in postmeropausal women with estrogen-receptor positive or urknown tumors. The recommended ones ione 60 mg tablet administered once per day. When used (of-libely) migate the estrogenic side effects of ambiolic/androgenic steroid use, male athletes and bodybaldness my use 30 mg to 60 me per day during second treatment.

#### Availability:

Toremifene citrate is widely available under the Fareston brand name. It is not commonly sold on the black market, nor is it a high profile item for counterfeiters.

# Faslodex® (fulvestrant)

#### Description

Fishestumi à a highly selective estrogen receptor autagenist (also classified sus mestrogen receptor downregalon). Il curet sia action in the body not by registique fre production of estiogen, that by preventing a from enerting activity in the body. It does this by histing anothele estrogen corpore in a corportier menter, making them unstable before circularly estrogens. This mode of action is very similar to Nobosket (transition citate) and Closal (chemphene citate), abridge timble them too agents fallestates does not here reside (chemphene citate), abridge timble entrope and the consistent of the consistent of the out as the first ipictuble estrogen autagenist to catch the attention of the athlect bodybuilding world. Although now body used how, when supplet it myo be an effective due for mitigating the sists effects of excess entropen caused by ambolic/intralogusic steroid use such as grococomists, the bulley and increased used restriction.

Feloratura is very potent as an articutogen, significantly more to than earlier medications. Bis Polosdee and Cleini Albrught turgues escapent as to expect and not signostication. It can still produce an environment of low estrogenicity on par with strong aromanes richarbion robe study for exemple, shows felorature to be as efficient in relative in residencing beast cancer patients who have already field with first line endocrine teamment good to significantly more efficiency than turnostic crime. 60° Studies investigating the physiological response to fidestearm note that the day actually downeygables estogen receptor concentrations. Furthermore, it also tends to downregulate projectome receptor concentrations. Furthermore, it also tends to downregulate projectome receptor concentrations for Flestearms does not some behaloud hard hard, and for this resons to felevolar produce four enoughput side effects related to estrogen artigorism such as he fishes, mod alternition, and to recept.

#### History:

Fibestrate was developed by AstraZence. It was approved as a prescription drug in the U.S. in 2002, and is sold under the Fastdacke brand name. The drug is indicated for the treatment of estrogen neceptory nositive breast cancer with disease prospession following intuitive and the state of the properties of the astronoidy. AstraZence has store expanded the market for Fastdact to include over one dozen countries, including Asprains, Belgiam Astra Canada and C

### How supplied:

Faslodex (fulvestrant) is supplied in pre-filled syringes containing 50-mg/mL fulvestrant, either as a single 5 mL or two 2.5 mL injections. The product must be refrigerated for storage

#### Structural Characteristics:

Fulvestrant is an estrogen receptor antagonist. It has the chemical designation 7-alpha-[9-(4,4,5,5,5-penta fluoropentylsulphinyl)nonyl]estra-1,3,5-(10)- triene-3,17- beta-diol.

# Side Effects:

The most common side effects associated with faboratural richale gustionitestimal disturbances such as muesa, vornitur, conseptions, abhormal pairs, and disturbance common side effects reduche healteds, back pair, but finders, and some fronte Less common side effects include and, loss of strength interpretations for compared sections, single bleeding, market pair, and low white development of mit reborn fixes, and should have reduced to the side of the pairs of the side of the compared pairs of the property with the property of the prop

# Administration:

Fubestura is FDA approved for the treatment of hormone receptor positive metastatic breast cancer in postmeropusual women with disease progression following artic-strongen therapy. The recommended dose is 250 mg administered intramscularly (battock) per morth, as either a single 5 ml igéction or two 2.5 ml injections. When used (off-labely to mirigate the estrogenic side effects of anabolic-indrodognic steroid use, mile athletes and bookbooklers may find a samira dose to be beneficial.

#### Availability:

Fulvestrant is available in more than one dozen countries. At the present time, all fulvestrant in circulation is Bloby to be of the Fashodex brand ranne. The drug itself is exceedingly expensive, and as a result is not widely traded on the black market.

# Femara® (letrozole)

# Description:

Letrook is a ron-strookla sleckrise third generation arountsee ribihot. The structure and activity of this compound are very similar to that of Arindrick (unstrook), and it is prescribed for similar medical purposes. More specifically, U.S. prescribing guidelines for lettenoite recomment of the usual for the treatment of postamropassas whereas when should be used for the treatment of postamropassas whereas when the structure of the cancer is responsive to estrogan recognitive and its in higher than the structure of the cancer is responsive to estrogan breast cancer. It is typically used as second law of relatent after an estrogeneous extension structure. It is typically used as second law of relatent after an estrogeneous extension structure and the first cancer of therepy depending on the cercumstances. Made the first cancer of therepy depending on the cercumstances when the first cancer of therepy depending on the cercumstances. Made effects associated with the use of arountable analous and other cercumstances are considered with the use of arountable analous analous and considered with the use of arountable analous analous and considered with the use of arountable analous analous analous and analous analous

Letroole expensests one of the never achievements in a long line of drugs trageting amounts withinkin in its immig them only other contemposal vortice lings developed to due, and has an effect significantly stronger than non-selective first generation arountsee inhibitors. Bike Tolse and (Asharin. The dosage of each halb of Formar is 2.5 milligams, which according to product information was sufficient to lower estrogen levels by an average of 75% during clinical stands. The days, lowered, peopens to done remembring the effective in lower does. The package insert for the product first convention that during clinical stadies doses as low use 1 and 5 milligamps produced 15% and 75% estrogen; reliablent, respectively in a format of the pool and the convention of the control of th

#### History:

The U.S. Foul & Dang Administration approved betweek for preceivious naise in 1907, where it is could by Nouveil naise the Fernam tend mern. Nouveilna also exclusively methods the drug in other nations, and mere than 70 matiens now earry letmode as an approved drug. The Fernam benth of his first disease interpretation workshoot, and is found in such markers as Augustria, Australia, Belgiant Board, Crinato, Chile, Cecho Republic, France, Germany, George, Hong Kong, Baira, Noubenhai, Veser-bande, Hall, Stand Hall, Stand Stand, Standish and Contact and Co

# How Supplied:

Letrozole is most commonly supplied in tablets of 2.5 mg.

#### Structural Characteristics:

Letrozole is classified as a non-steroidal selective third generation aromatase inhibitor. It has the chemical designation 4,4'(1H-1,2,4-Triazol-1- ylmethylene)dibenzonitrile.

#### Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes. joint nain weakness fatious mood changes depression high blood pressure swelling of the arms/less, and headache. Aromatase inhibitors may also decrease bone mineral density. which may lead to osteonomsis and an increase in fractures in suscentible nations. Some individuals may also respond to the medication with gastrointestinal side effects including nausea and vomiting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an aromatizable steroid such as testosterone enauthate is taken in conjunction with an aromatase inhibitor, suppression of HDL (good) cholesterol levels become significantly more pronounced. Since the estrogen receptor agonist/antagonist Nolvadex® generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovascular health

#### Administration:

Letrooks is FDA approved for 1) adjoint treatment of postmospassal worms with hormone receptor positive early besset career. 2 the extended adjoint trainment of early breast career in postmospassal women with nine received 5 years of adjoint trainment of early hereast, 3) first fee in centurist of postmospassal women with hormone receptor positive or treatment of advanced breast career in postmospassal women with disease progression fellowing artist-tonged transp. The recommended doos of between 6 or on Ering table administered once per day, without oragand to meals. When used (off-label) to nigitate the entropies side efficies of antabolical horgonic strend use or breast meak definition, using entropies that the contract of antabolical horgonic strend use or breast meak definition, using a finite contract of the contr

#### Availability:

Letrozole is most commonly sold under the brand name Fernara by the international drugmanufacturing firm Novartis. It is widely available at the present time.

# Fertodur® (cyclofenil)

#### Description:

Cycloferal is a non-steeroidal arti-estrogen that is used in the treatment of menstraal desturbances and anouslaup infertility (na milkipi too otable). It is so syniher in structure to Christifi & and Nobadeoff, and also works in the body as a mixed estrogen againstrategost. This drug is commonly used for foll-field purpose by milke holyblatifies and adulticis, typically at the conclusion of a steroid cycle for the purpose of increasing endogenous testiscenter beek. This is in an attempt to mirrine the regulier inpact that a period of low analogues lovels may have on the physique, which can include significant muck-tos. The day [14] of also commonly used for this purpose, but works by ministright exaction of literating hormone, not as an artis-entogen. He'd is typically looked at as a night exaction of literating hormone, not as an artis-entogen. He'd is typically looked at as a night exacting day used in the Grouph's of volus darken the attention are distribution. After sengang men and the proposed of the contract of the several works after the HCG has been removed (see PCLT-Dax Cycle Theory).

# Cyclofenil stimulates the release of testosterone via its anti-estrogenic action. The

Ispothalisms is one target size of this. By stratering with the britting of entrogen to its receptor in this may offen body, excluded libeds the negative declarick rishbolts recognite fourth by this sex hermone. The enhanced relates of granulchropin releasing homese (Caleff) may reads. It is a few principal of the principal of the principal of the control of the control of the control of the principal of the control of the

### History:

Cyclofenil was developed during the early 1960s, a time when other agents of the same class (such as tamoxifen and clomiohene) were being thoroughly investigated.

Cycledia was soon released as a procription drug agent, sold minky to increase the cluster of conception and to counter centain measurable suppriss. Allowagh the drug seemed to offer a good clinical effect without significant health concerns, it did not see extensive access sold consideration of the contraction of t

### How Supplied:

Structural Characteristics:

Cyclofenil is most commonly supplied in tablets of 200 mg.

# Cyclocia is most containing supplied in ables of 200 mg

Cyclofenil is classified as a selective estrogen receptor modulator, with both agenist and artagonist properties. It has the chemical designation 4.4\*(Cyclohexylidenemethylene)bis(pharnyl accetate).

# Side Effects:

Cyclorial appears to be self tolerated, with a low incidence of significant side effects. Common advance reactions incide low energine electations, someworft shales, (the flules), aldonimal discorrifert, musea/worfnitg, breast discorrifert, breadchet, and abnormal steries bedoing. Premeropasal women may be more assemptive to be fulseed as to be stronger effect entergenic disciprious can have on this population. In miles, the test-solvene boosting and accordance of the compression of

### Administration:

Cycloteal is most commody used (modically) to text accordancy inferrity. Thereposite protector (eccurrent a does of 200 mg free terms per day for 5 days, which is initiated must the start of the menstrual cycle. If Programmy is not achieved with the first cycle, it may be used for 3 or 4 cycles in textle In one interactive the days is also pion in lower does to treat merceposal computers. When used after steard administration (off-likely) to increase continue of the cycles in textle In one in the continue of the major and of the S weeks as part of a comprehensive post-cycle recovery program in place of Clerid (see PCT, Dest Cycle Theory). Similar does are used for except ministration proposes with convolve, although Noblack is usually given professor for the purpose. Seem allocate the cycles are a convolved in the propose collection as a consistent of the purpose. Seed of the cycles are a contracted with using cycles allow a contract for the grammer contraction.

uses an uni purpose a typicary non-tool tig per say an oro weeks. The some lawe reported this approach to be effective, many others find the drug too mild, especially in light of the effects of exogenous testosterone.

### Availability:

Cyclofenii is not widely produced. Availability is presently low on the international market. When located in the U.S., the drug is usually found in the form of Fertodur, made by Schering in Turkey. Counterfeits of cyclofenii drugs have not been a significant problem.

# Lentaron® (formestane)

# Description:

Formstare is classified as a selective inversible strovidal amountse inhibitor. This ager is straturally a derintion of andisonation, defining from this we liftering from the value of possion problemane only by the addition of a 4-judosol group. This group, however, is responsible for causing an inversible shartners between formstare and arountses when the two come into context with each other. This means that formstane will bond with the empire and never let it go, permenterly describing its as result. The curpous will need to be replaced, frough normal antition, before the body will recover its but entangant sparsitissing capacity. This may take several days or more flowing contention of theropy, General from end of classific general and the proposed of the content of the c

Because of fa potent estrogato-appressing action, formestate hus been used chically to trust broat enter protects in a narrher of courtesis relicified gradust, Germay, Switzerland, Spein, Australia, New Zealand, Ilaly, and Malaysia. It has been shown to be an efficitive option as a second less of defense and termording, an estergan receptor artangase, has fidad to elekt a positive response with patients, and produces an overall response statistically simits to mounting when administered as the first in femery, in terms of overall, potency, formestatus is not as storage as the selective fitted generation ribbition. Box Arrinales, (unstancible) or ferman (stronois), One soits, for example, rosts a 79% level of supersistent of surface I may deal for a present of assistance of surface I may deal for a present of a supersistent of estrogate heavy deal of the surface of the present of the immunicated fromestate control with Ferman used, but only a 50% seed for daypression with instrumental fromestate control with Certain and Nobaskas (turnoiden circus), formestatus is significantly more effective at blacking the effects of estions on the body.

# History:

Formstane was the first selective aromatuse irithibor to be developed as a prescription drug, first appearing its Tiance during the mid-1990s under the Lentanon Depot brand name. It was sold by Novaris, which markeded Lentanon Depot in note-loon contrains including Augustra, Austria, Beijarn, Barot, Carnale, Chie, Ceelt Republic, Dormard, France, Germag, Geneve, the Gong Indiana Ganel lay, Malapias, Natheriak, Portagel, Sodiana, Georgia, Ge

#### How Supplied:

Formestane is most commonly supplied in a sterile solution containing 125 mg/mL of drug in a 2 mL ampule.

#### Structural Characteristics:

Principal to the Mark and a second and the second to the s

Formestane is cassified a steroidal selective irreversible aromatase inhibitor. It has the chemical designation 4- Hydroxyandrost-4-ene-3,17-dione.

# Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes ioint pain, weakness, fatigue, mood changes, depression, high blood pressure, swelling of the arms/less and headache Aromatase inhibitors may also decrease hone mineral density which may lead to osteoporosis and an increase in fractures in susceptible patients. Some individuals may also respond to the medication with eastrointestinal side effects including nausea and vomiting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an ammatizable storoid such as testastorone enorthate is taken in conjunction with an aromatace inhibitor summession of HDL (good) cholesterol levels becomes significantly more pronounced. Since the estrogen recentor agonict/antagonict Nakadevili generally dose not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovaccular health

# Administration:

Formstare is indicated for the teatment of advanced breast cancer in postmeropausal women. The recommended dosage is 20 mg by intransactar injection offunction every two weeks. Although not a medically approved form of the drug, staties have demonstrated that a smile fael of colorogous appression can also be achieved with online of formstratuse. Due to poor bioxnailshifty, however, the dose needed is around 250 mg per dry. When used (off-livel) to miligate the estoque; sits effects of antibolicharinogeni steroid use or increase muscle definition, mile athletes and bodybuilders often take 250 mg every two weeks by nicetion, or 250 mgpc and results.

#### Availability:

Formestane is not widely available as a prescription drug, and consequently is rarely circulated in the athletic community.

Nolvadex® (tamoxifen citrate)

### Description:

Tamodine crate is a non-steroidal anti-estoparie ding used widely in chiral medicine. It is specifically a Selection Entropy-Recopering development of the trajumpt-fleeport family, and possesses both entropy anytate and antiquois properties. As such, it may at an attention of the contrast of the contr

The primary worsy among the athlicit body-building population is genecomastic, or the very unagifyly devolupment of femule beast issues in men. This can be first noticed by the appearance of sweding or a small large under the rippit. If if to progress, this can divelop into a large land-issue genecomation that may be an invesselled occurrence without surgery, and the state of the properties of the propert

Tamodine cintre also possesses the ability to increase production of FSH (folicle stimularies) hermone) and HI (Jacinize) phormon). This is accomplished by belong negative features distribution caused by estrogen at the hypothalums, which (in the actions of GirdH) fosters the relaces of the memorated pathaps, hermones. This is very similar to the function of Camidli and cyclotical Since a higher relaces of LH can similar the Logdigs colds in the constraint of the control of the co

Note that like some other triphosphelphes compounds, immotine citate can set as an estimage in the few - Eurogean action in the few - Eurogean action in the few is important in the regulation of serum choistened, and tends to support IDL (good) choistened synthesis and LDL (toud) choistened reductions. Some serool-sing shophidates are already calcium serious endings of the serious serious carbon serious endings of the serious serious carbon serious endings of the serious serious endines in the serious serious serious endines in the serious seri

# History:

Tamoxifen citrate was first synthesized in 1962 by ICI. It was made commercially available in the U.S. not long after, but was initially used to treat certain forms of female infertility, a numose for which tamoxifen citrate does not seemed ideally suited. In 1971, the first clinical trials evaluating the effectiveness of tamoxifen citrate in breast cancer patients were undertaken. Two years later noting the link between estrogen and breast cancer and the success of early trials. ICI pursued marketing the drug in the U.S. to treat breast cancer. It was not until 1977 that FDA approval for this use would finally be granted. Tamoxifen citrate was sold by ICI in a wide number of countries under the Nolvadex brand name (the company would later become known as AstraZeneca). A number of generics and other brands followed, presently too numerous to list. In 1998, the FDA approved expanding the indicated uses of tamoxifen citrate to include breast cancer prevention for women at high risk for developing the disease. In spite of continued clinical success with the drug for both cancer. treatment and prevention, in June 2006 AstraZenica finally discontinued the sale of Nolvadex in the U.S. A number of generic versions are still available in this country, however, ensuring easy patient access to the drug. Tamoxifen citrate is presently the most popular anti-estrogen used by athletes and bodybuilders.

#### How Supplied:

Tamoxifen citrate is most commonly supplied in tablets of 10 mg or 20 mg.

#### Structural Characteristics:

Tamoxifen citrate is classified as a selective estrogen receptor modulator, with both agonist and artagonist properties (also known as an estrogen agonist/angagonist). It has the chemical designation (Z)2- [4-(1,2-diphenyl-1-butenyl) phenoxyl-N, Ndimethylethamamine 2-hydroxyl-1,23- proparatrizathoxylate (1:1).

#### Side Effects:

Commo side effects associated with the administration of tumoxien critaria include links, signilia believing usquil dichings, usquilia believing too stronch, bandsche, light-haudschess, oderms, and har loss. Other listed advices teactions include skin rash reduced plated or white belood of court, visual distraturess, teatrie frouls, endormerious and plated or white belood of court, visual distraturess, teatrie frouls, distraturesis entering frouls, endormerious and engine levels, and increased trippicaries levels. An increased raiseless of endormerious career and settine sections with amoviden circuits.

Tamoxifen citrate may cause birth defects and should not be taken during pregnancy.

### Administration:

Timodon citate is indicated for 1) the tentiment of metastatic breast cancer in women and merc? a fullyinart beament of mode-regoint beast cancer for indove persists press arranger and radiation; 3) adjassant treatment of mode-positive breast cancer is poststraceptual women following beast staggery and radiation; 4) production in incidence of contrataent breast cancer in women with DCIS (Detail Carcinomi in Sian) following beast staggery and radiation; 4) protect cancer in women and high risk for breast cancer. In women and men with metastatic breast cancer, a dose of 10-20 mg is administent outless of any immigrate and constrained varieties and offsith offsit of the distribution of the contraction of the c

It is important to note that anti-estrogen use may slightly reduce gains made claring a steroid cycle, as many antiognationables steroids seem to eabth their most powerful anti-olic effects when accompanion by a sufficient level of enterings (Fig. 1) gains and constrained to the contract of the contra

# Availability:

Tamoxifen citrate is widely manufactured, and can be found in virtually every developed ration of the world. The drug is also commonly circulated on the black market. Given its relatively low price and high availability, counterfeit product do not appear to be a large issue

### Teslac® (testolactone)

# Description:

Testakence is a first generation non-selective steroidal morntaes ribilitor, used chically to true entogen-dependent breast cancer. In secent mode of action is urknown, but it is believed to ribilit the anomatuse engine in a noncompatitive and investrable marine. If so, the world he an archity that is very similer to that of Lettaron (formations.). This night also explain why coastion of the drug does not provide an immediate restoration of normal energies production. It does fromation, it is the service of the object to recover is normal energies upon the control of th

Albudy his stolutions is technically storoidal in structure, of offen to ambolio or anthogonic effect to is user. This because if does in process the train necessary to had and activate the anthogon necessor, must be made in 7-to-to-in-flowed progs, in face, it is a Dring is an extension of the contract of the contra

testolactone is used by athletes and bodybuilders not to increase muscle mass and performance, but to mitigate the estrogenic side effects caused by certain anabolic/androgenic steroids.

Note fur the level of aromates inhibition produced with testolectors is significantly lower turn fat produced by the never selectic field agrenation inhibition such as austroation, lemonib, and cuerustane. For cample, one study conducted in 1985 showed furt 1,000 mg of testolectors per day given to mie moral men for a period for day superposed serum estradial levels by 25%, Another using the same 1,000 mg does noted a 50% reduction after six days of use. These armbers see lower frum butte would be expected of the nower third agreeation agents given the substantial estrogen suppression figures they have produced during circlatal with with women.

# History:

Testolacione was first approved un a prescription ding by the FDA back in 1970. It was not any anti-astrogate drug, exhibitg a moderably promotered effect the filling by see and levels of high clinical success. As other more effective medications began to surface for the tenturner of breast cancert, estolacione would not see the success is developers likely planned for it. It would see production in sentl number of countries outside the U.S., most U.S. where the Testolacion is also consistent to the U.S. where the Testolacion and is contributed to the U.S. where the Testolacion and is contributed to the U.S. where the Testolacion and is contributed to the U.S. where the Testolacion and is the U.S. where the Testolacion and the Testolacion and the U.S. where the Testolacion and the U.S. where the Testolacion and the U.S. where the Testolacion are the Testolacion and the U.S. where the Testolacion are the Testolacion and the U.S. where the Testolacion are the Testolacion are the Testolacion and the Testolacion are the Testolacion are the Testolacion and the Testolacion are the Testolacion and the Testolacion are the Testolacion are the Testolacion and the Testolacion are the Te

# How Supplied:

Testolactone is most commonly supplied in tablets of 50 mg,

# Structural Characteristics:

Testolactone is classified as a steroidal noncompetitive irreversible steroidal aromatase inhibitor. It has the chemical designation 13-hydroxy-3-oxo-13,17-secoandrosta-1,4-dien-17-oic acid [day ]-lactone.

# Side Effects:

Common side effects associated with the use of an aromatase inhibitor include hot flashes. joint pain, weakness, fatigue, mood changes, depression, high blood pressure, swelling of the arms/legs, and headache. In 1999, the FDA officially added malaise to the list of possible side effects from this drug, reflecting something bodybuilders had noticed for some time: low estrogen levels can lead to lethargy, as this sex hormone plays an important role in the functioning of the central nervous system. Aromatase inhibitors may also decrease bone mineral density, which may lead to osteoporosis and an increase in fractures in susceptible patients. Some individuals may also respond to the medication with gastrointestinal side effects including nausea and vomiting. Aromatase inhibitors can harm the development of an unborn fetus, and should never be taken or handled during pregnancy. When taken by men (as an off-label use) to reduce estrogenicity during prolonged periods of steroid treatment, aromatase inhibitors may increase cardiovascular disease (CVD) risk by retarding some beneficial properties of estrogen on cholesterol values. Studies have demonstrated that when an aromatizable steroid such as testosterone enanthate is taken in conjunction with testolactone, suppression of HDL (good) cholesterol levels becomes significantly more pronounced. Since the estrogen receptor agonist/antagonist Nolvadex® generally does not display the same anti-estrogenic (negative) effect on cholesterol values, it is usually favored over aromatase inhibitors for estrogen maintenance by male bodybuilders and athletes concerned with cardiovascular health.

# Administration:

Testabetroes is FDA approved as adjustative therapy in the pullarite treatment of advanced or disseminated beats cancer in postureneapousal women when hormonal therapy is indicated. It may also be used in somen who were diagnosed as having that disseminated breast cancirors much mey remempassal, in whom overant fastion has been sibstepared terminated. The recommended dosage is 250 mg taken 4 times per explosed in the state of the proposed of

### Availability:

Testolactone is no longer commonly used in clinical medicine, and consequently is not manufactured on a large sale globally. Presently a small number of testolactone preparations still exist, but are not commonly diverted for sale on the black market given the very low demand for the drue in this novulation.

# Anti-Prolactin

# Dostinex (cabergoline)

### Description:

Cabengine is a selective departure receptor against. This agart is highly specific in its action, with a storage fifty for the departure. Directoper, and a softly for section, and a S-HT2-section receptors. Its man clinical use is for the teatment of principles of the section of the section

Productin is a somatoropic hormone, in the same family as harman growth hormone (somatopic). It is a single perigide hormone, contraining a chain of 199 autino acids. This makes it similer to (though slight) larger than) growth hormone, which is made of 192 autino the contract of th

Although this is almost never associated with runks, high levels of productin have actually been related to learning genocements in a very suml percentage is desired-sized guident. This disorder is other characterized by small flaid discharge fast becomes noticeable with the relation of the contraction of the contra

High products lovels (as would be associated with the need for calengalise) are not regularly documented in section-sing allithes, faither underscoving the relative incommon nature of this devote. We do know that estingen plays a stemilatory rule here, and likely is the key to increasing products careful in males. Welling "Older statisch, knower, allow appreciation are considered to the contract of the statistic products, and as the contract of the statistic products and as a strain hormous in relations, and not necessarily high entropy, may be the cause of laceting approximation. Scaming the medical books, there are the vastice over looking at products in lovel and strain of the statistic products and the statistic products and as a strain product in the statistic contracts are considered as a strain of the statistic contracts are contracted as a strain of the strain of

no consistent effect on protectin secretari.— A tima sen-administration study wan athletes, <sup>664</sup> and a fourth clinical with randrolone, <sup>665</sup> failed to show an increase in prolectin levels.

# History:

Cherupine was developed during the 1980s. The most popular trade rume for this agart is Destince, which is produced in the U.S. and muny often courties by the sign pharmacetral conglumenta Pharmacia. Domine retained market exclusivity on colorgistic in the U.S. for many years, but helvenes of 2005 and 2007 securities agarets: versions were in the U.S. for many years, but helvenes 2005 and 2007 securities agarets: versions were considered to a consideration of the Cherupine is widely available internationally, and can be found in more than 3-do-not different courties. Soutist the U.S. for Dostone trade nurse all dominates mort unders, and can be found in Augusties. Australia. Austria, Basines Busiless Brazil. Cannala. Chile. Casch. Republic, Dermark Finals, France, Germany, Geotec. Kings Republic, Carmada, Chile. Casch. Republic, Dermark Finals, France, Germany, Geotec. Kings Republic, Dermark Finals, France, Germany, Geotec. Mine Republic, Carmada, Chile. Casch. Africa, Singapore, Spain, Sweden, Switzerland, Tarkoy, United Kingdom, and Venezuck In addition to Dostone, colorogistic in marked-to dure at last 1-door offer trade names.

### How Supplied:

Cabergoline is most commonly supplied in tablets of 500mcg.

### Structural Characteristics:

Cabergoline is an ergot derivative with the chemical designation 1-[(6-allylergolin-8beta-yl)carbonyl]-1-[3- (dimethylamino)propyl]-3-ethylurea.

### Side Effects:

The most common side effects reported with cabergoine use include branderle, muses, and vourning, which covered in 26, 27, and 27% of prization (specietyle)) receiving the medication during one clinical trail/effo Other potential side effects include (but are not limited to) constpiction, of your data, beloarding pain, durineta, dizenses, verige, fingage, anarecti, anarectia, mulies, depression, arcomita, but flushes, heart pulphitations, lopotension, broast pain, and zuce, however muses and hackades were from sup primarier side effects. Muny side effects are close related, fairfur reason for starting off with the lowest possible therapeatic does and working part The prescribing information does not mentain data in as clear consequence of an overables, but if those list faither-traints, how blood pressure, and consequence of an overables, that if these list faither-traints, how blood pressure, and processing.

#### Administration:

When used medically to sithild productin secretion, calcergaine is given in an initial desage of 500 meg per work. If immy be batters in resign due one divide into 20 nerme does not separate days. The does may be accessed by 500 meg per work at mentity intervals sard as weak, although does to produce the production of the second by 500 meg per work at metalty intervals sard as week, although does to produce the production of the production of the second by the state of the production of the second by the state of the second by the state of the lower end of the thempoarie range are most commonly used. The user typically state with a dougs of 250mg per application (a half tablet) wise per week. This is used for the work, at which point the doague right be adjusted upwords to all fathet if recoded (1 mg words, at which point the doague right be adjusted upwords to all fathet if recoded (1 mg words) and the second table of the second to the sec

#### Availability:

Cabergoline is not widely used by bodybuilders and athletes, and consequently is not commonly traded in black market commerce. The drug itself is widely available in legitimate modified commerce.

### Parlodel® (bromocriptine mesylate)

# Description:

Bomocripire mesplate is a dopanizamient engra derivative with 12 dopanier receptor against and 10 dopanier receptor angainst and 10 dopanier receptor angainst activities. It is used most commonly as a probact inhibitor in cases of hypercolectionnia, a growth hormore supressunt in aconcegal (byth) does not required, and as an adjustive medication to levedops in the management of Parkinsor's disease. The structure and activity of first drug are very similar to that of callenging (Costonic), in the addition-oblophidating commants, between the management of particular desirable control of the control of

The most vocal proposent of bromceriptine use for fit loss is probably ligh McDraulf, author of the ordine z-Boick Bennereriptine. And Old Dary Bill Now Lies. In this book McDrauld describes how the drag can be used to resemble the metabolism, such that some of the normal physicial persposes to deling frieth height in some the loss of holy fits as the duration of desting accesses are insidered. At let of this Econes on leptin, a hormoze locked at as set of the thremstot, lefting such when labor height consuring (an "ani-structural" local price to the local position of the contract of

The human medical data concerning the potential role this drug might play in supporting oraging fit loss is encouraging. In cases where I was given while dieting, bromorepitue was complete of increasing just fit loss by a satisfactive, justificant depart and second to extend the duration in which the date was most effective. In one case, both placebo and tournest groups were morking around, however, considerable has significant armortion could get for the homorepitud group. However, considerable has significant armortion could get for the homorepitud group. However, considerable has significant armortion could get for the distribution of stops, is a common loss among these undertaking a cubic-rescribed died for the purpose of reducing body fit mass. Adrug that can prevent or delay this plateau may localish be able to increase the overall effectiveness of delating it many including.

#### History:

Bomorigine has been used widely in clinical medicine for its indicated used since the 1970s. It is also men more widely distributed thus its construptor medication cabergalies, which is used for a similar set of clinical indications. In the U.S., the most common brand mannes is Partfold, which is sold by Noramits. The drug is available in dozenous Courties, and is sold under a satisfully large arrather of different rate names including (to me it finited by a sold manner in the contraction of the contra

#### How Supplied:

Bromocriptine mesylate is most commonly supplied in tablets of 2.5 mg and 5 mg. The doses are expressed in terms of base bromocriptine, so each 2.5 mg tablet contains 2.87 mg of bromocriptine messylate.

#### Structural Characteristics:

Bromocriptine mesylate is an ergot derivative with the chemical designation (5°S)-2-bromo-12°-lydrosy-2°-(1- methylethyl)-5°-(2-methylpropyl)-ergotaman-3°,6°,18-trione methanesablorate.

#### Side Effects:

Bromocriptine can produce a number of unwanted side effects, the most notable being low blood pressure, dizzness, confision and nausea. These side effects do tend to be dose nebtted with the low mecommended doses used in brothwhildine are not likely to be much touble for many. Further, initial muses sometimes goes away after a couple of applications, once the user becomes accustomed to the dug. However, the strong incidence of any tarsoclorum side efficies should warrant discontinuing therapy, especially if blood pressure is becoming regardity affected (too low a drop). Less common advener exercious include survively of yrouth, edem, soizures, fittigue, headache, beltunge, usual congestion, rash, edvated liver exercious, and chances in unitary forumers.

### Administration:

When used medically to treat disorders murked by log-persplicationersis (typer secretion of production, an iritial classes of 1.25 mp to 2.75 mp end up of usual processors (and the production), and the production of the producti

### Availability:

Bomcrejtrie is produced in most developed coursies, including the United States where it is sold as a generic dang and under the Parkelo Heard name; Pottle brand name; product comes in the form of both 2.5 mg tablets and 5 mg capsales, with 100 doses per bothe. At the paramers, 10.0 mg capsales myo can tearly \$400. In some antiens, fits pixe may be as low as \$50 to 200 (5.50 to \$10.0 per dose) for generic and other brands of bromcreptire.

Low as \$50 to 200 (5.50 to \$10.0 per dose) for generic and other brands of bromcreptire, and the state of the brands of bromcreptire, and the state of the brands of bromcreptire.

# Appetite Stimulants

#### Periactin (cyproheptadine hydrochloride)

# Description:

Cynobepaties hydrocharide is a first-generiton prescription histanine and sontonia maragasti. This ding is most doing pion in the U.S. of the treatment of allegy-reducted symptoms, relating hyp fover, many mose, irritated oyes, hore, and aweding. It is also FDA approved for the treatment of analysphetic restorious caused by algorithms, colon as an adjust to injectually exprehenre (admental). The servotion inhibiting effect of this drug also joses t a tamega addity in stress genetic. This has led to a considerable manner of off-barbels are as successed with AIDS infection, cancer, or other debitating diseases. Cyprohytatine hydrocharidas is also und concession as madjust to growth hormore therapy includes to foster geneter nativest update and increases in lorse growth beyond what is mentally achieved with AIDS infection, cancer, or other debitating diseases. Cyprohytatine hydrocharidas is also under our consumer therapy in relative to foster geneter nativest update and increases in lorse growth beyond what is mentally achieved the consumer than the constraint of the proposition of the consumer than the constraint of the proposition of the consumer than the constraint of the proposition of the consumer than the constraint of the proposition of the consumer than the constraint of the manner than the constraint of the manner than the constraint of the manner than the constraint of the constraint o

Alhough this is a controversial use of the drug references to the apporties strainting properties of copposition follopsobles are absolutent in the medical lineature. One of the more detailed papers compares the appote increasing effects of cytolopsatates hydrochrisches to megators detactife. If me apport of 14 mm with which the associated with HIV infection. The other agent, megators, is a progastri that was approved by the FIDA in 1993 for the reasonner of america, caches, or weight loss in principares with AIDS. In this inestigation, cyprohydratine hydrochrische was about to have a similar level of brendf to FDM approved agent megastrol, with pattern excessing about 500 court anchies per day.

and gaining a moderate amount of weight with other medications. While the benefits were similar, the side offerts were not. The investigators reported that more than 50% of the patients taking megastrol suffered impotence during the investigation, while the prophesphatine hydrorchizing upon full not such side effects. Cyprohepstatine hydrorchizing upon that on such side effects. Separation placed bytechted may offer an effective alternative to megastrol therapy for many patients, especially those proce to negative side effects associated with this type of Domono manipulation.

# History:

Cyprobepadire Inducebaride is an early arti-listamine drug, and has been sold as a proscription medication in most developed materias for decades. It was introduced to the U.S. in 1961 under the Perucitally brand name by Merck & Co. This brand of cyprobepades phytochedrical was sold for many pases in the U.S. but was subtractly descortized by the produced and the sold for many pases in the U.S. but was subtractly descortized by form an article of the produced and the produced and the produced and the produced and the sold of the produced and the produced and the sold of the produced and the produced and the United Kinglom It is also add under dozens of other brand means careful the work, it plots higher and male specified properturations. A number of generic products are still sold in 1961 the 1962 and male specified preputations. A number of generic products are still sold in 1961 the 1962 and male specified preputations. A number of generic products are still sold in 1961 the 1962 and male specified preputations. A number of generic products are still sold in 1961 the 1962 and male specified produced and the produced and the produced preputations. A number of generic products are still sold in 1961 the 1962 and male specified produced and the produced as a deport-time mortice similar.

#### How Supplied:

Cyproheptadine hydrochloride is most commonly supplied in tablets of 4 mg.

#### Structural Characteristics:

Cyproheptadine hydrochloride is antihistaminic and antiserotonergic agent with the chemical designation 4- (5 H - diberzoo [a,d] cyclohepten - 5 - ylidene) - 1 - methylpiperidine hydrochloride sesouthydrate.

### Side Effects:

As a fire-generation anti-bisturine, cyproduptative lydrochloride my be ponce to producing a marker of riskel effects in the serv. The most common of which is solution or the closes "uni-bisturine lettings", which is common to these types of drugs. For some users, the treatment for the producing hydrochloride will produce will conside any potential as a greater than the contrastance of the producing and producing and potential as specified. recessible discontrastancing a best Order less common side effects of concern include, but are not limited to, disconsist distinct of them than of the contrastance and the contrastance and the contrastance of the contrastance of the contrastance of the contrastance and the variety distincts of the contrastance of the contrastance of the contrastance of the contrastance of the variety distincts of the contrastance of the contrastance of the contrastance of the contrastance of the variety distincts of the contrastance of the contrastance

#### Administration:

#### Availability

# дуанацииу.

Cyproheptadine hydrochloride is produced in a wide number of countries. Although it is not commody traded on the black market, high supply and the relatively benign nature of this drug loose controls) make it easily diverted for sale when needed. Given its low demand, counterfeiring of cyproheptadine hydrochloride preparations is not common.

It is interesting to also note that the Dominian steroid product, Anabolex, actually includes 1.5 mg of synochoptatine lydorchetric in each 3 mg methandrosterobere tablet, which was added by its developers to feditate irreased earlier intitude and weight gain during annihold therapy. The 2.1 min provided is optimal for a daily dose of 24 mg Diambol (as very common anarcus, as a would provide 1.2 mg/coprobaptatine photochietis (the measure common daily dose). This is the only common anabolic steroid product that includes correlated in the ordered time as an anabolic steroid product that includes

# Cardiovascular Support

# Lipid Stabil<sup>TM</sup>

#### Description:

Lipid Stabile is a chelesterol and candrouscolar health support supplement. It was specifically designed to be a foundation supplement for secretal users, and is expecting before project period. The ferminal specifically focuses on supporting four separatic areas of continuenciar learning 17 melectra. Life, Undersolary Represental Prod Chelestero, 25 trighcentics, and 40 reducing a continuenciar learning properties of the supplementation is an important and often convoloded part of rimanging cardiouscoaler below during endouscoaler belowing the continuenciar learning and the continu

Lipid Sabile contains a comprehensive blend of nearly one doesn natural ingedients. Each component in known to play an inportat noise in cardiovoscular bash and muny buse significant chiral support demonstrating beneficial effects on key health murkers. For example, gaing possible of some of the bedoods neighborst, and she no about noise in acrease IRL cholestorel, reading collection and in propose the attributed noise of exemption, and the propose for different neutral contradiation of the FCCO and high polyberial outports in some backborst inguiseless. And the gain, is also shown to reprove cholestoral and reduce contained street Enclaration of the Second other chirality stated in people in known of contained to the contained of the proposed properties and the gain of the contained of the properties of the properti

#### History:

Lipid Stabl was developed in 2008 by Molecular Nutrition (U.S.). The focus was specifically on designing a supplement that can help support cardiovascular health in antibolical/informer steroid users. The product can be found through international distribution, although it may possibly be considered a drug product in some regions with strict controls on herbal supplement product.

# How Supplied:

Each serving of Lipid Stabil is supplied in 3 capsules, and contains a blend of green tea extract (750 mg), garlic powder (600 mg), inositol hexanicotinate (400 mg), polygonum

cuspidatum standardized for resveratrol (200 mg), pantothenic acid (100 mg), phytosterol complex (100 mg), policosanol (10 mg), and selenium (200 mg).

# Side Effects:

Lipid Stabil is a natural dietary supplement and is not expected to have notable side effects.

### Administration:

For general cholesterol and lipid support or as an adjunct to anabolic/androgenic steroid use, Lipid Stabil is generally state at a desage of 3 capsales per day. Note that a natural product such as Lipid Stabil may help reduce cardionsecular toxicity, the carmot be reduct upon to completely definited postful cardio-resoluted manage from the above of anabolic-inadequesis second denge. Care should always be taken to motite at large-cet of theils them taking AAS

#### Availability:

Lipid Stabil is produced in the U.S. by Molecular Nutrition. It is available for export, and may be found in Canada, Europe, and other international markets.

Lovaza® (omega-3 ethyl esters)

# Description:

Lozza is a prescription comps-3-acid supplement which contains edgls clares of conceptaneous acid (EM) and docool-senson cair (EM) had its fundamentally miles to most over-the-counter fish of supplements, except that Lozza is highly particle to drug quality saturdates, from each with a high (FOP) concentration of compact 3-acids, and has good trought extensive clinical trials for a specified therapeut: use. Otherwise to the benefits of EMP-DEM Available to rependicable with an ogual neurora-yee for those on papersars. In the PEP-DEM Available to rependicable with an ogual neurora-yee for those on papersars and (~500 mg/dL). Clinical status showed trighceride reductions by as much as 45% with its use. <sup>57,000</sup> in Addition, this prescription approache to be not how to recome EDIC (apod), chelested beech improve lipoposition particle size and achelous distribution. <sup>601</sup> and reduce conditions that mentiling is now gained by 30% CE PANDEA supplements are communitation by an experiment of the contraction of

The mediumin of action of eiscosportaneois and (EPA) and deconductories acid (EHA) is in a fully understood. These orange-3 exists appear to cert their formoble properties over seram lipids frough a number of different but complimentary pathways. For one, EPA and DEA appear to be efficient a triavestige for group lepartic properties passe. When the nexuses the exerction of LDL closhstered and triply-certics. These orange-3 acids may also increases meta-chardinal personsoral mode-continue, rotate gate anability of fur againtering meta-decided and personsoral mode-continue, rotate-gate as anability of my acid for fail synthesis. We have a superior of the continue of the continue of the continue of the continue of the proposation. PLA and EPA also appear to the sevent field including reduced substitute availability, reduced pilely synthesis, and resumed paid breakfoots.

#### History:

The first prescription drug product containing omega-3 acids was approved by the U.S. Food and Dug Administration in 2001. It was sold in this market under the Ormocor brand name until 2007, when the manufacture, Relater Planmacoricals, changed the name of the product to Louze. This was done to eliminate any confusion with the blood clotting medication Amicar (antinocaprois acid). Louze is presently distributed in the U.S. by the international drug manufacture (Glossofmith Kine.

#### How Supplied

### LION SULDINGU.

Lovaza is supplied in soft gelatin capsules containing approximately 900 mg of omega-3-acids each. The doage consists mainly of eicosapentaenoic acid (465 mg) and docosalexaenoic acid (375 mg).

### Side Effects:

Lovaza is a natural dietary product and is not expected to have notable side effects. A small percentage of patients reported mild adverse reactions during clinical trials, including back pain (2.2%), flu symptoms (3.5%), infection (4.4%), pain (1.8%), againg sepectoris (1.3%), infection (4.4%), and alerted taste (2.7%).

### Administration

Loxan is prescribed in a dosage of 4 capsules per day for the treatment of very high trigheerinks. Given high cost and hired access, Loxan is not commonly store they IANS users, Instant, most accord users will artishner 4 gamp per day of a quality fall of the contract of the cost of the day containing the context. FIAN and IANA may ledy nodare entirescents toxicity, that carned be noted upon to completely definite potential context great metal and the cost of the cost o

### Availability:

High concentration omega-3 acid is marketed as a prescription drug product in the U.S. under the Lovaza brand name. Lovaza is also sold in select European and Asian markets. High quality fish oil supplements containing EPA and DHA are widely available over-thecounter in most markets.

### Diuretics

Description:

#### Aldactone® (spironolactone)

Spienobectore is an attaqueist of albotectore and is pharmacetically classified as a futuric. It is not by compression yielding dischostores behing to receptor so, expectally in the result has where albotectore is involved in sodium-plotession exclusings. The ding causes invested amounts of sodium-and water to be excreted and protession to be resident. Bits is different from some other agents of this class, such as a loop dareft like farsoments, which can significantly receive the executions of sodium water, and prosession in this light for a narrher of conditions including the renument of light blood pressure, edem related to congestive heart finites, leparaldostorous fine over production of ablostorous, and lippolarisms (drop of potensium, othen associated with other medical traversional), so the contraversional cont

#### History:

Spironolectone was developed during the late 1950s, and first saw widespread use in clinical medicine during the early 1960s. The drug filled an important need for a directic that does not deplete potassism; and therefore has a less dramatic impact on electrolyte balance in the body. In many regards it is looked at as a "safer" and "midder" directic compared to other agents in this general category, such as loop directics or thizacket, allowing this agent room

for murket stability and success. Today, spironolactore is widely distributed throughout most of the developed world. It is available in dozens of brand rames, the most commonly identified is probably Aldactons from Searle. It is also widely sold in mixed-agodient proparations alsogaside other directics. This includes foresmide, as seen in the product Lashector, or hydrochlorodinizale, in Searle's (also widely distributed) mixed directs product Aldactaside.

### How Supplied:

Structural Characteristics:

Spironolactone is most commonly supplied in tablets of 25 mg.

### Spirotolacione is most contribiny supplied in tablets of 2.

Spironoluctone is an aldosterone artagonist and diuretic. It has the chemical designation 17hydroxy-7alpha-mercapto-3-oxo-17alpha-pregn-4-ene-21-carboxylic acid y-lactone acceptate.

### Warnings (Dehydration, Elevated Potassium, Death):

The rissue of directic drugs for physique- or performance-relaturing purposes is characterized as a higherith periche. Directis mup produce a discretating less of debledation and electrolete instalance when administently without proper medical supervision. Many doubts have been associated with the misse of these drugs. It is also important to took that the superplanetation of potassism, other through pharmacerization or a difficient in posturing is gornally not advised while taking a posturan-purping distort the spirocolatone. Decessive postussion irake may cause hypertalenia, which may lead to condition resultant in an apposal by dark

### Side Effects:

Adverse neactions associated with spicrookerone administration my include gonceromatio, curranging diarrials, downsiens, lettings, breadules, skin irritation, reals, metal confisions, fewer, importance, loss of muscle coordination, mentalmal irregularities, vibrilation, and desporting of the vision. Spicrookeroleur has also been shown to easie turner in rate, Berese cancer lass been reported in some patients receiving spirorolectore, but no causal relationship has yet been established. Additionally, fits compound my colibit arillarial analoguier proporties, as both a weak inhibitor of audrogenireceptor binding and tectostome bisonofield.

#### Administration:

When used medically to teat hypotension, the infair commended dosage in adults is 50 mg to 100 mg per day in indicied dosses. It may lack to woweds for a menurum sepones to be achieved. The dosage may be adjusted later depending on the individual reads of the partier. When used of field-byld what bodyladies to increase much definition or althest to reduce weight before a weight-in a dosage of 100 mg per day in a single menting application; is most ormorms. This may be contriused of 7s to 5d says prior to the ever, and will define result in a harder and more defined appearance to the muscles (or substantial readson in body weight prior to reductions).

Woman are occasionally attaced to opionolactore for is effect as an arti-antrogen. It is sometimes used as a saley net at a pair when androgan levels have beenere problement during a cycle, and is used in an effort to reduce the risk of permaner witzzino symptoms. A chaogus of 25-57 in quality for 10 are used in so in the soft price purpose, and may be enough to ward off side effects while antropan levels decline (the steoral regimen terminaled). Euro-provincelactors in more effective at lovering endoprosa antropan levels than inhibiting antropan action, it is certainly not to be considered a cure-all remody for advantrous terminal states great formula affaires.

Size this compound is one of the milkest (prescription) distretic options, it is a common starting point for an endy competitor. Once finding with its effects, many attents to achieve a stronger level of water loss by mixing spironolaction with a thirdide or fitnesemide (Lassi). The goal is to provide strong water exceeds only this sea distimptionstain host bits than using the stronger distreties above. When mixed with hydrochherothizaide, for example, the 100 mg spironolaction dosage is often out in half and an equal amount of the thizaide is taken. The 50 mg/50 mg combination is reported to noticeably acrosses water excertion without demantic size effects. The potessians re-obseption secon with spinoncheone should be latalized or with the thinzide so pomission levels will not be as groutly affected. On the balanced or with the thinzide so possissant levels will not be a groutly affected. On the thin of the contract of the potential contract of the poten

It is important to note that while Lasix and Hydrodiaril appear to be more effective at inducing short-term water loss, they also have increased risks as compared to potassiumsparing directics, and should be approached with caution.

#### Availability:

Spironolactone is widely manufactured in both single ingredient and multi-ingredient drug preparations. Low cost and wide scale availability make this a poor financial target for counterfeiting.

### Dyrenium® (triamterene)

### Description:

Tienternes is an oral distret used medically to treat edeum. Edem may occur without known cause (distriptic edem), or be associated with her or taking disease, compassive hard failer, corticosteral/progent use, or the comprehension of abstances. Timaterous bare failer, corticosteral/progent use, or the comprehension of abstances. Timaterous bar preserving possions beeds. As the mare suggest, this drug produces a procuraced district effect without the potentian loss associated with fluidiselysop districts. The need for potentian supplementation is, therefore, reintened with this again the darset active of profession supplementation is, therefore, reintened with this again the darset active district effect without the profession supplementation is the contraction of the contraction o

Timitenes is stilled (off-likel) by bodybalden and affects to shed subcutanous water prior to a bodybalding conportion, or to make weight class adjustments in centil competities sports. Bodybaldines in particular roly heavily on the increased definition that can result when water tenerion is reduced. The highly defined "streddle" physique common to bodyballing today is nearly impossible to achieve without the use of districts. At the same transit, districts are found on the signal of the

### History:

Trientence for our extensive chiral use during the 1906. It was used bragly as a standardor agar at fine the stort on the corner would used again included modern contribution products with other district. George Telephone the contribution products with other districts. The proper districts is fruited and those against Henry, the other mode potent districts like thratical and those against Henry the production may be desired most soft of the modern in a nechocal or one orientated need for possissian supelumeration during themps. Spells implicitly reputation of triustreness are still sold, however, and can be found in several legislate the still a special possission of the still and the still a several form the still a several several several several contributions of the still and the still a several form the still a several sev

### How Supplied:

Trianterene is most commonly supplied in capsules of 50 mg and 100 mg.

# Structural Characteristics:

Triamterene is a potassium-sparing diaretic. It has the chemical designation 2,4,7-triamino-6phenyl-pteridine.

### Warnings (Dehydration, Death):

The risses of district dray(d) for playing- or porformuse-orbanicity purpose is characterized as in playing agreeds. Destricts may produce a file-florating bard of delightation and electrolyte installance when admissized witted proper medical supervision. Many deaths have been associated with the misses of these drags. It is also important to note that the supplementation of potassian, other through pharmecortains or a deef rish in postssian; a seemaly not advised while taking a postssian-squaring district like trinstructure. Excessive postssian inside may cause hyperkalenia, which may lead to cardiarespulsaries and possibly donth.

#### Side Effects:

Timterene use my be associated with electrolyse rischaine; including electrated or decreased prossions needs Sags of electrolyse irrelatures risched by morth, first, weakness, lettings, drossionses, rentleseness, muscle pain, muscle cramping, solares, rentleed urare volume, too blood pressure, and gestoresterized disturbaness. Other side effects may include museau, vorning, juanticie, blood platelet deficiency, ameria, zostenia (bodiaplo of metabole wasse products in the blood, rentl states, and other kidney disturbaness. Additionally, some me side effects characterized as hypersensishity sourcises consortiols for all complete, of meeting and consortiols for all complete on device meeting of meeting and consortiols for all consortal sides; in careful and consortions are all consortions and consortions are all consortions and consortions are all consortions are all consortions and consortions are all consortions a

### Administration:

When used medically to treat hypertension, the usal initial dosage in adults is 100 mg toice daily after mask (200 mger dos "). This may be increased, but should never exceed 500 mg per dos just total. Among bodybalders, this drug is commonly used for only a few drugs prior to a competition, adapting the dosage over the course to left in the sell of district effect. Since then a long being action, it is generally administend once per day. One capuals (100 mg) is usually latent he first first just from errors; with a must, and the effect judged over the next several hours. The dosage is usually increased by one capuals per day for 2 days at surgue and the sell of the contract treatment of the contract treatment

### Availability:

Trianterene is widely sold throughout the developed world. Although single-ingredient preparations containing trianterene are available, this drug is most commonly sold in multi-ingredient preparations targeting edems and/or high blood pressure. Low cost, modest demend and high availability make this drug a low-most traver for counterfeiting.

### Hydrodiuril® (hydrochlorthiazide)

#### Description:

Hydrochlorothiazide is a diaretic from the thiazide family, used medically for the treatment of edems and hypertension. This drug acts by reducing the reabsorption of electrolytes, thereby increasing the exerction of sodium potassium, chloride, and consequently water. In comparison to other diareties, Hydrodiaril is stronger than the potassium sparing agent.

AMERICANDO OPPURISACIONE, OU WORKER URAI HE ROP AGENT LEAST, (LEDOCHUSE), WHITE DESISSIM EXCEPTION IS HAND IN PRODUCED tHEN THE SEE WHY! LEAST, IN the use of a potassim supplement (or a potassim rich dief) may still be necessary with this product. This is usually dependent on the dose and duration is which the drug is administered. Clackim exception may also be pronounced with thiszides, but again, are weaker in this regard than Lesix.

The use of disturcts has been increasingly popular in a number of athletic displanes. For statutes, these dings are very popular among bolybalates who use them to she statutes, these dings are very popular among bolybalates who use them to she advantages of the statutes of the statutes

#### History:

Hydrochronizade was developed during the 1950s. Geon the widespread rature of diseases associated while pilothed pressure and destruct, and quickly actived large-scale acceptance and distribution. Hydrochronizatios became a finalmental famous flearing in the same of radicious, where treatment widely multible tools produced the same and distribution. Hydrochronization became a finalmental famous flearing in the same and admit and distribution of the distribution of different brant mere and generic products. Single-inspections (where Inputs and Section 1981) and the same and present production (where hydrochronization is the only active drugs are far outstandarded by programs of the production of the hydrochronization of the complex distribution of the same and th

#### How Supplied:

Hydrochlorothiazide is most commonly supplied in tablets of 25 mg and 50 mg.

#### Structural Characteristics:

Spironolactone is an aldosterone antagonist and diuretic. It has the chemical designation 17-llydroxy-7alphamercapto- 3-oxo-17alpha-pregn-4-ene-21-carboxylic acid y-lactone acetate.

### Warnings (Dehydration, Death):

The misuse of diaretic drugs for physique- or performance-enhancing purposes is characterized as a high-risk practice. Diaretics may produce a life-threatening level of dehydration and electrolyte inhalance when administered without proper medical suncevision. Many deaths have been associated with the misuse of these drugs.

#### Side Effects:

Hydrochimizacie use may be associated with electrodyse intulative. This may include postusian and sodium deficiency, as well as physochemic alkaside, sin increase in blood bizarbonia che to significant chinride bosto. Signs of electrodyse intulative reluded by morth, thist, undessors, feedings coloruses, reclassess, messele, messele, considerate disarbances. Other side effects may include reluciate appearing seators, reduced urine volume, low blood pressure, and gastronizatival disarbances. Other side effects may include reluciate appearing, musea, contingio, constpation, disarbance, transmission of side effects on particular effects, musea, contingio, constpation, disarbances proposed and control of side of the control of side of side of the control of side of the control of side of

### Administration:

When used medically to test hyportensors, the usual smit does in adults as 2.5 mg duthy given as a sigid does. The does may be invessed to 50 mg duith, often in two does of 2.5 mg. Note that duith does above 50 mg are often associated with marked neduction in serum possissian Afrikaes and bothyhalisers pointally use the drug (off-leide for very herf periods) of the adjustment. A comment possible is to a darknesse this drug once periods of the contract of the

If the application of hydrochlorolizatie is not producing the desired effect, many bodyhalakesthakest will cooke to dad nother distract (mild) before moving on the stronger loop agents. A combination of a postassian sparing distret: Bac Ablactone® (opironductous) and Hydrochtar it sequenced as practicating useful by many and as beloved to slightly balance on the calcium and potassian hoss associated with fie use of hydrochlorolizatic. The application and vocalization conductously, usual varieties with a 25 mg/25 mg application and vocking upwards.

It is important to note that the oversue of districts, saide from being potentially very diagnous, may reach in no much water loss. This can lead to flar, 'delithed' looking muscles. A higher district dosage, likewise, does not absups equate to invessed definition and muscalinty; it is usually regarded as good advice by those in the adhetic community to become familiar with the practice of using districts before using them dainty competition to the contract of the contract

#### Availability:

Hydrochlorothizzide is widely manufactured in both single ingredient and multi-ingredient drug preparations. Low cost and wide scale availability make this a poor financial target for counterfeiring.

### Lasix® (furosemide)

### Description:

Farounds belongs to a class of drugs known as loop districts, which cause the body to except water as set as possissins, soften, magestam cakerus, and chirtic. They are used most commody to text often and high blod pressure. Like other agents of this type, the contract of the contract

Addates and bodybaldens use directies for a couple of specific purposes, and usually for only briefperieds. Our pertire addates in sport with weight class restrictions may use fees drags to drop water weight, in an effort to make adjustments in their weight class standings. Seen the weight recondent is often also quite beginning to the control to the properties often also quite bodyweight considerably with directs, and be back to normal within hours after drug consistenably with directs, and be back to normal within hours after drug consists and relydation. This may provide a some competition advantages, allowing the is only offen to owned degree by the now near univocal nature of some form of 'thooping, we weight' practice in these sports. Bodybaldens may obly heavy on durates when preparing for a contest. Here, a drug like favourishe can efficiently lower subcutancous water concentrations, heights produced a more defined "ripport") but common to competities.

### History:

Farounds was developed during the early 1960s. Much of the intial research on this directive was conducted in Farope, mink foreinn guard labs. The day growed to be quite successful, however, and within a number of years guard workshoke anterior and a cooptime of the control o

#### How Supplied:

Furosemide is most commonly supplied in oral tablets of  $20~\text{mg},\,40~\text{mg},\,\text{and}\,80~\text{mg},\,\text{and}\,$  injectable solutions containing 10~mg/ml.

### Structural Characteristics:

Furosemide is an anthranific acid derived loop diuretic. It has the chemical designation 4chloro-N-furfuryl-5- sulfamoylanthranific acid.

#### Warnings (Dehydration, Death):

Furosenide is a highly potent diaretic, which can profoundly increase water exerction (directs) and talled to electrophe deposition. The misses of diarets drugs) like fromeomide for physique- or performance-enhancing purposes is characterized as a high-risk practice. Duratics may produce a like-frametaning best of debilidation and electropic inhalture when administered without proper medical supervision. Many deaths have been associated with the misses of these drugs.

### Side Effects:

Farsonici use mys le associated with decouple in relative. This my related to depletion of postularin (polystachem), a solurin (polystamin) and polystamin (polystamin) and calcium (polystachem), as well as hypothemic adalaties, an increase in blood biendroute due to significar (fairlier less Signs of describpt in trabuler include dym moth, this, weakness, futunge, drowsiteers, restlessenses, musele pain, musele cramping, science, rendeed uris voalane, the blood pressure, and gestoriestratel disturbances. Other side effects may include purceratis, juratice, associate, and and stometh irritation, cramping, durches, contrajone, musess, vonirigh, disturbances, anomic, decreased wheele of the blood pletted court, dermatile, enal, sain fairling and essentishy to light, the blood pressure, high thole danger levels (hypenglevnik), musel quantum wealness, the polystamin of the polystamin (polystamin and polystamin an

#### Administration:

When used medically to treat edems, it is often given only in a dose of 20 mg to 80 mg per day, which is laten one seing applications for the treatment of hypertensis it is generally recommended to administer 80 mg per day, which is given in two separate 40 mg applications speaced 120 bans parts Athlesse and bodybaleasts speaked just see its drug official high fire very heir periods (second days) of soater adjustment. The dosage and method of the affaire. Our labels see the most corresponding to the state of the affaire of the affaire will be affaire the affaire of the affaire will be affaire the second through the affaire of the affaire will be affaire the affaire of the affaire affaired to a show or competition. The inflind dosage is usually 30 mg to 40 mg, and the mixtures of the inflice ready occurs 80 mg in not not or minime to the effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds. 80 mg in nother to minime the side effects associated with the affaired proceeds.

drug, it is generally used for no longer than 4-5 days.

Note that since fanosemide has such a strong effect on electrolyte levels, it is generally consistend much aside to add a patients amonging agret like Allactione® (epitronhectors) than it is to keep increasing the amount of fanosemide used. Combination districts like this are widely produced as prescription medicines for this reason. The use of 50 mg. Aldactore® and 20 mg fanosemide is a common starting port, and is believed to have a roughly smirtle directive fields to 90 mg fanosemide, but whoth the same level of prossible to like the consideration of the product of t

Injectable foresemble solutions are considered to be significantly more powerful forms of the drug milligams for imaginary the regimes obstituces on the administrate distanceasted or situations), depending on the infoldant needs of the patient. The IV method is much more might acting and produces significantly large peak blood levels of the drug Glient hat the action of fasoestake can be noticed in a matter of seconds or mixtees when given by signiciant, the effect is eastly easier to judge and centred with the method of use, at least under romand conditions. Sience the spictous is much more powerful than the outal, however, is important to emphasize that the chooger must be considerably method in comparison of the conditional production in conditions and action, and state the production in comparison of the conditional production in conditions are considered in comparison of the condition of the condi

### Availability:

Furosemide is widely available, and is manufactured and sold under many different brand names, in many countries. No version of Laisi (or any other diarteix) is currently being counterfield on any large scale. Although it is doubtful these will circulate, make sure to never purchase the drug in 500 mg tablest. These are used only in severe medical conditions, and cortain a dosage that would fleel verve fight to a bealth or sexon.

## Endurance/Erythropoietic Drugs

Aranesp® (darbepoetin alfa)

#### Description:

Dubsportin alli is a synthetic devination of the harmat explresportin protein. It also less the same planumological action as reconfribent harmat englresportin (sportin alli), in the body, explresportin is normally released by the kidneys in response to hypoxa (low blood orogan levels). This in turn taggess hore marrow to acrosse not belond cell production, it such, this hormone is vital to the regulation of normal red blood cell concentration in the such as the total to use of the suggest orderposesies when the body is or ministrainey adoptant roll blood cell levels on in som It is FDA approved for the normater of surnita (low red blood cell court) associated with crober routal fails or chemotherapy.

Dubsports all differs from recordware larum explusopoieti (special all) misky in its duration of activity. This way retent materials is beles in the blood for approximately free times longs; causing it to have a much longer therapeatic window. This means that with durabportian all parties are required to daministe the product match be subquestly than they would apport all the invasion of the subsport of the subsport and its invasible product all the sunably injected on a schedule of three times per work, durabports all requires only one injection in the same time farm. His may endement parties confort considerably, especially when the patient is wisting the physician for routine drung administration.

Findame, earliests are highly attracted to derbeposite afts for the effect it has on roll blood cost production. It is no sever that the partice of blood depile has been popular with endamence sports. This procedure involves removing, concentrating, and storing a quantity of med blood cells from your combod pot her transferad letter. By adding the stored cells before an event (by then the body has restored the lost blood volame), the affalte has a much agonter concentration of roll blood cells. The blood should, Blowske imaged rough med efficiently, and the affalte may be given a significant endamence boost. This procedure cam be quite ricky, however, as blood products can be diffured to see and administer correctly. Darbeports afti is a drug that basically equates to chemical blood depring, and can achieve the same end result fully deprive red of concentrations) with the use of suspine medication.

### History:

Dubosporii alli was deceloped as a praccipito deng by Aragan. The U.S. Food & Dang. Administration fast approved 4 for such in 2011. Aragan is the world's largest biotechnologies company, and the same firm that first brought recombinant epidenzoidissi contents and a contract and a contract

### How Supplied:

Darbepoetin alfa is most commonly found in single-dose vials and prefilled syringes containing 25, 40, 60, 100, 150, 200, 300, or 500 mcg of drug,

### Structural Characteristics:

Darbepoetin alfa is a 165-amino acid protein that differs from human erythropoietin by the substitution of amino acids on the erythropoietin peptide backbone, which allows the addition of two additions N-tilked of diseasechanticle chains

### Warnings (Death, Viral Disease):

The misuse of darbepoetin alfa for physique- or performance-enhancing purposes is characterized as a high-risk practice. Like traditional blood doping methods, darbepoetin alfa can produce an abnormally high concentration of hemoglobin in the blood (polycythemia), which may result in heart attack, stroke, seizure, or death

Some forms of darbepoetin alfa contain albumin, a purified human blood product. Although effective donor screening and product manufacturing procedures are in place, it still carries a risk, though externely remote, for transmission of viral disease.

#### Side Effects:

Side effects associated with the use of dashpoperin alls may include the-like symptoms such as fewer, chills, headacher, muscle pairs, weakness, or dizerious. Such effects tent to be more pronounced at the initiation of therapy. Other adverse reactions may include infection, read, swelling of the side, museus, vonting, darriera, high blood pressure, low blood pressure, cough, tronchilis, or others in some instances distributed in the sheen associated with from the controllers, and the side of the controllers and the side of the side of the controllers and the side of the side of

### Administration:

Durbopoofin allá is indicated for the treatment of anemia associated with chronic renal fisher (CRF) and chemotherapy. The accommended starting dose for the treatment of amenia in adult CRF patients is 4.5 mrg/kg body weight, administered once per week as a single IV or SC injection. The dosage is subsequently adjusted based on changes in hematocirt. Healthy adulteds using darboportin allá for performance-enhancing purposes generally start on the very low end of the therapentic spectrum, and adjust according to changes in hermatocrit. This may ertail initiating therapy with as little as 0.5 megkeg of body weight once per week. Note that it is considered very important to maniter blood cell courts closely during the ertire inition of darbepoetin allia to help ensure hermatocrit is not allowed to increase to an unhealthy level.

#### Availability:

Durbepoetin alfa is not widely sold on the black market. Yet because of the high cost for enythropoissis stimulting agents like durbepoetin alfa, it is a high profile target for counterfeit drugs of this close have even infiltrated legistrate pharmocartical distribution channels, suggesting that care should be taken when purchasing this and smilter dure products.

### Epogen® (epoetin alfa)

### Description:

Eydropoien's a glocoprotein that is produced in the kidneys, and is responsible for strainting red blood of production. Epoch and his is palmenteating form of eydropoient, which was munificated using reconfirmer IDA technology. The compound is produced from animal ceds in which the gase coding for human eydropoients been insent. He biological activity and structure of optoria alia are indistinguishable from that of human eydropoient. Epoch and his suced to team may from of armine, factively strainting and entire eydropoient. Epoch and his suced to team may from of armine, factively strainting and entire eydropoient. Epoch replacements for left less effectively therming as could broad to a considerable of the efficiency of this day quickly much at a node replacement for left less effectively therming some star hand soft less.

Findames earlies are highly attracted to epocin alls for the effect it has on red blood cell production. It is no sever that the practice of blood deeping has been popular with endames, sports. The procedure incohes removing, concerning, and stering a quarity of red blood topy, the procedure incohes removing, concerning, and stering a quarity of red blood (by the rule body) has recorded the bath blood volume, the addition has an article greater concernation of red blood cells. The blood should, likewise, transport cogges more efficiently, and the affirm emply beginn significant endament boots. This procedure can be quarity (by however, as blood products can be difficial to sace and admissive correctly. Freprist all is a single that beautify against so chemical blood deping and can achieve the

### History:

Epocini and two advectoped by the biotechnologies firm August, and first introduced to the U.S. murket in 1984. The relusace of the only is required as a breadfrough in the tentumer of arcmin, which beforehand was being addressed manly with agents that infected or mospecifically trapsford oned productions, each as ownersholes, which may present a number of insomated side effects so the patient. Epochs and murked the development of the first due flat prediction and one flowershole that the contraction of the contraction

In 2002 the subctameous use of Egrec, which is sold only custice the United States, was liked to a run dissons called pure net-of calleys. This is a confinition where the body bees is ability to produce red blood cells. Those that suffer from pure net-of-call apilias issually become dependent on containal bold carriadisions for sarvial Cells to 200 people is likely Ejrecs were identified as developing this rune condition, far in excess of normal expected numbers. Internal insectingations by Johnson, Alchoon linked the high numbers to changes in the product that were made to satisfy European regulations limiting the use of albunin. According to a company polscaperson, a chemical nearction between the new stabilizer that resplicted albunin and the rubber stopper allowed organic compounds to beed into the visit. The company subsequently resplicated is equilial suppores with condition herbor prevent the

reaction. The incidence of pure-red cell anemia in patients receiving crythropoieth seems to have been reduced as a result, although warmings about this reaction remain on the prescribing information for all products sold in the U.S.

#### How Supplied:

Epoetin alfa is supplied as a dry sterile powder that requires reconstitution with sterile diluent before injection. It is most commonly found in single- and multi-dose ampules and vials containing 2,000-40,000 Units/ml.

### Structural Characteristics:

Epoetin alfa is a single chain polypeptide hormone containing 165 amino acids. It is identical in structure to the alobs obsoform of human exyltropositin

### Warnings (Death, Viral Disease):

The misuse of epoetin alfa for physique- or performance-enhancing purposes is characterized as a high-risk practice. Like traditional blood doping methods, epoetin alfa can produce an abnormally high concentration of hemoglobin in the blood (polycythemia), which may result in heart attack, stroke, seizure, or death.

Many forms of epoetin alfa contain abumin, a purified human blood product. Although effective donor screening and product manufacturing procedures are in place, it still carries an extremely remote risk for transmission of vital disease.

### Side Effects:

Side effects associated with the use of eportin ali my include fu-like symptoms such as force, chills, headeder, muscle pair, weakness, or dizzines. Set he fifects tend to be more pronounced at the relation of Berapy. Other side effects include reads, swelling of the side, rances, ventring, distribute, high blood pressure, hepselabilities (excess possissain in the blood), and irration in after side of ispection. In some instances epoced rails has been of the contraction of the side of the contraction o

### Administration:

Epochi and injectable solution is given by subcataneous or immenson isjection. The two paths of administration lawe goards differed effects on the blood level of the days, When given by IV faliation, pack blood levels of the drug gare needed within 15 minutes, and the meaning the path of the p

#### Availability:

Epoetin alla is a very expensive compound, and its use is additionally isolated to certain athletic fields. As such, it is not widely traded on the black market. Given the high cost of this drug, however, it is a lucrative target for counterfeiters.

### Provigil® (modafinil)

#### Description:

Modafial, Brown chemically as heurlydysluphinsplaceturike, is a certard stimular (posphorimitar). It is FDA approach for the tenturent of metospey (a disorder chemicatived by audient and uncontrollable attacks of deep sleep, metal faigue, or investigate for a marber of other use, is thatling the returnent of Arbientry disease, depression, and attaction defect disorder. Modafial belongs to a group of drugs known as Engineer's gread ourself), designed to promote a metal state of spilance and adverse. One of its known mechanisms is to work as an alphel—a democrepte agoint, certifig to also reads in alternation in its GI arbitary all democrepts are spilanced to all the control of the control of the control of the control of the democratic democratic and the democratic democratic democratic and democratic democ

The use of modifield as a stimulard has been shown in stadies to have many advantages over amphetamises. To begain with it is bedieved to have a much lower potential for abuse due to the fact that if produces a lower sense of cuprism. It also displays toop exprisar OSA to the fact that if produces a lower sense of cuprism. It also displays toop exprisar OSA interest to the contraction of the contraction of the contraction of the contraction in the contraction of the USA military, which is looking at it as an energy enfrance for piths and corridor soldiers that moved powers below profits of corridor to design below of the USA military, which is looking at it as an energy enfrance for piths and corridor soldiers that moved to operate for the greatest of the value of the contraction o

Recerts, mediafi has become a popular drug surrogs competite athletes. They use it not simply to 'stay syndrom' to at a performance critarize agar with the shi straintar and endurance-increasing properties. This type of use probably comes as a surprice to show who developed this city, as early reports agarded that this was a 'lind' admires drug without strong stimular properties that would improve athletic performance. Recent studies contradict this deemartment. A subject contradict this offermation. A subject contradict this commentation. A subject contradict to all confirmation in the subject of the subject to a group of 15 rathe volunteers. Three hors after ingestion, anothe concrise to accordance of on a year lenganteer at 15% Visituals (untrains anothe concrise was contained on a yeak enganeter at 15% Visituals (untrains marche) provides of the carries for significantly longs provides of first (~30%), and had guster corgan rathe at execution. They also reported lower subjective entire of previous forcer (REPs), which suggests that the increased performance was in part due to a significantly long provide of trains of the stay of the provides of

#### History:

Modafial was developed by Linfo Laboratories in Firance. It was approved for sake in the Linfo States by the FIsh is 1998, where was structured under the Provqillib Teard mure. Modafial is also found stereminously under this and several additional trade names sacisting Modafially, Vigile, Attenced, and Modasentilla, Attençable the dag appears to sake in the U.S. It is presently classified as a scholad IV cortroded substance, which places modafial in the same category as Vidian and Marce. This is resteed to limit is discussion for normedical purposes by placing considerable legal premises on as possession and importation. The modal applications for the day are fully broth, lowever realizing and importation. The modal applications for the day are fully broth, lowever realized production.

Modafial quietly became popular among competitive affaktes between 2000 and 2004, before the affatic bodies were source of the day, Is use as a performance-enterming agent was revealed to the public during the designer steroid (BALCO) depring searnful of 2004, however, when it was disclosed that many of the same affatices who tested positive for THX also used modafial The 10C quickly barned its use, and with the help of a marker of researchess a methodology for detecting the chemical in the true was developed. This test is now implemented as part of the standard Olympic level drug screening process. Most of the other international athlicit bodies have followed the IOC's lead in braining and testing for modaful. The drug has since lost all appeal as an "invisible" performance-enhancing agent, although is still being used by many athlicits that are not subject to random urine testing.

### How Supplied:

Modafinil is most commonly supplied in tablets of 100 mg and 200 mg each.

### Structural Characteristics:

Modafinil is a central nervous system stimulant related to adrafinil. It has the chemical designation 2- [(diphenylmethylbalfinyllacetamide.

### Side Effects:

Side effects associated with modafinal are commonly the result of its central narrows system strainfaing actions, and my include renoranees, anomain, sakinkess, exploris, sakinkess, exploris, and excitation. The drug may also produce gestrointestimal disturbances such as names, contraining, addominal pain, dry mouth, amoreia, and headache. Hypertension, heart pulpitations, or alternative and the action of the production of the production of the production of the pulpitations, or alternative and the action of the production of the pulpitation of the production o

#### Administration:

When used chinally to text excessive dayster sleepiness associated with merodepay or obsertate sleep aparent, the recommended does it pipels, 200 to 200 mg per day. This may be given in a single morning application, or in two diskeld does (morning and at midday). When used to ordave physical performance, the pipel affective does is in the range of 100–400 mg. This is offen given at lasst 2–3 bases pine to athletic corporation. Note that side effects on the dose deportal it. is often advised to an taxing modalital on the low end of the effective doesage range, and increase by 50–100 mg per application and a contently leady is desirable.

The areas in which this drug is applied are vost, and essentially include any sport focused on acrobic activity or enderance. It may also work well with from a unities focused on short requar losses of strength or speed (namerobic activities), such as sheep partiag pole-unsities; or long-interings. Modefini is not a popular drug name gloodybulders, as it holds little direct value for hading muscle or reducing body fit levels. Some, however, do fail it to be an other contents of the content of the

## Availability:

Modafinil is presently available in more than two dozen countries. The drug is not highly diverted for black market sale, however, and is not a lucrative target for drug counterfeiting oncentions.

## Fat Loss Agents - Sympathomimetics

Adipex-P® (phentermine hydrochloride)

## Description:

Phetermine Induchdoride is a sympathominetic stimulant of the amphetamine family. Like other amphetamine derivatives, it is categorized as an amonectic (apperties appressing) agent. Phetermine is commonly prescribed as a weight loss aid in obese patients. It is typically used for short periods of time (loss than 12 weeks), and as an adjunct to support and orangoing coercise and dieting regimen The main focus is to curb the desire to est, thereby reducing the total caloric intake. Although the data seems to vary from trial to trial, much of it supports at least a modest additional loss of fit mass with the use of phentermine hydrochloride. <sup>506</sup> Additets and bodybuilders use phentermine hydrochloride for the same purpose, typically when weight loss is required for physique remodeling or competition.

#### History:

Plentermine bydrochieride was first irroduced to the U.S. drug mutet in the 1970s. Buse plentermine was suitable in the U.S. as fir those as 1999. Plentermine flued tope born used as an appetre appressure, although the most notable attention to it came in the early 1990s, when the drug was sensestful paried with furtirantie carring destrates, Investigates that shown that this type of drug corribation was actually more effective at promoting weight too the time of the content of the process possible of the process of the proce

### How Supplied:

Phentermine hydrochloride is most commonly supplied in tablets and capsules of 18.75 mg and 37.5 mg each.

## Structural Characteristics:

Phentermine hydrochloride is a central stimulant and indirect-acting sympathominetic of the amphetamine family. It has the chemical designation 2-methyl-1- phenylpropan-2-amine (2methyl-amphetamine).

#### Side Effects:

Commo side effects associated with pleatermine hydrochloride include incomia, nersued blood pressure, minishility, neurosusses, and explorie. Less commo side effects include vision distributors, reduced blishi, confusion, diarbate, diziases, day month, handeshe, rangale hearthest, meanes, vonnifige, rath, and tiredenses. Presertemine is a CNS stimular with potential for fatal overdrots. Signs of overdrots may include rapid breating, force, such as the confusion of the confusion

#### Administration:

For optimal effectiveness, phararmine lydrochricite should not be taken with food. The usual said did not is one capisal or this (6.75 gml day), administrate before or 1-2 hours after breakfast. For some pointers a half of a tablet (18.75 gml day) may be adequate, while in other cases it may be adsolvable to give a find of a nabel (18.75 gml day) may be adequate, while in other cases it may be adsolvable to give a find of a nabel (18.75 gml twice day). When taken more than once per day, the second does should move be taken within 4-6 hours of skeep. The day is givenly used for 5-4 weeks as a trine, with length entanties of thereup, day of the control of day in a sinite observation and the length believed of side effects as the does exclude show of the normal flearnoset; more

### Availability:

Phertermine hydrochlordie is available in a number of different countries. It is not widely counterfeited. U.S. residents would not be advised to order the drug from overseas, however, since phertermine is a schedule IV controlled substance and carries similar legal restrictions as Valian and anabolic steroids. Many U.S. doctors who specialise in weight loss medications will readily disenses phertermine for cortrolled periods of weight loss.

### Albuterol (albuterol sulfate)

### Description:

Abatord saillate is a selective beta-2 adveragie agonis, very similar in structure and action to christorate United centrateural, bowers, althered is readily subtile as a prescription dung in the United States. It is also sold as substamol in a nurber of other countries, which is another generic manner for the ding. Abstracted is most commonly board in the form of a rescue influx, which is designed to disperse a measured amount of the drug mirediately and directly to the broachin these in times of circle (softem) antack.) This form provides the least amount of systemic date gather these in tests of circle (softem) antack.) This form provides the least amount of systemic date gather days possible, which is great for minimizing unusual systems does of the drug. These are the subject of retreets in the body-shalling and althoric communities, and they can provide significant bets-2 similation and measurable fit has throughout the body gene the right conditions. Now that a more comprehensive discussion of the benefits, activities, and side effects of bets-2 agosist drugs can be found under the clearband of large profits.

### History:

Abators shife was structured to the U.S. ding matter in 1980, old under the Vertoin brand rame. Abators shife has goon to be one of the mes opported rings in bistory for the menagement of acute asoften matters. As a result, many other companies have insected in the market. The elementary the elementary the control heard many is sill anables in the U.S. bower, the FDA has also also proposed a variety of other generic and heard mane from of the drug. Abatorsh silled the silled proposed as the proposed as a single proposed as a proposed as a single pro

#### How Supplied:

Albuterol sulfate is most commonly supplied in oral metered dose inhalers and tablets of 2 mg. 4 mg. or 8 mg each.

#### Structural Characteristics:

Abuterol sulfate (sabutamol sulphate) is a short-acting ,2- adrenergic receptor agonist. It is a racenic drug with the chemical designation (±) a1-[(tert-butylamino)methyl]-4- hydroxy-mxylene-a.4-ioli sulfate (2:1)(salt).

### Side Effects:

Commo side effects associated with alturered saftie irelate headerle, dežizass, glightheadenes, sommit, neuron, arcousses, souring, manes, orning, dimbra, and dry morth. Less commo but more serious adverse events relate altergis reactions (rush, hote, souring) of the jax, tongs, or fixer, or of fifted by bearings, clear pair, high blood pressure, and iregular hearboat. Abstends safties is a CNS stimular with potentia for faul conclude. Sign of conductor my kinds and pair beartings, blood pressure ingalants, irrigals hearboat, unconcisuousses, terribing, shaking, paria; extreme realissness, and source most avairation or distribes.

#### Administration:

The usual starting dougs for adults and children 12 years and older for the immegeneet of authins 2-6 an given or four inner per day. When used (off-likels) for fall so, an efficient done of historiol usually starts in the ranges of one to two 4 mg tablets per day (1 tablet X 1-2 applications). This is often increased slightly as the user becomes accentated to the day, perhaps to 4 mg three to four times per day. Individuals very sensitive to the stimular side effects of freat against usually start with the bower-dose trap tablets. Eith Endmittation intervals are spread out as everly as possible, so as to prevent overlap and sustain acche concentrations in the blood for a man chi of deep use possible. Althose and bodylatation in temperature with use of the day may include that probability the proposition of the side of the proposition of the starting of the startin As is noted with all bets agonitis, tolerance to the thermogenic benefits of this ding tracks to develop quick). If it is usualty noticed by the body temperature tenturing to normal pretented levels. Due to the potential side effects of fines drugs, it is not advised to will usually opt to disconstruct the damp for some time of weeks or frongs but let the body nestore is normal bets-adveraging receptors concentrations. More recently, the artifasturine Zaden (destribed) in boscome popular, which is a potent quegation of best-adveraging receptors, optically bets-2 receptors. This medication may enhance the temporarie effects, optically also also concerned to produce the scale exception of the collection of disconstructions of the collection of effects.

#### Availability:

Albuterol is a widely available and very cheap medicine. Counterfeiting is not a strong concern with this medication.

## Clenasma (clenbuterol hydrochloride)

### Description:

Clebaterol hydrochérick is an arti-softem medication that belongs to a broad group of drugs known as reprodumentatic. These drugs affect that symptotic reviews open mis a walk araber of ways, length medicated by the distribution of admonoscipers. There are sufficiently a superior of the distribution of admonoscipers. There are admit no beat and first endemapying they post nather. Desputing on the special efficient of these agents for the various receptors, they can potentially be used in the returnet of conditions such as suffern, legeratories, conflowescaler door, and printings, ingrain bandences and amphibition shock. The text Geochemic and Gillions The diverse many in which these often sufficie the bond doors ago of plot of describes put foresee many in which these often sufficie the bond doors ago of plot of describes put the

Most of the actions of catecholamines and sympathomimetic agents can be classified into seven broad types: (1) persphered excitatory action on certain types of smooth muccles such as them is bold vessely supplying the 8th Askey, and muccous aperiphend thibitory action on certain other types of smooth muccle, such as those in the wall of the gat, in the boundain tree, and in bod vesses supplying the 8th and muccle; (3) a cardiac excitatory action, responsible for an increase in heart rise and force of contraction; (6) metabolic actions, such as an increase in the net of glozognoshis in liver and muccle and liberation of free fatty acids from adjuses tissue; (5) endocrine caterius, such as more admitted free fatty acids from adjuse tissue; (5) endocrine caterius, such as notabilities of plays severition of justili, reminand palutary hormous; (6) CSS actions, such as neighostory stimulation and, with some of the drugs, an inverse in weldpless and psychostory artificial and activities in applicit, and (7) propagate, actions that result in either inhibition of secretariolism.

Chebiterol Inducedarida is specifically a solution beta-2 sympthomizate, primarly affecting only one of the time subsets of bette-response. Of practical intensit intensit in the fast that this drug liss little beta-1 stimulating activity. Stace beta-1 receptors are closely find to the candiac effects of these against, this allows clothaterol legislactivities to reduce recently active and the standard states with the state of the standard states. The standard states with the daily offent of the compared to non-section beta against. Gittel attales with the diagnose it is extensively effective as a bronchodiant, with a low level of near complains and high patient complaine. Chebraton legislactivities also exhibits an external high patient complaine. Chebraton legislactivities also exhibits an external high patient complaine control of the complaints of the complaints

In animal studies cleributerol hydrochloride is shown to exhibit anabolic activity, obviously an attractive trait to a bodybuilder or athlete. This compound is additionally a known

themagane; with ben-2 against like clumbated lydurchiede shown to directly strainlate fat cold and accentrate the breakdown of righterious to form free lay such. In effects, of the area makes clearbared lydorchiede a very popular fat loss drug arrong the strain makes clearbared lydorchiede a very popular fat loss drug arrong free fat loss of the contract of the contract of the contract process of the contract of the contract of the contract process of the contract of the contract process of the contract of the contract process of the contract of the contract of the contract process of the contraction. We also an appear then placed to the contract process of the contraction. We also an appear then have for clothest as to whether conjust of the broadcalistion. We also an appear then has been as for defents as so whether conjusted for broadcalistion. We also an appear then has been and to effect as also whether the contract of the contraction of the contract of the contraction. We also an appear to the free fat it builds marked, and use clearbared lydorchieder regularly as an off-section of adjusted analysis. On other, the MSSE report is confirming that addies a low seader valuable time and money on drugs that do not build market. The debate over clearbared lydorchieder's pertain analysis and the contracts today.

### History:

Chebaseo luptochieris Inu born svalishie as i berochollator for decades and a widely used in mung parts of the world. Altough the as god soldy-record and approach in a wide number of other countries, this compound has never boen made available for human use in the Utael Status. The feet that there are a marrier of similar effective andream selections aboutly approach by the FDA and oxidable may have something to do with this, as a travelagoriage the emperor of the new date approach goods. Because of the feet of the selection of the feet of the feet of the selection of the feet of the selection of the feet of the selection of the feet of the feet

#### How Supplied:

Clembuterol hydrochloride is most commonly supplied in oral tablets of 20mcg each. It is also supplied in oral syrups, injectable solutions, and for inhalation use.

#### Structural Characteristics:

Clembuterol hydrochloride is a long-acting selective ,2- adrenergic receptor agonist. It has the chemical designation 1-(4-amino-3 5-dichlorn-phenoli-2-(terribut/damino) ethanol

### Side Effects:

The possible side effects of clerbaterol hydrochloride include those of other CNS trimutars, and radde such occurrences as shally hands, incrunia, wenturing, increased blood pressure, and nauses. These side effects will generally assisted after a week or so of tues, once the term becomes accustomed to the drug. Clerbaterol hydrochloride is a CNS strainflaw plotterial for fail overtices. Spay of oversides may leaked rapid benefities, libide greaters, and the procession of the contraction of the contract

#### Administration:

When used for the munigement of radius, the most common claimd dose for adults is 20mg (1 table) visc pare day. Some paristent require up to diving; (2 tables) to tope of day. When using the drug (of-liable) for physique- or performance-orinaring purposes, proceedings of the process of the contraction of the cont

The drug will usually elevate the body temperature shortly after therapy is initiated. The rise in temperature is commonly .5 to 1 degree, sometimes a little more. This elevation is due to

one's body harning excises energy (draghy from 6t), and is usually not unconfroble. The marrier of connection days electrated pulsachietis is now used is usually dependent on the response of the individual To be clear, the atthictive benefits of this dings will only last for a limited time and then diminish, largely due to best-receptor devengation. By most accounts celebrated phytochietist seems to work well for approximately 4 to 8 weeks. During this proof, uses permally morned their body temperature or regarder tests. We are given some level of assurance that clerbatered hydrochietis is working by the temperature electrists. Once the temperature draps below to mental, receptor deveningations has reception to the control of the proposal proposal

Many bodybuilding competions orhance the fit barning effect of claritateral hydrochleride with the use of additional substances. Many have commended that when the drug is corribined with figured hermones, specificably the posteriol (Systeath), the dismergatic effect on the individual claritater and the control physique from the drugs. A claritater of the individual claritater a much more total physique from the drugs. A claritateral principal control and analysis of centre of the control when surge growth hermore, which is believed to orthoch each dependent and analysis of effect of HGH theory. Leady, knotifien has also been a popular adjust to claritate only a section of the control physical claritater in the trapelation between Leady for the control physical claritateral control and the control physical claritateral control physical claritateral control physical claritateral control control physical claritateral control physical claritateral control physical claritateral claritateral control physical claritateral claritateral control physical claritateral claritateral claritateral control physical claritateral clarit

### Availability:

Cleributerol hydrochloride is readily available on the international market. Although it is usually a very cheap drug in common source countries, allowing black market dealers ample opportunity to obtain legitimate drugs to divert for sale, cleributerol hydrochloride has been the subject of flow-level counterfeitine. A flow thines are important to note:

Clembuterol hydrochloride is not produced in the U.S., so avoid anything bearing a U.S. commany name.

Cleributerol hydrochloride should only be trusted when found with a proper brand name from a foreign drug maker. Spiropent, Novegam and Oxyflux from Mexico are the most common products in the U.S.

From Europe, the brand names of Spiropent, Broncoterol, Clenasma, Monores, Contraspasmin and Ventolase are popular.

Bulgarian cleributerol hydrochloride is also found commonly, but so are counterfeits. This is a slightly higher risk form.

### Ephedrine (ephedrine hydrochloride)

#### Description:

Fielderine is a structure thrus plan belongs to the group of medicine known as expendentiments. Secondary, it is not but multi-may float and best abovers; grainer (nour movements electrical parties) and major and from a formation of the contract of the composition of the contract of the composition and the grainer to make of the composition, a strong endogram, and major and contract the contract of the composition of the contract and contract of the composition of the contract and contract of the composition of the contract and contract contra

the force of skeletal muscle contractions.

### History:

Epichetine is a fairly off medication; and has been used in the United States for a number of model applications over the years including that of a stimular, appeles appressing, decongular, and hypotenson treatment associated with amendment fooling it is approved as an over-the-courter medicine, and also are gaining in recent press, in foresect, the contract of the contract in the United States are gaining in recent parts, thorever, that we seeker. Controls over-applications in the United States are gaining in recent parts, thorever, that we can avoid the case of the contract product, undergoond remnifications have been able to easily obtain it. A trend involving large volume ratal practices for OTC ophodrine products had been developing and must gates have recopcided with legislation cortrolling the safe of precursor materials like epichetine. In 2006, a federal law was prosed father restricting the record leeping requirement and saudable safe charmles for specifies in the United States. With the widespread accesses of metamprications contracting the correct leeping requirement and saudable safe charmles for specifies in the United States. With the widespread accesses of metamprications and calcium (and related curva), some total records of a state of the contraction of the contractions. In specific district resultation, as followed to the contractions of the contractions of the contractions of the contraction of the contraction of the contraction. States of the contraction of the contrac

## How Supplied:

Ephedrine (as ephedrine hydrochloride or ephedrine sulfate) is most commonly supplied in tablets of 25 mg or 50 mg each

### Structural Characteristics:

Ephedrine is a sympathomimetic amine related in structure to amphetamine and methamphetamine. It has the chemical designation (1R,2S)-2-(methylamino)-1-nhenylnorous-1-ol

### Side Effects:

Enhedrine can produce a number of unwelcome side effects that the user should be aware of. For starters, the stimulant effect can produce shaky hands, tremors, sweating, rapid heartbeat, dizziness, and feelings of inner unrest. Often these effects subside as the user becomes more accustomed to the effect of this drug, or perhaps the dosage is lowered. In general, those negatively impacted by caffeine would probably not like the stronger effects of ephedrine. The mental and physical state produced by this drug is also quite similar to that seen with cleributerol, so those who find little discomfort with that treatment should (presumably) be fine with this item (and vice versa). While taking this drug one may also endure a notable loss of appetite, usually a welcome effect when dieting. Ephedrine is in fact a popular ingredient in combination (prescription) appetite suppressants. The user may further notice headaches and an increase in blood pressure with regular use of ephedrine. Those suffering from thyroid dysfunctions, high blood pressure, or cardiac irregularities should also not be taking this drug, as it will certainly not mix well with such conditions. Ephedrine is a CNS stimulant with potential for fatal overdose. Signs of overdose may include rapid breathing, blood pressure irregularities, irregular heartbeat, unconsciousness, trembling, shaking, panic, extreme restlessness, and severe nausea, vomiting or diarrhea.

#### Administration:

The primer application for epithesise among bodybaldem and address (off-blob) is that of a central (fish-sols again. Here, the infinited and generally take fish ang is for trusps per dyduring a dieting phase of fraining, at a dosage of 25 to 50 m gper application. The widely touted stack of epithetic (25-50 mg), calliface (200 mg), and aspire (100 mg) (ECV-30) is shown to be extremely potent for fit loss, and is more commonly applied thus epithesis salme. In this combination, the epithesis and cultifice both at an article themrogensternians. The added aspirin also belays to right inpostage is by blocking the incorporation of acceta to first pass, a fit. Buildem upon an increase in body personne as a marker than concern the start of the added to the control of the combination of the combination of the combination of the combination is taken ? to 9 times daily, for several consecutive weeks. It is discontinued once the user's body temperature drops below to mornal, a clear sign free drogs are on longer working as desired. A breach of at least 4-6 weeks is usually taken so that this stackmy over eagint work at no sprimal level. Fipedarie is also used by some competitive athletes (reclaim geometitives) as a striniart before worknots or competition. The reading (digit) strength and energy increase may impose anaerobic performance and weight totals on major life. On this same rock, it is also believed by some to provide a mental edge, making the use rome energy fared abbret and to concentrate on the tasks ahead. A pre-event dose of 25-50 mg of epidedries is pipeally used for this propose. It is important to note that focus copraint is not used continuously as a pre-worknot or pre-event stimulart, as its effect wall drinninh as the body becomes accessment to the first, jun most statescape, these used value the order only 2 or 3 times per work, usually on flowed days personally "important" is individual would also be visite to have considered to the control of the formular fall for seven the control of the property of the property of the locations provide the optimal effect formular fall for seven the control of the formular fall for the property of t

### Availability:

Ephedrine is widely available in the U.S. and in a number of countries abroad. It is not commonly a target of counterfeiting operations.

### Meridia® (sibutramine hydrochloride)

### Description:

Shuramine hydrochloride is a selective serotoria and nonadrurulin re-spaske sibilitor used for the medical menagement of obesity. The pharmocetals is strended to be an adjust to a reduced calarie diet, which will help increase weight loss compared to that achieved with medicing food intime hydrochrolide is not advertised as a repail acting quite drug, but instead one that fasters slow, safe, and steady losses in fixt mass which are mutiated lotters better.

Sharamin bydecchariae centra a weight loss effect frought two distinct mechanisms. It has a marked ability to superso apprehe. Darig more stades, periors would reduce their distinct constraints and the survey include by as mechan 1.00 calories while taking this drug. Well in addition to a effects on calorie intale, substancine also straintside metabolism and abily calorie expenditure. As single 10 ray door law to be confirmation of a stronger board intention for all by a 150%, an their dealers are supersoned to be confirmed to the calories of the substancing systems, manyly brought he address apport of their 3 recopers actuation. While the adverages is specific manyly brought he address apport of their 3 recopers actuation. While the survey is supersoned to be supersoned to the survey of t

To get a betre idea of exactly how well sharamine lyducchiaride works, we refer to some of the clinical statistics on this agart. One meetingtion was contacted at the Kansas Fornation for Clinical Pharmacologie in 2001. Here a parcy of 22.2 chose patients were given their 20 mg of harbaminist or glocked once daily for 20 whose ly the conclusion of the state of the contraction of the contrac

## History:

Sbutramine Indrochloride received Food and Drug Administration approval for sale as a prescription weight-loss agent in 1998. It was developed and marketed by Abbott Laboratories, which sold the drug on the U.S. market under the brand ramm Merdia. The company also sold the drug in many international markets under the name Reductal Sbutramine control of wal influed beared of sales in the U.S. as it was removed from the market in October 2010 under FDA pressure, citing an increased incidence of adverse cardiovascular events. Abbott has since also withdrawn sibutramine from many markets worldwide. Note that sibutramine remains classified as a schedule IV controlled substance in the United States.

### How Supplied:

Sibutramine hydrochloride is most commonly supplied in capsules of 5 mg, 10 mg, and 15 mg.

### Structural Characteristics:

Sibutramine hydrochloride is a centrally-acting serotonin-norepinephrine reuptake inhibitor structurally related to amphetamine. It is chemically a racemic mixture of (+) and (-) entrationers of 1-(4-chlorophenyl)-N,N-dimethyl-a-(2- methylpropyl)-cvclobutanemethamamine.

### Side Effects:

The most common side effect with shatumine is an incruse in blood pressure, a runt that contrainfacture is use in pratest with high bode pressure or other enrifloweacher issues. Other common side effects include day mouth, sleepissorses, ritability, back part, storneth report, and conseptions, and of which inted to become reduced in imaginate as the one immediately famy of the true serious side effects or symptoms of toxicly occur, including exclement, realissorses, loss of consciouses, confusion, against weakness, therein, clammisses, mpil hearthoat, lurge propils, voniring, difficulty breathing, clear pairs, swelling of feet, ankless or legs, laring, discirational, depression, high fever, op pain, termor, or excession sensuing. Note that hereused inclease of caudiouscular overs has prompted the removal of this product from most markets. It is to large crossisted as saft product by

### Administration:

Sharamine Inducedurials is used for the management of obesity, rekading weight loss and marienance, and should be used in conjugation with a reduced-caber due fit. Bits drug lass been used with patients who have additional weight-related risk factors including controlled hypertension, disbeloss, and dop-splowing (high closlesston). The recommended starting domes for more patients is 10 mg corect day, which is to be algoined sprands to 15 mg affect to design for more patients in 10 mg corect day, which is to be algoined sprands to 15 mg affect commended.

#### Availability:

This drug is subject to limited availability following an FDA supported recall in the United States, and subsequently its removal from many other international markets for safety pressors.

### Zaditen® (ketotifen fumarate)

#### Description:

Kotofies is a artifistantive due that is used for the toutment of general allarge symptoms, contain alongs' contributions (relating conjustation), and the management of softens. When used for authen, the drug is not reguled as effective for tenting an immediate stack (it is not a rapid benechation), based, over time is use is associated with a relaction in the foregament, duration, and severity of stacks. It is usually prescribed as a ways is tercase the efficacy of other attention mediations. Likewise, dectofies firmative bussless specifies and cacking soften mediation program, and not replace the prescribing of immediate rescue devices such as an admit solder or budsless.

Kotatifon firmarato alloviatos alloras currentarno ha blackina historino H1 recontare a

reconstant annume activates among 3-julyacous 12 julyacous 12 julyacous 13 properly that is common to though of the arthbasine class. Its second and very unique mode of action, however, makes it useful in the treatment astime. Kortelin famunta: increases the concentration of both achieverage increases in the body (especially believe? receptors). Dises that stimulate beta-2 receptors are commonly prescribed as broachchildious, used to increase after those the gasen activate the constriction caused by sustram. While patertailly efficacious abuse, one key therappeain effect of lactoffen famunta is to increase the sensitivity of the body to drugs of the beta again cits.

The beat? Exceptor upregating properties of kotofien farantee make this drug of interest to the holyhalding and athies communistee. This is due to the strong role of the beat? exceptor in supporting fal loss. Although not a strong fal loss compound by heaft when taken a long in the major like it clerksteet of kettler finnesse may be received the terrogate file celectrical kettlerfort finnesse may be received the terrogate fall celectrical kettlerfort from the fall resident of seat disease the because beat-of the telectrical kettler for the fall seater for the

The ability of lexitón to potentiae the efficies of behr.2 aguist drags has been demonstrated in a mibro of crisial statistics. For campie, one easy phebled at 1900 demonstrated that when lexitónic and cleabrated was taken lexitonic production of the behavior of the statistic desired compared to the season of the statistics of the season of the statistics of the statist

### History:

Kentólin was globalised as a preccipiton medication by Novarán. It is presently precibed for slavejes, allajego conflictos, and (most commonly) the munagener of adrium in more than three-dozen countries around the world. The most widely available beard name is Novaria's Zadlen, which is old throughts more parts of Energe and Asia. In addition to generic forms of the medication, dozens of other brand manse can be fourth in mung different markets as well. Kentofic finaments in approach of such in the United States, but currently only as an ophthaline anti-slavey solution (Zadisto), not an enal alsoysisstimu medication. The chooge of kentoficia finaments in the product is also too bo for it to be considered useful for any other (off labels) purpose. Given the ready availability of kentofic influence in convenience with contribution of the contribution of th

The UK guidens on the chiral management of asthm consider lextofien to be inflictive for the management of this disease. There is a shirthedly conflicting date on the potential usefaless of Lextofien farmants for this purpose, with some studies reporting positive results and others showing an inspiritant effect. In Attouculy review of the lamp plobled on the Cochrane Database of systematic Reviews in 2004 concluded that it appeared to have some usefaless in controlling softman and wheesing in many childres, but the variability of the disease and response to the drug meant that these positive results could not be generalized for all asthm matrices.

### How Supplied:

Ketotifen furmarate is most commonly supplied in tablets of 1mg. This dosage is usually expressed in terms of the base, so each tablet actually contains 1.38mg of ketotifen furmarate.

#### Structural Characteristics:

Ketotifen furnarate is selective histamine H1 antagonist, anti-allergic, and anti-asthmatic agent. It has the chemical designation 4-(1-Methyl-4-piperidylidene-4Hbenzo [4-Stevelohene-1]-(2-Hb) one furnarate.

#### Side Effects:

Common side effects include dry mouth, appetite stimulation, weight gain, dizziness, CNS

stimulation, and drowsiness. These side effects are all commonly associated with strong antihistamine compounds. In rare cases severe allergic reaction on the skin or a urinary bladder inflammation called cystitis may occur.

#### Administration:

When used to reduce the frequency, duration, and secority of asthm attacks, keetafin finames is usually instant at a dosp got of my tope or pdy (or pg. 801). Horocostey, this may be increased to a minimum dosage of 2mg tokee per day (ong textal). Bodylvakkees and attacks well comment, or an dosage of 2mg tokee per day (ong textal). Bodylvakkees and attacks well comment, or and sought of the property of the per day (ong textal). Bodylvakkees and attacks well comment, or and sought of the property of the property of the property of property in the property of the property of the property of the property of the head-of-property of the property of the head-of-property of the property of the proper

### Availability:

Ketotifen famarate is widely available, and is sold under numerous brand names in many countries. Large scale counterfeiting of this medication is currently not known to be a problem.

Fat Loss Agents - Thyroid

Cytomel® (liothyronine sodium)

## Description:

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Bodybulkers and arthletes are attracted to forfopronies softem for its ability to increase metabolism and apport the breakdown of body fit. Most of shot in intied during context preparation or perods of "cating", the drug is usually said a significantly aid in the loss of fit, often on higher bods of calcer intelled mountain usual to significantly aid in the loss of fit in the context of the significant intelled and the significant in the loss of the significant intelled to the significant intelled and the significant intelled and the significant intelled and homeous like following soften in terrease the analose effect of seconds. While in these provides the significant intelled and the significant intelled and the significant intelled and provides the significant intelled and the significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant significant intelled and the significant intelled and the significant significant significant intelled and the significant significa

### History:

The first medication that included T3 was technically a thropic curract, first given to a patient with moucherm (as indisorder associated with prophropishing in 1891; 600 Narmal Hydroc with moucherm (as indisorder associated with prophropishing in 1891; 600 Narmal Hydroc with valve) used in medical particles from either mole) particle from the mole) particle from the object of the through the mountain the same showly sard giving usey to new synthetic through medications, namely holtpromies confurmed to construents that proper admiral estants. Although holtpromies sodium and beothyroxies sodium and levoltpromies sodium and levoltpromies sodium and levoltpromies sodium and levoltpromies sodium are both without particles and about the first (including a least the sodium and levoltpromies sodium are both without particles and about the time (including a least through holtpromies sodium and levoltpromies sodium are both without particles and about to this (including including a least through holtpromies sodium and levoltpromies sodium

significantly smaller portion of the global thyroid market. Given is more potent and last acting effect, however, liothyrorine sodium remains a popular thyroid drug with bodybuilders and athletes. Cytomellië is the most recognized trade mem for the drug in the U.S., where it is presently sold under the King Pharmaceuticals brand name.

### How Supplied:

Liothyronine sodium is most commonly supplied in oral tablets of 5 meg, 25 meg, and 50 meg.

### Structural Characteristics:

Liothyronine sodium is a synthetic form of T3 thyroid hormone. It has the chemical designation i-tyrosine o-(4- hydroxy-3-indon)orm/h-3-5-diaglo-monosodium salt

#### Warnings:

FDA regimes the following black box warning accompany prescription fooltpoories sodian products sold in the US. Though with quoid lammen activity, almoe or toughter with other theraperic agents, have been used for the treatment of obesity. In enthyroid patients, does with the range of help hormound requirements are netflectives for weight reduction. Larger does may produce serious or even life-frendering munifications of toxicity, particularly when given it association with sympathomicisc aimses such as those used for their when given its association with sympathomicisc aimses such as those used for their

### Side Effects:

Side offets are generally associated with overdosage, and may include headards, intribility, neuroscursos, sussing ingular headards, incrused bowed medigs or mustatul impalaries. Overdosage may also induce sheek; and may aggroute or tagger angin or congastics heat falses. Centure overoposages to ladfornes obstantial mylorabot symptom neurally associated with hyporthosation of the obstance of the obstance relief and see effects in remaily case to introducing various or contract and post of the contract of overoposation of the obstance is neurally case to immediately reduce or discontinue therapy with follynomies sediam Acute musice overdose more held fett obstantial.

### Administration:

When used to treat mild hypothyroidism, the typical recommended starting dosage is 25 meg daily. The daily dosage them may be increased by no more than 25 meg every 1 to 2 weeks. The established mirretanence dose is usually 25-75 meg per day. Once a day admiristration of the full daily dose is usually recommended. Although forthyrorine sodium is first acting, its effects may persist in the body for several days after discontinuance.

The usual protecol narrog bodybalders and athless taking follyonine sodiam to accelerate fit loss involves instituing is sue with a dosage of 25 mgp er day. This dosage may be be increased by 25 mge every 4 to 7 days, usually reaching a maximum of no more than 75 mgp er day. As in a medical setting, the interest of this slow balding is to help the body become adjust to the increasing thyroid hormone levels, and avoid sudden changes that may initiate side effects.

Cycles of forhymnie sodiam usually last no longer than 6 weeks, and administration of the drug should not be halted abruptly. Instead, it is discontinued in the same slow marner in which it was infaited. This usually entails reducing the dosage by 25 mg every 4 to 7 days. This tapering is done so that the body has time to readjust its endogenous hormone production at the exclusion of therapy, and to avoid the orest of side effects.

#### Availability:

Liothyroine is an old and widely prescribed medication. It can be found readily in most areas of the world, and is sold in a variety of different brand and generic forms. Counterfeiring is not a large-scale problem It is important to note than one should never purchase an injectable form of this drug. These are generally used as emergency room products only, with potentially very diangerous side effects if misused.

### Description:

Leodipowies sodium is a synthetically mundicated from of the materal throad homose testinoidepowine (Fe-1) Tipotal hormose are primary becomes he regading the body's metabole rate, and play a valid role in the body's utilization of protein, till, and which is characterized by instillater turnal production of flypod hormose. This my number to feel with a marker of symptoms including best of energy, lettings, weight gain, but loss, and changes in shit netters. Leodoprotes sodium is considered a dow-seeding medication, and my take up to 4 to weeks before full florespeeck levels are nearbed in the considered to the seeding the seeding the seeding the seeding the seeding the considered to the seeding throad the seeding the seeding the seeding the considered to the sential formout foresame for most cases of therefore the seeding the considered to the sentantal formout foresame for most easier of the order of the seeding through the seeding the se

The action of two dynomics and min is very similar to that of the popular dynoid preparation Cytembell (buddyness) soddism (Cytembell is slightly different infustance, however, being a sufficient levels of both 13 and 1-4 bytembell hormones; present in their body. 13 is considered the primury active form of flaptoid hormones; present in their body. 13 is considered the primury active form of flaptoid hormone, while 74 serves minkly as a reserve for 13, exerting most of its methods activity via conversion 130 in apprehend issuess. 13 is regarded as lawing an effect that is roughly four times stronger than that of 1-4 on a minglasm-few-inflagments is Lakewise, Cytembell is considered to be a none potent form for minglasm-few-inflagments is Lakewise, Cytembell is considered to be a none potent form for the contraction of the contra

Leodiposise sodiam is valued by mmy drug-using utilises and bodyheidess for it a dilip; to straintate the metallor anter and sproof the breaddown of Doby for stores. It is usually taken during a peried of calorie sentricine ("cating"), when the artiscitual is founced on fat loss or increasing much definition. It is often thought fur the use of throat drugs can support fit loss at a higher level of calorie raise fram would otherwise be possible without the drugs, adding to their proceder value among the commarkies. Analories sensors are generally used in conjunction with those hormous, but many before that the mathodiem have with the budge through the confidence of the distriction of the confidence of the

### History:

Leodynovies sodiam was the first orderic flyorid medication to be sod in the U.S., and was first introduced on morted in 1955 by First Laboratries as Significent. He drug has a long history of thempeate use in the U.S. and internationally, and for decades has been the most widely prescribed medication for the neutrneer of Jupophysics. The Synthroid brane has habstrashy been the most successful, with figures estimating that it retained 85% of total locally ordering the configuration of the soft of military international processing of the soft of the configuration of the

The Synthesis band lead'has a beg and at times controversial bissing. We for many years after its seption by Plett Lubouriers, Synthesi elegoid a structure morepely on the boothyrouse sodium musket. Genetic medications finally began taking a large share of leosthyrouse sodium anske gaing in the 10 Blook. In reports, Plett Lubourioris finaled a study at the University of California in 1986 which attempted to demonstrate that Synthesia had a higher thereprese take than its general contempts. The study was completed in that in higher thereprese takes than its general contempts, the study was completed in execution of a clause in its content requiring company approach before the university could publish its satch, A algabath one ver in publishent ensual Even after Fat Lubouriers was sold to Boots, and threateler Rosts sold to Rost, and thereafter Rosts sold to Rost, and thereafter Rosts sold to Boots, and the sold to the

#### How Supplied:

Levothyroxine sodium is most commonly supplied in oral tablets of 25 meg,50 meg,75 meg, 100 meh, 125 meg, 150 meg, 200 meg, and 300 meg.

### Structural Characteristics:

Levothyroxine sodium is a synthetic form of T4 thyroid hormone. It has the chemical designation L-3 3' 5 S', tetraiodothyroxine sodium salt

## Warnings:

FIA requires that the following black hox saming accorptory prescription followings softum produces both in the U.S. "Thyroid hormones, including bothsymous notion, either above or with other therapeutic agents, should not be used for the treatment of obesity or for weight loss. In cultipoly patients, does within the range of childy hormonal requirements are netflective for weight reduction. Larger does may produce serous or even the threatment metalstations of cookey, practically when given in association with symptomic mixer and analysis.

### Side Effects:

Side effects are generally associated with overdosage, and may include beadurel, intribulity, memoraness, sowingin ingular bendurel, increased bossel motificy or mustally interpretation of the production of the production of the conception bear little effect. Overdosage may also induce stock, and may agground on trigger anging or congagistic heart little. Choice incresponane to levelopments coline will produce spraptures mornilly associated with hyperhysidism, or the overpredaction of manual thysical hormans in the body. The occurrace of over-quotaristical date decisis normally associated with foreign the production of the decisis normally associated with hyperhysidism, or the overpredaction of decisis increasing least to immediately reduce or discorning therapy with levelsproxies sodium Acute mussion contribute may be life throughout the configuration of the contribute of the configuration of the contribute of the c

#### Administration:

When used to treat mild to moderate hypothyroidism, the average replacement dose of levoltyroxine sodiam is approximately 1.7 megkgdys. This equates to 100-125 megday per day for a 154b adult. The fill therapeutic dose may be given from the onset of therapy in otherwise healthy adult patients. Note that due to the long half-life of fevoltyroxine, the peak therapeutic effect at a siven dose may not be achieved for 4 to 6 weeks.

When used (of-libed) to accelerate fat loss by bodybadles and affates, the typical protected moless show badlay of the closure is not the body has made in two capita for the changing flyorid hormore levels, in rinductal will generally and with a low change of 27 the changing flyorid hormore levels, in rinductal will generally and will not be completely 250 mag. It is imported and usually be in the mage of 1001–150 mag and will medy second 250 mag. It is imported to nemerice that flyorid drugs are strong medications with significant side effect potential. Canation addicables all between the two consecutions of the contribution of the cont

#### Availability:

Although levothyroxine sodium is a widely manufactured drug, it is not as common on the black market as the stronger thyroid drug Cytomel®. Large scale counterfeiting does not appear to be a problem.

## Fatloss Agents - Other

#### Description:

DNP is one of the most controversial drugs in use by bodyhadden. This agert is not soil for human use anythere in the world at this firm, but is really somable as an indistant charrical. Armay other things, it is used as an intermediary for the production of certain dyes, for photographic development, as a fragical, in wood pressure—sentered to person ording, and as an isoccicide. It is technically classified as a poison. Although quite incorporation with the six of strong indistrictlerized uses, this cherical uses soil during the ent of potent medicine as a diet drug for humans. It is this property of dinitrophenol that remains of interest to some bodyhadders today.

Distrophenoi induces weight to sky uncoupling oxidatine phosphorylation, threely muckuly increasing the methods in and an lody temperature. While this is an entermely efficient way of producing mpid weight loss, there seems to be no ceiling to DNP's temperature increasing effect. Herein is perhaps in sort adiagnossi nat; it may allow body aptreparture to rise to level that can be duringing, even final. Whire Carl Matriereg much perhaps one of the enterts and most firms quates about the startege levels in the 1998, when he told of a physician who was "Remdy cooked to death" from using I. This was far from an isolated care, and deaths societied with DNP lime contrained over the deades. For exempt, a case, and deaths societied with DNP lime contrained over the deades. For exempt, a locate of the contrained over the deaded. For exempt, a locate of the contrained over the deaded. For exempt, a locate of the contrained over the deaded of the locate of the loca

### History:

The firshess properties of DNP were reportedly first moticed during World War 1, when convergift rum working with DNP in mattines plants started being substantial measured or weight. It did not take very long for this chemical to be identified as the cause. Soon after, it was packaged as a ding product, B. 19153. The more than 100,000 American lead about your 'poster motions' remoties that reckedd DNP in fact, DNP was the list synthetic drug that was occur used for weight enables in the cases. While the was solid, it was being worked as destricted as a new, soft, and effective way to get after typical trained from the DNP propularly, the date could be Dord in pleasured as a leason to the could be an expenditure, the start could be found in pleasured as a consequence of DNP propularly, the date could be Dord in pleasured as a leason the court of

While the drug may have worked for the intruded purpose, it was also introduced at a time before government review and approad of day assley, in the regular DNP hall some very strong shortcomings, and it dish't take leng for reports of side effects to began pouring in the day. The contract of the contract of the contract began content is been as for severy in the day. Durnerson steps of DNP bridged contained began contents in form as for severy France and hall, it was said to be happening with doses as fird as 100 mg dishy when taken for burg periods. Reports of times retirens ignificantly experienced from the contract was also highly unforceable saskly reports, the dang was soon paled. By 1938 it was off the sask highly unforceable saskly reports, the dang was soon paled. By 1938 it was off the sask to be a several contraction contraction. The contraction of the contraction

Audre's Note: I was bestart to over include a profile of distrophenol in this book, for fast in rigil erface someone with otherwise may not here become about it to use. I But distrance I decided it would be better to taking the Institute I and the I may be the I decided it would be better to taking the I besterial individual of the I besterial individual in an arealization may be an arealization may not an arealization may be made to the I besterial individual individua

#### How Supplied:

DNP is not supplied in a form prepared for human or veterinary consumption. It is available

as a research or industrial chemical only Structural Characteristics:

DNP (2.4-Dinitrophenol) is a cellular metabolic poison with the chemical designation 1hydroxy-2 4-dintrobenzene

#### Side Effects:

There are many potential side effects associated with DNP use including increased heart rate. increased breathing rate, nausea, elevated body termograture, incoming profine sweating rach skin losions docroacod white blood coll count, estaracts, come, and doath

#### Administration:

DNP is not approved for use in humans. Prescribing guidelines are unavailable. A common dose used among bodybuilders is reportedly 2mg per kg of bodyweight per day. This calculates to a dosage of 200 mg per day for a person of approximately 220 pounds of bodyweight. Note that this population tends to retain more lean muscle mass than the average (sedentary) person of the same bodyweight, which may substantially after the results and side effects of a given dosage. Admittedly, fat loss due to DNP use is highly rapid and extreme, with some people losing as much as 5 to 1 pound of fat weight per day. This can equate to a drop of 15 or 20 pounds in only a few weeks. Given the high risks associated with DNP use however it is usually taken for only a few weeks at a time. The strong incidence of side effects is also regarded as an indicator that the drug should be discontinued immediately. Note that most experts regard DNP as a drug with inherent dangers that far outweigh its notential benefits

### Availability:

DNP is not available as a human or veterinary medication in any part of the world Availability of products intended for human use is entirely in the underground realm, where products, dosages, and safety are not the subject of government approval.

## Lipostabil N (phosphatidylcholine/sodium deoxycholate)

#### Description:

Lipostabil N is an injectable medication that contains phosphatidylcholine (PPC), a natural phospholinid. Sodium deoxycholate (a bile salt) is also added (among other ingredients) to solubilize PPC in water. It was originally developed as an intravenous solution for the improvement of serum lipids, reduction of arterial plaque, improving liver values, and the prevention or treatment of blood vessel blockages by fat particles (fat embolism). It is approved as an intravenous drug in a number of countries, mainly in Europe. Lipostabil has also had a very popular off-label use over the past several years, namely as a localized fat loss agent. Clinics in many areas of the world including Brazil. Europe, and the United States have actually marketed this as a nonsuroical alternative to linosuction. In ocent wars bodybuilders have been paying some attention to this drug as well, using it as a cutting or finishing agent.

The mechanism behind Lipostabil's lipolytic (fat loss promoting) effect is unique. Upon injection, the solution acts as a detergent, causing nonspecific lysis (breakdown) of cell membranes. 698 The bile salt sodium deoxycholate is actually believed to play an important role here, and is therefore considered an active constituent of Linostabil for the context of this profile (it is normally considered an inactive ingredient). During this process the fatty acids stored in the cell membrane are released, which includes arachidonic acid. This will trigger the inflammatory cascade, benefiting lipolysis (the inflammatory system can be a powerful remodeler of body composition) but also causing unwelcome pain and swelling. Phosphatidylcholine itself also triggers the release of lipases used in the removal of fat. 699 All of this works together to dismantle localized fat stores, which are removed via the liver in the form of gall acids.

### History:

Lipostable first appeared as a medication during the 1950s. Although not approved for prescription use in the U.S., it is approved for medical use in a muther of other contribution. The most propair brand name is Lopostable No. No Notements, although it is also found as Lipostable Ford and paying is Lopostable Post and appealent off this deep first loss is generally viewed as an "tid-label" use of the medication, although it undoubtodly remains a highly proposal factor in the St. Some experience of the contribution of th

Network I polysis (an expansion of some 280 decises workshield that apports this use of Ilgorability sports one 18,000 contest incuments without suspected adverse events. Additionally, Dr. Hosengechansthure, the medical director of the Austrian client Therapy Contro Bell Lockbiles, has reported on bifurbin and german glamant bandiences (unless of first stess) values after selectaneous I postable up. <sup>200</sup> to see 2 this new modes of first recording is causing for stain. The resolve were in few with Yue, chowing an absenual charge in few values. Although we do not have a gener deal of data on this off-black use of I-postable with or the fourth of general postable up. <sup>200</sup> to provide and usages site sing (or mattal dung 8 you will is quite self-lipostable in presently sold in Germany, Spain, Italy, Caech Republic Hope Kope and South Africa.

#### How Supplied:

Linostabil is most commonly supplied in injectable annules containing 5 mL of solution each

## Structural Characteristics:

The primary active ingredients in Lipostabil are phosphatidyl-holine (phosphatidyl-Ntrimethyl-trlanolamine) and sodium deoxycholate (cholan-24-oic acid, 3,12-dihydroxy-, monosodium ath

#### Side Effects:

Potential side effects associated with subcutaneous Lipostabil injections include localized swelling, redness of the skin, burning, pain, tendemess, and bruising. Systemic side effects are reported in approximately 3% of users and may include diarrhea, nausea, dizziness, and interconventual blooding. <sup>701</sup>

### Administration:

The typical practice for using this drug to promote localized fit loss involves a series of subctanances injections. A total dosage of 1250-2500 mg is often used, which equates to 25-50 mt. of injectable solution. This dosage is disided into 20 or more separate smaller rigications. These are spaced throughout the problemantic area (quite commonly the abdominst or thighs), and area all given during the same office visit or application period. The drue is not administered on a duly base.

Lipotable lipotrons usually cause a significant amount of fullmutation in the new, which may sake a worke of reage to fully adside. When the influmntation does subside, however, it usually unseek a noticeable amount of fit loss. In a clinical setting, this procedure is order repeated a few times, so as to sciept the new and acheise the desired level of fit reduction. The current guidelies set forth by Newtork Lipolysis call for an 8-week break between remntant pericks. When that on coatio of a clinical setting, it is usually alwards any pile find it simply two painful to use, while others seem to tolerate the whole procedure extremely week.

As for the ultimate question of how well it works, it is difficult to give exact numbers, as few clinical studies have been conducted on this use of the drug. The anecdotal feedback is mixed. Many neonle who try it report positive results nearticularly for the removal of those has stableon mean of fit interfering with muscle definion. There do not seem to be many reports of dimutin weight loss, lowers, nor does it seem to be the "pharmacustical spoatcient" that some clinic describe it to be Regardies, the reports of wishle improvements in the sand muscle definition are consistent and compelling enough to be given credit. For those externelly overweight, this product is not Richy to perform well, but as a finishing tooch it may beld value.

### Availability:

Lipostabil is not a controlled drug in the U.S. or Europe, and as such is fairly easy to obtain on the black market or via mail order drug distributors. "Mesotherapy" clinics selling procedures with Lipostabil are also fairly common.

### Growth Hormones & Related

### Geref® (sermorelin acetate)

### Description:

Semeroin is a synthetic analog of endogenous growth hormon-releasing hormone (GRBH or GRB). Semeroin is a portion of this polypetic hormone, specially consisting of the first 20 of 8.4 minto acid structure. As for more states very clearly, the biological activity of GRBH is to strained the synthesis of growth hormone, which occurs in the platting gland. Studies hower down, however, that the GRBH peptide can be partly functed without searchings (6.4 fill studing adilly <sup>100</sup>. 2004.) you sale-deeped based on this research, and states the fill biological activity of GRBH with regard to increasing the orthogenous endoubtrance precision factor (GRBH or of GRBH with regard to increasing the orthogenous evolution for the control of the co

Semredia, as an acette salt, is used in clinical medicine for two primary applications. The first is to diagnose primary deficiency. The procedure involves measuring the seman ground hormone response over a 1-2 hear window following a single IV infaisor of semmetin acetta. The infaisor hopolumbrome tassis provide-hormone assis dataly ideal practicable acreases in the GHI level. The second common medical application is the treatment of growth hormone deficiency in children. As with necessitient growth hormone deficiency in children, and the contributing growth hormone medications (HEAL) semantipolis, semmedia acettet can provide the benefits of assistant electrisms in GR. mixing entrance (ICI+1 (Instain) Rec Gooden Factor) organ articarceased lance haptic. "Semmedia" is a very speech acitig dang, and has no officio or production. ILIT (SMI, mathr. controls) discover, glescage, or dipoid hormone becks. "Official beach to its therapeute."

Addeds are interested in semmenia nectate for the same reasons they use recombinar human growth homes. Among other things, growth hormone has arbicle and aricatabolic properties. An elevated Gel Iscel may apport new muscle issue growth, and also enfances strength, energy levels, and corrective issues; Gel is also a potent modulator office issue. The issue poster modulator office issue Till seals on potent modulator office issue, Till seals on potent modulator office issue, Till seals on potent modulator office issue in the institute of poster and performance-orizontary properties of growth hormone are widely accepted by the polyholding and comprehen adultic terminally and a visible admentise to low does it is slowly gaining acceptance among the commanday as a visible admentise to low does growth hormone injections. It is also presently popular in anti-signing medicine, where again it growth confirmations are proposed and and declerate the polyholding and the formation of a posteriol off in decleration.

When comparing this drug to recombinant harman growth hormore, semmedia acetate does appear to be less effective under normal therapetric conditions. During clinical statics, fewer potents on average seem to respond floworably to therapy in contrast to somutropia, and for those that do respond the improvements are often less pronounced. <sup>706</sup> Sill, it would be a mixture to average the contrast of the contrast o

Instance to Contact Scientification accounts on a vision intemperate, Operate, 100 Contings, Stateshow Enter Man State Mannership accordance rursed in significant increases in height velocity in children with GHI deficiency. The Furthermore, these improvements seem to be well sustained after one year of therapy, 788–790. Anticolos to GHRH do develop in some patients during extended therapy, and may impair the potency of the dang. 710 The full biological relevance of this authority reservition to towards contribution under a

an atmosphastic moves, reministrated for the restaurch of GH deliciony in clubture only. Produced studies tenting adults with somotopuse are lacking, One inscitation looked at the effects of 2 mg sermerein acutes per day for 6 weeks in a group of elsem hashly delay men (agod 6 to 7) with two creditating field: 1 leek. <sup>17</sup> Wisson measures of body corporation and performance were recorded at the beginning and end of the study. As a result for future, there was a significant reasons in norticum [61] output, area under peak GH release, and GH peak amplitude. This was accompanied by improvements in credit measures of strength and endirance including uppirt on, who although press, and addormal cruzely. Many often measures of day for a ready statistical significance, reflect and produced and the contraction of the contraction of trains.

Another investigation booked at the effects of sermerelin acetate in a group of HIV+ man with ipodystrophy. The HIV Ipodystrophy tends to be characterized by the abnormal disturbation of face data and paperssed levels of growth humane. Daving his investigation, 31 men agad 18 to 60 years were given In gar of sermerelin acetate or placebo by subscriancous rejection twice daily for 12 weeds. The purpose your content of the study was a significant reacease in searm KiF-1 in the semenoism acetate group (104 right. we for girlt.). This was accompanied by a forebulk increase in land who mays (+9 g ber, +3) gand adcrease in any change in other health markers including blood pressure, cholestend, traffection and produced to the study of the produced to the produce

Semreina acetate could be visced as offering a themposite advantage over recombinar gowth homors in some cases, that it is loss likely to reals in GH escens This is due to the fact that it relies on the body's own homore synthesis instead of coagenous supplementation. Thus, normal IGAT is Endest in Eribin 18, kely be less at natural intito the growth homore stimulating effect. This such, homore levels are more easily controlled with semeration actue. Under normal confirms, while sememed in Produce significant clearations in GH and IGAT, these levels should not exceed the high end of the normal large. <sup>11</sup>Gene this factors, semember accurate my be a more contributed option for some pariers. Studies seem to support this notion, finding many of the same playiously and produce and the studies of the studies

### History:

Semretin actate was developed during the early 1988s, and approved for prescription sate by the U.S. Food and Drug Administration in 1997. It was intraducted to motert unter the band name Geref Diagnosis: by the international biotechnologies firm Serons. As the more implies, it was primarily developed as a diagnosis: bot II was specifically used for evaluating potential plattary deficiency in CHI production. Gene in effect on growth homoralesels, however, the day was also approved by the FDA for the transmert of CHI deficiency in children Geref was never which prescribed, however, especially for uses relating to children GHI deficiency, where it was never able to compete with sometings for scenarios, children GHI deficiency, where it was never able to compete with sometings for a children GHI deficiency, where it was never able to compete with sometings for a children of the GHI deficiency of the GHI developed the competition of the children of the GHI developed the competition of the competition of the pharmacentral agreement in the United States, their is no longer routhflet has a such pharmacy learning that the competition of the

#### How Supplied:

Geref Diagnosti: was supplied in ampules containing day lopolitized semmorfin acetate, equivalent to 50 meg of semmorfin. This was reconstituted with a sterile dihaert (also supplied) before use. Generic compounded versions of this medication typically contain between 3.0 and 7.5 mg of dry lopolitized semmorfin acetate in a multi-dose vial Reconstitution before use is also recurried.

# Structural Characteristics:

Sermorelin acetate is the acetate salt of a synthetic 29-amino acid peptide (GRF 1-29 NH 2) that corresponds to the amino-terminal segment of the naturally occurring human growth hormone-releasing hormone (GHRH) with 44 amino acid in esidues.

### Warnings:

Sermorelin acetate should be used with care in epileptic patients. Obesity, uncontrolled hypothyroidism, hyperglycenia, or elevated plasma farty acids may impair the effectiveness of sermorelin. Therapy should be discontinued in patients treated for childhood GH deficiency once the eninhyses have closed.

## Side Effects:

The most common side effects to sermorelin acetate therapy are injection site reactions such as pair, redness, and swelling. During clinical trials, this occurred in approximately 17% of patients. Less common side effects include difficulty swallowing, tiching, dizzincss, flashing, beached; neurose avonition aborted areas of tester softsesses and sleviness.

### Administration:

When used medically for the teatment of idepathic growth hormore deficiency in prophential clikines with growth findare, semented nactient is admissioned by selectaneous sipecina rat a dosage of 0.03 mg per kg of body weight cone a day at bodium; Inspiction sites should be notated to avoid irration or the buildigh of care times. When used to evaluate philatory capacity in adults, a single intraceous inflation of 1.0 mg/kg body weight is distinisated in the mereing after an overlight fact. This is followed by 60-120 minutes of

When used for physique- or performance-enturing purposes, sommorth meetate is glornly subcataneous spicetim. It is yielday allomationed at a dosage of 102 to 10.5 mg per day (200-500 mgg), which is glorn before skep. Statles, however, do suggest that the drug is more effective when given twice day. <sup>201</sup> Therefore, is a folm predented to disked the total dayl dosage into two applications, one in the morning and one in the evening. Cycles of sumerical acutes usually last between 8 and 12 works. Some arrivaging practitions was of practice the medication for much longer periods of time, however, and cycles longs 24-66 hormore belowd with deriven estimation and conditional changes.

### Availability Trends:

Given its low financial value on global pharmaceutical markets, sermorelin acetate is subject to limited availability worldwide. The preparations most commonly found diserted for bodybuilding use include those made by private compounding pharmacies in the United States, and those sold by gray market research chemical supply companies.

### Human Growth Hormone (somatropin)

#### Description:

As a man sagests, harm growth hormore is an important mediator of the human growth process. This hormore is produced endospassely by the arteries printing plant, and easily as expectedly high levels during childhood. In growth-promoting effects are broad, and cam be protect, carebodynic, level, and mean temperature of the protect, carebodynic, lept, and mean temperature of conceives trained see growth of consective issues. Although vial to early development, human growth hormore is produced throughout adulthood. It is level and bological in declares with age, but confirst to approximate the confirst to approxim

(rhGH) is biologically equivalent to human growth hormone (hGH) of pituitary origin.

In a medical setting somatopin is used to help treat a setting of health condition. It is most mostly presented in cases of difficulty growth disouless that are characterized by randfacter growth homorous production. While issually not difficult controls, countraries are set in the control of the control

Somatopin is also somatimes prescribed to healthy men and women who are aging Grooth hormore looks tard to decline as we get older, and many physicians before that is before the first production of the control of the

Sommton) may be given by either abestureous or inturnacedar signistics. Dering claimstaken, he phermoconkorte properties of counterpost flowings but methods of the were determined. When given by substanteous rejection, sommtonia has similar but modernally higher level of bisconsibility (19% vs. 68%). The rate of ting metablosin flowings both mostes was also very virinke, with counterpoin displaying that life of approximately 3.5 hours also very virinke, with counterpoin displaying that life of approximately 3.5 hours select a result of the properties of the proper

A species amplies of sometopina earlier planes a hormone with a discress set of effects. It is another to skeld muck, down to its results but the size and transfer of coll (processes referred to as Inperceptly and Inperplasis, respectively). The hormone also seems to more growth-promring effects on all organs of these local results for some production and the size of the size

Growth hormore has both direct and indirect effects. On the direct side, the IGH protein attaches to receptors in mascle, bone, and adoptore fissues, senting messages to support another and logical field in the office of the intermed also directly increases glacose synthesis and a senting of the intermed and directly increases glacose synthesis code. The indirect effects of growth hormore are largely mediated by IGF-1 (results blee growth factor), which is produced in the level and vistualy all offers its sense in response to growth hormore. IGF-1 is also anabole to both muscle and bone, augmenting growth hormore. This includes increased hopping-risk first extention, precessed glacose consumption, and decreased glacoseconsarption, and decreased glacoseconsarption and decreased glacoseconsarption, and decreased glacoseconsarption, and decreased glacoseconsarption and decreased glacoseconsarption and decreased glacoseconsarption, and decreased glacoseconsarption and decreased glacoseconsarption and decreased glacoseconsarption and decreased glacoseconsarption and decreased glacoseconsarption.

Somatropin is considered to be a controversial anabolic and performance-enhancing drug in the realm of bodybuilding and athletics. The main issue of debate is the exact level of potential tental this substitutes carries. Write states with 1874 pattern in a wisting state tend to support potentially strong artificial and residuable properties, stated dominating these same effects in healthy abalts and athless are belong. During the 1980s, a large body of right amounted discussions of Gelf in hoophshafing circles which may have been faciled by the high cost of the dang and its very more ("goodt hommer"). It was come have been faciled by the high cost of the dang and its very more ("goodt hommer"). It was come having a contract of the cost of the cost of the danger and the state of the cost of the cos

#### History:

The first harm growth hormous proporations to be used in medicine were made from polarity ordines of Human regis. These are now controlly reflected to a cachier growth polarity ordines of Human regis. These are now controlly reflected to a cachier growth control and the polarity of the control of the cont

The FDA approved the first synthetic harms growth hormore drug in 1985. Synthesis produced a pure hormore without biological communitum, eliminating the possibility of CDI transmission. The drug approved was called somation (Protropin), and was based on a manufacturing technology developed by Gourcette in 1979. "Somation camer as an inportant time given the memoral of cadavor (GH by the FDA that same year. This hormore that the production of the protection of the production of the protection of the production of the production of pure synthetic somation; with the exact armin and called capture of modification growth formore." It was also discovered that the extrantal structure of somation makes a manufacturing the production of pure synthetic somation; with the exact armin and called experiment causes a much higher incidence of arthody reactions in partiest, which can make the gifted production of the production of pure synthesis contrastive short in partiest, which can make the gift and the production of the production of pure short production of pure specific production of pure specific production of production of pure specific production of pure specific production of pure specific production of pure specific production of production of pure specific production of pure specific production of pure specific production of production of pure specific production of pr

## How Supplied:

Somatropin is most commonly supplied in multi-dose vials containing a white hyphilized powder that requires reconstitution with sterile or bacteristatic water before use. Dosage may vary widely from Img to 24mg or more per visil. Somatropin is also available as a stabile pre-mixed solution (Nutropin AQ) that is biologically equivalent to reconstituted sometropin.

### Structural Characteristics:

Somatropin is human growth homone protein manufactured by recombinant DNA technology. It has 191 amino acid residues and a molecular weight of 22,125 daltons. It is identical in structure to human growth homome of fortiary origin.

### Storage:

Do not freeze. Follow package insert for storage information. Refrigeration (2° to 8°C, 35° to 46° F) may be required before and after reconstitution.

### Side Effects (General):

The most common advense reactions to ostumopoi therapy are joirt pain, headache, the like symptome, peripheral chem (under reoferin, and back pain. Less common advense reactions tacklet inflammation of macoon membranes in the nase (thinkis), dizheas, upper greated cheme, masses, new benes, capped larned syndrome, check pain depending, general collette, masses, new benes, capped larned syndrome, check pain depending, general median, sons, new benes, capped larned syndrome, check pain depending, general collette, masses, new benes, capped larned syndrome, for feet pain depending, general median, sons dependent pain and pain and pain and pain and pain and common pain and p

### Side Effects (Impaired glucose tolerance):

Somatropin may reduce sensitivity to insulin and raise blood sugar levels. This may occur in individuals without preexisting diabetes or impaired plucose tolerance.

#### Side Effects (Injection site):

The subcutaneous administration of somatropin may cause redness, itching, or lumps at the site of injection. It may also cause a localized decrease of adpose tissue, which may be commounted by the presented administration at the same site of injection.

### Administration:

Somtropis is designed for substantaneous or intranscular administration. One miligram of controlpis is capitalent to approximately 3 international links (3 LI). When used to treat adult onest growth hormore deficiency, the drug is commonly applied at a dosage of 0.05mg/lag per day. The capatas to modify 11 Ui as 3 Li per day for person of approximately 180-220 lbs. A long-term mintenance dosage is established after reviewing the matter's 13CF-1 lacks and clinical processors more firm.

When used for physique or performence-orbancing purposes, sometopin is usually administered at a dosage between III and if III per day Ce-II III being most common, III and the common of the dos commonly in the common orban orban

Other drag, are commody used in conjunction with contribution in order to clear a stronger response. Hippoid drag (standy 13 ray emritadushy common given the lowess effects of scentropies on flysiod beeds, and may significantly enthrance for loss during through Insalin is able commonly used with contrations, which form contexting error of the effects controlly of GP bridge proteins. I allowing for more (GF-1 actios)<sup>223</sup> growth between sord flack bowers (GF bridge proteins. I allowing for more (GF-1 actios)<sup>223</sup> growth between sord flack bowers (GF bridge proteins beds)<sup>223</sup> databolic anticipacity states with somethops, in an effort to maximize potential music-building effects. Anabolic strong in the contraction of the contra

### Availability:

Somatropin is produced by many different drug companies, and is distributed in virtually all developed countries. The most common brand names include Serostin (Serono), Saizen (Serono), Hurratrope (Eli Lilly), Norditropin (novo nodisk), Omnitrope (Sandoz), and Genotropin (Pharmacia).

Somatropin products are high value targets for drug counterfeiting operations. Many counterfeits are highly deceptive in nature, and have been found in both flict and legitimate drug distribution channels. Some counterfeit growth hommone products are made by nebbelino visk of IrCG which hear a very close visual resemblance to sematronin A home.

pregnancy test is sometimes used to help determine 2 HCG has been used to make a countrief it KH product. This test words by descring ICG in the urine. A few days into a countrief it KH product. This test words by descring ICG in the urine. A few days into a cock with sometime, the artifactual will be used. a 1-H U spicerup prior to bed. Upon ring the programsy test will be used, and a positive result will indicate that an ICG contrief it has the contribution of the contribution of a solid healthfood life. From this low roundred that contribution were most the few most of a solid healthfood life. From this low roundred that contribution were most the few most description.

### Increley® (mecasermin)

#### Description:

Measemin is huma insula ide growth factor-1 (GF-1) munification by recombinar DNA detechooy (infer). [GF-1 is the jump; mediate of the goods phenoming effects of human growth homene. As such measemin also can similate the growth of bree, muck, and internal organs. Is effects on delectal make are also strongly hypophosis, menting it causes an accuses in ord narrher. Unlike [GF] however, measemin has very storeg install-face (feets. It can aparty ordary by precising the paice of armin society, glacose, and fany acids, but lowers blood sugar levels so efficiently that it can infance severe hypophorenia from his choice possible and the control of first parish and measurement may promote his progressio, for an increase in the storage of body (if. His society) and all affects for its potential to appear the growth of society that the control of the procession of the pro

Measurem is most commonly prescribed for the restruct of severe primary IGF-1 deficiency (Primary (EFF). Bit disease is charactered by a filter to produce normal bests of IGF-1 due to insufficiencies in the growth hormore / IGF-1 axis (usually moshing GHI receptor, signing quitton, or (IGF-1) axis decisally moshing of IGF-1 receptor, signing quitton, or (IGF-1) axis decisally neither significant or even high levels of growth hormore, but their hocks do not respond to it with the sufficient production of IGF-1. Measurem may also be used for the returnent of patients who have developed arthoids to growth hormore therapy, in hoth instruces the patient is and CHI deficient, but does not respond properly in good the hormore, hereographically the consideration of the IGF-1 deficient production through the patients for IGH through, and reads a commonwealth of IGH descript, and reads a learner with GHI through, and reads a learner with GHI through, and reads a learner with GHI through and reads a learner with GHI through, and reads a learner with GHI through through the general content of the general

#### History:

The U.S. Food and Dug Administration approved Measuremin in August 2005. It is soil under the brand must herekes, termificated by Tercia lace, of Bellstone, Californi, Tercia licenses this technology from Generated, which was the first company to sell a synthetically munificated human gooth bornome product in the United States (Proteophy, Tercias's rife-II is produced human signature product in the United States (Proteophy, Tercias's rife-II is produced by a similar recombinate DNA technology, the process involves the process of the process

#### How Supplied:

Mecasermin (Increlex) is supplied in 4mL multi-dose vials containing 10 mg/mL.

### Structural Characteristics:

Mecasermin is human IGF-1 protein manufactured by recombinant DNA technology. It consists of a string of 70 amino acids and has a molecular weight of 7,649 daltons. Its amino acid sequence is identical to that of endogenous human IGF-1.

### Storage:

Do not freeze. Refrigeration (2° to 8°C, 35° to 46° F) required before and after reconstitution.

# Side Effects (Hypoglycemia):

The mst common adverse reaction to measuremin thurspy is hypoglycenia, which occurred on at last one occusion in 42% of parisers receiving the dup during clinical risk. Approximately 7% of patients roticed severe hypoglycenia, and 5% noticed hypoglycenia related lungs, as existen or loss of concisenoses. Steps of risk of the moderate hypoglycenia related lungs, disositous, blarted vision, depressive mood, dizieness, sosetting, pulpitation, trenze, restolencess, targing in the hards, for the post, or trappe, light-depotess, analyst occurrents, londarders, steps distributes, anasty, starred speech, rithality, almorated behavior, attendard momenta, and personally clauses. Fair of these swaring signs should occur, one should introducibly consume a food or drift, containing surple stages such as a carefly har of eartholythate clairs. Signs of severe hypoglycenia and health discrimination and hard for the control of the control

Mecasermin should never be taken before sleep or in higher than recommended doses. A meal or snack must be consumed within 20 minutes (before or after) of administration.

# Side Effects (Injection site):

The subcutaneous administration of mecasemin may cause bruising at the site of injection. It may also cause a localized increase of adipose tissue, which may be compounded by the repeated administration at the same site of injection. Rotation of the injection sites is recommended.

# Side Effects (General):

Other potential adverse reactions to measuremin therapy schale joint pain, growth of the troubs, strength, healther, dizziness, constraints, orange, heart pain, hearing loss, and hypertreply of the thymas glant. Mild elevations in serum NST, ALT, and LDH levels were found in a significant number of patients, the flow worse at association with hypotococcide of the first years of long-term flexupy in clicial trials, without decling result faction. Becausion in cholsestoral ant highesticate were also observed the reminent within the upper limit of normal values. Evidence of heart enlargement was constant including history and the contraction of the contr

### Administration:

Measurem is intruded for subcutaneous administration. The initiation of therapy mobiles close monitoring of blood glacous levels with all peoper ministenance does in established. The recommended starting does is .04 to .08 mg/kg (40 to .80 mg/kg) their closely. The older may be increased by .09 mg/kg per injection, reaching a ministrant of .12 mg/kg twice daily. The older may be approached all allows the subcutance of the subcutant of the subcu

Mecasermin is not widely used for physique- or performance-orlancing purposes. Common protocols of administration have not yet been established. Due to the potential for severe hypoglycenia, monimum doses armony bodybuldies and althetes are not likely to measurably exceed those supplied by therapeutic gaidelines. This drag will most fleely by taken in cycles lesting no longer than 8-12 weeds in an effort to minimize unwanted copyang growth of fit gain.

# Availability:

Mecasermin is approved for sale in the United States and Europe under the Increlex brand

### Description:

Sommton is a synthetically manifectured form of human growth hormore (Edil). It is exactly a variet of endogenous LEH protein, containing the same aspector of 191 aimso each, but with the addition of an ereal nation each, meaning the contrast period excellent and excellent endogenous contrast to excellent a medium frame growth hormore. As an Edil reduction, corgan. It also plays a role in protein, carbohydrate, legit, and mixed metabolism. In a mendical setting, contraste is used to next edil-lent with growth framework growth from the contrast to the contrast of the contrast to the contrast of the c

Abough counteren is considered expiculent to harmin growth herence, it is not a manal promotion the harmin body. Here my increase the chares for devologing arthfoliot so goods hormone design tentimes to general the models work by being with the growth hormone investigation. 2014 of the children tracted with one more devologing arthfoliot so goods hormone after one year. <sup>268</sup> In a strinkly configured investigation involving the administration of sometropic for year, one by It in "protest produced some arthfoliot so growth hormone. <sup>278</sup> It is important to note that in both studies the arthfoliot year-forms were not storage and did not depart on substantially definish the ability of the days to be storage and did not depart on substantially definish the ability of the days to be an very want powerings (CSF) of patients taking sometern Still, the convex 191 mino and configuration of semmings in consideration meet destable to use

# History:

Sommton was approved for sale in the U.S. is 1985. It was the first synthetic grouth hormone medization analishe workshies, produced with a mandlateing process called linkaisin Body Technology. <sup>23</sup> The technology insolves sensiting the DNA exceeding for the EIGH protein to exceeding a GHz EGO based to the state of synthesis provide intermed. EIGH was made into a medication was a contraction of the contraction of the Cost based of the synthesis growth hormone, EIGH was made into a medication was called, was burned in the U.S. in 1985 the to the high procedure of a mer tearn-legisle disease in patients. Sommton was approved for sale that same year, giving Generated a system recomposite or the state of the system of the contraction of the contrac

### How Supplied:

Somatrem is most commonly supplied in multi-dose vials containing a white hyphilized powder that requires reconstitution with sterile or bacteriostatic water before use. Dosage may vary from 1 me to 10 me per vial.

# Structural Characteristics:

Somatrem is a polypeptide (methionyl human growth hormone) manufactured by recombinant DNA technology. It has 192 amino acid residues and a molecular weight of 22 256 datous

#### Storage:

Do not freeze. Refrigeration (2° to 8°C, 35° to 46° F) required before and after

# Side Effects (General):

The most commonly reported advene reactions to committen therapy include carpial transities, beyondome, accessed growth of ravie (most and bridments), agreements, and parameters. Note that the side effects of sometimen will generally mirror those of sometimenty of the side of sometimenty cause disberts, are correctly to visible thickening of the boots, most readily the fact, forcheath, lancks, jow, and elebory, and enlargement of the internal capture, reachily the fact, forcheath lancks jow, and enlargement of the internal capture. The properties of the capture of the capture of the properties of the capture of th

### Side Effects (Impaired glucose tolerance):

Somatrem may reduce sensitivity to insulin and raise blood sugar levels. This may occur in individuals without preexisting diabetes or impaired plucose tolerance.

# Side Effects (Injection site):

The subcutaneous administration of somatrem may cause redness, itching, or lumps at the site of injection. It may also cause a localized decrease of adipose tissue, which may be commounded but the pre-stated administration at the same site of injection.

### Administration:

Sommtons ágion by sub-taincous or intransacular specians. One miligano of sommtonis equivalent to approximately 3 International Units (EIU). Muse used to teast children with growth failure due to growth hormone deficiency, the drag is applied at a dosage up to Oslimgka per day. The equates to a manurant for oughly follul per day for a particular of approximately 100 lbs. A long-term maintenance dosage is established after recieving for approximately 100 lbs. A long-term maintenance dosage is established after recieving for approximately 100 lbs. A long-term maintenance cover in the contract of the c

When used for playsique or performance-enhancing purposes, sometrem is usually administened at a dosage between III and 6 IU per day (2-4 II Usbeing most common). The drug is commonly cycled in a similar marner to a moleclic/antioogenic steroids, with the length of intake generally being between to weeks and 24 weeks. The ambole effects of this drug are less apparent than its lipstific fall tools properties, and generally take longer periods of time and higher does to marified themselves.

### Availability:

Somatrem is available in the United States under the Protropin brand name which is distributed by Roche. In Europe and most maiors the vast majority of hGril is the correct 1911 amino acid sequence somatropin. Somatrem can be found in some markets, however, most commonly in Asia, where it tends to sells for a substantially lower price than somatronin

# Hypoglycemics

# Glucophage (metformin hydrochloride)

### Description:

Metformin hydrochloride is an oral anthyperglycemic medication. It is prescribed for the management of Type-II diabetes, sometimes also referred to as marue onest diabetes since it ends to develop later in life. The drug is typically utilized when dietary management and exercise alone have not been able to control the progress of the disease, yet rigicable insulin is not an appropriate option. While the main activity of metformin HCL is the increased and the progression of the disease. utilization of glacose, it does not directly mine the action of issulia. While its precise mode of a caction is utilizons, it is uterestored to encluse the output of glacose by the live, decrease the intestinal absorption of glacose, and increase insulin sensitivity in certain organs and peripheral issues. Use of this agent will buser the patient's blood stager, though its activity or makes it less filedy to cause a dangerous state of hypoglycemin if the dosage is accidentally missiated (a concern with injectable is unifain).

Insula mujutation is common in sports due to the biological actions of this hormore. Insular is involved in nativest storage, helping to bramport aims out, off, they sock, and carbolyduttest (glecoso) into various cells. In the case of muscle cells, multin also hellants collected multiples (protein) instructions, insular and ins

# History:

Medicinis has a long and finity complicated bistory in medicine. This drug was for sportheaded in 1979 drug with a neiso of other higamates. Atthogs some of as beneficial properties used exhibited at the time, it was not risinly subject to human trial. "If not all in the research book for many free deceades after its central. The first clinical angle in exhibiting is therepartic potential as a gloose lowering medication was finally initiated in 1995 (50°Phone trials were very successful, and metalium is sus-school for clinical development, it was given the rate name Glacophage, which mendates to "glacose entail". This of course, refers it is ability to help the body dispose of body dispose. Glacophage went on to be the most recognized trade name for metalium HCL. This trade name is still being used to mutest the drug dody, presently the Bristal-Meyer, explaid Company.

Metfernis was not initialy the glocos-lowering agent of choice among chiciaus. During the 1990s, it had been standed aleagued perhaterian and banfami, which had also been selected for commercial dovelopment. Metfernism was initially pashed saids in favor of these two desays, both of which had prouses to be significantly more potent at lowering blood saids to the properties of the properties of the properties of the properties and badfernis were producing unacceptable high in Suckeres or labels existins, and often fail methods. Glosself charactered by a rapid drop in pilt. By the close of the 1970b, most good exhaustered by a rapid drop in pilt. By the close of the 1970b, most good exhaustered by a rapid drop in pilt. By the close of the 1970b, most growth pilt.

The structural and pharmacological similarly of metiferius to phenformia and baderius held back it desiral potential for many same. Sometimes were which concerned that this agree would also present unforceable risks. While lack acidsois is a legitimate concern. It occurs for the present of the present of

### How Supplied:

Metformin hydrochloride is most commonly supplied in oral tablets of 500, 850, and 1000 me each.

### Structural Characteristics:

Metformin is a synthetic derivative of the natural antidiabetic agent guanide. It is specifically the 1.1- dimethylated bizuanide variant.

### Warnings:

In rate cases, the use of metforms HCL is associated with heir actions, an other heart and the method desorber motivoing tumour gother factors in a frenze in heisted beeds (heir) and a prosonered decroses in blood pH (actions). This risk increases with conflictors such as special, deblymatic, one sees shedwidt rather, but its sufficiency, and prapiriment, and acte congestive heart faince. Symptoms Officie actions include making, muscle pain, respiratory defineses, downstrase, and absorbing distincts. Laboratory shormalizes include to perfect the action of the contraction and are found in the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction of the contraction and are found to the contraction of the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction of the contraction and are found to the contraction an

### Side Effects

Commo side effects of materima H.T. therapy richale distribe (33.2%), musculvoring (25.5%), flutheres (12.4%), weakness (29.5%), flutheres (12.4%), weakness (29.5%), indigentes (17.4%), almebrari of (19.4%), and breakness (27.5%) indigentes (17.4%), admental content of the production of the product

### Administration:

The coal absorption rate of metricums HCL is show, with the body taking approximately six houses to absorbed and distribute each does. Described releases (200) formalistics are also made, which further desloy the absorption of metricums HCL. Estended releases tables should made the state of the control of

When used for physique or performance-enhancing purposes, the typical protects of is to like 850 mg concer or two-part of life fairing polisions in desired, it is highly liketin 1-2 losses to be the excrete, so that the drug can line to passed the change three enhys stages of recovery. It is highly corrons to use these carefulness as appelled the change three enhys stages of recovery in the high corrons to take a carefulness as perfect as the property of the

### Availability Trends:

Metromis is readly available given its videopread use in clinical medicine. It is sold under meny bond marea, so both a standation and confinition medicinion. Bolyhalides and athletes tent to list their use to presputations containing only metformis. While the drug is not a thick seem to list their interest by seriod courtefeiting or undergood manufacturing operations, is in the target of many other general days courtefeiting period and the containing on the sold. As seen it is bould not be assumed that all pendaged their growthes blend as menformis are legitimete. Care should be taken to crease that all products blend as menformis are legitimete. Care should be taken to crease that all products bearing this impaction those bone canciled through legitime pharmocracial charmen pharmocracial charmen.

### Description:

Insulin is peptide hormore produced in the lists of Langerhuss in the parceness. The release of this hormore in the human body is most closely ted to blood glaces level, although a marrher of other factors including purcentic and gastroinstrail hormores, amino acids, farty acids, and factors bodies are also moboted. The min bological role of malin is to percent the intracellular titlation and storage of amino acids, glacose, and fatty acids, while simultaneously shifted pure boundaries of glocose, protein, and it is in most mobility of the standard proteinstrains and the simultaneously shifted graph boundaries of glocose, protein, and it is in most mobility proceived to people with diabetes, a methodic disorder characterized by hyperapheronic fleigh blood signally while assist targets many different organs in the body this hormore is both anabolic and anti-catabolic to skeletal muscle fissos, <sup>122, 123, 124, 124</sup> a fact that explains the inclusion of pharmoreacial instina in the result of additions and bodylated and anti-catabolic to different organs in the relation of addition and bodylated in stanta five factor of addition and only of additional bodylated in a stanta for a factor of additional organic and anti-catabolic and anti-catabolic organic and organic and anti-catabolic anti-catabolic anti-catabolic anti-catabolic anti-catabolic anti-catabolic anti-catabolic anti-catabolic anti-catabolic anti-cataboli

The use of isadin to improve performance and body composition on the a little tricty because this bettmen can also promote materie stages in fat cold. This, between it as the intermed can be provided in the control of the control of the control of the regimen of interne weight training and a dist without excess colories and fit intels can enable made to above a much higher affinity for protein and placous storage in muscle is opposed to fing acid storage in adoption) cells. This is specially true in the post-curvice enhanced absorptive state, where results sensibily in skeletal matech has been shown to increase supfacturily our beautiful results of the beautiful training and the control of the control beautiful training and the control of the control of the control of the control of the harmon, it is level, capable of producing grapf and in retended muscle gains. The muscles therem, and the overall coulse of forbears are control occurred to control of the control occurred and the control of the control occurred to the control of control occurred to the control of control occurred to the contro

The fact that results use convox be detected by witholds his control for place in the drug requires of many affects and produced model hopkhaders. Note that there has been some progress in drug detection, especially with the analogs, but to date regain risulfs is sail considered a "said" fage Institus is done used in convisations which or "cortest said" frauge like harms growth hormore, throat endestines, and low done testostence spictors, and supplier can have a darms electron of the early "splenges and performs without fast of single care have a darms electron of the endestines, and low done testostence spictors, and supplier can have a darms electron of the end "splenges" and performs without fast of fast that results contributed with anti-lockin/unloopie; servides can be a very syntepsistic contribution. This is because the two activities appear an analosie start frough different reclusions. Insulin strongly enhances the transport of natures into mack cells and shibles protein breakforms and the analosis kernelly disregate the analosis of the protein breakforms and a markode kernelly disregate the analosis.

As mentioned, the usual medical purpose for insulin is to trust different forms of diabetes. More specifically, the humm body may not be producing enough insulin (Pspel diabetes), or may not recognize residu well at the cell see although some level is present in the blood ((Pspel diabetes), and previolence, required to inject residu on a regular basis, as they are left without a sufficient level of this hormore. Along with medication, the inhibitual will meal to constating market belong dispose levels and regular their sages intakes. Together with lifestly in-rediffications such as regular exercise and developing a balanced dart, and the constant of the constant

### History:

Insulin fist became available as a medicine during the 1920s. Credit for the discovery is most appropriately given to Caradian physician Feld Barting and Caradian physician Feld Barting and Caradian physician Feld Barting and Caradian physician Feld Receive treatment of dischess. There work sterred from an else in stillay proposed by Barting with as a syong doctor theoreted that an active center could be must from airmal selection and the state of the stat

in his work. A coin flip determined who would work with Banting, and he was eventually paired with graduate student Best. Together they made medical history.

The first insulin preparation they produced were made of crude purceastic extenses taken medges. At one point the supply of blorestory arisins was cellusted, and desperate to continue their research, the pair actually began taking story dogs to supplement their particles apply. Since the continue the first point of the Point of the Continued to the Continued to

And reinidy declined the assistance in the loops that they could work out production issues on their own, Barring and his tourn worked with E-ILDy & Co. to develop the first meaproducted malint medicts using their aimine function incivingue. Their production reaches producted malint medicts using their aimine function incivingue. Their production reaches 1922, the same year Barring and Mackeds own the Nobel price That same year. Nortisk Intendishontonium sus Strated by Danish sceiterist Augusta Korgly, who despensibly wanted to bring back an insulin runnificating technique to Dermutk to treat his wife, who wanted to bring back an insulin runnificating technique to Nortisk, the world'statute of the strategies of the strategies of the strategies of the strategies of the world belief understood of Findiance at E-ILD & Co.

The early insulin medications were fairly impure by today's standards. They opically contained 40 units of animal residing per millite, in contrast to today's accepted standard concentration of 100 units. The large does needed with these early box-concentration drags were not very confinetable for patient, and injection-sel reactions were not uncommon flavy also contained significant protein imparises that would associately associated and patient protein imparises that would associate and large beforehead were focal with a sew death sentence following a diagnosis of dishotes; Eli Illy and Nowo Nordisk improved the parisy of their products in the conting years, but no migor improvements in insulin technology developed until the mid-1930s, when the first longer-acting issuit, portunities began to arrive first longer-acting issuit, portunities began to arrive.

The first Regar-acting chung made use of proteinine and zine to delay the action of risusin in the body, extending the activity care and redesige the number of duals pieceious required for many patients. Diebbed Proteinine Zire Insulin (PZD), the preparation would live an effect integra to large 3-45 hours. Neurall Proteinine Hagsdom (PVI) Plansis, due Downson Isosphane insulin, followed, reaching market by 1950. This preparation was very similar to PZP randin cucypt their could be mixed with regalar studies without disturbing the relationship of the protein protein and the protein protein protein and the protein protein and the same syring with NPH issulin, providing a hybrids relates patient characterized by an early peak effect due to the centar issulin and an extend active to local or by the NPH.

In 1951 the Letter issuline began is surface, which included sembents, bets, and when-letter proposations. The amount of rize used in each work quick exploration with distinct and lenge-string pharmacolarizers, Unifice previous Insulins, this was also achieved without from Phil Insulin over to a sight norming does of Lette insulin, the land health of the control of the new drugs as a big advance in insulin medications (Rough some would still require an excentige does with a letter insulin to matterial of thought some would still require an excentige does with a letter insulin to matterial of thought some would still require an excentige does with a letter insulin to matterial of court does be thought produce levels during the 24-loss protectly. Up to this point the insulin drugs made by the large pharmacontical delicency schedules are would come for matter? 2 years.

A substitute of the control of the contro

In 1974, chromatographic purification techniques allowed the manufacture of animal insulin with extremely low impurity levels (less than 1 pmoll of protein impurities). Novo was the first to release a drug made with this technology, which it called monocomponent (MC) [Conf. DEL 1984 to accessed a configuration of the protein purity of the configuration to the amoun zet any mor received a versant cross y sugar reas. meant, meny receiving to the side protein point foreid upon chemical analysis. This advance findly significat, would be ident book in 1975, Cho-Geg produced the first synthetic insidii preparation (CGP received for the synthetic produced for the synthetic produ

The FDA has approved a surkey of other issulf and go combinations over the years, ixelding works to be plant is mitted backed at the difficilly generate of replant and hoper-acting resident. More necerify, we have also seen the FDA approval of FB Liky's repré-acting issulfa analyge thereby the resident of the plant and Applied from a survey of the resident and Applied from the resident and applied from the resident and applied from the resident and approved and sold in the U.S. and other mixens, it is imported to exclusive mixens are also under investigation at this time. With the large variety of different resident medications approved and sold in the U.S. and other mixens, it is imported to exclusive the resident and approved and and the U.S. and other mixens, it is imported to exclusive the understand that "resident an extraction of the plant and the proposed and approved and and the U.S. and other mixens, it is imported to exclusive a continue to expend as new agents are developed and associately stend. Today, it is continued that 55 milkin people use some form of speciable insulin on a regular looks to mixend that mixend and the proposal and also the source of manner.

### How Supplied:

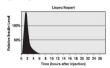
Plemenocinical insulin coruss from one of two basic origes, minul or synthetic With arimal source insulin, the formes is extensed from the paracruss of either jape or coves (to bulk) and proposed for modical use. These preparations are father disclosed into the consequence of the contract making their way into the preparate drug. Specifically called biosynthetic, synthetic insulin is produced by a recombinant IDAN procedures saint to the process used to mutualization produced by the contract of the contra

There are a variety of synthetic issulfs available, with each possessing uringe properties relating to speed of ossest, peak and durintion of activity, and concentration of dose. This therapetic variety may allow physicisms to take a treatment program for issulfs-dependent durinties that allows for the least marcot of dost ipactions and the greatest beef of partiest conflict. It is important that one should be some of the individual activity of any residun drug before attempting is so. Doe to the differences between preparations, it is also michally advised that externe care be taken whenever a physician attempts to switch an insulfadeependent dubiety profess from one from first miles medication such as

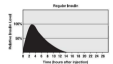
Below is a list showing the distinctions between popular forms of biosynthetic insulin.

# Short-acting Insulins:

Hamalogió Benultu Lisproi: Hamalogió is a short-scritg artulog of harma irsufin, specicially the Logisti2p ProtEgiva junto go frisatio rectaed when the artino acids is positions 28 and 29 are receivent. It is considered equipotent to regular soluble instalion on surf-to-surf basis, but with more mpil activity. <sup>28</sup> The cents of long action following scheatments admiration in approximately 15 mirates, and is peak effect is reached in 30 to 90 mirates. Has no atol duration of action between 3 and 5 bruse. Insufili legios is usually used as a supplement to a longer acting insufin product, providing a fist-scring medication that can be taken before or immediately after males to mirate the body's matural sainte response. Many athletes believe that its short window of effect mines it an itsel insufin medication for plession or performance-ordanic grapusces, are most of a section can be Novologie (Insulin Agrost): Novologie is a bott-seitig analog of huma issidii ceased when the minso del produce at position RSs in repulsed with supportion cit. The costs of time gas critical field in the second time and the cost of the second in a support time to a programmely 15 minutes, and its pask effect is recarded in 1-3 hours. Insulin input is usually used as a supplement to a longer acting insulin product, providing a field-seiting mondaction that can be taken before or introducing older most beach before or introducing older most intention between the submitted insulin response. Many athletes before that its short weaknow of effect must a find insulin medication for the policy—to preference endemanding paperses, as a most of in action can be



Hamulin-Fi. "Regular" (results ligit leaf tall to human mufin. Also sed as Hamilin-Fi. (Schibb)) is some mutect, this product consists of rize-minin regulation disorder in the third. There is nothing added to solve the release of this product, so it is generically referred to as soluble human leadin. This diet quotes not supply and the as a short cartaint or effect. The once of thing action following selectaneous administration is 20-30 matter, and its peak effects is reached in 1-30 hours. It has not all charaction of action between 5 and 8 hours. The product of the control of the co



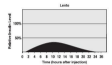
# Intermediate- and Long-acting Insulins:

Hamularis-N. NPH Instalin isophanes): A crystaline asspension of istalin with protamine and zine to delay te release and extend its action. Instalin sophume is considered intermediate length issalin. The onset of drug action following subsctaturous administration is approximately 1-2 Duxer, and its peak effect is reached in 1-10 Duxer. It has a total duration of activity leating more than 1-4 Duxer. This type of instalin is not commonly used for physiqueor performance—entrapressors.

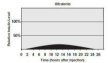
		NPH
Lovel	100%	
e Insulin		
£	50%	



Hamulards-L, Lente (medlum zinc suspension): A crystaline suspension of isaalin with zinc to delay is relase and extent is a critical. Hamilarl. Is considered an intermediate length isaalin. The cone of drug action following subentaneous administration is approximately 1–3 looses, and its past effect is nearboth in 6-14 hours. It has a total duration of activity lasting more than 20 hours. This type of insulin is not commonly used for physique- or performanceendronousment rumous.



Hamulin<sup>8</sup>-U. Ultralente (prolonged zine suspension): A crystaline suspension of insidn with zic to delay is release and extent d sacrior. Hamilto-U is considered a long-acting insulin. The onset of drug action following subcutaneous administration is approximately 6 hours, and is peak effect is reached in 14-18 hours. His set a total duration of activity isting 18-24 hours: This type of issulin is not commonly used for physique- or performanceerhancement purpose.



Lanton (malatin glazpine): A long-acting analog of human installa Institution glazpine is created whethe nations and aspuragied a profictors (All s beipedout by glacitic, and two arginisms are added to the Cetemins of the institution B chain. The croset of drug action following softenances administration is approximately?-1-2-brase, and the drugs controlled to how no significant peak (it is designed to have a very suble release pattern throughout the characteristic administration and carbon, limited grade last between 20-24 hours in the hody following subscataneous injection. This type of itsulfu is not commonly used for physique-or performance-relateurent purposes.

	Glargine	
lu Level	s	
gree programme son		
£ 30		

# Biphasic Insulins:

Humulin® Mixtures: These are mixtures of regular soluble insulin for a fast onset of action, and a long- or intermetiate-acting insulin for a prolonged effect. These are labeled by the mixture percentage, commonly 1090, 2080, 3070, 4060, and 5050. Mixtures using Humulou as the raudi-actine insulin are also available.

### Warning: Concentrated Insulin

The most common forms of issidis come in a concentration of 100 fl. of hormone per millite. These are desired as "L-10" programations in the U.S. and may other regions. In addition to this, however, these are also concentrated forms of soalin available to printers that require lighter closes and a rune concentrated controlled control to U-00 or 500 IU per milliter, are also sold. These are identified as "U-500" preputations, and are available by perception only. It can be externed duagnosco not fire thereating to replace a U-100 insidi product with a U-500 product without ranking the necessary dosing adjustments to compress for the general endocrocartistics. Given the guarred directly accurately measuring effects doses; C-15 III) with a drug of such high concentration, U-100 programations are used almost exclusively for physique- and reformance-channels.

# Side Effects (Hypoglycemia):

Hypogleomia is the primary danger with the use of rankin. This is a dangerous condition that coccurs who though glosco levels that look. It is a common and potentially find reaction experienced at some time or another by most medical and memodical ranks users, so it belows to be a superior of the common and the common a

It is also very important to note that you may notice a tendency tog at shope dair injecting amount. This is an entire way opportune of people-germa, and a clear seg ship to user should be consurring more carbohydrates. One should absolutely noted the temporate to tog to skep at the consumer of the people of the

# Side Effects (Lipodystrophy):

The subcutaneous administration of insulin may cause a localized increase in adipose tissue at

the site of injection. This may be compounded by the repeated administration of insulin at the same site of injection.

# Side Effects (Allergy to Insulin):

In a small percentage of users, the administration of readin may cause a localized alerge. This may relate irritation, swelling, taking analize rechess at the site of ispection. This often subsides as therepy contracts, in some instances in may be due to an integer to an ingulent, or in the case of ariminal results, a specific containant. Less commen, but posterially more valued to the contract of the pressure. In more bestures of the only the place, avointing uniform a robotic on the pressure in more bestures of the only the place avointing uniform and the contract of the pressure in more bestures of the contract of the contract

# Administration (General):

Gion that there are varying forms of results available for medical use with differing a pharmacolative patterns, saw vell as products with different deap concentrations; is extensively important that the user be familiar with the dosage and actions of any specific insign preparation by strend to use so the pack-effect, total fine of effect, total dosage, and carbolydrate intake can be closely monitored. Rapid-acting insign preparations (Novolog, Harmage, and Harmafiel) are the most popular choices for physque- or performance-orbancing papeose, and the subject of the dosing information presented in this could be considered to the contract of the c

# Administration (Short-acting Insulin):

Shot acing forms of results (Novolog, Humiles, H

Insulin dosages can way slightly among athletes, and are often dependent upon factors like body weight, reads resistivily, activily level (i.e. and the use of other days). Most uses choose to administer results insteadiledly after a workout, which is the most opportunistic time of the days to use this elig. Among bodybodistics, dosages of regist ir most [Hamilton 1904] are usually in the range of IIU per 15-20 pounds of the modolyweight. I'UIU is perhaps the more common dosage. This amortant mpt is adjusted downward delightly for seas of the more office. The fair contrast was such grown bodyweight guidelines, and instead start at a low dosage with the interiors of gradually working up to a round dosage. For example, on the first day of result therapy occur may begin with a dose as low as 2 IIU. Each connectative post-workout application find soages gript be in received by IIII. Until the user determines a confrictable range. Many feel this is safer and much more tailweigh of the trainformation was a supplication of the proteing a door, as some fifth by to before adolphy more or less reside that way to be the artificial than strainformation of the protein adoless, as one fifth by to before adolphy more or less reside the protein adoless, as one fifth by to before adolphy more or less reside the protein adoless and complete the protein adoless a some fifth by to before adolphy more or less reside the protein adoless and complete the protein adoless and one of the protein adoless and of the protein adoless and the protein adoless and other three or the protein adoless and the protein adoles

signly ingle relative/paterieties, as read unappy a soroni to on lower sectional, or under collar resistance to familiar.

One must also remember that it is very important to commar embedydrates for several houses following institute. Core should generally follow the rule-of-flatim for figuring a lates 10-15 genes of simple carbolydrates per III of readin injected (with a minimum immediate into 10-15 genes of simple carbolydrates per III of readin injected (with a minimum immediate into 10-15 genes generalles of does) limit is useful to 10-15 genes useful sea for similar and 10-15 genes angulates of does) limit is useful to 10-15 genes useful sea for similar of 10-15 genes angulates of does) limit in useful to 10-15 genes and 10-15 genes an

transcent manuscripts was among never a founce as surgin, region scripts at least on unexpected drop in glacone levels in notice of Many affects will also take creative monologistic with their carbohydrate drist, since the institute of the carbohydrate drist, since the institute of the carbohydrate drist, since the since and the care a good read and in the market. So of maints after ignoring institute or an advantage of the carbohydrate in th

# Administration (Intermediate-acting, Long-acting, and Biphasic Insulins):

Intermediate-acting, long-acting, and hiphasic insulin are designed for substaneous injection intermension injection mile came the days to be released to pupils, potentially resulting in hypodycenum. Following substantaneous injection, the injection size should be left adocument for method or present the design democration to trainful, the long-action and to reduce the present the design democration to trainful or beginning to the contract the

### Availability:

U-100 insulins may be dispensed from pharmacies in the United States without a prescription. This is so that an insulin-dependent diabetic will have easy access to this lifesaving medication. Concentrated (U-500) insulin is sold by prescription only. In most regions of the world, high medical use of the drug leads to easy access and low prices on the black market.

# Hypotensives

# BP Stabil TM

### Description:

BP Stubi à a blood prossure mungament product, it à net phermecutical based, but instead contrain nutural dietury appelents that apport fe finciscionig of the civolatory system (it à classified as a dietury supplement), in definite no to-event viames and minenda recessary for optimal contrain a runnier of components with substantial clainal support including headerm, "in curvection," metalement of components with substantial clainal support including headerm, "in curvection," metalement, "in sufficient," in such conjust," in Placific in general confusion, and in the confusion of the confusion o

BP Shall does not reduce blood pressure simply by increasing water too (distrets), Instead, it takes a milk-tool proposed to supporting circulatory bash! Most findamentally, it was designed to help maintain optimal functioning of the cadebulant. The exhibition is a seal designed to help maintain optimal functioning of the cadebulant. The value of the cadebulant is a cherical formal to the cadebulant in the cadebulant is a calculated by the careful cadebulant of the careful cadebulant is not healily and reaches, the system can become rigid or self. This would increase bodo pressure and restrictive when transporting blood further away from the heart. The loss of endothelial statesthy; can even lead to a state of child all presentates, definall by pressure clearlists in 16th of pressure careful of the

потты гапде.

# History:

BP Stabl was released in 2010 by Adeceder Nation (Island States). It is an expunsion of the Health Stabl product inc, which contains several products designed to support the turinge necks of Prodybulkers and adulests undergoing aggressive training or supplementation program. BP Stabl is closeful as a desay, pedial supplement, and a seal is a soubble program. By Stabl is closeful as desay, pedial supplement and a seal is a soubble program. By Stabl is a seal of the stable program of the stable program

# How Supplied:

BP Stabil is supplied in capsules containing a blend of hawthom extract, querectin dehydrate, nattokinsse, garlie, green tea extract, girkgo bibba extract, olive leaf extract, CoQ10, notassistrummenssium and calcium and visumins A. B3 (riacin). B6 (novosidine). C. and D.

### Side Effects:

BP Stabil is a natural dietary supplement and is not expected to have notable side effects. Individuals should monitor their blood pressure closely during use to make sure normal healthy levels are maintained.

#### Administration:

For blood pressure support, IPP Stabil is taken at a dosage of 2 capsaks per day. A dosage of up to 6 capsaks per day my be taken during periods of heightened activity or or supplementation. Note that a natural product such as IPP Stabil may hely support healthy blood pressure levels, but carnot be redied upon to replace the need for pressure medications, especially if a state of clinical physertension has developed. Care should be taken to research up monthly observed the contraction of the states of o

# Availability:

BP Stabil is produced in the United States by Molecular Nutrition. It is available for export, and may be found in Canada, Europe, Australia, and some other international markets.

Catapres (clonidine HCL)

### Description:

Choidine is a centrally acting ables-agoint lapotensis agast. It is most commody prepared as a lapticachiest is after use in tables and capsals, although it is not appelled in ordinal lagadit, rijectable solutions, and transdormal patches. Choidine straintiests alpha-adenocupous in the thirt, which reduces sympathics could be from the central across to five more casaly. Choidine is widely prescribed for the memorar of lapotension (and the contraction of the translet. Bodybaldiers pixelly use this drug for the same purpose, namely to content blood pressure which may be acrossed to the contract blood pressure which could use Cradine is also commonly prescribed for off-tiled purposes including the treatment of alcohol, opinic, and incoine addiction, continue.

Clinical studies show that cloridine can also produce a significant increase in the serun growth homme beet. This occars via stimation of hypothatine GBHH (drowth Hommer Releasing Hommer) relaces which in turn supports the secretion of growth hommer from the pintary. "I close the known anabodic properties of growth hormore, feature secretions specifically chosen by bodybalkers (when a blood pressure mediation is secretions specifically chosen by bodybalkers (when a blood pressure mediation is a target togetable or noduce near formation or oblov corresionism innoveneurs. This may be a target togetable or noduce near formation or oblov corresionism innoveneurs. This may be due to is effect on GH being door fixed. Studies agagest that in many patients, clearline will not road in assistant divences in GH with contrasted days  $me^{-6/247}$ . The percentilg literature even warms that is GH clearting properties are acute, and do not result in chronic electations with long-form use. Chairfarm why for of whate for both opersoare mumagement of their GH being between the order of the off-thick properties. It is not widely accepted as a GH stimulating days for bothyladder.

Oral closidire is considered a fast acting medication. Its blood pressure lowering effects are pipically noticed within 30 to 60 minutes after ingestion. Peak blood levels are achieved within 3-3 bloors after administrator, and the drug is metabolized with a half-size and produced and the produced of the produced of the produced of the produced administrated at least roice per day in order to markini theraportically effective blood levels. Food does not amorar to relatence the relatmoscharicies of closelation.

The absorption of cloridize from transformal patches may be significantly delayed in comparison to an labelse, and may the 2-3 days before peak bodd tools are arisined. Tolerance to the blood pressure lowering effects of cloridize may develop over time. It may be necessary to periodically revealute the dosage during prolonged theory. Cloridize may be used above, but is also commonly contributed with a second analypertonise agent when a stronger blood-messure beavierine effect is monitor?

# History:

Custine for superord in clinical states during the law 1960s, <sup>20</sup> and was approved by the US. Food and Dup, Administration for prescription sales in 1944. <sup>24</sup> One the years, its founds salety profile and level of efficacy have allowed to become a seroundst common fine-fine-fine points for the content of Dependencin The most saleshy those trades must be changing a content of the profile of the content of the potention. The most saleshy those trades must be changing a content of the profile of the content of the potential of the content of the c

# How Supplied:

Clonidine hydrochloride is most commonly supplied in oral tablets of 0.1, 0.2, and 0.3 mg each.

### Structural Characteristics:

Clonidine is an alpha-agonist hypotensive agent derived

Clonidine is an alpha-agonist hypotensive agent derived from imidazoline. It has the chemical designation 2 6- dichlors-N-2-imidazolidin/denehenzeramine

### Warnings:

Abrupt discontinuation of clonitine may result in side effects including rebound hypertension, nervossness, againin, headache, and of termor. In rare instruces clonitine withdrawal has been associated with serious cerebrovascular accident and even death. Patients should not discontinue clonitine treatment without consulting their physician. When discontinuing clonitine, it is usually advised to slowly reduce the dosage over a minimum period of 2-4 days to reduce withdrawal symptoms.

### Side Effects:

Commo side effects associated with clorifien treatment reduct day moth (40% of patients during clinical statistics), downsines (35%), declines (16%), consistention (10%), and sociation (10%), and sociation (10%), and sociation (10%), about mile hard best (congestive hort failings, fewer, toreused sensitivity to alcolo, shormed hard best (congestive hort failing, edition; increases, parasothesia (trigding of the skin), hair loss, rash, abdorniral print, americal, happatins, vorning, decreased libela, creeted sylataction, frequent weight out strategies are the right (notarity, kg cramps, masch pair, blamed vision, and dyness or barring of the eyes. Some side effects appear to be dose-dependant. Taling more than the proceeding amount (overlossing) may result in serious advents energiate through the control of the control o

nypotnerma, neart irregularities, coma, or death.

# Administration:

When used medically to tent hypotension, the dosage must be tailered to the individual needs of the princt. The usual arisist strainty dosages in adds is 1, mg gioen troke dully, mering and evering (2 mg per day). This dosage may be irrecused orace per work by 1 mgloby until the proper ministrance dosages is achieved. An effective disrupacttude of the effective disrupaction of the effective disrupaction of the effective disrupaction of the effective disrupaction of processing of proces

### Availability Trends:

Clonidine is widely sold throughout the developed world. Low demand, low cost, and high availability make this drug a low-profit target for counterfeiting. Counterfeits are unlikely to be a simificant problem.

### Liver Detoxification

# Essentiale forte N (Compound N)

# Description:

Essertiak forn N is the trude rums for a liver-support supplument distributed in Europe by Acestré Platurus. While this regarded as mendication is more regions, a factally contains a selection of intural viamins and phosphologist. Licevoise, in muny areas, is challege to bland concept of the contract viamins and phosphologist. Licevoise, in muny areas, is challege to bland complex of the viamins, viamin E and phosphologist florns of looks in Chinece and olic aced. Essertiak forts N is used widely in Europe to treat cases of hepsite deplications, such as those canced by forterin factories, allergy, desputies, or order descents. Essertial florn N is of risterest to attend using bodylosakes and athless for it a shipty to rotate the level of fore relative to the contraction of the contraction of the contraction of the contraction of the composite florns of 1-10 and the contraction of the composite florns of 1-10 and the adoleton.

The min mechanism of action with Essertial frets N appears to be ficused on the supply of arrivolutes and building bolest necessary for the rapior of dimagned cells. Bis product contains mainly polyarostaneted phospholophis (mostly phosphatis)chioleo), and a complex of B and E viantine, Prosphatis[chioleo is electrified as a membrane pilt, and is a ley component of the Essertial frets. N formâla. This phospholopid is important to the integrity of cells, adding both decibility and strength. Phosphatis[chioleo is loss) been destribed as an important supplement for the lore, supporting normal leparic fit metabolism and cowall lore health. It is also believed to be an important anticidual for feer and purcentia vellens. Bit additional viantines in the Essertial frets. N formâla nee Redy included to increase the articidated and regenerative properties of the mediation.

Secribs fort N was the first product shown to migate the hypotoxic effects of ambolic standards are included in a fixed standard. The increasing in a decided stand, the first standard standard standards are standard standards and the standards are standards and the standards are standards are standards as the standards are standards are standards as the standards are st

increased hepatic enzymes' levels suggest a relationship between abused AAS and hepatic cel damag. However, when AAS were taken with [Essertiale forts N]... the hepatotoxic effect appears to be attenuated. While this one study does not assure that steroid fore toxicity can be completely eliminated, it does lend strong support for the use of Essertial forts N with heratoxic arabolic strends.

# History:

Essentiale fotte N has been sold in Western and Esstern Europe for musty years, where it is distributed by Aventis (Kennely Mbone-Poulers Rore). This compound has been approved for the memory of highly. Been seations and often begand dysfarctors. European for the contract of the contract of the contract of the contract of the regard to her beath. It did not each the anterior of addess in the Usted States and Condo that 2006, blowers, when the resistanciation into its effects with stated alteres was published in the Clinical Tracticopie journal State then this matual medication has been retrieved in the Clinical Tracticopie journal State then this matual medication has been retrieved in the Clinical Tracticopie journal State then this matual medication has been retrieved in the Clinical Tracticopie journal State then this matual medication has been retrieved in the Clinical Tracticopie journal State than the state of the contraction of the contraction of the clinical state of the contraction of the clinical state of the contraction of the clinical state of the cl

# How Supplied:

Essentiale forte N is supplied in soft gelatin capsules containing vitamin B1 (6mg), vitamin B2 (6mg), vitamin B2 (6mg), vitamin B2 (6mg), vitamin B2 (6mg), vitamin B1 (6mg),

### Side Effects:

Essential forte N is a natural vitamin and supplement medication and is not expected to have notable side effects

# Administration:

When used medically to treat hepatic dysfunction, the most common recommended dosage is 2 capsales 3 times daily with meals. Bodybulders and athletes using this medication to reduce the hepatic stress of onal stored use will hypically follow the same medical prescribing gaidelines, and will taken the product for as long as the hepatotoxic steroids are administered.

### Availability:

Essential forte N is widely available in Western and Eastern Europe. It is a relatively inexpensive supplement, and is not a high interest target of counterfeit manufacturing

# operations.

# Description:

Lis-25 as in-held medicine used widely in Europe and Asia to support metabolic and kee helds. While is one occurities this product is regarded as a drig of contains all matter inguidents including captures spirous, terminals arjun, cichrain riphots, achillea midelalum, solumaringum tumming faller, and casio occidentals it is specifically used in the prevention or treatment of Praptisis, alciolic liver disease, early liver cirricosis, proxim energy malarization, loss of appetier, melation and chempthemp-relaxed here damage, as an adjust to hepatotoxic diseg, and to support metabolism during considerance or protograf films. As the first time better of liver more odial suggest, overall for health for primary factor of this product. Body-haldkers and athletes use Liv-22 as a way to reduce the level of strain placed on the few by hepatotocia enabloical incidents restords.

Numerous medical studies have been conducted on Liv-52 in recent years, many of which involve its ability to protect the liver from damage by alcohol or other toxins. 751 752 753 754 One investigation in particular looked at how the herbal medication affected the breakdown

of account in the body, stowerg time it notatiny accessed as execution, even to the point of being adds to reduce and buy largours opprious after bring of string, <sup>175</sup>5 Andrest study investigated what underlying mechanism might be insoled in Lis-52's ability to protect the been against alcohol toxicy. It demonstrated that one mechanism involved a specific ability to show the rate of glantilinear deplication. <sup>286</sup> This may be very important to the stood-using attacks, as glantifarious deplication is looked as as a direct number of here steen with C1 ability adjusted outs. Note that while these studies bent support for the use of a natural transply like that the strength of a process flow of the strength of the process flow of the strength of the strength

### History:

Li-52 is an herbal product that has is roots in syavedic medicine, mag cold formof Hindu science and medicine of intent elements. Li-52 is numelicated by the Himilitya Drug Ci., it Bushiny, India, and soss first struckaced to the global medice in 2002, the Stoning processor and the science of the struckaced to the global medice in 2002, the Stoning processor actually exhibited Li-52 as a pulmental product, which is believed to be the first time in herbal remody was adopted as a prescription due; in Western Bergon, 2011, Li-52 and advantage of the science of the struckaced and the science of the scie

### How Supplied:

Liv-52 is supplied in capsules containing a 450 mg of a blend of caper bush (capparis spinosa), arjuna (terminalia arjuna), wild chicory (cichorium intybus), yarrow (achillea millefiliam), black nightshade (sobraum nigrum), tamarisk (tamaris gallica), and negro coffee (cassia occileratilis).

# Side Effects:

Liv-52 is a natural herbal supplement medication and is not expected to have notable side effects

### Administration:

For general liver support or as an adjunct to hepatotoxic pharmaceuticals, Liv-52 is generally taken at a dosage of 1-2 cansules 2 times per day.

### Availability:

Liv-52 is produced exclusively by the Himalaya Drug Company, and is distributed widely in many areas of the world. In some regions, including the United States, the product is marketed under the Liver-Care® trade name.

Liver Stabil TM

# Description:

Lies Stable à a lor support supplement. It contains more than a donne materal impredient designed to help ponces and obtously the lever, and my be specially useful directly exposure to hepatotonic substances such as coal anabolezandragueis steroids. At the foundation of his formation are several reliciously stated inspections sainte to those Doral In-Ilarida fold medicare, has been the subject of mortium states solvening or compreted the level (separatus armos-termelense and alumine armos-termelense) and bilinthis beek after took exposure. <sup>200</sup> Jupin as loss used, and gain his been substance administration, <sup>200</sup> Various (scalibles after the substances and the substance administration, <sup>200</sup> Various (scalibles) and stress meteors following hepatotonic substance administration, <sup>200</sup> Various (scalibles) secretion. To Phosphatidylcholine (a key constituent of cellular membranes including liver cells) and a combination of key vitamins round out the Liv-52/Essential forte-like base of the Liver Stabil formula.

Ljid Sagel expands on its foundation with for additional well-statistic for beath regioneties. The most procedure of these are N-coedy systems and L-glataflone. Both matriers are important to ministring onesping for health, and may be especially useful in concurring hapting faintiness depletion. We mell need comment with on AIAS toxicty. Note, mak finds central standardized for adjunction contains concurring the contract standardized for adjunction contains on of the most extensively statisfical rapports are developed to the contract the contract of the contract the contract the contract that the contract the contract that the contrac

# History:

Lier Sahl was developed in 2008 by Molecular Nutrition (Linde States). The ficus was specifically on designing a supelment fluer can help support the Pathal in users of hapatotosi substances such as cell amboliciantiosgenic steroids. The formals was intended to target several law general forestimating larger and levation and hapatic enspire elevations, glantifinese depletion, suffarmation, and the secretical and hapatic enspire elevations, disturbine thereinteed individuolism, most noteby in Carnada, Astonifia, and centain parts of framespec larger laws for this some regions with sinker controls of the control of

# How Supplied:

Liver Stabil is supplied in capsules containing a blend of N-acetyl cysteine, milk thistle extract, wild chicory (cichorium intylus), L-ghtathione, wasabi japonica, arjura (terminalia arjura), phosphatidylcholine, yarow derdillea millefolium), vitamin B1, vitamin B2, vitamin B6, vitamin B1, vitamin B1, witamin B1, wi

# Side Effects:

Liver Stabil is a natural dietary supplement and is not expected to have notable side effects.

### Administration:

For general liver support or as an adjusct to hepatotoxic pharmaceuticals. Liver Stabl is a taken at a dosage of a capsakes per duy. A dosage of type to 6 capsakes per duy my be taken taken at a dosage of capsakes per duy my be taken my help reduce the level of Sev toxics, but an antural product such as Liver Stabl is my help reduce the level of Sev toxics, but cannot be redul upon to completely eliminate potential durings from the abuse of hepatotoxic drugs. Care should always be taken to manifor her health when taking few toxic substances.

### Availability:

Liver Stabil is produced in the U.S. by Molecular Nutrition. It is available for export, and may be found in Canada, Europe, Australia, and some other international markets.

### Reductase Inhibitors

# Avodart® (dutasteride)

# Description:

Datastrick is an iribitor of the S-alpha reductase cropne. Reductase iribitors are designed to present the convension of Isotastron to is true an integracy convergent DETI (diplydrottostorous). DETI is implicated in a number of disorders in men including male pattern having some disorders. Datastrick is specifically approached free treatment of symptomic broagn prostate chappends in (BFI). While datastrick is smaller in statute and action to instructive, if all continuous for a format in the statute and action to instructive, if all continuous free agreement in the statute and action to instructive, if all continuous free agreement in the statute and action to instructive, if all continuous free agreement in the statute and action to instructive in the statute and action in the statute a

The DHT sinhing effects of datastride make this drug of some interest to bodyhalders and adulties, particularly betwee concerned with the androguic component of relassions related staroids. Datastride is capable of reducing the androguic side effects produced by DHT concension, changing be profile of testostrove days measurably Provided moderate does of testostrove are being used, the result can be a substrating reduction in the occurrence of only shirt and zero. From the proof is reducing the provided moderate does not be substrated to the provided moderate does not be a substrated in the process of the latent has no selective type. 2 tribbite, and the provided of the provided of the provided of the provided provided to the provided of the

In terms of overall potency, a study published in the Journal of Clinical Endocrinology and Matcholom (Most 2004) of therefor potential endocrinology and Matcholom (Most 2004) of therefor potential endocrinology and the studies of the potential counterpart, finanterial.<sup>500</sup> In this investigation 399 miles suffering from being protestic hypotrophy were senserabled and sequential to the organization groups, and necessity disably or placebo, for a period of 24 weeks. Over the 24- week period, the disasterate group roaded the storages level of DHF sithstorn. The beneficial effects of the ding also occurred over a wide range of diseages. For example, a 3 mg dish pomest caused 58-8% shall be in DHF fireds, when more of 5-8% of the distoration of the state of the stat

This was in great contrast to the 5 mg finasteride group, which noticed only 70.8% inhibition. Researchers also noted that there was significantly more of a variation in the results of the finasteride group, with some patients noting DHT suppression in the range of ronly 50-55%.

Just as three can be benefits to lowering 5-alpha neductuse activity by way of less anthogogic side effects, there can also be some disabolists. For one, a strong and length if legislation with necessary and the side of th

### History:

Datasterile was first described in 1997.76 It was developed by the U.S. based planmeutrical company GlosoSmithKine. It was approved by the FDA in November 2001, and introduced to market the following year by Gloso under the Avodant trade manu. GlosoSmithKine also markets the drug in a number of other countries in Europe and South America under the same trade rune.

### How Supplied:

Dutasteride is supplied in soft gelatin capsules containing .5 mg each.

# Structural Characteristics:

Dutasteride is a synthetic 4-azasteroid. It has the chemical designation (5·,17,)-N-{2,5 bis(trifluoromethyl)phenyl}-3- oxo-4-azaandrost-1-ene-17-carboxamide.

# Warnings (Pregnancy):

This drag must never be taken during programsy. Be source that disasteriske cam be absorbed through the sids. Wintern who are, or might become programs, should never handle disasterisk capsules. The DHT blocking action of disasterisk can cause severe developmental profession to an unborn made fient, seen in very small amounts. Unlabered disasterisk can also be recovered in the semen. It is utknown if the drag can be absorbed exhibited to a single property of the disasterisk can also be recovered in the semen. It is utknown if the drag can be absorbed exhibited can also be recovered in the semen. It is utknown if the drag can be absorbed admission of the desired of the desi

### Side Effects:

The most common adverse reactions to datasteride therapy are impotence, reduced libido, and difficulty ejaculating, Cynecornastia was also noted during clinical trials, but occurred in less than 1% of patients. Some patients have also developed allergic reactions to the drug, including raish, tehing, edorm, and hives.

### Administration (General Considerations):

Reduction: elibbitors current completely protect against authorganic sale effects such as settori-in-hazden lairs, soo, shy skin, and came. Reductions: elibbitors lessen these side effects by reducing, not eliminating, the level of antiogenic activity in the sisk and sculp. Authorganic and anabolic effects are both mediated by the same receptor, and there is presently no suglarous its completely separate filmes two properties. Displantics solutions as also not training therefore does not forfer complete protection anaiset this side effects, by Delf additional therefore does not forfer complete protection anaiset this side effects.

Reduction inhibitors are only applicable with testistence, methylactoristence, and dampyrentience. These there daing are consecuted to stranger "lighty" destrations by the analyses the entire of the control of the stranger of the reduction engine. Numberloom and some of its derivatives become weather upon interaction with this copper, as their "lighty," metablosis but the analyses, more proper very peoply. Reduction: inhibition may introlly their analogusic side effects. Methandrostroubre and boldence undergo concession to stranger "solahe medical entirelistics, but at such mall looks that reduction inhibitions level the effect on their analogusicity. Most other synthetic analosis stansists are undificated bythe reductions common and reduction inhibitors.

# Administration:

When used medically for the treatment of symptomatic benign prostatic hyperplasis (DPRI), transactive its taken in a dosage of 5 ang (1 expasily per day. When used by bodyniklers and address to reduce the antiogenicity of testosterone, methylestosterone, or flaxopresserone, distinctively is commonly taken in a dosage of 5 mg (1 expasily) once every 1-2 days. The drug is typically administered for as long as the offending steroids are also taken.

### Availability:

GlavoSmithKline distributes this drug in the U.S., Europe, and South America under the Avodart trade name. Additionally, a number of other brands can be found in different markets including Austria (Avobe, Zyfetor), Greece (Daugen), India (Duprost), Netherlands (Daugen), Portugal (Daugen), and Spoint (Daugen).

# Proscar® (finasteride)

### Description:

Frustrick is an inhibitor of 5x-roductuse, which is the enzyme responsible for converting textosterow in to DHT (displotecontensory). Bits or que miléctury roduce the serim concentration of DHT, thereby minimizing the susmeated analogues: effects that result form is present. The effect of the sings is quite rapid, suppressing earn III off concentration is to the properties of the propert

Finastrick is a specific ribbitor of the Type-II Sa reduction errors. There are extually two someones of reduction in the human body, labeled as Type-II and Type-II Type-II Sa-reduction is predominarly found in the fore and schoocine glaris of this skin Type-II Sa-reduction is predominarly found in the fore and schoocine glaris of this skin Type-II Sa-reduction is predominarly found the time of the third this school is the skin Type-II Sa-reduction of the sk

Finastrici is considered a highly specific drug, as a line lift spillower effect on the other homenes in the body line an offlinity for an analogous cortisogue, controperios, and therefore does not olibit any antiooganic, antionganic, estropanic, or anti-estroparic properties. It has no apprecible impact on cricating legisles of cortical, thyrotic ristanting homene, or thyrouse, nor does a appear to alser HIL/LDL cholesteed levels. Changes in Intensing homenes, therefore the control of the contro

### History:

The first release of finisterile in the U.S. was under the hand mure of Procure! (Merch), which was approach by the FDA in 1902. It was specifiedly going approach fir see by patients with beings postute hepspelasis (prostate enlargement). In December 1907, the Food and Drug Administration again approach firsterile, this time for a different propose, Food and Drug Administration again approach firsterile, this time for a different propose proposed in particular to finish probability of the proposed product of the proposed proposed for the proposed proposed for the proposed proposed for the proposed first proposed for the proposed proposed for the proposed for th

# How Supplied:

Finasteride is most commonly supplied in tablets of 1 mg and 5 mg.

# Structural Characteristics:

Finasteride is a synthetic 4-azasteroid. It has the chemical designation 4-azaandrost-1-ene-17-carboxamide, N-(1,1- dimethylethyl)-3-oxo-,(5-,17,)-.

### Warnings (Pregnancy):

This drag must never be taken during programs; Finsterrice can be absorbed through the sist Women sho are, or might become programt, should never hundle broken or uncould finisheric haltes. The DHT blocking action of finisheric can cause severe developmental programment of the programment of the programment of the programment of the second of the second is substantial valued to the programment of the programment of the second of the second is substantial valued to the programment of the programment of

# Side Effects:

Adverse reactions commonly associated with the short- term (1 year) use of finasteride

include impotence (8.1%), decreased libido (6.4%), decreased ejaculate volume located. (3.7%), ejaculation disorder (.8%), genecomstin (.5%), breast tendemess (.4%), and rash (.5%).

### Administration (General Considerations):

Reduction elimbitors current completely protect against antiogenic side effects such as testioni-instead mile soo, soly skin, and care. Reductions elimbitors besom these side effects by reducing, not eliminating, the level of antiogenic activity in the side and scale. Antiogenic and antiolic effects are both mediated by the same receptor, and there is no very presently larger than the completely separate libes they proported by desirate testions as also not unguite productions. The completely receptor is a side of the completely protection against this side effect. In the control of the complete protection against this side effect.

Reduction sirbilities are only applicable with testistence, methylastistence, and that discourstence. These free drugs are contented to stronger "lighto" destrainties by the reductions engine. Nandrokine and some of is derivatives become weaker upon interaction with this engine, as the "lighto," metablosis bid the anthoganic receptive two proofs. Reductions ribilities may intend free intelligence and considerable with the anthoganic side effects. Methandrostrowhere and boliconee undergoo concession to stronger "solay metable anthoganic side effects. Methandrostrowhere and not an anthoganic side effects on their anthoganic side of the stronger side of the stronger and the stronger side of the s

# Administration:

When used medically for the tentiment of male pattern had too (ant/oupcretic alspecia) in mrs., the recommended doosage is lings per day. When used for the unstartent of being prostatic hyporphism (BPH). S mg per day is usually administered. When used by body-dulated and allottes to reduce the analogousity of fector-storest, methylessectionser, or hyporphism of the second property of the contract explosion of the property of the spirally administered for as long as the officing sectorial are also tulean Stoce DHT righthetic and lesson strength and possibly market gains during studentser, methylessectories, or flassymsterrore therapy (spon the positive actions of antiqueness on the automatically section, a "late only both necessity" approach is usually lead with

### Availability:

Finasteride is widely available in most regions of the world. The most prominent brand names in commerce are Prosear® (5 mg) and Propecia® (1 mg), although a number of other brand and generic forms of the drug can also be located.

# Testosterone Stimulating Drugs

hCG (human chorionic gonadotropin)

### Description:

Hamus Chroixe Coundstopin (ICG) is a prescription medicates containing obvinive grandstopin obstanced from a natural (Iranium) eight. Chroixe grandstopin is a polyspeptic hermore normally favor in the famile tody during the early mortles of receiving the production of progestorous, a preparay-sensiting hornore. Chroixe sensessing the production of progestorous, a preparay-sensiting hornore. Chroixe grandstopin is present in significant amounts only during programs; and is used as an anticater of pregnanty by standard over-the-counter pregnancy sets the Biodo levels of chronic grandstopin become mixtuodile as easy as seven days after routdine, and the hornore level will deligo grandsally strift the post of better. Although it possesses minor FSH-like (Follicle Stimulating Hormone) activity the physiological actions of chorionic gonadotropin mainly mimic those of the gonadotropin luteinizing hormone (LH). As a clinical drug, hCG is used as an exogenous form of LH. It is typically applied to support oxidation and pregnancy in women, most specifically those suffering from infertility due to low concentrations of gonadotronine and an inshifty to oxilate. Due to the ability of LH to stimulate the Levdio's cells in the testes to manufacture testactorine hCG is also used with mon to treat hungarous/detronic hungarous/sigm a disorder characterized by low testosterone levels and insufficient LH outnut. The drug is also used in the treatment of prepulsertal eruntachidism, a condition in which one or both of the testicles have failed to descend into the scrotum. HCG is used by male athletes for its ability to increase endogenous testosterone production, generally during or at the conclusion of a steroid cycle, when natural hormone production has been interrupted. History:

Charionic consideration was first discovered in 1920 767 and was identified as a pregnancy hormone approximately 8 years later. 768 The first drug preparation containing chorionic consideration came in the form of an animal nitritary extract, which was developed as a commercial product by Organon. Organon introduced the extract in 1931, under the trade name Pregnon. A trademark dispute forced the company to change the name Pregnyl however, which reached market in 1932. Pregnyl is still sold by Organon to this day. although it no longer comes in the form of a pituitary extract. Manufacturing techniques were introduced in 1940 that allowed the hormone to be obtained by filtering and purifying the urine of pregnant women, and by the late 1960's were adopted by all manufacturers formerly using animal extracts. Over the years the process and manufacturing protocols have been refined, but hCG is made in essentially the same way today as it was decades ago. While modern preparations are of biological origin, the risks of biological contaminants are said to

he low (although cannot be completely excluded).

Early on the indicated uses for charionic consideration preparations were much broader than they are presently. Product literature from the 1950's and '60's recommended the use of these drups for among other things, the treatment of uterine bleeding and amenorthea-Froehlich's syndrome, cryptochidism, female sterility, obesity, depression, and male impotence. A good example of the wide uses of charionic gonadatronin are illustrated in the preparation Glukor, which was described in 1958 as being, "Three times more effective than testosterone. For tired young men in male climacteric. For tired old men in male senility. Beneficial in impotence, angina and coronary heart disease, neuropsychosis, prostatitis, [and] myocarditis." Such recommendations, however, reflect an era less tightly regulated by onserment agency and less relient on proven clinical trials. Today FDA-approved indications for hCG are limited to the treatment of hypogonadotropic hypogonadism and cryntocridism in men, and anosulatory infertility in women

hCG was once widely used for the treatment of obesity. The trend seemed to have become popular in 1954, after a paper was published by Dr. A.T.W. Simeons claiming that chorionic consideration was an effective adjunct to dieting. According to the study nations were able to effectively stave off hunger with severely low-calorie diets provided they took the hormone injections. Dubbed the Simeons diet, people all across the country were soon subjecting themselves to severe calorie restriction (500 calories per day) and taking hCG injections. Soon after, the hormone itself became the main focus for fat loss. In fact, by 1957 it was said that hCG was the most commonly prescribed medication for weight loss. More recent and comprehensive investigations, however, refute that there is any anorexic or metabolic advantage to the use of hCG.769 In 1962, the Journal of the American Medical Association had already been warning consumers about the hCG-inclusive Simeons diet. stating the more basic fact that severe calorie restriction, which causes the body to sacrifice muscle and organ tissue to obtain necessary protein, was more hazardous than obesity itself. By 1974, the FDA had had enough of the hCG fat loss claims, and mandated the following statement to be included with all prescribing literature. "HCG HAS NOT BEEN DEMONSTRATED TO BE EFFECTIVE ADJUNCTIVE THERAPY IN THE TREATMENT OF ORESITY THERE IS NO SUBSTANTIAL EVIDENCE THAT IT INCREASES WEIGHT LOSS BEYOND THAT RESULTING FROM CALORIC RESTRICTION THAT IT CAUSES A MORE ATTRACTIVE OR 'NORMAL' DISTRIBUTION OF FAT. OR THAT IT DECREASES THE HUNGER AND DISCOMFORT ASSOCIATED WITH CALORIERESTRICTED DIETS." This warming

HCG has no significant thyroid-stimulating activity. This is specifically pointed out because

persists on all product sold in the U.S. today. In spite of this warning and evidence to the contrary, some clinics still promote the use of hCG for dieting purposes.

Haman Chroine Gendedoppi is a widely popular drug preparation today, owing to the fart it remains an indispensable part of outlaint heapy for irrung causes of framile that it is remains an indispensable part to outlaint heapy for irrung causes of framile to the control of the

# Structural Characteristics:

Chorionic goradotropin is an oligosaccharide glycoprotein composed of 244 amino acids. It has an alpha subunit that is 92 amino acids long and identical to that of lateinizing hormone (ILH), follicle-stimulating hormone (FSH), and thyroid-stimulating hormone (TSH). It has a beta subunit that is unious to hCG.

# How Supplied:

Harman Chorionic Goradotropin is widely available in various harman and veterinary drug markets. Composition and dosage may vary by country and manufacturer, but typically contain 1,000, 1,500, 2,500, 5,000, or 10,000 interminend unlist (III) per dose. All forms are supplied as a hyphilized powder, requiring reconstitution with sterile dilatent (water) he/for use

# Administration (General):

Haman Chorionic Gonadotropin is generally given by intramusecular (IM) injection. The subcutamous route is also used, and has been deemed to be roughly equivalent therapeutically to IM injections. <sup>700</sup> Peals concentrations of chorionic gonadotropin occur approximately 6 hours after intramuseluri nejection, and 16 to 20 hours after subcutamous.

### Administration (Men):

injection.

When used to tent Ipapograndscoppis Ipapograndsm, currest FIA-approved protectors recommend effert as ofter 4-wede prosgnar or a long-term prongan lasting up to 1 year, depending on the infloidant most of the pattert. Proceeding guidelines for short-term use recommend that 600 is 1000 mins to be gain 5 mins a vasele. 50 a works, followed by the contraction of the to be administered 3 stress weedly for 6 to 9 meetins, after which point the dosages in reduced to 2,000 units 3 times weedly for an additional arrowth Bodybudens and additions to 1000 offert on cycle, in an effort to mariant institution from the contraction of the contract

### Post-Cycle:

Harma Chronic genderops in solen used with other medications as part of an in-depth Post Cycle Theory (PCT) program Genesic on restoring enlargence testosteroes production more rapidly at the end of a stead cycle. Bestoring enlagonous testosteroes production in a special concern at the condision of each cycle, it sime where subscrain physique. But min concerns the action of corteol, which in many ways is blundered out by the effect of arthogram. Cortisel sends the opposite message to the musics time testosterone, or to breakdown protein in the cell. Let urchecked by a low best of testosterone, or and can quickly simp mark of your new marker massage. Protects the contents control can quickly simp mark of your new marker massage. Protects the control control can applied by marker of your new marker massage. the pose-cycle the OLEA Spectrumy can be the administration of 2000-2000 Units every critic or 3rd day, taken for no longer than 2 or 3 weeks. Thread fix too long or at no high a doos, the drug may actually function to desensitize the Leydig's cells to Interitoring hormone, further Indicing a return to hornestassis. For a more comprehensive wire of ECGs note in a proper hormonal-recovery program places refer to the Post Cycle Therapy section of this book.

### On-Cycle:

Bodybalders and athlete may also administ Flamm Chronic Conschoroph troughts a stord cycle, in an offer to avoid selection trapply and the reading reduced ability to respond to 114 strainals. In effect, this practice is used to avoid the problem of storiate analysis, instead of rejing to cornect it kine on when the cycle is one. It is important to remarker that the dosage needs to be carefully monitored with this type of use, as high levels office of the contraction of the contractio

These on cycle IcG protocols were developed by D. Abin Criste, a well-known figure in the unti-sign and hemore-oxplications field, for see with its indiscontence replications that the unti-sign and hemore-oxplications field, for see with its indiscontence replications that the complex is a common countrie compliate of puttiess respective of the ministence of normal analogous levels. De Criste's IcG programs designed to alleviate this concern in a meaner that is acceptable for longar-time to Ecr for lose intensional proceeds the IcG does in relation to a prescribed teachistence replication program. De Criste's program of the Criste's process of the Criste's program of the Criste's program of the Criste program of the Cris

### Administration (Women):

When used to induce oxidation and pregnancy in anoxidatory infertile woman, a dose of 5,000 to 10,000 units is administered one day following the last dose of memotropins. The trining is specific so that the hormone is given precisely at the right moment in the oxidation cycle. Human Chorionic Gonadotropin is not used by women for physique- or performance-refunctive numeral.

### Availability:

When we find IrCG, we see it is always packaged in 2 different visikarmpules (one with a reported produced must be reported produced must be reported produced must be reported produced must be reported produced by the reported produced produced by the reported produced produced produced produced for later use. Make sure your product must be descriptor. Harmon Cortonic Grounderprops in which your mediatent, and easily obtained on the black market. To date, counterfeits have not been much of a cornern, although a couple of oddities have more done of unfail in malfe loss visik).

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# Glossary

# AAS - Ambolic/Androgenic Steroid

Acute - Short duration. An acute effect is one that occurs rapidly, not after long-term administration

Aerobic - Refers to a process where oxygen is used to generate energy in the muscles from carbohydrates, fats, and protein, in the form of ATP (adenosine triphosphate). Long distance numing is an example of an aerobic activity.

### Adipose - Fat tissue.

properties

Adrenocentor - A type of receptor in the body involved in the regulation of heart rate. metabolism, and thermogenesis. Stimulated by endogenous catecholamines such as epinephrine (adrenaline), norepinephrine (noradrenaline), and dopamine,

Agonist - A substance that initiates a biological response. An estrogen agonist acts as an estrogen in the body.

Ampule - A glass container that holds a single dosage unit of a liquid drug product. An ampule must be broken open before use.

Anabolic - A process that involves the building of tissues such as muscle and bone. AAS are most valued by bodybuilders and athletes for their anabolic properties.

Anaerobic - Refers to a process where energy is generated in the muscles from carbohydrates in the form of ATP (adenosine triphosphate) without the use of oxygen. Weight lifting is an example of an anaerobic activity.

Androgenic - Refers to the musculinizing properties of a substance. AAS stimulate male libido, secondary hair growth, acne, and male pattern hair loss via their androgenic

Antagonist - A substance that inhibits a biological response. An estrogen antagonist blocks the action of estrogen in the body.

Arrhythmia - An irregular heartbeat. Arrhythmias may be life threatening or benign in nature.

Atherogenic - Promoting the formation of plaque deposits on the walls of arteries.

Atherosclerosis — A progressive cardiovascular disease characterized by the buildup of plaque deposits in the arteries. This may obstruct blood flow, causing heart attack or stroke.

Bacteriostatic – Inhibits the growth of bacteria. Bacteriostatic water contains ingredients that prevent bacteria from contaminating the liquid.

Contraindicated - Not advisable for use. A contraindication is a condition that would prevent someone from using a particular drug product.

Diabetogenic - Increases blood sugar.

Diastolic - The phase of blood circulation where the pumping chambers of the heart (ventricles) are being filled. Pressure is at its lowest during the diastolic phase.

Downregulate — To reduce in number. Some cellular receptors downregulate with high levels of drug stimulation, inducing tolerance. Anabolic/androgenic steroids generally do not cause a reduction in respective androgen receptor concentrations. Classic downregulation does not occur.

Edema - The retention of excess water in the circulatory system and/or body tissues.

Endogenous - Occurring naturally within the body. Testosterone produced by the testes is an endogenous hormone.

Epiphyses – The growth plates at the end of long bones. The epiphyses plates regulate increases in linear height during development, and fise at maturity preventing further linear growth.

Erythropoiesis – The process in which red blood cells are produced in the body. Anabolic/androgenic steroids can stimulate erythropoiesis.

Esterified – Refers to a steroid compound that has one or more fatty acids attached to the molecule, usually to slow its release from an injection site. Testosterone expionate is an exterified from of festosterone.

Exogenous - Caused by an agent outside the body.

Glycemic - Relating to blood sugar levels.

Hepatotoxic – Liver toxic. All c-17 alpha alkylated anabolic/androgenic steroids are considered henatotoxic.

Homeostasis - A state of equilibrium among physiological processes.

Hyperplasia - Growth that occurs via an increase in cell number.

Hypertrophy - Growth that occurs via increases in existing cell size.

proinflammatory state, and increased risk of cardiovascular disease.

Anabolic/androgenic steroids produce growth through hypertrophy.

Metabolic Syndrome — A poorly defined common metabolic disorder characterized by abdominal obesity, an atherogenic lipid profile, insulin resistance, elevated blood pressure, a

Multi-dose Vial — A vial with a rubber stopper on the top, designed to be pierced repeatedly by a needle (for multiple uses).

Pathological - Involving or caused by physical disease.

Peripheral – Near the surface of the body. Psychoactive drugs are often regarded as centrally acting (brain/central nervous system), while hormones like anabolic steroids affect both central and peripheral tissues (such as muscle and skin).

Prognostic - Serving to predict the likely outcome of a disease.

Pulmonary - Related to the lungs.

Recombinant – Refers to a synthetic manufacturing technology that involves the splicing of genes or DNA segments and inserting them into a cell culture in order to replicate a specific protein. Recombinant DNA technology is used to manufacture many protein-based drug products including human growth hormone, insulin-like growth factors, and human insulin.

Selective - Describes a drug with a very specific effec, and little spillover into other biological systems.

Subcutaneous - Located beneath the skin and above the muscle

Supraphysiological – In excess of normal biological levels. High doses of testosterone produce supraphysiological levels of hormone in the blood.

Supratheraneutic - In excess of normally defined theraneutic levels.

Systemic - Affecting the entire body through general circulation.

Systolic - The phase of blood circulation where the pumping chambers of the heart (ventricles) are actively pumping. Pressure is at its highest during the systolic phase.

Upregulate — To increase in number. Usually relating to cellular receptor concentrations. Anabolic/androgenic steroids can increase respective androgen receptor concentrations, possibly increasing sensitivity to androgens.