The andropause: fact or fiction?

Nicholas Burns-Cox, Clive Gingell

The menopause is a condition with a proven pathophysiological aetiology, leading in some women to unpleasant symptoms and bone loss over the ensuing years. The relatively sudden onset of ovarian failure causes amenorrhoea, infertility and oestrogen deficiency. There is no analogous process in the male. However, the term andropause is being increasingly used to describe a collection of symptoms including lack of energy, depression, decreased libido and erectile difficulties occurring in middle-aged or elderly men with a testosterone level at the lower end of normal range. The term andropause, like menopause, implies a state of hormone deficiency secondary to gonadal failure and the similarity of the two terms tends to give the andropause some unproven credibility. The incidence of erectile dysfunction increases with age and it is tempting to correlate this with the relative decrease in testosterone seen with age. It is therefore likely that androgen supplementation will be used to try and boost testosterone levels in middle-aged men with erectile difficulties who are not hypogonadal. There is, however, no evidence that the increase in erectile difficulties with age is related to decreasing testosterone levels in these eugonadal men. In this article we review the action of testosterone, the changes that occur with aging and the likely benefits and risks of supplementation.

Androgen production and aging

Testosterone is a C19 steroid and 0.24 μmol/day are secreted by the Leydig cells of the testicle. Its production is regulated by a negative feedback loop involving luteinising hormone and luteinising hormone-releasing hormone (LHRH), and forming the hypothalamic–pituitary–gonadal axis. Androgens are also secreted by the adrenal cortex in men (0.002 μmol/day), mainly as androstenedione but the quantities are of little clinical significance in comparison with the amount secreted by the testes. Testosterone is metabolised by the action of 5-alpha-reductase to to the biologically active androgen dihydrotestosterone and by aromatase activity to oestradiol. Secretion of testosterone begins in the foetus with a peak in the male at 12 weeks. There is another peak after birth and a low level equal to the female remains until puberty. At puberty increased pulsatile secretion of luteinising hormone leads to maturation of the Leydig cells and hence to increased testosterone production.

The action of the testosterone is firstly androgenic leading to the secondary sexual characteristics of the adult male. It also has an anabolic effect on receptive tissue leading to increased muscle bulk. The great majority of testosterone (80%) is bound to sex-hormone-binding globulin (SHBG) and to a lesser degree to albumin and cortisol-binding globulin. Only about 2% of total testosterone remains free, and it is this free portion that is thought to be of biological importance. There is debate as to which fractions are bioavailable but it is generally considered that the non-SHBG bound fraction is biologically active. The other measure of bioavailability is the Androgen Index which is the ratio of total testosterone to SHBG multiplied by 100, and is comparable to the free testosterone concentration. In males, there is a progressive decrease in testosterone levels with age (table 1). However, there is a concurrent increase in SHBG and therefore a greater decrease in the concentration of the free testosterone, which is reported to fall at a rate of 1% per year. The levels of dihydrotestosterone responsible for many of the actions of testosterone at the cellular level do not show a decrease with age. Although the decrease in testosterone is seen in healthy elderly men, a greater decrease in testosterone is seen in people with concomitant illnesses, which are very common in this age group.

There is evidence for some degree of testicular failure with advancing years. There is a decrease in the number of Leydig cells, impaired testicular perfusion, decreased steroid response to stimulation by beta-human chorionic gonadotropin, and a moderate rise in luteinising hormone levels. But there is also evidence that the decrease in testosterone is not all of primary testicular...
Table 1  Influence of age on plasma testosterone, free testosterone and SHBG levels in man

<table>
<thead>
<tr>
<th>Age (years)</th>
<th>n</th>
<th>Testosterone (nmol)</th>
<th>SHBG (10^-8 M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>18–29</td>
<td>105</td>
<td>23.1 ± 6.8</td>
<td>4.2 ± 1.2</td>
</tr>
<tr>
<td>30–49</td>
<td>30</td>
<td>21.4 ± 7.3</td>
<td>4.5 ± 1.4</td>
</tr>
<tr>
<td>50–59</td>
<td>24</td>
<td>17.0 ± 8.0</td>
<td>5.5 ± 1.3</td>
</tr>
<tr>
<td>60–69</td>
<td>63</td>
<td>16.3 ± 6.6</td>
<td>6.0 ± 1.4</td>
</tr>
<tr>
<td>70–79</td>
<td>63</td>
<td>16.5 ± 7.3</td>
<td>7.3 ± 1.8</td>
</tr>
<tr>
<td>80–89</td>
<td>60</td>
<td>15.0 ± 5.7</td>
<td>7.3 ± 2.0</td>
</tr>
</tbody>
</table>

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origin, the hypothalamic–pituitary–gonadal axis is also implicated. There is loss of the circadian rhythmicity of serum testosterone\(^\text{18}\) and increased sensitivity to negative feedback by the sex hormones on gonadotropin secretion.\(^\text{19}\)

However, although there is a decrease in both total and free testosterone with age, the decrease is gradual and levels do not fall outside the normal range.\(^\text{7}\)

The variability between individuals is great and some elderly men have the same levels as young men, indicating that, unlike the ovary after the menopause, the testicle continues to secrete significant quantities of testosterone throughout life. In contrast to the post-menopausal ovary, the testicle continues gamete production and fertility persists life-long. Although a loss of fertility potential is not a typical finding in the elderly, with increasing age the sperm count is reduced due to a decrease in Sertoli cells\(^\text{15,20,21}\) and some decrease in motility is reported.\(^\text{1,22}\) Nevertheless, it seems that the zona-free hamster egg fertilizing capacity of the spermatozoa of elderly men is similar to that of young men. Although there is no sudden degeneration of the testicle analogous to that of the ovary, there is still a definite progressive decrease in the ability of the testes to produce spermatozoa and testosterone. The reduction of fertility in the elderly male rarely becomes of clinical significance. The decrease in testosterone secretion, however, is thought by some clinicians to be responsible for a group of symptoms ascribed to the andropause. These include nervousness, insomnia, hot flushes, mental and physical tiredness, irritability, reduced libido and erectile dysfunction. There are age-related changes, including decreased muscle mass and body hair, and disturbance of bone density and haemopoiesis\(^\text{8}\) have also been described as being due to testosterone deficiency.

**Hormone replacement and supplements**

The case for hormone replacement in men and children who are truly hypogonadal is generally accepted as beneficial.\(^\text{23–25}\) There are a variety of types and modes of administration of testosterone. The currently available compounds for androgen replacement are shown in table 2. Some of the early oral preparations (eg, methyltestosterone and Danazol) are alkylated at the 17-

Table 2  Summary of the current available testosterone compounds for androgen replacement

<table>
<thead>
<tr>
<th>Type of testosterone</th>
<th>Name (Trade name)</th>
<th>Route of administration</th>
<th>Dose (mg)</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Testosterone esters</td>
<td>Testosterone</td>
<td>oral capsules</td>
<td>40-120</td>
<td>once daily</td>
</tr>
<tr>
<td></td>
<td>undeconoate (Restandol)</td>
<td>oral</td>
<td>25, 40</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mesterolone</td>
<td>oral</td>
<td>3–4 times a day, reduced for maintenance every 2 weeks</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Testosterone</td>
<td>intramuscular injection</td>
<td>40</td>
<td></td>
</tr>
<tr>
<td>propionate (Sustanon)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Testosterone</td>
<td>−</td>
<td>implant</td>
<td>600</td>
<td>6-monthly</td>
</tr>
<tr>
<td>Testosterone</td>
<td>(Testop)</td>
<td>transdermal on scrotal skin only</td>
<td>3.6</td>
<td></td>
</tr>
<tr>
<td>Testosterone</td>
<td>(Andropatch)</td>
<td>transdermal patch placed on any skin which does not cover a bony prominence</td>
<td>2.5</td>
<td>once daily</td>
</tr>
</tbody>
</table>
hydroxy position, making the androgen more resistant to hepatic metabolism. However, these oral preparations often give unpredictable serum levels of testosterone and are more likely to cause side-effects, including cholestatic jaundice, and can rarely lead to hepatic failure (haemorrhagic liver cysts). For these reasons they are no longer available in the UK for androgen replacement. Alternatively, testosterone can be esterified and given as an oral agent daily (eg, mesterolone), a depot injection (eg, testosterone propionate) every two weeks or as a surgically implanted pellet every six months. These result in a sustained level of plasma testosterone. The availability of testosterone patches in the UK will provide a more physiological and ‘user friendly’ delivery system for patients requiring androgen replacement therapy and has been shown to be very effective.

Support for hormone supplementation comes from trials on men who have moderately low, age-related levels of testosterone. Potentially advantageous increases in lean body mass and insulin sensitivity have been shown, as have increases in fibrinolytic activity. Androgens have been shown to affect the balance of blood lipids. Men generally have lower levels of high-density lipoproteins (HDLs) and higher levels of low density lipoproteins (LDLs) than premenopausal women and the administration of testosterone in hypogonadal men further decreases the HDL levels. However, the type of testosterone administered determines the effect on lipids. The orally active 17-alkylated testosterone preparations cause a markedly lower HDL concentration and raised LDL concentration; this is not seen with the parenterally administered testosterone esters. Indeed, Tenover et al reported an 11% decrease in LDL with testosterone ester supplementation; the significance of this finding for cardiovascular disease is unclear, however. Conversely there have been reports of associations between low testosterone and increased adiposity, insulin resistance, low fibrinolytic activity, and hepatic enzyme induction. The importance of these associations is unclear. Some authors consider that the low testosterone may be the result of ill health and not a contributing factor to it. The effect of aging on sexual function and its relationship to androgen production is difficult to quantify. Certainly there is a decrease in sexual activity with age but many factors are involved, including psychological (eg, fear of failure, boredom with partner) and organic (eg, decreased penile sensation and impaired blood supply to the penis) ones. Korenman et al showed that the levels of both testosterone and gonadotropin are similar in potent and impotent elderly men. Both groups show an impaired response to LHRH stimulation as compared with younger men. The authors concluded that hormonal changes and impotence were two independent conditions in older men.

Individuals complaining of decreased well-being, loss of sexual interest and erectile difficulties have symptoms that are common, nonspecific and frequently associated with stress. It has been reported that well-being and sexual interest can be increased by testosterone supplements. However, of the 10 men who objectively showed a small improvement, only three felt that the therapy was adequate. There is also clinical evidence from trials that androgen supplementation has no beneficial effect in men with erectile dysfunction who are eugonadal. With regard to erectile dysfunction, there is evidence that the normal process is androgen-independent. Hypogonadal men have been shown to have a normal erectile response to erotic stimuli. Further, a group of sexual offenders on the anti-androgen cyproterone acetate showed no decreased erectile response when shown erotic films. There is little evidence to date to support the concept that androgen supplements in eugonadal men are of any significant benefit.

The role of testosterone in prostate cancer

It is known that prostate cancer is androgen-dependent and that the principal therapy of advanced disease is the removal of androgen stimulation. This is achieved by medical or surgical castration or the administration of anti-androgen, alone or in combination, so-called maximum androgen blockade. It is likely therefore that the administration of androgen to a man with prostate cancer will stimulate activity in what is mainly a hormone-dependent tumour. Certainly in the first 10–12 days after the administration of an LHRH analogue the serum testosterone rises and so-called ‘tumour flare’ results, in which bony metastases become more painful, paraplegia may be precipitated, and ureteric obstruction may occur. It is therefore likely that giving testosterone to men with latent cancer within the prostate may promote the development of active clinically evident disease. It is consequently of paramount importance to check the plasma prostate-specific antigen (PSA) level before the administration of testosterone to any man greater than 50 years and to check this regularly on...
treatment. Unfortunately, there is no clear cut-off point, but a rising PSA on treatment should arouse suspicion. In a study of 33 men with low serum levels of free or total testosterone aged 45–75 years, with normal age-adjusted PSA levels and negative digital rectal examination, transrectal ultrasound-guided biopsies of the prostate were undertaken. 49 Six (18%) were discovered to have cancer, five of whom were aged between 62 and 65 years old with a PSA of less than 3 ng/l. These results raise serious concerns about the use of androgen therapy in men over 60 years old. The authors speculated that low androgen levels may have falsely lowered the PSA into the normal range in some of these men. They concluded that prostate biopsy should be mandatory prior to testosterone therapy in men over 60 years old and strongly encouraged men in between 50 and 59 years old.

Conclusion

Evidence-based medicine demands that the extent of the risks be known and the benefits quantified by controlled trials. The symptoms of fatigue (mental and physical), loss of energy, depression and decreased libido which constitute the so-called andropause, can readily be explained by stress and there is no convincing, scientifically valid, placebo-controlled study that shows any benefit for testosterone supplements in these patients.

References

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Postgrad Med J 1997 73: 553-556
doi: 10.1136/pgmj.73.863.553

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