The aging male project

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Abstract

With an increasing life expectancy and a decreasing reproduction rate, the population structure changes. A Jenapharm R & D program investigates the endocrinology of aging men. In men, a decrease in production of sex steroids and other hormones with age can be observed. The typical patterns of daily rhythmicity become less distinct. This is part of a very complex picture in which not only isolated hormones are involved, but also the influence of hormones on each other. Many factors from the external and internal environment mediated by neurotransmitters constantly affect the highly sensitive hormonal balance. Therefore, aging has also been defined as "the gradual dysfunction of homeostatic processes". Declining testosterone (T) levels are involved in 'andropausal' symptoms in men: loss of libido, erectile dysfunction, insulin receptor resistance, obesity, osteoporosis, disturbances of lipid metabolism, myocardial and circulatory disturbances, impaired well-being and mood. Data are derived from studies in hypogonadal men treated by T replacement. In such patients under T treatment libido increases, fat mass decreases, muscle strength, bone mineral density and erythropoiesis increase. Whether the symptoms of andropause in aging men could successfully be treated by T substitution remains to be investigated. Negative effects of T, especially on the prostate and the cardiovascular system, are under discussion. There is increasing evidence that low T levels seem to be a risk factor for both the prostate and the cardiovascular system. Jenapharm's new testosterone undecanoate formulation for intramuscular injection can be administered every three months. T levels remain within the physiologic range. No supraphysiologic peaks occur. In women, estrogens have beneficial non-genital effects. Studies concentrate on synthetic estrogens for men without feminizing properties such as gynecomastia and reduced testicular size. Several derivatives of 17-alpha estradiol have been synthesized some of which show selectivity for the central nervous system. CNS effects have been demonstrated in female and male animals. Cardiovascular protection by estrogens has been shown in animal and human studies. Atherosclerotic plaque size was reduced after estrogen injections in cholesterol-fed rabbits. Phytosterogen-fed monkeys had lower total cholesterol and LDL cholesterol and higher HDL cholesterol. Apart from atherosclerotic lesions, coronary artery vascular reaction was improved. Some of these experimental findings were confirmed in human studies in postmenopausal women with and without estrogen replacement. Whether all of the described estrogenic effects can be seen in men remains to be investigated. (Med J Indonesia 2001; 10: 127-33)

Keywords: aging, andropause, testosterone, estrogens

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With an increasing life expectancy and a decreasing reproduction rate, the population structure changes in a dramatic way. In 2000, we see for the first time in history an equal distribution in percentages of under 15 yr olds and over 60 yr olds in the developed countries. It is predicted that in Japan, for example, as soon as in 2010 more than one third of the population will be older than 60 years. According to the calculations of United Nations, the same change will occur in the less developed regions with a delay of about 50 years. The latest statistics indicate that this may happen even sooner.

For these reasons, scientific investigation of the phenomenon of aging not only from a social and financial, but also from a biological and medical point of view needs more and more attention. Jenapharm and its mother company Schering are primarily focused on the endocrine aspects of aging. Looking back at several decades of research and developmental work in the area of female menopause (leading to hormone replacement therapy, HRT), it is now our goal to learn more about the endocrinology of aging men.

In both genders, a decrease in production of sex steroids, DHEA, growth hormone, and melatonin can be observed. An anatomical equivalent is the reduction of size of the pituitary gland with age. Levels of corticosteroids remain stable. This leads to a relative hypercortisolemia. Not only total hormone production decreases but also the typical patterns of daily rhythmicity become less distinct. These findings become even more complex by the fact that the three axes of hormonal regulation involving hypothalamus, pituitary gland and target organs, influence each other. The CRH-ACTH-corticoidoestrogen axis seems to have a suppressive effect on both the GHRH-GH-IGF I and the GnRH-FSH/LH-sex steroid axes. In addition, many effects from the external and internal environment mediated by neurotransmitters in the central nervous system constantly affect the highly sensitive hormonal balance. Therefore, aging has also been defined as "the gradual dysfunction of homeostatic processes."

**TESTOSTERONE**

Declining testosterone levels are involved in a variety of symptoms in men: loss of libido, erectile dysfunction, insulin receptor resistance, abdominal obesity, osteoporosis, disturbances of the lipid metabolism, myocardial and circulatory disturbances, impaired well-being and mood. Hajjar et al. showed that perceived libido improved significantly in older hypogonadal men after long-term testosterone replacement. Serum testosterone concentrations in obese (BMI > 28) men were lower than in non-obese (BMI < 27) subjects. Several investigators showed that in hypogonadal men receiving testosterone treatment body fat decreased. At the same time, bone mineral density increased. Thus, testosterone replacement in hypogonadal men is of crucial importance in the prevention of osteoporosis and the postponement of the fracture threshold within the life span.

Although the symptoms of andropause may not be exclusively testosterone dependent, testosterone plays a crucial role. It would certainly be too soon to draw the conclusion that therapeutic administration of testosterone can improve or cure all these symptoms. Numerous studies will be necessary to prove the potential benefits of testosterone treatment in aging men.

For a long time testosterone was associated with negative effects, especially on the prostate and the cardiovascular system. This opinion needs to be revised. Testosterone does not cause prostate carcinoma but enhances the proliferation in approximately 80% of prostatic carcinomas that are already present. There is not a single case described in which testosterone replacement in hypogonadal men has caused prostatic cancer. There is no evidence that testosterone can cause growth of the prostate exceeding what is considered normal. The contrary could be true because both benign prostatic hyperplasia (BPH) and prostatic cancer occur increasingly at an age when testosterone levels decline.

There are also new findings concerning the cardiovascular system. Whereas it had been assumed for a long time that testosterone supports atherosclerosis, there is now evidence that the incidence of coronary heart disease often coincides with low testosterone levels. Low testosterone levels seem to be a risk factor for both the prostate and the cardiovascular system. Jeppesen et al. showed that in patients with acute ischemic stroke those with higher testosterone levels had a better 6 months survival rate than those with lower testosterone levels. In more than a million users of androgenic anabolic steroids in the U.S. within a period of 15 years (1976 - 1991),
only 26 cases of fatal cardiovascular events have been reported.\textsuperscript{13,14} It would definitely be too soon to come to a final conclusion but it should be kept in mind that old prejudices need to be questioned.

Different causes of the decline of testosterone levels with age have been discussed. One possible cause could be the change of intratesticular regulatory factors leading to an increased conversion of pregnenolone to progesterone instead of testosterone in the Leydig cell. Another hypothesis is a failure of appropriate GnRH secretion by the hypothalamus.\textsuperscript{15} Decreased androgenicity of physiologic serum testosterone concentrations may also lead to symptoms of andropause despite of impaired testosterone production and secretion.

Testosterone levels show very sensitive reactions to exogenous stress factors. For example, experimentally induced metabolic stress led to a significant decline of testosterone levels within less than one hour.\textsuperscript{16} Strollo et al. reported that astronauts had significantly lower testosterone levels inflight as compared to preflight.\textsuperscript{17} Even winning and losing small amounts of money in gambling had almost immediate effects on testosterone levels.\textsuperscript{18}

Jenapharm is currently developing a testosterone ester for intramuscular injection, the well-known testosterone undecanoate. This formulation has major advantages compared to preparations that are currently available: It can be administered every three months with testosterone levels remaining within the physiologic range (Figure 1). There are no supraphysiologic peaks as observed with testosterone enanthate.\textsuperscript{19-22} The ups and downs of testosterone levels are held responsible not only for mood swings that seem to be quite significant in many patients, but high peak levels may also be the reason for an elevated hematocrit.

In the near future, we will concentrate on the identification of dissociated or tissue specific androgens. These are steroids that will ideally have all of the desired androgenic properties but none of the less desired. There are already compounds which are in principle dissociated or tissue-selective androgens, e.g., anabolic androgens with a three-fold stronger effect on the muscle compared to native testosterone but with less than half of the genital effects. In addition to its selective properties, such a molecule should be orally available to avoid injections.

**ESTROGENS**

There are controversial results and opinions whether estrogen levels in men decline with age.\textsuperscript{23} Whereas some investigators claim that estrogens remain stable,\textsuperscript{24-27} more and more papers show an age-related decline in estrogen secretion.\textsuperscript{28} Our own experience shows that the correct measurement of estrogens
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depends significantly on the hormone essays employed. Of the circulating estrogens in men, 75 – 90% arise from peripheral aromatization, mainly in adipose tissue, of both testosterone to 17-beta-estradiol (E2) and androstenedione to estrone (E1). The testes synthesize the remaining 10 – 25%.

From the experience with hormone replacement therapy (HRT) in peri- and postmenopausal women it is well accepted that estrogens have beneficial effects on the cardiovascular system and on the central nervous system. We have established a program to investigate whether estrogens could have similar effects in the male gender. From studies as well as therapeutic experience in patients with hormone-dependent prostatic carcinoma who have been treated with high doses of estrogens it is known that ‘natural’ estrogens such as 17-beta-estradiol can lead to feminizing effects, e.g. gynecomastia and reduction of testicular volume. Similar findings have been reported from studies in male-to-female transsexuals who also receive high doses of estradiol. Therefore we are working with synthetic estrogen compounds that show a reduced genital effect compared to native estradiol.

A lead compound displaying this particular property is 17-alpha-estradiol. Based on this molecule we have synthesized a number of derivatives for our further studies (Figure 2 and 3). Some of these substances show a remarkable selectivity for the central nervous system. Several mechanisms of the neuroprotective functions of estrogens are being discussed: Genomic effects mediated by transcription factors, antioxidant activities, and modulation of neuronal membranes.

Figure 2. Effects of chronic estrogen administration on the retention of conditioned avoidance behaviour in ovariectomized rats I

Figure 3. Effects of chronic estrogen administration on the retention of conditioned avoidance behaviour in ovariectomized rats II
CNS effects of estrogens have been demonstrated in vitro in dopaminergic neurons using cell cultures. Experimentally induced oxidative stress caused significant cell death which could be prevented by preincubation with both 17-beta- and 17-alpha-estradiol. The effects of estrogens on the CNS could be shown in a number of animal experiments. For instance studies in which cognitive functions of animals including the ability to learn as well as memorise are assessed before and after estrogen treatment. These effects could be shown in female and male animals. Estrogen-treated male-to-female transsexuals showed improved learning scores compared to a control group awaiting treatment. The Max Planck Institute of Psychiatry in Munich sees several parallels between estrogens and Alzheimer's disease.

A human model has been described by the group of Carani from Modena, Italy. They have identified a patient with an aromatase deficiency. This patient cannot metabolise testosterone to estradiol. Treatment with estradiol significantly changed several psychological parameters and aspects of sexual behavior.

In this patient, it was also very well demonstrated that estrogens are of crucial importance for the bone. When he was diagnosed at approximately age 30, his epiphyses were unfused. The man did not respond to testosterone treatment but to treatment with estradiol. Other studies have demonstrated the role of estrogens in idiopathic osteoporosis and, since most of the subjects investigated had physiological testosterone levels but low estrogens, it can be hypothesized that aromatase deficiency of varying degrees could be the cause.

Cardiovascular protection by estrogens has been shown in animal experiments and human studies. Atherosclerotic plaque size was reduced after estrogen injections in cholesterol-fed rabbits (Figure 4). In rabbits and monkeys, feed containing phytoestrogens in soya products showed the same effect. Moreover, the phytoestrogen-fed animals had lower total cholesterol and LDL cholesterol and higher HDL cholesterol. These findings could be confirmed in humans.

Apart from atherosclerotic lesions, coronary artery vascular reactivity (vasodilatation) was improved in estrogen-fed monkeys. Some of these experimental findings were confirmed in human studies in postmenopausal women and men with and without estrogen treatment. Again, different possible mechanisms of rapid vasomotor effects of estrogens are being discussed: (1) nonreceptor mediated effects such as direct effects of E2 on membrane function or ion channels; (2) effects mediated by a novel, as yet undescribed estrogen receptor; and (3) effects mediated by one or both of the classical Ers acting in a novel manner.

More recently, musculoprotective effects of estrogens have been described in animal experiments in rats. From in vitro studies in peripheral blood mononuclear cells, a stimulatory effect of estrogens on immunoglobulin production has been reported.

Whether all of the described estrogenic effects can be seen in men remains to be investigated. We hope to be able to start our first human studies with non-feminizing estrogens within the next two years.

REFERENCES


