Clinical implications of altered thyroid status in male testicular function

Implicações clínicas das alterações tireoidianas na função gonadal masculina

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ABSTRACT

Thyroid hormones are involved in the development and maintenance of virtually all tissues. Although for many years the testis was thought to be a thyroid-hormone unresponsive organ, studies of the last decades have demonstrated that thyroid dysfunction is associated not only with abnormalities in morphology and function of testes, but also with decreased fertility and alterations of sexual activity in men. Nowadays, the participation of triiodothyronine (T3) in the control of Sertoli and Leydig cell proliferation, testicular maturation, and steroidogenesis is widely accepted, as well as the presence of thyroid hormone transporters and receptors in testicular cells throughout the development process and in adulthood. But even with data suggesting that T3 may act directly on these cells to bring about its effects, there is still controversy regarding the impact of thyroid diseases on human spermatogenesis and fertility, which can be in part due to the lack of well-controlled clinical studies. The current review aims at presenting an updated picture of recent clinical data about the role of thyroid hormones in male gonadal function.

Keywords
Testis; thyroid hormones; spermatogenesis; reproduction; fertility

INTRODUCTION

In mammals, altered thyroid status is known to adversely affect many organs and tissues. Nevertheless, for many years, the impact of thyroid disorders on male reproduction remained controversial. Early studies in the 1950’s demonstrated that testes were essentially independent of thyroid hormone effects (1,2). For this reason, the potential of thyroid hormone in the modulation of male reproductive function was not determined.

However, in the past two decades, clinical studies have demonstrated that thyroid hormone plays an important role in testicular development and function.
It is now established that T3 regulates the maturation and growth of testis, controlling Sertoli cell and Leydig cell proliferation and differentiation during testicular development in rats and other mammal species (3,4). The efficiency of spermatogenesis, reflected by daily sperm production in adulthood, correlates to the total number of functional Sertoli cells established during prepubertal life (5). Furthermore, changes in thyroid hormone levels during early testis development have been shown to affect testicular maturation and reproduction later in life (6). These data, in conjunction to the findings that thyroid hormone receptors are present in human and rat testes from neonatal to adult life (7,8), confirm that thyroid hormone plays a key role in testicular development. The presence of iodothyronine deiodinases, enzymes that modulate the concentration, and thus the action of thyroid hormones in different tissues, were also recently identified in the rodent testis from fetal to adult life (9-11). Clinical literature indicates that most patients with thyroid hormone disorders experience some kind of sexual dysfunction, which improves or normalizes when patients become euthyroid (6,12,13). Hence, although thyroid hormone was not historically viewed as a major regulator of the male gonad, it is clear now that it has critical effects on the testis, especially during development period (14). This review intends to present an updated view of clinical studies regarding the effects of thyroid hormones on testicular function.

**THYROID HORMONES**

The thyroid hormones, thyroxin (3,5,3',5' L-tetraiodothyronine or T4) and the more potent triiodothyronine (3,5,3'-L-triiodothyronine or T3) are synthesized in the thyroid gland. Most of the hormone released is in the form of T4, as total serum T4 is 40-fold higher than serum T3. The main pathway for the production of the bioactive form, T3, is via outer ring deiodination of the prohormone T4 by iodothyronine deiodinases (15). This activation reaction is catalyzed by types 1 and 2 deiodinases (D1 and D2) in peripheral tissues. Both T4 and T3 may also be inactivated by inner ring deiodination to 3,3',5-triiodothyronine (rT3) and 3,3'-diiodothyronine (T2) predominantly by type 3 deiodinase (D3), and to a lesser extent by D1 (16). Deiodination is the most important pathway of thyroid hormone metabolism not only in quantitative terms, but also because it accounts for most of the circulating T3 (~ 80%) in humans (17).

The serum concentrations of thyroid hormones are regulated by a negative feedback system that involves the hypothalamus, pituitary and thyroid gland (H-P-T axis). Both the hypothalamic thyrotrophin-releasing hormone (TRH) and pituitary thyroid-stimulating hormone (TSH) secretions are negatively regulated by circulating thyroid hormones (18). The majority of thyroid gland–released hormones circulate in blood in order to carry proteins such as thyroxin binding globulin (TBG), albumin, and thyroid binding prealbumin. Only 0.03% of total serum T4 and approximately 0.3% of total serum T3 circulate as free (unbound) hormones.

Although thyroid hormones may exert their effects on a number of intracellular loci (16), their actions on target tissues are predominantly mediated by specific nuclear receptors (TRs) able to bind to regulatory regions of target genes modifying their expression (19). Two separate genes, TRα and TRβ, encode thyroid hormone receptors (TRs). Similarly to other nuclear receptors, TR consists of a DNA-binding domain, ligand-binding domain and amino-terminal transactivation domain. The remaining non-hormone binding TR isoforms lack portions of the DNA-binding and/or ligand-binding domains, and their functions are still unclear (20). In the nucleus, TRs recognize and bind to specific DNA sequences termed thyroid hormone response elements (TREs), located in the promoter region of target genes, and activate or repress transcription in response to T3 binding (Figure 1).
In order to interact with specific nuclear receptors and generate a biological response, thyroid hormones have to cross cell membranes. It was originally believed that thyroid hormones, due to their lipophilic nature, enter target cells by passive diffusion. Currently, however, there is growing evidence indicating that T4 and T3 cross the plasma membrane by carrier mediated mechanisms (20,21). Several membrane transporter families have been identified, however only monocarboxylate transporter (MCT) 8, MCT 10 and organic anion-transporting polypeptides (OATPs) demonstrate a high degree of specificity towards thyroid hormone (22). The OATPs form a novel family of transporter proteins that have been detected in several tissues, including testis, in rodents and humans (22-28).

**THYROID HORMONE AND TESTICULAR FUNCTION**

The testes are mainly comprised of tightly coiled seminiferous tubules, which are supported by loose interstitial connective tissue in which the steroidogenic Leydig cells are located (29). Inside the basement membrane of the seminiferous tubules, there is a columnar epithelium composed of germ cells and somatic Sertoli cells. It is well established that the number of Sertoli cells presented at puberty is closely correlated with both adult testicular size and sperm output, which is crucial for future male fertility (5). Although several factors are presumed to play a role in proliferation and maturation of Sertoli cells (29,30), T3 is likely to represent a major hormonal signal to Sertoli cell proliferation during testicular development, and ultimately affecting the establishment of the adult Sertoli cell population.

Testes perform two major functions: production of gametes (spermatozoa) and androgen synthesis. The anterior pituitary participates in the control of both of these functions through its secretion of gonadotropins, with follicle stimulating hormone (FSH) acting directly on Sertoli cells and luteinizing hormone (LH) on interstitial Leydig cells (28). Testosterone, the main androgen produced by the latter cells in response to LH, diffuses into the seminiferous tubules, where it is essential for spermatogenesis. Both testosterone and FSH are required for initiation and maintenance of spermatogenesis.

Thyroid hormones are also known to affect Leydig cells proliferation and function (4). Several studies have shown that altered thyroid status has remarkable effects on Leydig cell differentiation in experimental animals (4,30). Likewise, evidence of direct actions of thyroid hormones on Leydig cell steroidogenesis has been demonstrated in different studies (31,32).

**Thyroid hormone disorders and androgenic hormones**

Although gonadotropins are the major regulators of testicular function, there is controversy with regard to the relationship between hypo- and hyperthyroidism and luteinising hormone (LH) and follicle-stimulating hormone (FSH) levels in men. Prepubertal males affected by primary hypothyroidism have been reported with normal (33,34) or, more frequently, elevated FSH and LH (33-37) and total serum testosterone in the prepubertal range (33,36-41). The variations in the levels of LH and FSH have been reported in different studies on adult hypothyroid patients and results are somehow conflicting. Wortsman and cols. (42) investigated eight adult hypothyroid male patients aged 37 to 77 years. All patients had evidence of hypogonadism; five were hypergonadotropic, and the remaining three, hypogonadotropic. Other authors describe elevated (43) or even normal gonadotropin levels in these patients (12,44). Basal serum gonadotrophin concentrations are usually normal in hyperthyroid males. LH and FSH response to exogenous GnRH seems to be significantly greater in thyrotoxic patients in comparison to patients who were rendered euthyroid (44,45). It has been suggested that the LH elevation could be secondary to changes in sex steroid binding and peripheral metabolism, alterations in the hypothalamic-pituitary feedback, or due to the direct effect of thyroid hormones per se at this level (45). All these changes are fully reversible with restoration of the euthyroid state, and require no other specific treatment.

Several studies have shown a drop in the circulating level of testosterone in overt and subclinical hypothyroidism (12,40-44), whereas other studies (46), measuring free testosterone, did not find any change in its level in hypothyroid patients. Jaya Kumar and cols. (43) studied endocrine function of eight males with primary hypothyroidism during the hypothyroid phase and after achieving euthyroidism with T4 replacement levothyroxine therapy, and showed a subnormal testosterone response to hCG leading to reduced steroidogenesis, probably resulting from a testicular lesion induced by hypothyroidism. In contrast, Velazquez and Arata (47) observed normal testicular response of hCG in hypothyroid males. In rare cases of severe prolonged primary hypothyroidi
dism, pituitary hyperplasia may occur causing multiple pituitary hormone deficiencies, including gonadotropin and corticotropin deficiencies (45). Low T3, high TSH and hyperprolactinemia caused by hypothyroidism could directly act on the Leydig cells in suppressing steroidogenesis (42,43,46). All the above changes are fully reversible with restoration of euthyroid state (12). Hypothyroidism is also associated with a decreased serum total testosterone, dehydroepiandrosterone (DHEA), DHEA-sulfate and pregnenolone sulfate (12).

An increase in SHBG is a consistent feature associated with thyrotoxicosis, and leads to an increase in circulating levels of total T4 and reduction in the metabolic clearance rate of testosterone. However, the plasma level of free testosterone is usually maintained within the normal range, which is in keeping with the lack of clinical consequences of the noticeably elevated levels of total testosterone found in thyrotoxicosis (12). Peripheral conversion of androgen to estrogen is enhanced in thyrotoxicosis, probably due to changes in peripheral blood flow (47) rather than a direct effect of thyroid hormones on the aromatase complex. An increase in the production rate of estrogens is also observed in some men with thyrotoxicosis, although it is unclear whether this is due to increased production of adrenal androgen precursors (specifically Δ4A) or to other mechanisms (48). Furthermore, serum progesterone was reported to be higher in hyperthyroid than in euthyroid males (49), while mean basal testosterone bioactivity was lower in patients as compared to controls (50). Thyrotoxic males often present clinical features compatible with exposure to increased estrogen bioactivity (gynecomastia, spider angiomas, and a decrease in libido) (51). Whether this results from alterations in estrogen metabolism is a direct effect of hyperthyroxinemia is unknown.

Thyroid hormone disorders and male reproduction

Thyroid failure in the pre-pubertal period is associated with testicular enlargement as well as alterations in sexual hormones (6,40). Hypothyroidism initiated in infancy may occur in association with macroorchidism without virilization, although the pathogenesis remains uncertain (6). The longer the hypothyroidism persists, the greater is the degree of damage to the testes (6,52). When adequately treated with thyroid hormone, however, boys with congenital hypothyroidism progress through puberty normally and at the appropriate time (53-55).

Morphological changes may be observed in pubertal and adult men testes with chronic untreated hypothyroidism, although the results are conflicting. Griboff (56), investigated five hypothyroid patients and all demonstrated normal sperm counts, whereas other authors (4,34,37,57) found morphological and spermatogenic alterations in testicular biopsies of pre-pubertal and adult hypothyroid patients. Hypothyroid prepubertal testis may present a preponderance of tubular compartment with early onset of spermatogenesis and no increase in the number of Leydig cells. Adult testis, however, may present fibrosis and hyalinization of tubular walls, fibroblastic proliferation, peritubular and interstitial fibrosis with sparse Leydig cells, as well as reduction of tubular diameter, interstitial edema and tubular basal membrane stickiness.

The potential role of thyroid hormone on mammalian spermatogenesis has been addressed by several studies (58). Corrales Hernandez and cols. (59) studied spermatogenesis in ten treated hypothyroid patients. Changes in seminal and sperm profile after discontinuation of hormone replacement were not marked, although some decrement in motility was reported. Apparently, post-pubertal hypothyroidism does not cause severe seminal alterations. Another study designed to investigate the correlation of thyroid dysfunction and semen parameters in infertile men, however, analyzed 305 patients and did not find any correlation between thyroid dysfunction and semen parameters, but between elevated thyroid peroxidase antibody (TPO-Ab) and panthozooospermia and asthenozoospermia (60). These results, conversely, were not confirmed in a recent cohort of subfertile men (61). Recently, Meeker and cols. (62) found a positive association between free T4 and sperm concentration in a group of 388 men with mean age of 36 ± 5.4 years.

The effects of hyperthyroidism on male fertility were the topic of some studies. Early clinical observations have described oligospermia and loss of spermatozoa mobility in hyperthyroid men (63). The same findings were observed by Kidd and cols. with hyperthyroid patients and low total sperm counts (64). In the early 1990’s, Hudson and Edwards (65) assessed testicular function in men with Graves’s diseases and found that the percentage of forward progressive motility was significantly lower than control values. Seminal abnormalities were corrected when patients became euthyroid. Another study that evaluated the effects of Grave’s disease on male reproduction showed that thyrotoxic male presented several changes on semen parameters, such as: asthenospermia, hypospermia, oligospermia,
necrospermia and teratospermia, as well as complaints of impaired sexual function (49). A normalization of 85% of seminal alterations was observed when euthyroidism were reached. A recent controlled prospective study (66) showed that 50% of the 23 thyrotoxic males studied presented sexual dysfunction associated with decreased libido, improved after treatment. These patients also showed lower sperm densities and motility, whereas no differences were found on sperm morphology. Again, following treatment of thyrotoxicosis, sperm density and motility improved, but sperm morphology did not change.

Erectile dysfunction and loss of libido or impotence are not uncommon in males with thyroid dysfunctions (13,67). Carani and cols., conducting a clinical trial that evaluated 34 patients with hyperthyroidism and 14 with hypothyroidism, both initiated in adulthood, demonstrated that 12.8% of these men showed some degree of sexual function deterioration (14). The most common symptom in hyperthyroidism were precocious ejaculation, while hypothyroid patients referred a greater number of complaints such as erectile dysfunction, ejaculation delay, loss of libido and impotence. This study showed that most patients with thyroid hormone disorders experience some sexual dysfunctions that may be reversed with the normalization of thyroid hormone levels (14).

Erectile dysfunction were found in about 80% of patients with hyper- or hypothyroidism in a controlled study that investigated the impact of thyroid hormone alterations on male sexual health (67). After treatment, the proportion of erectile complaints were the same in both patients and control groups. Thyroid hormone alterations on male reproductive function are summarized on table 1.

**CONCLUSION**

In the past decades, it has become clear that thyroid hormone plays an important role in Sertoli and Leydig cell proliferation and function, also influencing spermatogenesis, sperm motility and ultimately fertility. Disturbance of the normal euthyroid state affects the morphological and functional development of the testis. Thyroid hormone is also likely to contribute to normal spermatogenesis and metabolic processes in the adult testis, but these aspects are not completely understood at the present. Nevertheless, despite the gaps in our knowledge, the data herein reviewed may provide considerable evidence to conclude that thyroid hormone plays an important role in the human testicular development and function.

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<tr>
<th>Table 1. Effects of hypo and hyperthyroidism on male gonadal function</th>
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<tr>
<td><strong>Hypothyroidism</strong></td>
</tr>
<tr>
<td>Prepubertal testicular volume and function ↑</td>
</tr>
<tr>
<td>Early onset of spermatogenesis</td>
</tr>
<tr>
<td>Sperm counting Normal or ↓</td>
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<tr>
<td>Sperm motility ↓</td>
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<tr>
<td>Sexual function Impaired</td>
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<td>Erectile function ↓</td>
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**REFERENCES**

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