Symptomatic late-onset hypogonadism but normal total testosterone: the importance of testosterone annual decrease velocity

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Late-onset hypogonadism (LOH) is a relevant condition from an epidemiological perspective, since it has an estimated prevalence of 2.1% (1). In recent years, this issue has gathered growing interest, due to the possible adverse effects which testosterone replacement therapy (TRT) may expose to. Notably, LOH arises in aging, a period of life wherein cardio-metabolic diseases are highly prevalent (2) and the risk for cardio- and cerebro-vascular events is not negligible (3).

LOH seems to be due to a central (hypothalamic-pituitary) or testicular dysfunction, which may concomitantly occur in some cases. This editorial aims to discuss the management of patients with symptomatic LOH but with total testosterone (TT) levels in the lowest quartile of the normal range.

Before deepening into the pharmacological issue, some premises have to be made.

First of all, prior to make LOH diagnosis, a proper evaluation of the clinical conditions leading to LOH-like symptoms (e.g., anemia, hypothyroidism, hypocortisolism, mood disorders) should be accomplished, although sexual dysfunctions (erectile dysfunction, decreased libido) show a high specificity for LOH (4). Accordingly, the European Male Aging Study (a cohort study of community-dwelling middle-aged and older men in Europe) found that only sexual symptoms (poor morning erections, decreased libido, and erectile dysfunction) are associated with TT and free-testosterone (FT) concentrations (5). In addition, validated questionnaires might represent a potentially useful diagnostic tool for differential diagnosis (6), despite their use is not fully recognized by the current guidelines (7).

Second, TT serum measurement is usually done by chemiluminescence immunoassay (CLIA) in the clinical practice, although radioimmunoassay (RIA) or liquid chromatography-tandem mass spectrometry (LC-MS/MS) are the only two assays mentioned by the Endocrine Society (ES) guidelines. Specifically, the last ES guidelines define LC-MS/MS as the assay offering the highest specificity, sensitivity and precision for TT (especially in the low range) than do most immunoassays (7). Currently, the diagnosis of LOH is made for TT serum levels lower than 264 ng/dL (9.2 nmol/L). To confirm the diagnosis, at least two T measurements are needed (7). FT measurement should be performed by the equilibrium dialysis method or calculated if this method is not available in men whose serum TT concentration is around the lower limit (e.g., 200 to 400 ng/dL) of normal range (7).

Actually, the concept of “Testosterone Annual Decrease Velocity” is not considered, despite data from
the Massachussets Male Aging Study (MMAS) showed a testosterone annual decrease in the adulthood of 1.6% per year of age (8). Testosterone decrease begins when bone mass peak occurs. It is highly heterogeneous, especially in case of metabolic (e.g., obesity) or chronic diseases, that associate with a higher annual decrement rate. T decrease may hypothetically be responsible for symptoms based on the degree of the intra individual annual variation. Therefore, TT measurement starting from the adulthood would be of great utility to assess whether a greater than expected decline occurs in aging.

Few markers of Leydig cell dysfunction, which may anticipate TT decline have been proposed. In particular, insulin-like 3 (INSL-3), a Leydig cell-derived glycoproteic hormone, may represent a good candidate. Similarly, the rise of luteinizing hormone (LH) in the presence of normal serum TT identify a “subclinical hypogonadism” (9), whose treatment is not codified by none of the current available guideline. Alteration of testicular function occurring in aging may also be assessed by non-hormonal methods, such as ultrasound scan, which can usefully detect testicular volume decrease or changes into echo texture, due to a progressive rarefaction of the testicular parenchyma compared to that of the fertile age.

Some polymorphisms in the androgen receptor (AR) are known to influence the AR responsiveness to testosterone, affecting signal transduction intensity. Particularly, the number of CAG trinucleotide repeats is one of the most studied. A low number of repeats is associated with a higher AR activity, thus raising the risk for prostate cancer (10). This polymorphism is not taken into account in the clinical practice.

Moreover, some features may resemble LOH, such as (I) lower prostate volume (evaluated by ultrasound) in comparison with aged-matched peers or low total PSA levels, an index of prostate volume and androgenization (11), (II) apparently idiopathic low hematocrit levels, suggested by the role of TT on erythropoiesis (TRT is contraindicated in case of pre-treatment hematocrit levels of 48% or when it is >54% during TRT) (12).

Finally, patients with apparently idiopathic primary osteoporosis (secondary forms, e.g., those due to hyperparathyroidism, hypercortisolism, etc. have to be excluded) and hypovitaminosis D should also be checked for LOH. In particular, hypovitaminosis D may hide a central hypogonadism, due to the LH-induced hydroxylation of vitamin D at the testicular level (13).

On the basis of the above-reported considerations, would it be reasonable to consider TRT in the following cases characterized by TT levels within the lowest quartile of the normal range?

(I) Hypogonadism-related symptoms and Testosterone Annual Decrease Velocity higher than expected;
(II) Hypogonadism-related symptoms, decreased testicular volume and echo tessiture rarefaction;
(III) Hypogonadism-related symptoms and low INSL-3 levels;
(IV) Hypogonadism-related symptoms and high number of CAG trinucleotide repeat of the AR;
(V) Hypogonadism-related symptoms, low prostate volume and/or low PSA values compared to aged-matched peers;
(VI) Hypogonadism-related symptoms and (idiopathic) low hematocrit values;
(VII) Hypogonadism-related symptoms and idiopathic primary osteoporosis.

In all of these combinations, the exclusion of previously mentioned diseases causing LOH-like symptoms has to be carefully accomplished prior to plan TRT.

An experimental proposal may be to start TRT using half dose of that usually used for hypogonadism. Any possible side effect has to be timely detected adopting a shorter follow-up, which may be done after 6, 12 and 24 weeks [or 3, 6 (or 9 according to the clinician’s judgement) and 12 months and at least annually thereafter as currently suggested for hypogonadal patients in TRT by the current guidelines (14)]. In addition, transdermal preparations could be preferred to injective ones when a low dosage is required. Furthermore, the interruption of the administration of this latter preparation more rapidly allows a decrease of serum testosterone concentrations compared to the injective one. The improvement of symptoms after the beginning of TRT would confirm LOH diagnosis. Conversely, if no benefits ensue, TRT should be suspended.

Finally, to overcome the pitfalls of the current clinical practice, TT measurement from the age of 20 should be done (for example, every 5–10 years) in all men to assess testosterone annual decrease velocity (Figure 1).
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Footnote

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