Recent advancement in the treatment of boys and adolescents with hypogonadism

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Abstract: Clinical manifestations and the need for treatment varies according to age in males with hypogonadism. Early foetal-onset hypogonadism results in disorders of sex development (DSD) presenting with undervirilised genitalia whereas hypogonadism established later in foetal life presents with micropenis, cryptorchidism and/or micro-orchidism. After the period of neonatal activation of the gonadal axis has waned, the diagnosis of hypogonadism is challenging because androgen deficiency is not apparent until the age of puberty. Then, the differential diagnosis between constitutional delay of puberty and central hypogonadism may be difficult. During infancy and childhood, treatment is usually sought because of micropenis and/or cryptorchidism, whereas lack of pubertal development and relative short stature are the main complaints in teenagers. Testosterone therapy has been the standard, although off-label, in the vast majority of cases. However, more recently alternative therapies have been tested: aromatase inhibitors to induce the hypothalamic-pituitary-testicular axis in boys with constitutional delay of puberty and replacement with GnRH or gonadotrophins in those with central hypogonadism. Furthermore, follicle-stimulating hormone (FSH) priming prior to hCG or luteinizing hormone (LH) treatment seems effective to induce an enhanced testicular enlargement. Although the rationale for gonadotrophin or GnRH treatment is based on mimicking normal physiology, long-term results are still needed to assess their impact on adult fertility.

Keywords: anorchia, cryptorchidism, delay of puberty, disorders of sex development, DSD, Kallmann syndrome, Klinefelter syndrome, micropenis

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Introduction

The concept of male hypogonadism is often associated with low testosterone production by the testes. This notion derives from adult endocrinology; indeed, during adulthood, testosterone is the most conspicuous testicular hormone because of its circulating levels and target organ actions. Conversely, in paediatric endocrinology, basal serum testosterone determination is helpful only during the first months following birth and after mid-puberty. During the rest of infancy and childhood, serum testosterone is physiologically below the detectable levels using classical assays, and anti-Müllerian hormone (AMH) and inhibin B are more adequate biomarkers to initially screen testicular function (Figure 1).

The ontogeny of the hypothalamic-pituitary-testicular axis: its importance for the diagnosis of male hypogonadism

The hypothalamic-pituitary-testicular (HPT) axis is undoubtedly the pituitary axis that shows the most remarkable changes throughout life (Figure 1). The definition of male hypogonadism should take into account this ontogeny and reflect the inability of Sertoli, Leydig, and/or germ cells to fulfil their functions. Primary hypogonadism, called hypergonadotrophic hypogonadism in adult endocrinology, reflects a primary defect of testicular components—Sertoli, germ and/or Leydig cells—that can affect all cell populations concomitantly or initiate with the impairment only one of them and affect the others secondarily. Secondary
or central hypogonadism, called hypogonadotrophic hypogonadism in adult endocrinology, reflects a gonadal dysfunction that results from a hypothalamic-pituitary insufficiency.

The prenatal period: consequences of foetal-onset hypogonadism

During the embryonic and foetal periods, the testes differentiate before the gonadotroph is functional and secrete AMH and testosterone, which play essential roles in sexual differentiation of the genitalia. Testosterone, secreted by Leydig cells in response to placental human chorionic gonadotrophin (hCG) in the first trimester of gestation and by foetal pituitary luteinising hormone (LH) thereafter, drives the differentiation of the Wolffian ducts into the epididymides, vasa deferentia and seminal vesicles. Upon transformation to dihydrotestosterone (DHT) by the action of the enzyme 5α-reductase in target tissues, it induces the formation of the prostate and the virilisation of the urethra and the external genitalia in the first trimester and the increase in the size of the genitalia, as well as testicular descent in the second and third trimesters of gestation. AMH, produced by Sertoli cells independently of follicle-stimulating hormone (FSH) action in the first trimester of foetal life, provokes the regression of Müllerian ducts, the primordia of the uterus and Fallopian tubes. Subsequently, FSH stimulates Sertoli cell proliferation resulting in testicular enlargement and increased secretion of AMH and inhibit B. Note that the Sertoli cell population represents the major testicular component until the onset of puberty (Figure 1).

Foetal onset male hypogonadism has different clinical consequences according to the time of onset during the embryonic/foetal stage and to the testicular cell populations affected (Table 1). Very early foetal-onset hypogonadism can only be primary, since the gonadotroph is not yet functional. Androgen deficiency may result in a complete lack of virilisation and, thus, a female phenotype of the external genitalia at birth in the case of complete gonadal dysgenesis, a condition characterised by whole gonadal dysfunctions and named dysgenetic disorder of sex development (DSD), or of Leydig cell steroidogenic failure, characterised by specific cell dysfunction with normal Sertoli cell activity and known as non-dysgenetic DSD. In these cases, newborns are assigned female and will not be discussed further in this review. Alternatively, the gonadal disorder may be partial, resulting in ambiguous genitalia with different degrees of virilisation, which may lead to male gender assignment at birth.

Foetal-onset hypogonadism during the second or third trimesters, irrespective of its primary or central origin, is associated with a completely male phenotype of the external genitalia. Nonetheless, the androgen insufficiency may lead to micropenis and cryptorchidism. When the condition is of central origin, the FSH insufficiency results in reduced Sertoli cell numbers, leading to microorchidism and low serum AMH and inhibit B. If congenital hypogonadism goes unnoticed during the first 3 to 6 months of postnatal life, androgen insufficiency cannot be detected during childhood, and low Sertoli cell biomarkers and genetic tests are more helpful for the diagnosis (Figure 1). At the age of puberty, lack of pubertal signs prompts the assessment.

Childhood: male hypogonadism may remain unnoticed

As already mentioned, gonadotrophin and testosterone serum levels decline to very low or even undetectable values after the age of 3 to 6 months in the normal boy (Figure 1). Therefore, the male hypogonadism established during childhood may be challenging not only in the case of central hypogonadism, because gonadotrophins and testosterone cannot be lower than in normal boys explaining why the term ‘hypogonadotrophic’ may be misleading at this age, but also in face of cases of primary hypogonadism, because serum gonadotrophins show normal prepubertal values in a significant number of cases and the term ‘hypergonadotrophic’ may be confusing. A clear example is Klinefelter syndrome, the commonest aetiology of primary hypogonadism in males.

Adolescence: pubertal delay

The reactivation of GnRH secretion in the hypothalamus leads to a progressive increase in LH and FSH pulsatile secretion. FSH triggers again Sertoli cell proliferation, inducing a moderate but perceivable increase in testicular size, from 2 to 4 ml as measured by comparison to Prader’s orchidometer, or from 1.0 to 2.7 ml as measured by ultrasonography. Concomitantly, LH acts on Leydig cells to induce a progressive
increase in intratesticular androgen concentration during Tanner stages 2 and 3 of pubertal development, which provokes Sertoli cell maturation, reflected in a decrease in serum AMH. Then, Sertoli cells cease to proliferate and become capable of supporting adult spermatogenesis, the main responsible for the dramatic increase of testicular volume during puberty to reach an adult volume >15 ml by orchidometer or >17.5 ml by ultrasound. Inhibin B levels represent a useful biomarker of Sertoli cell maturation and adequate spermatogenic progression. Serum testosterone levels increase progressively from Tanner stage 3 to reach adult values in Tanner stage 5. The gradual increase in serum testosterone is associated with the development of secondary sex characteristics, especially penile growth which occurs from Tanner stage 3 onwards, and a timely bone age maturation to attain adult height.

When there is a whole gonadal dysfunction, due to either primary or central hypogonadism, the lack of testosterone rise results in the lack of signs of pubertal development. For primary hypogonadism to be associated with complete lack of development of secondary sex characteristics, the testicular disorder must be severe, for example, complete testicular regression syndrome or gonadal removal. Otherwise, Leydig cell function often remains sufficient to
provoke some signs of androgenisation, even if Sertoli cells are severely affected, for example, in many cases of partial gonadal dysgenesis or longstanding cryptorchidism.\textsuperscript{17}

Apart from the already mentioned congenital form, central hypogonadism may be acquired during childhood, for example, due to tumours or other lesions of the central nervous system affecting the hypothalamic-pituitary region or to functional states such as chronic or acute illnesses that impair the HPT axis.\textsuperscript{5} The severity of the condition may lead to a complete lack of pubertal onset or to a normal or delayed onset with incomplete progression.

In those cases where there is no personal or family history that could drive the diagnosis, constitutional delay of puberty is the commonest cause of lack of pubertal signs in a boy reaching the age of 14 years, that is, that corresponding to more than 2 standard deviations from normal pubertal age onset.\textsuperscript{32,33} The differential diagnosis may prove difficult: basal LH and testosterone levels are usually uninformative and several dynamic tests have been developed with controversial results.\textsuperscript{34} Although neglected in males, the FSH-gonadal axis seems to be more informative.\textsuperscript{35–37}

### Pharmacotherapy for male hypogonadism

**Medicines used**

**Androgenic drugs.** The most frequently used formulations in adolescents are testosterone esters, such as enanthate, cypionate and propionate,
available as oil-based compounds for slow-release IM injections. Testosterone enanthate is available in 200- or 250-mg ampoules or ready-to-use syringes. It has a half-life of 4.5 days and is cleared 90% through the kidneys and 10% through the bile (10%). Testosterone cypionate is available in 100-mg or 200-mg vials, and its half-life is 4 days.38 A mixture of 100 mg decanoate, 60 mg isocaprate, 60 mg phenylpropionate and 30 mg propionate exists in 250-mg ampoules.39-41 IM testosterone undecanoate has not been used in paediatric patients owing to its long-acting period.42

In adolescents >12 years-old, the anabolic and androgenic effects induced by replacement therapy are expected to mimic the timing and tempo observed during normal pubertal development. In other words, androgen supplementation should start at low doses with progressive increases so that adult testosterone serum levels are attained 2 to 3 years later. Thus, with a progression of approximately one Tanner stage every year, a precocious early closure of the epiphyses is avoided. The most commonly used protocols with testosterone enanthate or cypionate consist of an initial dose of 50–100 mg every 4 weeks.43,44 Due to the pharmacokinetics of these formulations, circulating testosterone attains supraphysiological levels in the first week after IM injection and subphysiological levels in the fourth week (Figure 2),45,46 with interindividual variations.47,48 The dosage is progressively increased in 50- to 100-mg increments every 6 to 12 months to mimic the evolution of serum testosterone levels observed during pubertal stages,28 to reach the adult dose of 250 mg every 4 weeks after approximately 2 to 3 years.

Transdermal testosterone formulations are the most frequently used in adults,53 but like IM formulations, they have not yet been approved for use in paediatric patients. Only few studies are reported in the literature in prepubertal boys.

While the initial oral androgen formulations showed hepatotoxicity,53,54 testosterone undecanoate capsules avoid hepatic metabolism, and pilot studies have been reported in adolescents.55,56 Mucoadhesive tablets, designed to provide sustained testosterone release as it hydrates after inner cheek adhesion,53 have not been used in children or adolescents. Similarly, no experience exists in paediatric patients with a testosterone gel for intranasal administration featuring easy delivery and low risk of transference.53,54

Side effects commonly described for androgenic drugs used for long periods include erythrocytosis, irritability, acne, persistent erections, and gynaecomastia due to aromatisation to oestrogens.57 One particular concern in paediatric ages is the potential acceleration of bone age, which compromises adult height. Early pubarche can also be observed. Oral testosterone undecanoate shows lower risk for erythrocytosis than injectable formulations.58 It should be kept in mind that the beneficial effects that systemic androgen therapy has on most organs contrast with their negative effects on the HPT axis. Indeed, exogenous androgens usually reach circulating levels that inhibit the gonadotroph. Because of the reduced secretion of LH, intratesticular androgen concentration remains low, spermatogenesis cannot go through meiosis and spermiogenesis, and testicular volume persists small.27,59

DHT, also known as stanolone or androstanolone, is the most potent natural androgen and cannot be aromatised to oestrogens. It is commercially available in some countries as 2.5% gels for
DHT is typically applied directly to the external genitalia in patients with micropenis and is indicated in patients with nondysgenetic DSD due to $5\alpha$-reductase deficiency.

Nandrolone and oxandrolone are non-aromatisable drugs with a predominant anabolic action and are thus used when androgenic effects need to be avoided. Nandrolone exists in 1-ml vials at 200 mg/ml for IM injections. Oxandrolone is available as oral tablets of 2.5 and 10 mg. In patients with Klinefelter syndrome, positive effects have been reported on psychosocial and visual-motor functions as well as on cardiometabolic health markers after oxandrolone administration. The main side effect is hepatotoxicity, though rarely serious or irreversible. Hepatic peliosis hepatocellular neoplasia have been described after long-term administration. In paediatric patients, even if its androgenic effects are reduced, oxandrolone may provoke precocious genital development. Selective androgen receptor modulators (SARMs) with different androgen receptor binding capacity are under investigation.

Aromatase inhibitors, such as letrozole and anastrozole, repress the metabolisation of androgens to oestrogens and have been tested to boost androgen levels in patients with preserved androgen synthesis, that is, constitutional delay of puberty.

Gonadotrophin formulations. Several preparations containing hCG, LH or FSH, alone or in mixture, have existed in the market and have been used especially in adult reproductive endocrinology. Here, I will focus on those formulations used in paediatric patients with hypogonadism.

Human chorionic gonadotrophin (hCG) has been available for several decades as solutions for IM or SC injection containing between 500 and 10,000 IU, obtained from the urine of pregnant women. LH and hCG bind the same receptor on Leydig cells, but hCG has a longer half-life and can be given once or twice a week according to the therapeutic aim: in boys with cryptorchidism, hCG is usually administered at 500–1000 IU per week for 5 weeks, whereas in adolescents with delayed puberty the usual dosage is 1000 IU twice a week.

LH and FSH are also present in preparations obtained from urine: human menopausal gonadotrophin or menotropin (hMG) contains FSH, LH, and hCG, whereas immunological-based technologies have allowed to obtain highly purified urinary FSH preparations. However, almost no experience has been published in paediatric male patients with these preparations. Conversely, in the last two decades, recombinant gonadotrophin formulations have been developed, including r-LH, r-hCG, r-FSH, long-acting r-FSH (corifollitropin alfa), and a mixture of r-LH/r-FSH.

Two r-FSH preparations, follitropins alfa and beta, are very similar and only differ in the technical approach using Chinese hamster ovary (CHO) cells resulting in slight differences in posttranslational changes. Nonetheless, all recombinant follitropins obtained in CHO cells differ from the natural gonadotrophins in some of the attached glycans, which may affect their bioactivity. In an attempt to solve this problem, recombinant folitropin delta was generated using a human cell line. Also more recently, corifollitropin alfa has been developed; it consists of r-FSH fused to the carboxyl-terminal peptide of the beta-subunit of hCG, which conveys a longer plasma half-life. Recombinant LH (lutropin alfa) and r-hCG are produced in CHO cells. Although gonadotrophins may be given as IM or SC injections equivalently, SC administration is preferred by the overwhelming majority of patients and supports long-term adherence to treatment.

GnRH is available in a few countries for pulsatile administration using a SC pump delivering the drug at a rate of 25 ng/kg per pulse every 120 min.

Pharmacotherapy in neonates and infants
Irrespective of the aetiology of impaired gonadal hormone production, newborns or infants with male hypogonadism may require treatment for micropenis and/or cryptorchidism. However, those with central hypogonadism have a more favourable fertility potential, and treatment options should consider future spermatogenic development as an aim.

Treatment of newborns/infants with DSD. Newborns with dysgenetic DSD, including partial testicular dysgenesis, asymmetric gonadal differentiations (also known as mixed gonadal dysgenesis) and ovotesticular DSD, assigned male usually present with micropenis, hypospasias and
Cryptorchidism. While hypospadias can only be repaired surgically, micropenis and cryptorchidism may be subject to hormonal treatment. For penile enlargement and enhancement of scrotal tumour, the usual practice consists in the administration of 2 or 3 courses of IM testosterone enanthate 25–50 mg every 3–4 weeks. Longer courses may provoke bone age advancement and pubic hair development and should therefore be avoided. Although infrequent, side effects related to androgen excess are erections and acne; non-specific side effects are pain and infections in the injection site. In some countries, testosterone is available for percutaneous treatment: 0.2 g of 5% testosterone cream (i.e. 10 mg of testosterone) applied onto the phallus daily for 1 month showed efficacy (9-mm increase in penile length) and safe (no significant advancement in bone age). DHT gels, which are available in certain countries, may be even more efficacious in patients with DSD, and especially in those with 5α-reductase deficiency. Treatment consists of gel application onto the penis twice a day at a daily dose of 0.1–0.3 mg/kg/day, with caution not to exceed 5 mg/day, for a maximum period of 6 months.

If LH levels are not too high, hCG treatment could be tried to induce testicular descent when the gonads are palpable in the lower part of the inguinal canal. However, this is infrequent and surgical orchiopexy is usually performed.

Treatment of newborns/infants with primary hypogonadism and completely virilised genitilia. Patients with testicular regression syndrome or testicular torsion occurring in the second half of gestation usually present at birth with micropenis and nonpalpable gonads. Once the lack of testicular tissue or its extreme scarcity is ascertained by the finding of undetectable or extremely low AMH or inhibin B in association with elevated FSH, together with low or undetectable testosterone and elevated LH, androgen replacement is indicated following the same protocol as that described above for patients with DSD.

Cryptorchidism may be associated with primary hypogonadism or with eugonadism. Expectant behaviour during the first year is suggested by many authors, given that the testes may finalise their descent to the scrotum. However, others point to the importance of an adequate hormonal milieu for germ cell development during the neonatal activation period, also known as mini-puberty, and are prone to hCG or GnRH treatment without delay. Three systematic reviews showed that the efficacy of hormone therapy is related to testis position—the lower, the better—rather than to age at treatment. Several protocols exist; hCG given IM or SC 500–1000 IU once a week for 5 weeks is the most widely used, whereas native GnRH or GnRH analogues have also been used, but are not available in most countries.

Treatment of newborns/infants with central hypogonadism. Like patients with other forms of hypogonadism, those with central hypogonadism are brought to medical attention for micropenis and/or cryptorchidism. Micro-orchidism usually is underestimated or overlooked. These signs might represent a red flag for ruling out other pituitary hormone deficiencies, which would need to be treated as a priority given their vital roles, for example, cortisol and thyroid hormone deficiencies.

Testosterone treatment has been classically used off-label to induce penile growth, as explained above, and the age at which treatment is installed does not seem to be critical. More recently, a few studies have adopted a physiology-based rationale to test gonadotrophin replacement during the first 6 months of life, in order to mimic the neonatal activation of the HPT axis. Combined treatment with recombinant FSH plus LH or hCG, or GnRH or a GnRH analogue, was expected to induce Sertoli cell proliferation—thus resolving micro-orchidism—and adequate germ cell development, together with Leydig cell androgen production promoting testicular descent and penile enlargement. Initial case reports showed that SC injections or SC pump infusions of recombinant LH and recombinant FSH led to a significant increase (twofold to fourfold) in testicular volume and hormone (AMH, inhibin B and testosterone) levels, followed by an enlargement of penile size (from ~1 cm to 3–4 cm). LH was administered at 20–40 IU twice a week or 56–75 IU/day, whereas FSH doses were 21.3 IU twice a week or 67–75 IU/day. More recently, a case series study reported the results of recombinant LH 50 IU/day plus recombinant FSH 75–150 IU/day given through a SC pump, showing and increase in testicular size and penile length in the 8 patients included, with complete testicular descent in 6 of them.
Pharmacotherapy in childhood

Continued hormone replacement therapy does not seem necessary during childhood, based on the knowledge of the normal ontogeny of the HPT axis (Figure 1). Nonetheless, in boys with Klinefelter syndrome androgenic treatment has been assessed even though hypoandrogenism has not been unequivocally observed. In placebo-controlled trials, oxandrolone oral administration at a daily dose of 0.05–0.06 mg/kg for 24 months to boys aged 4–12 years showed improvements in patients’ cognition and behaviour, as well as in their motor and visual capacities. Clinically non-significant adverse events, such as minor advancement in bone age and decline in serum HDL, were more frequently observed than early pubarche and an increased risk of early testicular enlargement; HPT axis hormone levels were not affected. These results cannot be applied to patients with higher grade sex chromosome aneuploidies, such as 48, XXXY, 48, XXYY or 49, XXXXY, in whom testosterone treatment resulted in an earlier and persistent suppression of testicular hormone production.103

One study assessed the effect of r-FSH treatment on 3 boys with central hypogonadism. The underlying rationale was that FSH is important to provoke Sertoli cell proliferation before pubertal maturation induced by intratesticular testosterone. Subcutaneous r-FSH, administered for 12 months at 1.5 IU/kg 3 times a week, induced testicular enlargement and increased serum inhibin B levels, reflecting Sertoli cell stimulation, suggesting that FSH treatment is an option before pubertal maturation to induce Sertoli-cell proliferation with a potential of an enhanced sperm-producing capacity in adulthood.

Pharmacotherapy at pubertal age

Irrespective of the aetiology, androgen insufficiency needs to be treated at the age of puberty in order to tackle the lack of development of secondary sexual characteristics and the impaired growth spurt. Indeed, boys with hypoandrogenism maintain a childlike body aspect and become progressively shorter than their peers, which usually lead to psychosocial distress. Androgen therapy may provoke androgenic and anabolic effects on target tissues. Androgenic effects include the development of male secondary sexual characteristics as well as of male sebaceous gland activity and hair growth pattern, whereas anabolic effects include greater skeletal muscle mass and bone metabolism.57,65

Treatment of patients with DSD or delayed puberty due to primary hypogonadism. As discussed for newborns, some forms of DSD reflect a primary gonadal insufficiency. In many cases, especially those due to dysgenetic DSD with increased risk of gonadal tumour development, orchiectomy is performed before pubertal age. Therefore, androgen replacement is needed to induce the development of secondary sex characteristics, growth spurt and bone mass accrual typical of puberty. The classic protocol of IM testosterone administration is used, starting at 50 mg every 4 weeks when bone age is at least 12 years-old, with progressive increases to reach 250 mg every 4 weeks approximately 3 years later. The same considerations are applicable to patients with primary hypogonadism not related to DSD but severely affecting testicular androgen secretion, such as testicular regression syndrome. Other forms of primary hypogonadism, for example, related to long-standing cryptorchidism, orchitis, chemotherapy, pelvic radiotherapy or Klinefelter syndrome, usually do not require androgen supplementation to induce pubertal changes, but may require it when testosterone levels are below normal. In these cases, adult doses can be used directly, especially if near-adult height has already been achieved. A particular controversy exists regarding the initiation of androgen therapy in adolescents with Klinefelter syndrome. Testosterone production is most often within the normal range, which makes most authors recommend watchful waiting even if LH levels are elevated, limiting androgen replacement to those cases with low serum testosterone and clinical signs of hypogonadism.5,107 Others are prone to earlier exposure to androgen treatment based on the observation of improved physical and neurocognitive outcomes. A clinical trial with a 1% testosterone gel showed that daily administration starting with 0.5 g and a progressive increase up to 5 g per day resulted in a two-fold increase in serum levels of testosterone and DHT, with no side effects.109

Treatment of patients with constitutional delay of puberty or central hypogonadism. Patients with
functional hypogonadism usually do not require hormonal treatment since normal activity of the HPT axis is re-established once the underlying condition is resolved. Conversely, pharmacological treatment may be needed in boys with constitutional delay of puberty and is absolutely necessary in patients with central hypogonadism. Interestingly, approximately 20% of patients diagnosed with central hypogonadism may show a spontaneous increase in testicular volume and serum levels of gonadotrophins and testosterone when treatment is discontinued in adulthood.

**Androgen therapy.** In boys with a well-established diagnosis of central hypogonadism, treatment should not be delayed beyond the age of 12 years to avoid the psychosocial burden and the negative effect that the delay in exposure to sex steroids may have on the skeleton and pubertal growth spurt. Similarly, treatment is initiated even if a differential diagnosis between constitutional delay of puberty and central hypogonadism could not be solved. Although all current treatments are off-label, the protocol of IM testosterone administration with progressive increase already described for primary hypogonadism is the most widely used. The doses (~50 mg every 4 weeks) used at the beginning are too low to inhibit the reactivation of the HPT axis in boys with constitutional delay of puberty, so that testosterone administration can be stopped if testicular volume reaches 4 ml. A frequently used alternative is to give testosterone for 6 months, then discontinue it for 3 months and watch whether testicular volume progresses. In a few cases, the patient feels satisfied with the androgenic effects and is not willing to discontinue testosterone administration. Treatment discontinuation can then be delayed until full development has been attained, approximately 3 years after initiation of treatment. In this case, a longer washout (~6 months) is needed to confirm that no spontaneous development occurs, given that the full dosage of 250 mg every 4 weeks provokes the inhibition of the HPT axis. While monitoring the potential side effects of androgen treatment on haematocrit and liver function is not critical when therapy is used for short periods like in boys with constitutional delay of puberty, the standard monitoring used in adults should be applied in boys receiving longer therapy.

In boys with constitutional delay of puberty, oral testosterone undecanoate has been tried for up to 15 months, at a daily dose of 10 to 160 mg. Serum testosterone showed physiological levels for pubertal onset, in association with the beneficial effects of secondary sex characteristics and height velocity, while no excessive acceleration of bone age was noted. Smaller case series or controlled trials had also been previously reported. Other androgen formulations have received little attention. A gel of 2% testosterone administered at 10 mg/day has shown good results in a small cohort of boys with constitutional delay of puberty. Testosterone patches have shown variable efficacy and patient adherence.

**Aromatase inhibitors.** Aromatase inhibitors, especially third-generation ones such as anastrozole and letrozole, have been used in the last two decades to increase adult height in boys, by delaying bone age progression. However, a recent controlled clinical trial used letrozole to induce puberty in boys with constitutional delay. Boys 14 years-old or older willing medical treatment and exhibiting the first signs of puberty received either letrozole 2.5 mg/day orally or testosterone 37.5–75 mg IM every 4 weeks for 6 months. Treatment with letrozole resulted in higher serum LH, FSH, testosterone and inhibin B, as well as greater testicular volume increase, suggesting that letrozole induces the activation of the HPT axis in boys with CDGP.

**Gonadotrophins and GnRH.** As proposed for newborns and infants with central hypogonadism, a more physiological option, aiming to mimic the normal development of testicular function during puberty, is the administration of gonadotrophins. Several treatment protocols have been reported, including hCG alone or in combination with r-FSH, hMG or GnRH. Initially, hCG treatment of adolescents with central hypogonadism showed a comparable effect to IM testosterone on secondary sex characteristics. However, pulsatilie administration of GnRH has showed better efficacy as compared to hCG in adolescents with central hypogonadism, underscoring the complementary role of FSH. In fact, other studies showed the beneficial effect of FSH or hMG addition to hCG. Nonetheless, to mimic physiological chronology, FSH proliferative effect on Sertoli cells should be induced before testosterone provokes Sertoli cell maturation inducing mitotic arrest. Indeed, studies with r-FSH priming prior to hCG treatment showed enhanced testicular growth and increased circu-
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lating concentrations of AMH (Figure 3) and inhibin B (Figure 4). Subsequent addition of hCG induced testosterone secretion by Leydig cells and Sertoli cell maturation, as reflected in AMH decline. The increase in inhibin B levels is useful to monitor sperm production. A randomised clinical trial in adolescents with central hypogonadism showed better results in patients pre-treated with r-FSH in terms of sperm production. Typically, r-FSH is given 75–150 IU SC daily or every other day to attain target serum FSH levels of 7–9 IU/L during for 2–6 months, followed by combined r-FSH plus hCG starting at 250 IU weekly and progressively increasing by 250–500 IU weekly every 6 months to finally attain 1500 IU 3 times a week. Alternatively, hMG is given instead of r-FSH at 75–150 IU 2–3 times per week. FSH priming may be especially needed in patients with small testes (e.g. volumes less than 5–6 ml), as compared to those with pubertal arrest or partial forms of central hypogonadism, who have larger testes volumes and are likely to only require hCG. Once full development has been obtained, treatment may be switched to testosterone administration for maintenance until the patient desires paternity.

Pulsatile GnRH treatment is also a physiological approach in adolescents with central hypogonadism. GnRH is administered in a pulsatile manner using mini-infusion pump with an initial dose of 25 ng/kg per pulse every 2h, with a subsequent titration to attain target serum testosterone. Although gonadotrophin and GnRH treatments have a physiology-based rationale in patients of pubertal age, whether fertility outcomes are improved as compared to using the more classical androgen therapy and waiting to administer these agents in adulthood still needs evidence.

Concluding remarks and research agenda
All treatments used in paediatric patients with hypogonadism are off-label. Testosterone administration IM is the most frequently used therapy in order to provoke genital enlargement in childhood or the full development of secondary sexual characteristics and growth spurt in adolescents. While this is the only possibility for patients with primary hypogonadism, the administration of gonadotrophins or GnRH may represent a more physiological therapy in boys with central hypogonadism. Clinical trials with long-term follow-up are needed to assess whether gonadotrophin treatment yields better results than initial androgen replacement followed by gonadotrophin treatment in adulthood when fertility is sought. Other possibilities based on recently developed technologies, such as Leydig cell or spermatic development in vitro, represent stimulating alternatives.

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