ABSTRACT

Masculinisation of internal and external genitalia during foetal development depends on the existence of two discrete testicular hormones: Leydig cell-secreted testosterone drives the differentiation of the Wolffian ducts, the urogenital sinus and the external genitalia, whereas Sertoli cell-produced anti-Müllerian hormone (AMH) provokes the regression of Müllerian ducts. The absence of AMH action in early foetal life results in the formation of the Fallopian tubes, the uterus and the upper third of the vagina. In 46,XY foetuses, lack of AMH may result from testicular dysgenesis affecting both Leydig and Sertoli cell populations: in this case persistence of Müllerian remnants is associated with ambiguous or female external genitalia. Alternatively, defective AMH action may result from mutations of the genes encoding for AMH or its receptor: in this condition known as Persistent Müllerian Duct Syndrome, testosterone production is normal and external genitalia are normally virilised. Finally, AMH may be normally secreted in intersex patients with defects restricted to androgen synthesis or action, resulting in patients with female or ambiguous external genitalia with no Müllerian derivatives. (Arq Bras Endocrinol Metab 2005;49/1:26-36)

Keywords: Androgens; Cryptorchidism; Germ cells; Intersex states; Persistent Müllerian Duct Syndrome; Sertoli cells

RESUMO

Hormônio Anti-Mülleriano nos Distúrbios da Determinação e Diferenciação do Sexo.

A masculinização dos genitais internos e externos durante o desenvolvimento fetal depende da existência de dois hormônios testiculares distintos: a testosterona secretada pelas células de Leydig conduz à diferenciação dos dutos de Wolff, do seio urogenital e dos genitais externos, enquanto que o hormônio anti-Mülleriano (HAM) produzido pelas células de Sertoli provoca a regressão dos dutos de Müller. A falta de ação do HAM no início da vida fetal resulta na formação de tubas uterinas, útero e terço superior da vagina. Em fetos 46,XY, a falta do HAM pode resultar de disgenesia testicular, afetando tanto as células de Leydig quanto as de Sertoli: nesse caso, a presença de remanescentes Müllerianos está associada a genitais externos femininos ou ambíguos. Por outro lado, distúrbios na ação do HAM podem resultar de mutações em genes que codificam o HAM ou seu receptor: nessa afeição, conhecida como síndrome da Persistência dos Dutos de Müller, a produção de testosterona é normal e os genitais externos são virilizados normalmente. Finalmente, o HAM costuma ser secretado normalmente em pacientes com intersexo decorrente de defeitos restritos à síntese ou à ação de andrógenos, resultando em indivíduos com genitais externos femininos ou ambíguos com ausência de derivados de Müller. (Arq Bras Endocrinol Metab 2005;49/1:26-36)

Descritores: Andrógenos; Criptorquidismo; Células germinativas; Estados intersexuais; Síndrome da persistência dos dutos de Müller; Células de Sertoli
THE EXISTENCE OF ANTI-MÜLLERIAN hormone was first suggested by the pioneering experiments performed by the French scientist Alfred Jost in the 1940's. In early stages of development in mammals, foetuses of both sexes have two pairs of unipotential internal genital ducts: the mesonephric (Wolffian) ducts and the paramesonephric (Müllerian) ducts, and bipotential primordial of external genitalia. Testicular differentiation from the gonadal ridge, occurring in humans during the 7th week of foetal life, drives the fate of internal and external genitalia. First Jost grafted testicular tissue to castrated foetuses and observed that Wolffian ducts gave rise to epididymes, vasa deferentia and seminal vesicles while Müllerian ducts regressed. In subsequent experiments, he noticed that a crystal of testosterone propionate was capable of masculinising Wolffian ducts in the castrated foetus but did not affect Müllerian ducts, which formed the Fallopian tubes, the uterus and the upper third of the vagina. This proved that a testicular product different from testosterone, that he named “hormone inhibitrice” or “Müllerian inhibitor”, was responsible for the regression of Müllerian ducts in the male foetus (1) (figure 1).

However, the identification of the “Müllerian inhibitor” did not prove easy. A test for the detection of testicular anti-Müllerian activity was developed in 1969 (2) but it was only until 1978 that the first evidence was produced indicating the glycoprotein dimeric nature of this testicular factor (3). Finally, AMH could be purified to homogeneity in 1984 (4), and in 1986 the human and bovine genes were isolated and sequenced (5,6). Along its history, the Müllerian inhibitor received several names, of which anti-Müllerian hormone (AMH) and Müllerian inhibiting substance (MIS) are the two most widely used nowadays. In the last 10 years, the use of molecular techniques has allowed the identification of AMH receptors and its signalling pathways, as well as the understanding of the mechanisms involved in the regulation of AMH expression, which will be discussed hereafter. All this knowledge has helped understand the responsibility of AMH in disorders of sex differentiation. As already shown by Jost’s pioneering experiments almost 60 years ago, the testis has a determinant importance in foetal sex differentiation, via two independent pathways. Leydig cells secrete androgens, necessary for the masculinization of Wolffian ducts, the urogenital sinus and external genitalia, whereas Sertoli cells secrete AMH, required for the regression of Müllerian ducts (figure 2). With this simple scheme in mind, the comprehension of intersex states can be more easily achieved.

![Figure 1. Experiments of foetal endocrinology that enabled Alfred Jost to suggest the existence of a testicular factor, different from testosterone, responsible for the regression of Müllerian ducts in the male.](image-url)
AMH: The Protein And The Gene

AMH is a 140kD homodimeric glycoprotein (3). Human AMH is synthesised as a 560 amino acid pre-cursor with a 24-25 amino acid leader containing a 16-18 amino acid signal sequence and a putative 7-8 residue pro-sequence (5) (figure 3). The carboxyl-ter-minal region of AMH shares homology with that of members of the transforming growth factor-β (TGF-β superfamily). Most members of this family require proteolytic cleavage at a site 110 amino acids from the carboxyl terminus to be active. While the full-length AMH molecule is active in organ culture, a cleavage site 109 amino acids from the C-terminus releases a more active fragment (7). However, the cleaved N-terminal domain interacts and enhances the activity of the C-terminus (8). While AMH can be experimentally cleaved using plasmin (8), PC5 and furin, two members of the pro-protein convertase family, have been reported to be able to process AMH in embryonic rat testes (9).

AMH is encoded by a 2.75kb gene divided into 5 exons (figure 3), characterised by a high GC content. The human AMH maps on chromosome 19 p13.3 (10). While the mouse and rat promoters contain an almost perfect TATA box, the human AMH promoter lacks consensus TATA or CCAAT box elements (5). AMH transcription of the human gene has been shown to contain a functional initiator (Inr) element that is specifically recognised by transcription fac-tor TFII-I (11). Cloning of 3.6kb of the 5’-flanking sequences of the human AMH gene allowed to identify a major transcription initiation site and three minor sites, a putative oestrogen response element at –1772 and an Sp1 binding site at –303 (12). The more recent finding of the existence of the SAP62/SF3A2 house-keeping gene, encoding for a spliceosome protein, at –789 of the human AMH ATG codon (and –434 in the mouse), has generated doubts about the real length of the AMH promoter (13) and abridged research studies to the proximal promoter sequences (less than 400bp), where binding elements for SOX/SRY proteins (14,15), SF-1 (16,17) and GATA factors (17,18) have been identified (figure 3). However, recent stud-ies have shown that more distant sequences are required for the normal chronology of expression of AMH in the testis (19,20).

The Ontogeny of AMH Expression

In the mammalian foetus

AMH is one of the earliest Sertoli cell-specific proteins expressed by the gonad (21). As soon as testicular cords begin to assemble in the foetal gonad, AMH expression is triggered. AMH is secreted by the human testis from the 8th week of amenorrhea and provokes irreversible Müllerian duct regression, which is completed by the end of week 9 (22). After that time, Müllerian ducts become insensitive to AMH, which highlights the importance of a tightly regulated chronology of its secretion by the foetal testis. Although AMH is no longer active on Müllerian ducts, its expression by Sertoli cells remains at high levels until puberty, which indicates that the end of the critical window of Müllerian duct regression is dependent on the expression pattern of AMH type II receptor (reviewed in ref. 23).

In the postnatal testis

Except for a transient decline in the peri-natal period, testicular AMH secretion is maintained at high levels until puberty, when Sertoli cell maturation is charac-terised by a decreasing AMH activity (24,25). There-fore, determination of serum AMH levels is useful to assess Sertoli cell maturation (26-28). During pubertal development, AMH expression fains in coincidence with the increase of androgen production by Leydig cells and the onset of germ cell meiosis (24,25,29,30).

In the ovary

Ovarian granulosa cells, the homologous to testicular Sertoli cells, also produce AMH but with several differ-ences: AMH expression only begins at the peri-natal
period (25,31), remains low throughout reproductive life and becomes undetectable after menopause (32). Gonadal AMH secretion shows a clear-cut sexual dimorphism in prepubertal ages, when serum AMH is significantly lower in females; in adults, serum AMH is similarly low in both sexes (26,32).

**AMH Action on Target Organs: Receptors and Signalling Pathways**

The AMH signalling pathway began to be unravelled in 1994, when the specific AMH receptor type II was cloned (33,34). This specific AMH receptor is present on the cell membrane of target organs and is responsible for ligand binding. It subsequently recruits a non-specific type I receptor in order to transduce its signal. Three different type I receptors are considered to mediate AMH response in target cells: ALK6, named BMPRI-B (35), ALK2, named ActRI (36,37), and ALK3, also known as BMPRI-A (38). The intracellular transduction pathways involved after type I receptor recruitment by the specific type II AMH receptor seem to vary according to the target cell (reviewed in ref. 39). AMH receptor type II is a single transmembrane protein with serine/threonine kinase activity, that is encoded by a 8kb gene composed of 11 exons and mapping to chromosome 12q13 (40).

AMH receptor type II has been identified in the mesenchymal cells surrounding the epithelium of Müllerian ducts in both sexes. Its expression follows a cranial-to-caudal chronology that explains why the cranial-most end of the ducts, that are near the testes, regress first and the more distant portions of the ducts regress later (41). After AMH binding to its receptor in mesenchymal cells, paracrine-mediated mechanisms trigger epithelial cell apoptosis and epithelial-mesenchymal transformation finally resulting in complete Müllerian duct regression in the male. The absence of AMH (e.g. in the normal female foetus) or of its signalling mechanisms (e.g. mutations of the AMH receptor) results in the stabilization of Müllerian ducts. AMH receptors are also present in granulosa cells of the ovary and in Sertoli and Leydig cells of the testes. In different experimental conditions, AMH has been shown to inhibit Leydig cell differentiation and steroidogenesis as well as granulosa cell response to LH and FSH, but whether it plays any essential role in gonadal physiology still needs to be determined (42,43).

**The Regulation of AMH Expression**

Owing to its sex-specific and time-restricted requirement during foetal development, AMH expression needs to be tightly regulated (reviewed in ref. 44). Uncontroversial proof exists indicating that SOX9 binding to the proximal AMH promoter is essential for the initiation of AMH expression in early foetal development (45). SF-1 (16), GATA-4 (18) and WT-1 (46) enhance, while DAX-1 impairs (46), AMH transcription.

As already mentioned, testicular AMH production remains high during foetal life and childhood and is downregulated at puberty. The decline of AMH production by Sertoli cells is related to the stage of pubertal development: the most significant decrease in serum AMH is observed between stages II and III of pubertal development (29), in coincidence with the increase of intratesticular testosterone concentration (figure 4): the decline in AMH production is a marker of the elevation of intratesticular androgen concentration, which

![Figure 3. The AMH gene and its 3-kb promoter (top) and the AMH immature and mature proteins.](image-url)
inhibits AMH production at puberty. Independently from androgen action, meiotic germ cells seem to play a role in downregulation of AMH expression at puberty (30,47). Interestingly, AMH is not down-regulated by testosterone during foetal life and in the first months after birth owing to the lack of expression of the androgen receptor in Sertoli cells (30). The physiological androgen insensitivity of foetal and neonatal Sertoli cells explains, thus, the transient coexistence of high concentrations of androgens and AMH (figure 4). In the absence of the negative effect of androgens and meiotic germ cells (e.g. in foetal and neonatal periods), FSH upregulates testicular AMH production. On one hand, FSH induces Sertoli cell proliferation and, on the other, it enhances AMH transcription in each Sertoli cell through a cAMP-PKA mediated signalling pathway involving transcription factors AP2 and NFB, which bind to specific response elements in distant sequences of the AMH promoter (20).

Abnormal AMH Secretion or Action
When AMH is not produced between the 8th and the 9th weeks of foetal life, or if the AMH receptor pathway is not capable of transducing AMH signalling, Müllerian ducts differentiate to form the Fallopian tubes, the uterus and the upper vagina. Absence of AMH action is the normal situation in the XX foetus: although the AMH receptor and its signalling pathways are present in Müllerian ducts, the foetal ovaries do not express AMH. In the XY foetus, the absence of AMH action results in an abnormal persistence of Müllerian ducts. This can be an isolated situation, known as the persistent Müllerian duct syndrome, or a disorder in which the existence of female internal genitalia is associated with defective virilisation of the Wolffian ducts, urogenital sinus and external genitalia.

The Persistent Müllerian Duct Syndrome (PMDS)
The uncontroversial and unique role of AMH in foetal sex differentiation can be deduced from the phenotype of patients with AMH or AMH receptor mutations (48) or of transgenic mice with manipulations of AMH expression or signalling (49-51). It seems clear that the absence of AMH activity in early foetal life results in one conspicuous phenotype, the persistence of Müllerian ducts. No other growth factor or hormone is capable of replacing AMH in this function.

For PMDS to be considered as a possible diagnosis in an XY patient, a *sine qua non* condition is that the external genitalia, urogenital sinus and Wolffian ducts have normally differentiated, accounting for normal androgen activity. Therefore, PMDS patients are genotypic and externally phenotypic males with cryptorchidism, sometimes associated with inguinal hernia. The presence of Müllerian derivatives is usually not suspected and discovered at surgery. Two anatomical forms of PMDS have been described. The most common one is known as transverse testicular ectopia: one testis descends into the scrotum dragging the ipsilateral Fallopian tube into the inguinal canal (a condition known as *hernia uteri inguinalis*) and pulling the uterus and the contralateral Fallopian tube together with the testis, which becomes located in the abdomen. More rarely, PMDS presents as bilateral cryptorchidism, the uterus is fixed in the pelvis and both testes are embedded in the broad ligament in ovarian position. These clinical variants are not genetically determined and may occur within the same sibship (52). In PMDS, the testes are abnormally mobile because they are not anchored to the scrotum (53). The spermatic cord is usually very short because the vas deferentia are embedded in the mesosalpinx,
lateral uterine wall and cervix. Lack of proper communication between the testis and excretory ducts and difficulties at orchidopexy probably explain why fertility is rare in PMDS patients (55).

Two biological variants of PMDS can be distinguished: those with normal AMH production (AMH-positive) and those with impaired or null AMH secretion (AMH-negative). The AMH-negative variant is due to mutations in the AMH gene, resulting in complete lack of AMH production or secretion. These patients have very low or undetectable serum AMH concentrations because (56), in rare cases with AMH mutations, serum AMH concentration may be normal for age; these mutations impair bioactivity but not secretion (57). The AMH-positive variant is generally due to mutations in the gene coding for AMH receptor type II (AMHR-II); as expected, these patients have a normal AMH serum concentration for their age (58) (figure 5). Serum testosterone and response to hCG are normal in all cases. Female relatives of AMH-resistant patients who share their genetic background are phenotypically normal and fertile.

As could be expected the inheritance of PMDS is according to an autosomal recessive pattern: affected patients are either homozygotes or compound heterozygotes. AMH gene mutations have been reported in 47% of PMDS families: 35 different mutations, for the most part missense type, have been detected along the gene, with no hot spots (figure 6). AMHR-II mutations have been found in 38% of PMDS families: 25 different mutations of AMHR-II, mostly missense, were spread out over the whole length of the gene (figure 6). Five mutations were recurrent, with a deletion of 27bp in exon 10 present in 45% of families of this group (48). In 15% of the families studied to date, no mutation of either the AMH or the AMHR-II genes was detectable and no large DNA rearrangements were seen by Southern-blotting. Associated diseases, such as jejunal atresia, lipoatrophic diabetes and vitamin D resistant rickets, lymphangectasia and other malformations were present in approximately half the cases. Unexplained PMDS may reflect mutations in the proteolytic enzyme involved in AMH processing or in components downstream of the type II receptor in the AMH transduction cascade; alternatively, in view of the high incidence of associated defects, unexplained PMDS may be part of complex malformative syndromes with no relationship to the AMH pathway.

Persistence of Müllerian derivatives associated with ambiguous external genitalia

Based on the endocrinology of foetal sex differentiation (figure 2), it can be easily deduced that a combined defect of androgen and AMH secretion should be suspected if Müllerian derivatives are associated in an XY patient with ambiguous or female external genitalia. This phenotype excludes the diagnosis of PMDS and genetic analysis of the AMH or AMHR-II genes is unnecessary. Since both Leydig and Sertoli cells are affected, gonadal dysgenesis is the most probable aetiology. Depending on the degree of gonadal dysgenesis, testosterone and AMH levels might be from low to undetectable (27). This usually correlates to the anatomic phenotype: low testicular hormones are observed in 46,XY patients with ambiguous external genitalia and rudimentary Wolffian and Müllerian

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**Figure 5.** Serum AMH levels in patients with Persistent Müllerian Duct Syndrome (PMDS). The shaded area indicates the normal serum AMH range, as assayed by the AMH/MIS ELISA® (Immunotech-Coulter-Beckman).
derivatives (a condition commonly referred to as dysgenetic male pseudohermaphroditism). Undetectable AMH and no response of testosterone to hCG is usually observed in 46,XY females (pure gonadal dysgenesis, also known as Swyer syndrome).

In 46,XX patients bearing a uterus and with very low or undetectable AMH levels, the most probable diagnosis is congenital adrenal hyperplasia (59). However, most often the diagnosis is made early in the neonate owing to adrenal insufficiency.

In patients with rudimentary Müllerian derivatives with XX karyotype or with mosaicism, the degree of Müllerian structure development is indicative of the amount of AMH produced by the gonads. AMH acts as a local factor on the homolateral Müllerian duct. Therefore, an ovotestis with a scarce testicular component is usually associated with a homolateral hemi-uterus and Fallopian tube, but these structures are very hypoplastic or even absent if the ovotestis is composed of abundant testicular parenchyma. Similarly, in patients with asymmetric gonadal differentiation (mixed gonadal dysgenesis), a Fallopian tube and hemi-uterus are present on the side of the streak gonad; contralaterally the degree of testicular dysgenesis will determine the development of Müllerian derivatives.

Absence of Müllerian derivatives in XY patients with ambiguous external genitalia

Once again, based on the endocrinology of foetal sex differentiation (figure 2), the absence of Müllerian ducts in a 46,XY patient indicates that the gonads are not dysgenetic, since Sertoli cells have correctly differentiated and secreted sufficient amounts of AMH (27). In these cases, referred to as non-dysgenetic male pseudohermaphroditism, only the androgen pathway is affected, and the probable diagnoses are defects of the LH receptor, of steroidogenic proteins or of the androgens receptor or its coactivators. When the impairment of androgen secretion or action is severe, a female external phenotype results together with a smaller vagina and the absence of uterus and tubes. These patients may only present at puberty owing to primary amenorrhea.

Clinical Utility of AMH Measurement in Intersex Patients

Except for the first 7-14 days of postnatal life, when endocrine testicular activity seems to be low, serum AMH levels reflect the amount of functional Sertoli cells (i.e. testicular parenchyma) in prepubertal boys, before they decline owing to the inhibitory effect

Figure 6. Mutations found in patients with Persistent Müllerian Duct Syndrome (PMDS).
exerted by androgens and meiotic germ cells (24,28). Therefore, in most cases serum AMH can be used to estimate what might have happened in foetal life (one should be aware that this is an extrapolation which could overlook, for instance, a late onset of AMH secretion after the 9th week, in which case high AMH would coexist with persistent Müllerian ducts).

Normally high serum AMH indicates that Sertoli cells are quantitatively and qualitatively normal: the predictive value of serum AMH for the existence of normal testicular tissue is higher than that of testosterone response to hCG (60). In 46,XY patients with female or ambiguous external genitalia, the androgen pathway is clearly impaired. The possibilities are numerous and androgen response to hCG may not always elucidate the diagnosis. For example, a low response to hCG can be observed in gonadal dysgenesis but also in patients with LH receptor or steroidogenic enzyme defects. Serum AMH determination can be very helpful, since levels are low in testicular dysgenesis but normal to extremely high in patients with mutations in the LH receptor or steroidogenic proteins (27) (table 1). Another difficult differential diagnosis is that between androgen insensitivity owing to a mutation in the androgen receptor and a defect in DHT owing to a mutation in 5α-reductase, serum AMH may also be helpful here. AMH is downregulated by intratesticular testosterone in normal conditions. This inhibition is absent, resulting in high AMH, in patients with androgen insensitivity (27) (table 1). Conversely, in patients with 5α-reductase deficiency, AMH down-regulation by testosterone occurs normally (61).

In 46,XY normally virilised boys with nonpalpable gonads, normal serum AMH clearly indicates the existence of testes in ectopic position (62,63). Undetectable AMH would indicate anorchia (or vanishing testes, if previously present), with the sole and rare exception of cryptorchidism owing to AMH-negative PMDS. Normal AMH also suggests that FSH activity has been adequate during foetal and neonatal periods, whereas low AMH may suggest congenital hypogonadotropic hypogonadism (64) (table 2). Conversely, high AMH has been reported in a patient with Sertoli cell hyperplasia due to an activating mutation of the Gsα subunit involved in the FSH receptor signalling pathway (65). Declining serum AMH is indicative of pubertal development of the testes. An early decrease of AMH is observed in patients with central precocious puberty or with testotoxicosis (table 2); serum AMH recovers normal values after treatment (29). Finally, persistently high levels of AMH in boys of pubertal age might indicate delayed puberty if androgens are low or mild androgen insensitivity if androgen levels have risen. Hypogonadotropic hypogonadism seems to be characterised by low levels of both androgens and AMH in patients with micropenis or cryptorchidism but no ambiguous genitalia (table 2).

| Table 1. Serum AMH and testosterone in 46,XY intersex patients. |
|----------------------------------|------------------|
| **Testosterone**                 | **AMH**          |
| Low or Undetectable Low or Undetectable | Normal or High | Gonadal dysgenesis 5α-reductase defect |
| Normal or High                   | LH receptor defects Steroidogenic protein defects | Androgen Insensitivity |

| Table 2. Serum AMH and testosterone in 46,XY boys with normally virilised external genitalia. |
|----------------------------------|------------------|
| **Testosterone**                 | **AMH**          |
| Low or Undetectable Undetectable | Anorchia PMDS (AMH mutation) |
| Low                              | Hypogonadotropic hypogonadism Normal or precocious puberty |
| Normal (high for pubertal age)   | Delayed puberty Bilateral Cryptorchidism PMDS (AMH-II mutation) Mild PAIS |
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