MEDICINE

P uberty is a sensitive phase of physical, mental, and social development for both girls and boys. A thorough acquaintance with the normal course of puberty is necessary. Any deviation from it, though it will be viewed with great anxiety by the young patient, can represent either a normal or pathological variant of pubertal development. The physician should be able to provide the young patient with accurate information and see him or her through the process of puberty in a reassuring manner.

If a normal variant is present, the treating physician can help the patient and his or her parents with thorough counseling. Rarely there is a need for a time-consuming and expensive diagnostic evaluation. If the child’s pubertal development is pathological, the cause of the pubertal disorder must be found by specific diagnostic testing, and the necessary treatment must be initiated.

The learning objectives of this article are to acquaint the reader thoroughly with:

1. Normal pubertal development and its temporal course;
2. Normal variants and pathological disorders of pubertal development, and their etiologies.

This article is based on a selective review of the literature. No clinical guidelines on this subject are available.

Normal puberty

The hypothalamic-pituitary-gonadal axis undergoes an active phase during fetal and neonatal development and then enters a resting phase that lasts for the rest of childhood till puberty. Puberty begins with an activation of the hypothalamic-pituitary-gonadal system (figure). The influence of the hypothalamic hormone GnRH (gonadotropin releasing hormone), the gonadotropins LH (luteinizing hormone) and FSH (follicle-stimulating hormone), and the sex steroids estradiol or testosterone brings about the manifestations of puberty, both external

Hormonal control of puberty

Puberty begins with activation of the hypothalamic-pituitary-gonadal system.
(breast development, genital enlargement) and internal (uterus, ovaries, testes). Pubic hair develops independently of the activation of the hypothalamic-pituitary-gonadal pathways, largely through the effect of androgens secreted by the adrenal glands (adrenarche).

The different phases of external pubertal development in girls are conventionally designated as Tanner stages B1 through B5 for breast development and PH1 through PH6 for pubic hair growth. For a detailed description of the Tanner stages, the reader can refer to standard textbooks of pediatrics and pediatric endocrinology (1, 2). Tables 1 and 2 show the timing of the various stages in girls and boys, respectively.

The mammary gland grows from a breast bud that can be palpated under the nipple (Tanner stage B2) to a fully developed female breast (Tanner stage B4 or B5) over a period of 3.6 years, on average (3, 4).

Ultrasonography reveals an increase in uterine volume, at first without, and then with, a visible layer of uterine mucosa ("mucosal reflex"). The ovaries develop follicular cysts. Multicystic ovaries with more than six cysts can already be seen in the early stages of puberty (5). The first menstrual period (menarche) occurs at an average age of 13.4 years, according to the longitudinal data obtained by Largo et al. (3). In 2006, the German Health Interview and Examination Survey for Children and Adolescents (Kinder- und Jugendgesundheitsurvey, KiGGS), using the status quo method, found the median age at menarche to be 12.8 years (6). The mean age at menarche is highly correlated within families, between monozygotic twins, and within ethnic groups (7). Emotional factors and the nutritional state are also important.

At first, the menstrual periods are irregular and consist mainly of anovulatory cycles; in 80% of girls, the menstrual periods become regular and ovulatory within five years after the menarche.

In boys, pubertal development begins with an increase in testicular size, which can be gauged using the Prader orchidometer (8); the normal values for testicular size

<table>
<thead>
<tr>
<th>TABLE 1</th>
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<tbody>
<tr>
<td><strong>Chronological age at the onset of pubertal development in girls (Tanner stages). Mean, standard deviation (SD), and normal range of the initial development of signs of puberty (−2SD to +2SD)</strong></td>
</tr>
<tr>
<td>Parameter</td>
</tr>
<tr>
<td>PH 2</td>
</tr>
<tr>
<td>B 2</td>
</tr>
<tr>
<td>PHV</td>
</tr>
<tr>
<td>Menarche</td>
</tr>
</tbody>
</table>

*B = breast development; PH = development of pubic hair; PHV = peak height velocity, i.e., the time when longitudinal growth during puberty is fastest

*modified from (3)
during puberty are as given by Zachmann (9). Over the further course of puberty, pubic hair appears (Tanner stages PH1 through PH6), as does facial hair, the voice breaks, and penis size increases (Tanner stages G1 through G5) (4) (table 2).

Spermatozoa first become detectable in specimens of boys’ spontaneously produced morning urine at a mean age of 13.4 years (spermarche) (10). As the testes become larger in the ensuing years, the maturation of spermatogenesis is completed.

**Psychosocial and psychopathological aspects of pubertal development**

Major mental and emotional changes accompany the physical changes of puberty. A basic structuring of the personality (identity) takes place; the child separates itself emotionally from its parents and experiences social orientation outside the family. These mental and emotional maturational processes do not necessarily occur in close parallel with the young person’s physical development.

During puberty, the body comes to be perceived differently. Feelings of self-worth or self-doubt can arise, as can feelings of shame and emotional vulnerability. An increasingly strong pursuit of autonomy leads the young person to retreat into his or her own living space and establish a separation from other family members, despite a persistent dependency on them. Social orientation to the peer group takes place. The manner in which the individual copes with these developments is influenced by the perceptions of others. Family members, friends, teachers, and the entire social environment no longer see a child, but rather an adolescent. These processes often manifest themselves in the adolescent’s behavior as emotional lability or insecurity, or else as rebelliousness or aggression. In the phase of social maturation, the adolescent breaks away from the family, assumes personal responsibility and social roles, and contends with the larger society’s norms and values (11).

Other physical changes, such as the weight gain accompanying puberty, can lead to emotional disturbances because they conflict with the Western ideal of slimmness. This development can cause especially girls to suffer from eating disorders (anorexia nervosa, bulimia) or depression (13, 14).

**Normal variants of puberty**

The diagnostic evaluation of disorders of puberty necessarily involves the differentiation of pathological disorders from the normal variants of early or late puberty (table 3). Normal variants include constitutional delay or acceleration of growth and puberty (pubertas tarda, pubertas accelerata), early breast development (premature thelarche), early development of pubic hair (premature pubarche), and pubertal gynecomastia.

In constitutional delay of growth and puberty (pubertas tarda), the spontaneous onset of puberty occurs more than two standard deviations later than normal mean age of onset—or, alternatively, later than the 97th percentiles—for the Tanner stage in question (see tables 1 and 2) (15, 16). For example, in a girl with delayed puberty, breast development to Tanner stage B2 occurs when she is more than 13.3 years old; in a boy with pubertas tarda, a testicular volume of 3–4 mL is first reached when he is (12, 13). On the other hand, late pubertal development can lead transiently to a negative self-image. Emotional disturbances such as expansive behavior and alcohol and drug consumption may arise as potential compensating mechanisms.

**TABLE 2**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Mean (years)</th>
<th>SD (years)</th>
<th>Normal range (years) (–2SD to +2SD)</th>
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</thead>
<tbody>
<tr>
<td>G 2</td>
<td>11.2</td>
<td>1.5</td>
<td>8.2</td>
</tr>
<tr>
<td>Testicular volume (≥ 3 mL)</td>
<td>11.9</td>
<td>0.9</td>
<td>10.1</td>
</tr>
<tr>
<td>PH 2</td>
<td>12.2</td>
<td>1.5</td>
<td>9.2</td>
</tr>
<tr>
<td>Spermarche</td>
<td>13.4</td>
<td>—</td>
<td>11.7</td>
</tr>
<tr>
<td>PHV</td>
<td>13.9</td>
<td>0.8</td>
<td>12.3</td>
</tr>
</tbody>
</table>

G = genital development; PH = development of pubic hair; PHV = peak height velocity, i.e., the time when longitudinal growth during puberty is fastest

*Modified from (4) and (10)

**Psychosocial aspects of puberty**

The mental and emotional changes of puberty include basic structuring of the personality (identity), separation of self from parents, and social orientation outside the family.

**Normal variants**

Normal variants include constitutionally delayed or accelerated growth and puberty, premature thelarche, premature pubarche, and pubertal gynecomastia.
more than 13.7 years old. Growth is markedly slower than usual for age, and height falls from the patient's previously maintained percentile to a lower one. The patient's adult height is often (17), but not always (18), in the target height range. A suspected diagnosis of constitutional delay of growth and puberty can be definitively confirmed only when puberty and the pubertal growth spurt finally do occur spontaneously, more than two standard deviations later than the normal mean age.

The mirror image of constitutional delay of growth and puberty is constitutional acceleration of growth and puberty. The onset of puberty (breast development, testicular enlargement) occurs more than two standard deviations earlier than the normal mean age of onset. The hypothalamic-pituitary-gonadal axis is activated earlier than usual, but not before age 8 in girls (breast development) or before age 9 in boys (testicular enlargement) (differential diagnosis: precocious puberty).

In girls with premature thelarche, breast tissue develops early—sometimes in the first two years of life, sometimes later, and occasionally even in the neonatal period—and then persists until the true onset of puberty (19). Premature thelarche is an incomplete form of pubertal development without any other signs of early puberty, such as a pubertal growth spurt or marked acceleration of skeletal age. In rare cases, premature thelarche undergoes a transition to precocious puberty (20).

Premature pubarche is a further normal variant of pubertal development in which pubic and/or axillary hair develop in a girl before age 8 or in a boy before age 9 (21). The serum levels of adrenal androgens are normal, and there is no androgen-induced growth spurt. The skeletal age corresponds to the chronological age or is only mildly accelerated. Premature pubarche is to be distinguished from premature adrenarche with elevated levels of adrenal androgens, the adrenogenital syndrome, and an androgen-secreting tumor.

Some degree of pubertal gynecomastia can be observed in about 50% to 90% of boys (22). In most cases, breast tissue is palpable only as a small induration under the nipple that regresses spontaneously in 6 to 18 months. Treatment is not indicated.

In rare cases, the gynecomastia is more pronounced and persists longer, so that severe psychosocial problems ensue that may necessitate medical treatment, e.g., with tamoxifen (23), or surgical intervention. Only in exceptional cases is pubertal gynecomastia due to increased estrogen production or other hormonal causes, chromosomal anomalies such as Klinefelter syndrome (47, XXY), or medications such as spironolactone. Pubertal gynecomastia is to be distinguished from lipomatosis in an overweight male adolescent.

### The pathology of early puberty

Pathological conditions in which puberty occurs early are divided into gonadotropin-dependent disorders (true precocious puberty) and gonadotropin-independent disorders (precocious pseudopuberty). A further possibility to be considered is pathological adrenal dysfunction leading to the isolated, premature appearance of pubic hair (premature adrenarche) (table 3).

### Gonadotropin-dependent early puberty (true precocious puberty)

By definition, true precocious puberty is initiated by premature activation of the hypothalamic-pituitary-gonadal axis. Its prevalence is estimated at 1:5000 to 1:10 000. It is five to ten times more common in girls than in boys (24, 25).

Pulsatile GnRH secretion begins as in normal puberty. In girls, breast tissue develops under estrogenic influence. In boys, the testes enlarge first, and then the penis enlarges under the influence of testosterone. The clinical course resembles that of normal pubertal development. Pubic hair usually appears later, as a result of stimulation by androgens derived from the adrenal gland and from the testes or ovaries.

### Delayed puberty

When growth and puberty are constitutionally delayed, puberty spontaneously begins at a time that is more than two standard deviations later than the mean time of breast development in girls or increase in testicular volume in boys.

### Pubertal gynecomastia

Pubertal gynecomastia of varying extent is seen in 50% to 90% of boys.
An early pubertal growth spurt occurs at about the same time as the external signs of puberty appear. Because of the simultaneous acceleration of skeletal development, the individual’s final height is often reduced, sometimes to the point of short stature, i.e., height below the third percentile.

The diagnosis of true precocious puberty must be considered when the initial signs of puberty appear in a girl under age 8 or a boy under age 9. The diagnosis is then confirmed by the clinical findings, including the Tanner stages and growth spurt, acceleration of skeletal growth, the results of GnRH testing, and the findings of gonadal and uterine ultrasonography. In the GnRH test, the stimulated LH/FSH quotient at 30 minutes is greater than 1. The estradiol level (in girls) or the testosterone level (in boys) is markedly elevated for chronological age, but corresponds to the current pubertal stage.

In girls, ultrasonography reveals a multicystic ovary with more than 6 follicles of diameter 4 mm or more (5). The uterine volume increases, and endometrium is produced (a “uterine mucosal reflex” is visible).

The cause of precocious puberty remains unidentified in 80% of girls and 40% of boys. Aside from these idiopathic cases, precocious puberty can also result from organic lesions in the hypothalamic and pituitary area, primarily in boys (e1). Hypothalamic hamartoma, glioma, astrocytoma, and germinoma can cause precocious puberty; it can also occur in children with internal hydrocephalus or other lesions of the central nervous system, such as an earlier episode of meningitis or traumatic brain injury or prior radiotherapy to the head. Magnetic resonance imaging of the brain should be performed in order to search for a possible organic cause.

In true precocious puberty, treatment is indicated because of the major psychosocial stress on the affected child resulting from the very early appearance of signs of puberty, the frequent (and generally wrong) assumption by others that the child possesses a correspondingly early mental and emotional “maturity,” and the risk of reduced adult height due to disproportional acceleration of skeletal age. The treatment involves the administration of GnRH analogs that suppress the effects of the elevated gonadotropins LH and FSH through down-regulation of the pituitary GnRH receptors (24). Physical examinations during treatment reveal the slowing or cessation or sometimes even a return of the prepubertal stage; on biochemical testing, the gonadotropins LH and FSH as well as the sex steroids estrogen or testosterone are detectable only in very low concentrations, or not at all. In doubtful cases, a GnRH test can be performed during the trough just before the next scheduled GnRH injection, in order to determine whether the gonadotropins have been adequately suppressed. If basal and/or stimulated LH and FSH are measured in higher concentrations, then GnRH analogs should be given at a higher dose or at shorter intervals. The pubertal stage, height, and skeletal age of the patient should be monitored over the course of treatment (24).

The treatment of true precocious puberty should be terminated when it is time for normal puberty to begin, and when it can be expected that the patient will attain an optimal adult height. The decision to end treatment should be taken by general agreement between the physician, the child, and the parents. Puberty will then run its course spontaneously, the duration of puberty depending on the stage that had already been attained by the time the treatment for precocious puberty was discontinued. The follow-up studies on treated patients have not revealed any treatment-related disturbances of the hypothalamic-pituitary-gonadal axis (e2, e3). When treatment is begun early, the patient’s adult height lies in the range predicted before therapy was begun (e4).

**Gonadotropin-independent early puberty (precocious pseudopuberty)**

Precocious pseudopuberty arises, by definition, before and independently of the maturation of the hypothalamic-pituitary-gonadal axis (e5). The appearance of secondary sexual characteristics is due to the increased production of female or male hormones. Estrogens induce isosexual pseudopuberty in girls and heterosexual pseudopuberty in boys; conversely, androgens induce isosexual pseudopuberty in boys and heterosexual pseudopuberty in girls. The hypothalamic-pituitary-gonadal axis is suppressed by the abnormally elevated secretion of androgens or estrogens.

Precocious pseudopuberty has many different causes. It can be due to external factors, such as the therapeutic or accidental ingestion of estrogens or androgens, or a large number of other conditions. These include tumors, disturbances of steroid biosynthesis, and congenital syndromes (e5).

Hormone-secreting tumors of the central nervous system, the adrenal gland, the liver or other organs can be responsible for the development of precocious pseudopuberty. Germ-cell tumors secrete human chorionic gonadotropin (hCG), which, in turn, stimulates the LH-receptors of the testes (for example), which then

**Pathological forms**

Pathological forms of puberty include gonadotropin-dependent precocious puberty, gonadotropin-independent precocious pseudopuberty, and premature adrenarche.

**The time of diagnosis**

The diagnosis of precocious puberty should be considered when girls develop the first signs of puberty before age 8, or boys before age 9.
produce testosterone. Tumors of this type can arise in the gonads, the central nervous system (pineal and pituitary glands), the liver, the retroperitoneal space, or the posterior mediastinum, which are the sites of origin of the sex-determining cells during embryonic development (e6).

Adrenal tumors can produce androgens as well as cortisol and thereby induce iso- or heterosexual precocious pseudopuberty in addition to the clinical signs of Cushing syndrome (e7).

Leydig cell tumors must also be considered in the differential diagnosis of precocious pseudopuberty. Physical examination often, though not always, reveals a palpable asymmetry of testicular size. In unclear cases, ultrasonography, magnetic resonance imaging, and/or testicular biopsy must be performed.

A special case is that of familial, gonadotropin-independent precocious pseudopuberty ("familial testotoxicosis"), a disorder in which an activating mutation of the LH receptor causes early Leydig cell maturation and increased testosterone production (e8). The disease is transmitted in an autosomal dominant inheritance pattern affecting boys only. In the affected boys, signs of puberty appear at the age of 1 to 4 years.

Another very special case is that of McCune-Albright syndrome, which is characterized by three clinically evident phenomena (e9). First, the characteristic, jagged café-au-lait spots become visible on the skin. Later, polyostotic fibrous dysplasia of the bones develops, leading to pathological fractures that can result in severe osseous deformities. Puberty occurs early as the result of repeatedly arising ovarian cysts, which can cause withdrawal bleeding even in early childhood.

Patients with McCune-Albright syndrome have a somatic mutation in the alpha subunit of the G protein, leading to continuous activation of adenylate cyclase and thereby to the pathological secretion of a variety of different hormones. In addition to precocious pseudopuberty, the patient may suffer from hyperthyroidism, Cushing syndrome, and increased growth hormone secretion.

Hereditary disorders of adrenal steroid biosynthesis are transmitted in an autosomal recessive pattern. The associated enzyme defect in androgen, glucocorticoid, and/or mineralocorticoid synthesis produces a clinical condition that presents with premature adrenarche (e10). The enzyme defect can be diagnosed by means of a steroid profile and/or ACTH test in combination with molecular genetic analysis. 21-hydroxylase deficiency is the most common cause, leading to the simple virilizing form of the adrenogenital syndrome. A 3ß-hydroxysteroid dehydrogenase defect should be included in the differential diagnosis.

Pathological absence of puberty
When puberty does not occur spontaneously, no development of secondary sexual characteristics is observed. If pubic hair develops, this is usually due to the secretion of adrenal hormones and does not imply activation of the hypothalamic-pituitary-gonadal axis. The concentrations of the pituitary hormones LH and FSH are low when the disturbance has its origin in the hypothalamus or pituitary gland (hypogonadotropic hypogonadism); they are high when the cause is ovarian or testicular failure (hypergonadotropic hypogonadism). In either case, the level of the gonadal hormone, estradiol or testosterone, is low. Follicular maturation or sperm production does not take place.

Further diagnostic evaluation is needed if no breast development has yet occurred in a girl aged 14.5 years (mean + 3 standard deviations) or if the testes have not reached a size of 3 mL or more in a boy aged 14.6 years (mean + 3 standard deviations). Even at this late age, the major differential diagnosis is constitutional delay of growth and puberty.
Hypogonadism, whether it is tertiary (hypothalamic), secondary (pituitary), or primary (gonadal), can be either congenital (e11) or acquired (figure). The classic example of congenital hypogonadal hypogonadism is Kallmann syndrome, in which hypogonadism is characteristically accompanied by hyposmia or anosmia (box 1, rightmost column in the figure). Kallmann syndrome is due to an impairment of the normal migration of the GnRH neurons from the region of the olfactory nerve to the ventral hypothalamus. Mutations in the KAL1 gene on the short arm of the X chromosome (Xp22.3) are responsible for the X-chromosomal recessive form, while mutations in the FGFR1 (fibroblast growth factor receptor 1) gene on the short arm of chromosome 8 (8p11.2–p11.1) are responsible for the autosomal dominant form (e12). In addition, mutations in the prokineticin-2 gene, the prokineticin-2 receptor gene, and the nasal embryonic LHRH factor (NELF) gene have been described as rare causes of Kallmann syndrome (e12–e14) (box 1).

Hypogonadotropic hypogonadism can also manifest itself as a disturbance of GnRH secretion without any abnormality of the sense of smell (e15–e16). Mutations in the GnRH receptor account for about 6% to 13% of all autosomal recessive forms of hypogonadotropic hypogonadism (e17). Further hypothalamic causes of absent puberty are listed in box 1. Inactivating mutations in the GPR54 (G-coupled protein receptor 54) gene are located on chromosome 19 (19p13.3) (e18).

The term “functional hypothalamic hypogonadism” refers to a usually reversible dysfunction of the hypothalamic-pituitary-gonadal axis that can arise in the setting of anorexia nervosa, during situations of severe stress, or when the affected person participates in very intense physical activity, including sport.

A number of different disorders can be the cause of secondary (pituitary) hypogonadotropic hypogonadism (see “pituitary hypogonadism” in the figure and box 2). Congenital developmental abnormalities of the pituitary gland usually cause a complex deficiency of multiple pituitary hormones; therefore, the diagnosis of hypogonadotropic hypogonadism is often preceded by that of a growth hormone deficiency, pituitary hypothryoidism, and/or pituitary ACTH deficiency. The associated genetic defects affect the transcription factors HESX1, PROP1, LHX3, LHX4, and others, which are responsible for the normal development of the pituitary gland during embryogenesis. Magnetic resonance imaging may reveal hypoplasia or aplasia of the adenohypophysis, a rudimentary or absent pituitary stalk primordium, and/or ectopy of the neurohypophysis.

The pituitary hormones LH and FSH consist of a common alpha-subunit and a specific beta-subunit. Mutations in the latter or in the receptors for LH and FSH cause hypogonadotropic hypogonadism (e19). Moreover, radiotherapy of the head for the treatment of leukemia and brain tumors can cause either secondary or tertiary hypogonadism.

The classic congenital gonadal disorders associated with hypergonadotropic hypogonadism are chromosomal anomalies such as Ullrich-Turner syndrome (45, X) in girls and Klinefelter syndrome (47, XXY) in boys (45) (see “hypergonadotropic hypogonadism” in the figure). Acquired forms are due to autoimmune diseases, radiotherapy, or chemotherapy. These patients have significantly elevated LH and FSH levels because of the lack of the negative feedback mechanism that normally results from the estrogens or from testosterone, as well as from the inhibins A and B, and that normally inhibits secretion of the hypothalamic-pituitary hormones LH.

Pathology of absent puberty
Hypogonadism is divided into hypogonadotropic hypogonadism (of hypothalamic or pituitary origin) and hypergonadotropic hypogonadism.

Kallmann syndrome
The classic type of hypothalamic hypogonadism, in which this disturbance is combined with anosmia, is called Kallmann syndrome.
and FSH. On clinical examination, the patient is found to be in the prepubertal Tanner stages B1 and G1, with testicular volume less than 3 mL. Gonadal ultrasonography reveals a small uterus without a mucosal reflex. The secretion of adrenal steroids is unimpaired in most cases, and pubic hair therefore appears in the normal fashion.

Any type of hypogonadism can arise either as the complete form of the disease, in which all signs of puberty are absent, or as an incomplete form, in which partial functioning of the hypothalamic-pituitary-gonadal pathway is reflected in some degree (usually incomplete) of external pubertal development. In some cases, such disturbances can be difficult to differentiate from constitutional delay of growth and puberty (pubertas tarda).

The treatment of hypogonadism consists of substitution therapy with estrogens/gestagens or with testosterone. It begins with low doses of estrogens or testosterone, which are slowly raised to the full substitution dose while the clinical findings are monitored (development of secondary sexual characteristics). For further details, the reader is referred to textbooks of pediatric (2) and adult endocrinology (e21).

**REFERENCES**


**Classic disorders**
The classic forms of hypergonadotropic hypogonadism are Ullrich-Turner syndrome (45, X) in girls and Klinefelter syndrome (47, XXY) in boys.

**The treatment of hypogonadism**
The treatment of hypogonadism begins with a low-dose substitution therapy with estrogens/gestagens in girls or testosterone in boys.


Further Information on CME

This article has been certified by the North Rhine Academy for Postgraduate and Continuing Medical Education.

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For volume 21/2009 we plan to offer the topic “The Differential Diagnosis of Food Intolerance.”

Solutions to the CME questionnaire in volume 9/2009:

Schrem H, Barg-Hock H, Strassburg CP, Schwarz A, Klempnauer J: Aftercare for Patients With Transplanted Organs: 1c, 2d, 3d, 4a, 5b, 6d, 7e, 8a, 9b, 10e
Please answer the following questions to participate in our certified Continuing Medical Education program. Only one answer is possible per question. Please select the answer that is most appropriate.

**Question 1**
Which of the following is a normal variant of pubertal development?
- a) Premature adrenarche
- b) True precocious puberty
- c) Pubertas tarda
- d) Precocious pseudopuberty
- e) Kallmann syndrome

**Question 2**
Which of the following clinical findings generally indicates the presence of true precocious puberty?
- a) Development of the first signs of puberty in a girl aged 8 or older
- b) Development of the first signs of puberty in a boy aged 9 or older
- c) Unilateral testicular enlargement in a boy aged 11 or older
- d) Onset and continuation of breast development in a girl under age 8
- e) Decreasing velocity of growth in a boy in whom puberty has begun

**Question 3**
Which of the following biochemical laboratory parameters indicates the presence of true precocious puberty in a six-year-old girl?
- a) An elevated estradiol level and suppressed LH and FSH levels
- b) A stimulated LH-FSH quotient greater than 1 in the GnRH test
- c) Isolated elevation of DHEAS levels
- d) Low estradiol levels
- e) Elevated hCG levels

**Question 4**
What would you do if you found isolated, bilateral mammary gland enlargement in a neonate?
- a) Nothing
- b) Gonadal ultrasonography
- c) Skeletal age determination
- d) GnRH testing
- e) HCG testing

**Question 5**
Which of the following characterizes hypergonadotropic hypogonadism?
- a) Low LH and FSH levels
- b) Elevated testosterone or estradiol levels
- c) Anosmia, as in Kallmann syndrome
- d) An increase in testicular volume before age 5
- e) Elevated LH and FSH levels in adolescence and/or adulthood

**Question 6**
Which of the following data or findings are consistent with the diagnosis of a normal variant of pubertal development?
- a) The appearance of signs of puberty at a time corresponding to the 10th to 90th percentile for normal pubertal development
- b) Testosterone levels corresponding to Tanner stage 2 of puberty
- c) The appearance of signs of puberty in a six-year-old girl
- d) The appearance of signs of puberty at a time within two standard deviations of the mean for normal pubertal development
- e) The appearance of signs of puberty at a time later than two standard deviations from the mean for normal pubertal development

**Question 7**
What is the first sign of pubertal development in boys?
- a) Increasing size of the penis
- b) Deepening voice
- c) Acne vulgaris
- d) Increasing size of the testes
- e) Spermarche

**Question 8**
What is the first sign of pubertal development in girls?
- a) A growth spurt
- b) Deepening voice
- c) Breast development
- d) Vaginal secretion
- e) Acne vulgaris

**Question 9**
What is meant by "premature thelarche"?
- a) The isolated appearance of pubic hair
- b) A normal variant of pubertal development
- c) Premature development of the breasts in association with a "pubertal" growth spurt
- d) A preliminary stage of precocious puberty
- e) Breast development beginning at age 12

**Question 10**
Which of the following can cause precocious pseudopuberty?
- a) A hormone-secreting tumor of the central nervous system
- b) Pituitary hypoplasia
- c) Activation of the hypothalamic-pituitary-gonadal axis
- d) Klinefelter syndrome (47, XXY)
- e) Ullrich-Turner syndrome (45, X)
CONTINUING MEDICAL EDUCATION

Disorders of Pubertal Development

Jürgen Brämswig, Angelika Dübbers