Congenital Adrenal Hypoplasia and Isolated Gonadotropin Deficiency

Gad B. Kletter, Jerome L. Gorski, and Robert P. Kelch

Congenital adrenal hypoplasia with gonadotropin deficiency is a rare X-linked recessive disorder that usually manifests with symptoms of adrenal insufficiency early in infancy. Adequate replacement therapy with glucocorticoids, mineralocorticoids, and salt has resulted in an increased survival. Slow growth and failure to undergo sexual maturation during the adolescent years usually ensues, secondary to hypogonadotropic hypogonadism. The X-linked congenital adrenal hypoplasia locus has been mapped to region Xp21.3–p21.2. Interstitial deletions of the X chromosome overlapping this region have been observed to cause complex clinical problems, with adrenal hypoplasia as a prominent component. Within a family segregating the disease, there is a 50% risk of having an affected male and a 50% risk of having a carrier female; considerations of genetic heterogeneity, possible chromosomal abnormalities, and prenatal diagnostic studies warrant medical genetic evaluations. The following case presentations illustrate the clinical spectrum of this condition. (Trends Endocrinol Metab 1991; 2:123–128)

• Case Presentation 1

A 6-week-old infant was admitted to a local hospital because of weight loss and severe failure to thrive. Gestation and delivery were normal; birth weight was 4600 g. After receiving intravenous (i.v.) hydration he recovered and did well until ~3 years of age, when he began to require frequent hospitalizations for recurrent vomiting, hyperkalemic, and hypotonic dehydration, lethargy, and weakness. The diagnosis of primary adrenal insufficiency was made and replacement therapy with hydrocortisone succinate and 9-a-fluorohydrocortisone was begun. Occasionally because of seizures and unresponsiveness associated with hypoglycemia and hyponatremia. Two brothers had died suddenly at the ages of 13 months and 13 days, respectively. A maternal grandmother who died in early childhood, and a sister of that grandmother had two sons who died in early childhood from "hypotonic dehydration."

At age 16, when he initially came to our institution, he was a thin, short boy with a eunuchoid body habitus. Physical examination revealed a scant amount of pubic hair and minimal scrotal thinning. Pituitary gonadotropin responses to synthetic GnRH (2.5 µg/kg) were low [mean basal luteinizing hormone (LH) 3.9 IU/L, maximal increment 2.9 IU/L; and mean basal follicle-stimulating hormone (FSH) 1.9 IU/L, maximal increment 1.6 IU/L]. Serum testosterone was 2.1 nmol/L, and bone age was 13 years.

He was admitted to the Clinical Research Center for detailed evaluation of gonadotropin secretion and responses to pulsatile administration of synthetic GnRH (0.025 µg/kg per dose every 2 h for 5 days). His response to synthetic GnRH was minimal (Figure 1): serum LH a subunit did not increase (data not shown). He was started on therapy with depot testosterone enanthate, and full masculinization ensued.

• Case Presentation 2

A 7-year-old boy was referred because of vomiting and increasing fatigue over the past 3 months. He had not experienced weight loss or diarrhea. Family history was noncontributory. On admission, his weight was 30.2 kg (90th percentile for age) and his height was 130 cm (80th percentile). Physical examination revealed increased skin pigmentation, especially of the elbows, knees, knuckles, and gingivae. Serum Na+ was 121 meq/L and serum K+ was 5.2 meq/L. Serum cortisol was 193 nmol/L (normal, 28–662 nmol/L) and did not increase after stimulation with adrenocorticotropic hormone (ACTH). Urinary 17 hydroxysteroid excretion was low. Plasma ACTH was 6300 ng/L (normal, < 200 ng/L). Primary adrenal insufficiency was diagnosed, and therapy with mineralo- and glucocorticoids was initiated. Evaluation of a possible autoimmune endocrinopathy included determination of all of the following antibody titers, all of which were undetectable: antiadrenal, antiislet cell, antimitochondrial, anti-smooth-muscle, antiparietal cell, antithyroglobulin, and antimicrosomal. Seven years later, failure to develop secondary sex characteristics prompted further evaluation. At 15½ years of age, his external genitalia showed no pubertal changes: there was a scant amount of pubic hair, and the phallus and testes were prepubertal in size. Bone age was 13 years; serum gonadotropin concentrations were somewhat low (FSH 4.2 and LH 1.6 IU/L); serum testosterone was undetectable. The diagnosis of hypogonadotropic hypogonadism was suspected, and he was admitted for evaluation of gonadotropin

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secretion before and after 5 days of exogenous GnRH given intravenously every 2 h (0.025 μg/kg per dose). The results are depicted in Figure 2. Pulsatile administration of synthetic GnRH resulted in an increase in FSH secretion, but no change in LH secretion. He then received human chorionic gonadotropin (hCG), 2000 U intramuscularly (i.m.) every 3 days for 8 weeks, which resulted in an increase of serum testosterone from undetectable to 2.1 nmol/L, a poor response. Subsequently he received therapy with gradually increasing doses of the long-acting testosterone ester, testosterone enanthate (up to 200 mg i.m. every 4 weeks), with induction of masculinization. At 20 years of age, his physical examination revealed Tanner stage-3 penile development and pubic hair. Testes were small, measuring 3 cm on the longer axis, and the scrotum was poorly developed.

- **Case Presentation 3**

An infant boy presented at 2 weeks of age with frequent vomiting, weight loss, dehydration, and apathy. His case history was presented before (Virdis et al. 1983) and only a brief summary is outlined here. His brother died suddenly at the age of 2 months; an autopsy revealed interstitial pneumonia, but the adrenal glands were not evaluated. Birth weight of this infant (case 3) was 4300 g. Laboratory evaluation revealed hyperkalemia and hyponatremia. Primary adrenal insufficiency was diagnosed, based on low concentrations of urinary 17-hydroxysteroids and 17-ketosteroids, and plasma aldosterone, cortisol, and 17-hydroxyprogesterone, as well as unresponsiveness to ACTH stimulation. Therapy was started with hydrocortisone succinate and a mineralocorticoid (fludrocortisone). He grew along the third percentile until 12 years of age, when his relative growth slowed further. Skeletal maturation had been 4'/2 years at 9'/2 years of age, and the delay in skeletal maturation increased progressively; at 12'/2 years of age, his bone age was 9'/2 years. Serum androgen concentrations [dehydroepiandrosterone (DHEA), DHEA sulfate, androstenedione, and testosterone] were measured periodically beginning at 12 years of age and were low. At 14'/2 years, slight development of pubic hair growth was noted, but no further pubertal changes developed. Between 14'/2 and 16 years of age, he had a slight increase in serum testosterone, but serum testosterone fell to within the prepubertal range by age 17 years. Results of a synthetic GnRH stimulation test (2.5 pg/kg i.v.) suggested gonadotropin deficiency, as LH and FSH concentrations did not increase. He was admitted to the Clinical Research Center for a 5-day GnRH stimulation protocol during which intravenous bolus injections of GnRH (0.025 μg/kg per dose) were given every 2 h. His plasma gonadotropin responses are depicted in Figure 3. Plasma immunoreactive and bioactive FSH concentrations increased, but LH concentrations as well as LH α subunit (data not shown) remained low. At 18 years of age, treatment with a depot preparation of
testosterone esters was begun, and secondary sex characteristics developed over the ensuing months.

## Discussion

Idiopathic congenital adrenal hypoplasia is a rare disorder with an overall estimated frequency of 1:12,500 live births (Kelch et al. 1984). Two distinct forms have been identified, based on the histologic appearance of the adrenal glands. The miniature adult form, also called anencephalic or secondary form, is characterized by small adrenal glands with normal architecture. There is a well-defined permanent zone and a greatly reduced fetal zone. This form is believed to be secondary to ACTH deficiency, such as seen in anencephalic infants, infants of mothers who received therapy with glucocorticoids during pregnancy, or infants with congenital anomalies of the hypothalamus or pituitary gland. This secondary form is usually sporadic, although few cases of autosomal recessive inheritance have been reported (Kelch et al. 1984; Prader et al. 1975). The observed association of achalasia and adrenal hypoplasia in a subset of familial cases suggests that autosomal recessive congenital adrenal hypoplasia may be heterogeneous (Allgrove et al. 1978).

The primary, or cytomegalic, form is characterized by the absence of the permanent zone of the adrenal cortex; the cytomegalic cells are larger than normal fetal adrenal cells and have nuclear inclusions due to cytoplasmic invagination. The primary form of congenital adrenal hypoplasia is an X-linked disorder usually manifesting as adrenal insufficiency early in infancy, with low serum concentrations of glucocorticoids, mineralocorticoids, and androgens. Failure to respond to ACTH stimulation is characteristic. As illustrated by all three case presentations, hypogonadotropic hypogonadism is noted at the expected time of pubertal maturation (Prader et al. 1975; Virdis et al. 1983; Hay et al. 1981; Kelly et al. 1977; Golden et al. 1977; Lippe and Golden 1978; Petersen et al. 1982).

Clinical signs and symptoms of infants with adrenal hypoplasia include poor feeding, failure to gain weight, hyperpigmentation, vomiting, diarrhea, vascular collapse, and sudden death. Dehydration, hyponatremia, hyperkalemia, acidosis, and hypoglycemia are common biochemical findings characteristic of combined glucocorticoid and mineralocorticoid deficiency. These signs and symptoms are indistinguishable from those seen in male infants with the salt-wasting form of congenital adrenal hyperplasia due to 21-hydroxylase deficiency. A clue for differentiation between these two possible diagnoses in a male infant is the size of the genitalia. In infants with 21-hydroxylase deficiency, the testicles are normal in size and the phallus may be larger than normal owing to elevated plasma androgen concentrations. In contrast, the testes and phallicus are usually small in infants who have adrenal hypoplasia (Kelch et al. 1984). With prompt diagnosis and replacement therapy, survival has increased greatly (Kelch et al. 1984; Virdis et al. 1983). Although the majority of patients present in infancy, some patients similar to case 2 do not present with classic signs and symptoms of adrenal insufficiency until the later childhood years (Hay et al. 1981, case 2; Kelly et al. 1977; Golden et al. 1977).

In a male infant, a presumptive diagnosis of congenital adrenal hypoplasia may be reached by determination of serum concentrations of cortisol, aldosterone, 17-hydroxyprogesterone, 17-hydroxyprogrenolone, and DHEA and its sulfate, all of which are low. Plasma ACTH concentrations are increased. Demonstration of failure to increase serum concentrations of the above-mentioned steroids following administration of ACTH strongly supports the diagnosis (Kelch et al. 1984; Kelly et al. 1977; Petersen et al. 1982). Distinguishing patients with congenital adrenal hypoplasia from those with hereditary ACTH unresponsiveness, although sometimes difficult, is usually made clinically, as the latter patients do not have significant mineralocorticoid deficiency nor are they gonadotropin deficient (Kelch et al. 1972).

Replacement therapy with hydrocortisone succinate, fluocortisone, and salt (NaCl) should be initiated promptly and monitored carefully (Kelch et al. 1984; Hughes 1988). Careful attention to growth velocity and rate of skeletal maturation is essential since, unlike the 21-hydroxylase-deficiency form of congenital adrenal hyperplasia, there is no useful biochemical marker for adequacy of treatment of patients who have congenital adrenal hypoplasia.

As gonadotropin deficiency is commonly associated with the primary form of adrenal hypoplasia, the patients must be followed closely for physical signs of pubertal maturation (Kelch et al. 1984; Petersen et al. 1982; Zachman et al. 1980). When a patient with primary adrenal hypoplasia shows no signs of sexual maturation by 13–14 years of age, hypogonadotropic hypogonadism should be suspected. Low serum concentrations of LH, FSH, and testosterone are usually found, and the responses of serum gonadotropin concentrations to a single GnRH bolus are usually low (Dunkel et al. 1985). A stimulation test with hCG, performed by administration of hCG 2000 U i.m. twice a week for 4–6 weeks, and determination of serum testosterone concentrations before and after the hCG therapy are recommended to evaluate testicular function before therapy with androgens is started (Dunkel et al. 1985). The majority of reported patients had normal testicular responses to hCG (Kelch et al. 1984; Virdis et al. 1983, case 1; Hay et al. 1981; Kelly et al. 1977; Golden et al. 1977; Lippe and Golden 1978; Petersen et al. 1982; Zachman et al. 1980; Gordon et al. 1984; Kikuchi et al. 1985).
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critical period of adrenal development and also to permanent gonadotropin deficiency. 

There is little evidence to support this theory, however, and it seems unlikely that ACTH deficiency is involved in the pathogenesis as ACTH secretion is nor-

mal or increased in patients with pri-

mary adrenal hypoplasia. Also, this theory does not explain the finding of patients who have a delayed onset such as in case 3 and others (Hay et al. 1981; Kelly et al. 1977) who, based on clinical data, were presumed to have had normal function of the adrenal gland until the childhood years. Moreover, normal adrenal function was documented by deter-

mination of normal urinary excretion of adrenocorticosteroids in a patient who had a positive family history and who developed adrenal insufficiency at the age of 10 years (Golden et al. 1977; Lippe and Golden 1978). Similarly, Sills and colleagues (1983) reported a case of a male infant who demonstrated a progres-

dive decline in adrenal function: at 2 months of age, he had a normal cortisol response to ACTH, but his adrenal func-

dion declined and, by 3 years of age, he had no response to ACTH. 

Whether the hypogonadotropic hy-

pogonadism in patients who have con-

genital adrenal hypoplasia is the result of pituitary or hypothalamic dysfunction has not been settled. Initial reports of the association of congenital adrenal hy-

poplasia with isolated gonadotropin deficiency documented poor gonadotropin responses to a single dose of synthetic GnRH in the majority of the patients (Kelch et al. 1984; Prader et al. 1975; Virdis et al. 1983; Hay et al. 1981; Kelly et al. 1977; Golden et al. 1977), but not all (Petersen et al. 1982). This did not differentiate primary pituitary from hypothalamic dysfunction, as most pa-


tients who have isolated gonadotropin deficiency, due to GnRH deficiency, as well as patients who have primary pitui-

tary dysfunction, respond poorly or not at all to a single dose of synthetic GnRH (Dunkel et al. 1985). 

To try to differentiate between hypo-

thalamic and pituitary gland dysfunc-

function, more prolonged administration of GnRH is required (Kelch et al. 1984; Barkan et al. 1985). Several recent re-

ports have suggested pituitary gland dysfunction as the primary defect, based upon lack of gonadotropin responses to prolonged, pulsatile stimulation with synthetic GnRH (Gordon et al. 1984; Kikuchi et al. 1987; Bovet et al. 1988). Gordon and associates (1984) described striking differences in responses to GnRH between a patient with Kallman's syn-

drome, whose LH and FSH rose to within the normal range after 1 week of therapy, and a patient with congenital adrenal hypoplasia in whom only mini-

mal increases of LH and FSH were found after 16 weeks of therapy. Serum testosterone concentrations increased in the patient with Kallman's syndrome, but not in the patient with adrenal hypoplasia. Similar observations of failure to produce pubertal LH and FSH secretion in patients with X-linked adrenal hy-

poplasia after prolonged treatment with pulsatile GnRH have been reported by others (Kikuchi et al. 1987; Bovet et al. 1988). The possibility of primary pitui-

tary dysfunction is further supported by the finding of Marsden and Zakhour (1978) who reported cytomegalic changes in the pituitary gland of an infant who died of congenital adrenal hypoplasia. 

In contrast, Partsch and Sippell (1989) reported successful induction of puber-

tal concentrations of LH, FSH, and testosterone 2–3 months after initiation of therapy with pulsatile synthetic GnRH, suggesting hypothalamic dysfunction as the principal defect in their patient with congenital X-linked adrenal hypoplasia. At least initially, patients with severe endogenous GnRH deficiency respond to exogenous GnRH with the preferential secretion of FSH (Barkan et al. 1985). This may be important, as the patients studied by Kikuchi et al. (1987) had increased serum FSH concentrations and normalization of FSH responses to synthetic GnRH although serum LH hyporesponsiveness persisted. Thus, the increases in serum FSH noted in our cases (Figures 1–3), as well as others, support the possibility of a primary hypothalamic abnormality (Kruse et al. 1984). Finally, in case 2, serum bioactive FSH responses to synthetic GnRH in-

creased after pulsatile GnRH therapy similar to findings in patients with idi-

opathic hypogonadotropic hypogonadism (Padmanabhan et al. 1988). 

Differences in gonadotropin responses of patients with congenital adrenal hy-

poplasia to the prolonged administra-

tion of synthetic GnRH might be attrib-

uted to differences in the treatment protocols, but this seems unlikely, since the frequency of pulsatile injections var-

ied only between 90 and 120 min and the dosage range was 0.05–0.2 μg/kg per dose (Bovet et al. 1988; Gordon et al. 1984; Kikuchi et al. 1987; Kruse et al. 1984; Partsch and Sippell 1989). Alternati-

vely, since histologic or precise genetic confirmation of the diagnosis has been unavailable in patients who have re-

ceived pulsatile GnRH, the syndrome of congenital adrenal hypoplasia with iso-

lated gonadotropin deficiency may be genetically heterogeneous. Thus far, no more than one affected individual from an involved family has received pro-

longed stimulation with synthetic GnRH. It will be important to determine whether GnRH responsiveness differs among af-

ected boys within a kindred and whether different families have different genetic abnormalities. 

Appropriate genetic counseling is an important component of the overall man-

agement of families with an affected child. To assist in differentiating be-

 tween the autosomal recessive and the X-linked forms of congenital adrenal hypoplasia, a detailed pedigree of the proband's family should be obtained,
with particular attention to the previous occurrence of neonatal or unexplained deaths, similarly affected individuals within the maternal lineage, and the possibility of consanguinity. Since adrenal insufficiency can precede neurologic symptoms by months to years in individuals with X-linked adrenoleukodystrophy (ALD), a distinctly different X-linked disease mapped to Xq28 (Mandel et al. 1989), pertinent laboratory assays, such as serum long-chain fatty-acid quantification (elevated in ALD), may be indicated to assist in making an accurate diagnosis. In families with a documented affected male infant with X-linked adrenal hypoplasia, the recurrence risk for a subsequent male infant may be as high as 50%, with a similar risk for a female sibling of being a gene carrier.

Hammond et al. (1985) suggested that the locus for X-linked congenital adrenal hypoplasia and that for glyceral kinase were located on the short arm of the X chromosome within the region Xp11.2-p21, based on the findings of an interstitial Xp deletion with breakpoints at Xp11.2 and Xp21 in a phenotypically normal mother of a male infant who died at 36 h of cytogemal adrenal hypoplasia with glyceraluria (indicating glyceral kinase deficiency) and ornithine transcarbamoyltransferase deficiency. The molecular genetic analyses of multiple unrelated affected males with observed cytologic and submicroscopic interstitial deletions have localized the X-linked congenital adrenal hypoplasia locus to region Xp21.3-p21.2 (Bartley et al. 1986; Francke et al. 1987; Mandel et al. 1989) between two anonymous DNA sequences, DXS67 and DXS268, with a gene order of Xpter-DXS67-AH-GK-DXS268-DMI.

OTC-Xcen (Xp telomere, Xpter; adrenal hypoplasia, AH; glyceral kinase, GK; Duchenne muscular dystrophy, DMD; ornithine transcarbamoyltransferase, OTC; and X centromere, Xcen) (Goonewardena et al. 1989). These studies indicated that it may be feasible to use polymorphic DNA markers closely linked to the adrenal hypoplasia locus to provide prenatal diagnostic studies to families with a previously affected member.

Not infrequently, patients with X-linked adrenal hypoplasia have been found to be affected with additional X-linked diseases such as Duchenne muscular dystrophy, glyceral kinase deficiency, ornithine transcarbamoyltransferase deficiency, and mental retardation as a consequence of having an interstitial deletion within Xp21 disrupting a number of neighboring, or contiguous, genetic loci (Hammond et al. 1985; Bartley et al. 1986; Francke et al. 1987). These patients present with a variety of clinical problems depending on which contiguous genes have been disrupted; the various clinical syndromes resulting from different overlapping deletions have been referred to as contiguous deletion syndromes (Schmickel 1986; Emanuel 1988).

Because X-linked adrenal hypoplasia is not uncommonly a consequence of an X-chromosome deletion, affected patients should be evaluated for clinical signs of other diseases mapped to Xp21 and, when appropriate, diagnostic studies, including a karyotype and urine organic acids, should be obtained.

In a male infant from an affected kindred, the finding of normal results of adrenal function at birth does not rule out the possibility of X-linked adrenal hypoplasia, as onset of symptoms may be delayed in some patients (case 1; Hay et al. 1981; Kelly et al. 1977; Golden et al. 1977; Silts et al. 1983), and thus careful follow-up is indicated.

## References


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