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P W Speiser, ..., M I New, P C White

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#### Research Article

Genotyping for 10 mutations in the CYP21 gene was performed in 88 families with congenital adrenal hyperplasia due to 21-hydroxylase deficiency. Southern blot analysis was used to detect CYP21 deletions or large gene conversions, and allele-specific hybridizations were performed with DNA amplified by the polymerase chain reaction to detect smaller mutations. Mutations were detected on 95% of chromosomes examined. The most common mutations were an A----G change in the second intron affecting pre-mRNA splicing (26%), large deletions (21%), lle-172----Asn (16%), and Val-281----Leu (11%). Patients were classified into three mutation groups based on degree of predicted enzymatic compromise. Mutation groups were correlated with clinical diagnosis and specific measures of in vivo 21-hydroxylase activity, such as 17-hydroxyprogesterone, aldosterone, and sodium balance. Mutation group A (no enzymatic activity) consisted principally of salt-wasting (severely affected) patients, group B (2% activity) of simple virilizing patients, and group C (10-20% activity) of nonclassic (mildly affected) patients, but each group contained patients with phenotypes either more or less severe than predicted. These data suggest that most but not all of the phenotypic variability in 21-hydroxylase deficiency results from allelic variation in CYP21. Accurate prenatal diagnosis should be possible in most cases using the described strategy.

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### Disease Expression and Molecular Genotype in Congenital Adrenal Hyperplasia due to 21-Hydroxylase Deficiency

Phyllis W. Speiser, \* Jakob Dupont, \* Deguang Zhu, \* Jorge Serrat, \* Miriam Buegeleisen, \* Maria-Teresa Tusie-Luna, \* Martin Lesser, \* Maria I. New, \* and Perrin C. White \*

\*Department of Pediatrics, Division of Pediatric Endocrinology, Cornell University Medical College, New York 10021; and <sup>‡</sup>Division of Biostatistics, North Shore University Hospital-Cornell University Medical College, New York 10021

#### **Abstract**

Genotyping for 10 mutations in the CYP21 gene was performed in 88 families with congenital adrenal hyperplasia due to 21-hydroxylase deficiency. Southern blot analysis was used to detect CYP21 deletions or large gene conversions, and allele-specific hybridizations were performed with DNA amplified by the polymerase chain reaction to detect smaller mutations. Mutations were detected on 95% of chromosomes examined. The most common mutations were an  $A \rightarrow G$  change in the second intron affecting pre-mRNA splicing (26%), large deletions (21%), Ile-172  $\rightarrow$  Asn (16%), and Val-281  $\rightarrow$  Leu (11%). Patients were classified into three mutation groups based on degree of predicted enzymatic compromise. Mutation groups were correlated with clinical diagnosis and specific measures of in vivo 21-hydroxylase activity, such as 17-hydroxyprogesterone, aldosterone, and sodium balance. Mutation group A (no enzymatic activity) consisted principally of salt-wasting (severely affected) patients, group B (2% activity) of simple virilizing patients, and group C (10-20% activity) of nonclassic (mildly affected) patients, but each group contained patients with phenotypes either more or less severe than predicted. These data suggest that most but not all of the phenotypic variability in 21-hydroxylase deficiency results from allelic variation in CYP21. Accurate prenatal diagnosis should be possible in most cases using the described strategy. (J. Clin. Invest. 1992. 90:584-595.) Key words: steroid 21-hydroxylase deficiency • CYP21 • allelic variation

#### Introduction

Congenital adrenal hyperplasia due to 21-hydroxylase deficiency is among the most common inborn errors of metabolism. Steroid 21-hydroxylase is a microsomal cytochrome P450 required for synthesis of cortisol and aldosterone but not for synthesis of sex steroids (1). The gene encoding the active adrenal 21-hydroxylase, CYP21, has been characterized (2, 3). Located on the short arm of chromosome 6 in the midst of the class III HLA region, the gene is closely linked to the highly polymorphic genes encoding HLA-B and DR (4, 5). Mutations in the CYP21 gene are almost always either deletions (6) or transfers of deleterious sequenes (7, 8) from the adjacent pseudogene, CYP21P, to the active gene, CYP21, in a process termed gene conversion (9, 10).

Address correspondence to Dr. Phyllis W. Speiser, Room N236, New York Hospital-Cornell University Medical College, 525 East 68th Street, New York, NY 10021.

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There are three major disease phenotypes. In the classic salt-wasting form, both cortisol and aldosterone synthesis are severely impaired; adrenal overproduction of androgen precursors leads to pre- and postnatal virilization, which is most apparent in affected female newborns with ambiguous external genitalia. In the classic simple virilizing form, cortisol synthesis is impaired, but aldosterone synthesis is normal to elevated (11); these patients also undergo pre- and postnatal virilization. In the nonclassic form, a subtle defect in cortisol synthesis can often only be detected during stimulation with corticotropin; aldosterone synthesis is normal and adrenal-derived androgen precursors are moderately elevated (12). In females with the nonclassic form of 21-hydroxylase deficiency, prenatal virilization is not observed.

The molecular genetic basis of this disease has been thoroughly investigated (for reviews see references 9 and 10). Table I provides a summary of the known mutations, their prevalence, and effects upon enzyme activity as determined by in vitro mutagenesis and expression. The strongest associations reported to date have been deletion of CYP21 in patients with the salt-wasting form (13–16) and a point mutation in the seventh exon, which results in a conservative amino acid substitution, Val-281  $\rightarrow$  Leu, in patients with the nonclassic form of 21-hydroxylase deficiency (17). The most frequent non-deletional mutation found in classic patients is a point mutation in the second intron, which activates an aberrant splice acceptor site (18); this mutation has been detected in patients affected with either the salt-wasting or simple virilizing forms of the disease (19, 20).

In this report, we genotyped a large number of affected patients and members of their nuclear families for deletions and nine other discrete CYP21 mutations, allowing assignment of specific mutations to the respective maternal and paternal chromosomes. Patients were categorized into one of three mutation groups depending on degree of predicted enzymatic compromise and clinical and hormonal statuses were retrospectively examined. We sought to determine how closely the predicted behavior of various CYP21 mutations was reflected by in vivo measures of 21-hydroxylase deficiency in each of the three adrenal zones. Conversely, we asked whether the clinician could be expected to predict the genotype based on the clinical diagnoses of salt-wasting, simple virilizing, or nonclassic 21-hydroxylase deficiency. Thorough clinical evaluation of patients genotyped in this study suggests that most but not all of the phenotypic variability in 21-hydroxylase deficiency results from allelic variation in CYP21.

#### **Methods**

Patients. A total of 88 families of diverse ethnic backgrounds were studied. In 90 patients from 79 of these families, CYP21 mutations were identified on each of the two chromosomes. Clinical data are

Table I. Functional Mutations in CYP21 and Activity of Expressed Mutant 21-Hydroxylase Enzyme

Designation	Site	Prog/ 17OHP NL Activity	Function	Allele frequency (reference)*
		%		
Group A: No enzyme activity				
Deletion	Exons 1–8	0/0	No gene product	20% (6); 33% (13); 35% (14); 11% (16); 19% (20)
Del 8 bp	Exon 3	0/0	Frameshift; no activity	3-10% (7, 18-20); 1% (36)
Cluster	Exon 6	0/0		17% (20); 3% (36)
Ile236 → Asn				
Val237 → Glu				
Met239 → Lys				
Phe306 Insert T	Exon 7	0/0	Termination before heme- binding site	1% (36)
Gln318 → term	Exon 8	0/0	Δ mRNA stability; termination before heme-binding site	3–7% (8, 18, 19)
Arg356 → Trp	Exon 8	0/0	?∆ Substrate binding	14% (36)
bp656 A → G	Intron 2	?	Activates new splice acceptor	42% (19); 26% (20); 31% (36)
Group B: Severely impaired enzyme activity				
Ile172 → Asn	Exon 4	2/2	?Δ ER membrane orientation	5-10% (7) (19) (20)
Group C: Moderately impaired enzyme activity				
Pro30 → Leu	Exon 1	30/60	$?\Delta$ ER membrane orientation	17% (32) (NC patients only)
Val281 → Leu	Exon 7	20/50	?Δ Tertiary structure	84% (20) (B14+ NC patients)

<sup>\*</sup> Allele frequencies are from studies in classic patients unless otherwise stated.

tabulated only for patients from the latter group. Five patients had been previously reported as completely genotyped. Patients were born between 1949 and 1991; 74% were regularly treated at The New York Hospital-Cornell University Medical College (NYH-CUMC), and 59% were followed at our institution from infancy or early childhood. Medical records and laboratory tests were thoroughly reviewed for those patients not diagnosed in our institution. To assign the CYP21 mutations to maternal and paternal haplotypes, patients were preferentially selected from families where both mother and father were available for study. Thus, both parents were genotyped in 67 families, in 10 additional families one parent was genotyped, and in the remaining 2 families the patient's child was genotyped, allowing assignment of the patient's two mutations to each of his two chromosomes. HLA-B14 homozygotes were excluded to avoid including an excessive number of Val-281 → Leu homozygotes.

Hormone assays. Adrenocortical steroid hormones in serum were measured by radioimmunoassay (21-23). Patients were diagnosed according to published reference data derived from this laboratory (24) by measuring the basal and/or corticotropin-stimulated serum 17-hydroxyprogesterone in the absence of glucocorticoid treatment. Urinary aldosterone was measured as the acid-hydrolyzable conjugate, aldosterone-18-glucuronide (25). This metabolite of aldosterone represents a constant percentage of secreted aldosterone (26, 27). Plasma renin activity (PRA)<sup>1</sup> was determined by the method of Sealey and Laragh with modifications (28).

Phenotype evaluation. Two methods were used to assess disease severity. First, the clinical diagnosis of salt-wasting, simple virilizing, or nonclassic disease was assigned by the primary pediatric endocrinologist (M. I. New for patients diagnosed and treated at NYH-CUMC) based on history, physical examination, presence or absence of electro-

lyte abnormalities, and hormonal data. Genital ambiguity in females was graded according to the scale of Prader (29). Patients who had overt evidence of sodium depletion, i.e., hyponatremia, hyperkalemia, dehydration, and/or shock (adrenal crisis), were classified as salt wasting, and among them basal serum 17-hydroxyprogesterone, where available, was usually > 500 nmol/liter. Those with no evidence of sodium depletion, but markedly elevated basal and corticotropin-stimulated 17-hydroxyprogesterone levels (usually > 300 nmol/liter), were said to have simple virilizing disease. Nonclassic patients had lower levels of 17-hydroxyprogesterone and no evidence of prenatal virilization (12).

Presence or absence of adrenal salt-wasting crisis and basal sodium and potassium levels were recorded at the time of diagnosis. When the medical records indicated that there had been no electrolyte abnormalities the serum sodium and potassium were recorded as normal in the phenotype score. Efficiency of the adrenal zona glomerulosa in synthesizing aldosterone, a function dependent on 21-hydroxylase, was determined in selected patients by measuring the ratio of PRA/urinary aldosterone-18-glucuronide standardized for body surface area. Higher ratios of PRA to aldosterone were assumed to be, at least in part, reflective of increasingly severe 21-hydroxylase deficiency. These studies were always performed with adequate glucocorticoid replacement (dexamethasone, 2 mg/d for  $\geq$  2 d, or hydrocortisone, 15-30 mg/m<sup>2</sup> per d chronically), no mineralocorticoid supplements, and under conditions of dietary sodium deprivation (10 mmol/d for 2-5 d). The majority of patients studied were over the age of 5 yr. Sodium balance, expressed as total daily sodium excretion divided by intake (thus, the higher the number, the more negative the sodium balance), was recorded for the same period. Several patients were studied at two different ages.

Because several of our patients did not precisely fit all criteria for one of these three categorical clinical diagnoses, we used a second mode of phenotype evaluation: an objective rating scale was devised to in-

<sup>1.</sup> Abbreviation used in this paper: PRA, plasma renin activity.

Table II. Phenotype Score for Patients with 21-Hydroxylase Deficiency

	Phenotype score								
	0	1	2	3					
Serum Na (mM)	Na 135-145	Na 125-134	Na < 125						
Serum K (mM)	K 3.5-5	K 5-6	K > 6						
PRA/ALDO	< 1	1–10	> 10, < 50	≥ 50					
Na BALANCE	≤ 1	1–4	≥ 4						
17OHP 0' (nM)	< 3	3–30	> 30-300	> 300					
17OHP 60'	< 6	6-60.5	60.5-605	> 605					
Ptriol (µmol/d)	< 3	3–15	15-30	> 30					
Genitalia (Prader)	nl	1–2	3	4–5					

Score = total point score/number of data points available  $\times$  100.

For each parameter listed, a subscore of 0 indicates a normal value, and disease severity increases as subscores and total score increase. A maximal subscore of 2 (serum Na, serum K, Na balance) or 3 (PRA/ALDO, 170HP, Ptriol, and Prader score) indicates a severely abnormal value. As an example of phenotype score calculation, patient 1 in mutation group A1 (Table V) received a subscore of 2 each for serum Na and K of 117 and 10 mmol/liter, respectively; PRA/ALDO of 3.7 yielded a subscore of 1; Na balance of 2.4 warranted a subscore of 1; the Prader score was 4, for which she received a phenotype subscore of 3. Thus, patient 1 had an adjusted total phenotype score of 9 based on five available data points:  $(9 \div 5) \times 100 = 180$ .

clude the several measures of the in vivo expression of 21-hydroxylase deficiency (Table II) described above. The total score for each individual was divided by the number of data points available and multiplied by 100; only patients with four or more data points were included in the statistics. For patients diagnosed in the years before the advent of radio-immunoassay for serum 17-hydroxyprogesterone, 24-h urinary pregnanetriol measurements were recorded. Both the clinical diagnosis and the phenotype score were compared with the patients' genotypes.

Genotype evaluation. Peripheral blood leukocyte DNA samples were obtained as described (30). An alternative protocol that avoids phenol extraction (31) was used in a minority of samples.

Southern blot analysis. This procedure was performed as described (6), using the restriction enzymes Taq I and Bgl II to determine CYP21 gene copy number. Deletions of CYP21 were identified by the absence (or by decreased intensity in heterozygotes) of the Taq I 3.7-kb and Bgl II 12-kb hybridizing bands. Large gene conversions were detected by the absence or decreased intensity of the 3.7-kb Taq I fragment without any apparent abnormality in the Bgl II digest. To identify smaller mutations not detectable by Southern blotting, the following procedures were employed:

(i) Polymerase chain reaction. Genomic DNA was amplified in two segments by the polymerase chain reaction using primers that selectively amplify CYP21, not the highly homologous CYP21P gene (19) (Fig. 1). Fragment 1 represents a 677-bp segment extending from exon 1 (primer sequence 5'-TGGAACTGGTGGAAGCTCCGG-3') through the 8-bp deletion in exon 3 (primer sequence 5'-AGCAGGGAGTAGTCTCCCAAG-3'; underlined nucleotides represent the 8-bp deletion typically found in CYP21P). Fragment 2 is an ~ 1.5-kb fragment extending from the 8-bp deletion in exon 3 (primer sequence 5'-TTGTCCTTGGGAGACTACTCC-3') through the eighth intron (primer sequence 5'-GCTCGGGCTTTCCTCACTCAT-3'). Amplification reactions were carried out as described (32).

(ii) Allele-specific oligonucleotide probe hybridizations. Serial hybridizations were performed with end-labeled, allele-specific normal and mutant oligonucleotide probes (32) representing point mutations that may be introduced into CYP21 from CYP21P by gene conversion (Table III; Fig. 1). The presumption was made that DNA samples that amplified with both sets of polymerase chain reaction primers did not have a homozygous gene conversion involving the 8-bp deletion in the third exon. A heterozygous gene conversion within the third exon was suspected if a patient was apparently homozygous for a given mutation, but only one parent was heterozygous for the mutation. Homozygous gene conversion at the 8-bp deletion was suspected if DNA from the

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patient failed to amplify, yet Southern blot analysis showed a normal restriction pattern in both parents. These unusual cases were further analyzed by constructing separate CYP21-specific oligonucleotide primers not centered on the 8-bp deletion (20). Mutations were designated A, B, or C according to decreasing order of severity as described below (see Results; Tables IV and V).

Statistical analysis. Differences between basal and stimulated 17-hydroxyprogesterone levels among mutation groups A, B, and C were detected using the Kruskal-Wallace test. Analysis of variance was used to assess differences in phenotype score between mutation groups. Discriminant analysis (PROC DISCRIM; SAS Institute Inc., Cary, NC) was used to classify patients into each of the three mutation groups based on the phenotype score or clinical diagnosis. Classification cutoff points were computed from this analysis. Classification error rates were computed based on prior probabilities that were equal to the observed proportions of each of the three mutation groups. Since several patients' phenotype scores were outliers, the discriminant analysis was also performed using a rank transformation. The 56 individuals with four or more data points for phenotype score were ranked from 1 to 56 (with midranks assigned to tie scores) and these ranks were used as the dependent variable in the analysis. Cut-off points were computed for

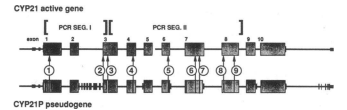


Figure 1. Functional CYP21 mutations and PCR amplification strategy. Sequence differences between the CYP21P pseudogene and CYP21 active gene known to cause clinical 21-hydroxylase deficiency are shown by the circled numbers: 1, exon 1, Pro-30  $\rightarrow$  Leu; 2, intron 2 A  $\rightarrow$  G; 3, exon 3, 8-bp deletion; 4, exon 4, Ile-172  $\rightarrow$  Asn; 5, exon 6, cluster of mutations (Ile-236  $\rightarrow$  Asn; Val-237  $\rightarrow$  Glu; Met-239  $\rightarrow$  Lys); 6, exon 7, Val-281  $\rightarrow$  Leu; 7, exon 7, Phe 306 +T; 8, exon 8, Gln-318  $\rightarrow$  term; 9, exon 8, Arg-356  $\rightarrow$  Trp. Other vertical lines represent differences between CYP21P and CYP21 that are apparently not functionally significant. The CYP21 active gene was specifically amplified in two segments as described in text.

Table III. Oligonucleotide Probes Used in Hybridization Analysis of Patient Samples

Designation	Sequence					
P30 wild type	5'-CTCCACCTCCCGCCTCTTGCC-3'					
P30L mutant	5'-CTCCACCTCCTGCCTCTTGCC-3'					
bp656-A wild type	5'-AGCCCCCAACTCCTCCTGCA-3'					
bp656-G mutant	5'-AGCCCCCAGCTCCTCCTGCA-3'					
I172 wild type	5'-TGCAGCATCATCTGTTACCTC-3'					
I172N mutant	5'-TGCAGCATCAACTGTTACCTC-3'					
I236/V237/M239 wild type	5'-AGGGATCACATCGTGGAGATGCAGCTGAGG-3'					
I236N/V237E/M239K mutant	5'-AGGGATCACA <u>A</u> CG <u>A</u> GGAGA <u>A</u> GCAGCTGAGG-3					
V281 wild type	5'-GGAAGGGCACGTGCACATGGC-3'					
V281L mutant	5'-GGAAGGGCACTTGCACATGGC-3'					
F306 wild type	5'-GGTGAAGCAAAAAACCACGGCC-3'					
INSERT p F306 mutant	5'-GGTGAAGCAAAAAAACCACGGCC-3'					
Q318 wild type	5'-CAGCGACTGCAGGAGGAGCTA-3'					
Q318ter mutant	5'-CAGCGACTGTAGGAGGAGCTA-3'					
R356 wild type	5'-CTGCGCCTGCGGCCCGTTGTG-3'					
R356W mutant	5'-CTGCGCCTGTGGCCCGTTGTG-3'					

Bases underlined in boldface represent differences between wild type and mutant probes.

the ranks, and then the ranks were "inverted" to yield the corresponding cut-off point in phenotype score units.

#### **Results**

Phenotypic evaluation. By clinical diagnosis, a total of 53 salt-wasting, 25 simple virilizing, and 12 nonclassical patients were studied. The distribution of salt-wasting versus simple virilizing classic patients is compatible with the  $\sim 2-3:1$  proportion expected based on the results of newborn screening programs and case surveys (33). Nonclassic patients were deliberately underrepresented in this study because of previously published data demonstrating that the Val-281  $\rightarrow$  Leu mutation found in association with the HLA-B14,DR1 haplotype accounts for  $\sim 75-80\%$  of nonclassic mutations (17, 20).

Table IV. Mutation Groups of Patients with 21-hydroxylase Deficiency

Group	Mutations	Possible genotype combinations	n	Clinical diagnosis
$A_1$	DEL	$A_1/A_1$	47	SW
	DEL Ex 3	$A_1/A_2$	5	SV
	Cluster Ex 6	$A_2/A_2$		
	Phe-306 $\rightarrow$ +T			
	Gln-318 → term			
	$Arg-356 \rightarrow Trp$			
$A_2$	Int 2			
В	Ile-172 $\rightarrow$ Asn	A/B	4	SW
		B/B	16	SV
			2	NC
C	Val-281 → Leu	A/C	2	SW
	Pro-30 → Leu	B/C	4	SV
		C/C	10	NC

NC, nonclassic; SV, simple virilizing; SW, salt wasting.

All females with the salt-wasting form of the disease had both clitoromegaly and urogenital sinus, Prader stage III-V, requiring corrective surgery. Females affected with simple virilizing disease had clitoromegaly, but did not necessarily have a urogenital sinus, and were classed as Prader stage II-IV. Mild clitoromegaly, or Prader stage I, and slight webbing of the labia minora were the only signs of genital virilization in females affected with the nonclassic form of 21-hydroxylase deficiency.

All patients had elevated serum 17-hydroxyprogesterone measurements (24) and/or urinary pregnanetriol >  $8 \mu \text{mol/d}$ ; however, levels measured during treatment were not recorded here or used in the statistical analysis. Classic 21-hydroxylasedeficient patients had basal serum 17-hydroxyprogesterone levels ranging from 151 to 2604 nmol/liter in the absence of glucocorticoid replacement therapy. Basal 17-hydroxyprogesterone levels were significantly higher in salt-wasting compared with simple virilizing patients (mean $\pm$ SD: 1,239 $\pm$ 719 vs.  $526\pm412$  nmol/liter, P=0.002). There was one patient classified as salt-wasting with basal serum 17-hydroxyprogesterone level < 200 nmol/liter in the neonatal period. All other defects of steroid biosynthesis had been excluded. This patient had severe electrolyte derangements and subsequently had a 17-hydroxyprogesterone level above 400 nmol/liter while on treatment during a period of stress.

Patients affected with the nonclassic disorder had 60-min corticotropin-stimulated serum 17-hydroxyprogesterone levels that were statistically different from those of classic simple virilizing patients (mean $\pm$ SD: 324 $\pm$ 167 vs. 1,217 $\pm$ 433 nmol/liter, P = 0.003).

The ratio of plasma renin activity (ng/liter per s) to aldosterone 18-glucuronide (nmol/m² per d) under conditions described above ranged from three to infinitely high (ratio arbitrarily designated 100 in Table V) among young salt-wasting patients, with the normal ratio being maximally 0.1 in sodium-deprived healthy adults (34). One sibling pair studied at ages 30 and 36 yr showed relatively low PRA/aldosterone ratios. All but one patient categorized as salt wasting showed evidence of hyponatremia, hyperkalemia, and/or hypovolemic shock in

Table V. Phenotype and Genotype in 21-Hydroxylase-deficient Patients by Group

						Mutatio	n group A <sub>1</sub>	and $A_2$ ( $n = 52$	<u>:)</u>						
	Clinical		Mu	tation	Zona fasciculata function			Zona glomerulosa function							
ID	Dx	Prader stage	Maternal	Paternal	17OHP 0'	17OHP 60'	PΤ	PRA	Aldo	PRA/Aldo	Na balance	Age	Serum Na	Serum K	PF SC
					nmol	/liter	μmol/d	ng/liter per s	nmol/m²/d		O:I	yr	nmo	l/liter	
Subgrou	up A <sub>1</sub>														
1	SW	4	DEL	DEL				31	8.3	3.7	2.4	19	117	10	18
2	SW	M	DEL	conver									127	8.8	_
3	SW	4	CL6	Q318ter									116	8.7	_
			V281L F306+T Q318ter R356W	R356W											
4	SW	4	DEL	×3	2057								133	8.6	22
5	SW	4	I172N	CL6	2037								116	11	
J	3**	•	CL6 F306+T Q318ter	CLO									110	11	
6	SW	4	×3	$\times 3$	988	1168							131	7.1	24
7	SW	M	DEL	Q318ter			8	48.3	8.3	11	8	17	113	9	18
8	SW	4	conver	DEL			-						137	6.6	_
9-a	SW	4	conver	conver	1840			18.6	4.2	4.4	5.5	18	130	5.8	16
10-b	SW	4	conver	conver			27	34.4	1.7	20.2	4	19	129	7.2	21
11	SW	M	Q318ter	CL6 R356W	1819	1837							135	6.3	20
12-a	SW	M	×3	DEL			47	32.8	10.1	3.3	1.3	15	120	7	17
	2		,				• • •	29.4	9.7	3.0	7.5	20			
13-b	SW	M	×3	DEL				27.6	10.3	2.7	8	20	120	9.8	16
13-0	5**	141	7.5	DEL				22	2.9	7.6	8	24	120	7.0	10
14	SW	4	DEL	F306+T Q318ter R356W				22	2.7	7.0	Ü	24			
15	SW	4	DEL	×3	2390	3026							131	7.2	24
16	SW	3	×3	×3	847										_
17	SW	M	×3 I172N CL6 V281L	DEL	1589									8.8	_
18	SW	4	DEL	$\times 3$				13.9	2.2	6.2		1	117	7.3	20
19	SV	2	1172N	$\times 3$	1292	1522	12.8	5	23	0.2	1	1.5		4.1	ç
			CL6					18	30	0.6	1.5	7			
								7	9.1	0.8	4.5	13			
20	SW	M	R356W	R356W				187	2	94	3.5	4			_
21	SW	4	×3	×3	187	262		21	1.8	11.9		0.1	125	8.6	20
22	SW	4	conver	conver	560	805		4.4	ud	> 100	2	0.3		6.8	22
23	SW	M	DEL	DEL	200	555			uu	- 100	~	0.5	133	5.7	_
24	SW	M	DEL	×3	454								133	5.7	
25	SW	4	DEL	//3	454										-
Sub-G	roup A2														
26	SV	M	R356W	i2	1579	1846		20.1	30.3	0.7	1	3	135	6	11
27	SW	na	i2	DEL											_
28-a	SW	4	DEL	i2				6.7 20.3	0.4 3.8	18.8 5.3	1.7 4	13 19	118	5.9	1
29-b	SW	M	DEL	i2				42	1	42	7	28	134	7.2	17
20	SW	4	i2	DEL	817								126	8.4	22
30	-												120	0.7	

Table V. (Continued)

				_		iviutati	on group A	$A_1$ and $A_2$ $(n=5)$							
	Clinical		M	futation	Zona fasciculata function				Zona glomerulosa function						
ID	Dx	Prader stage	Maternal	Paternal	17OHP 0'	17OHP 60'	PΤ	PRA	Aldo	PRA/Aldo	Na balance	Age	Serum Na	Serum K	PI SC
					nmo	l/liter	μmol/d	ng/liter per s	nmol/m²/d		O:I	yr	nmo	l/liter	
32	SW	4	i2	i2				1.2	ud	> 100	0.1	15	129	6.2	18
33	SW	M	i2	i2	1490								115	9.3	
34	SV	M	i2	×3	218	608		5.6	125	0.05	0.8	2	137	4.7	8
35	SW	4	i2	DEL			6						134	9.8	17:
36	SW	na	DEL	i2	523									8	_
37-a	SW	M	DEL	i2				60.6	3.3	18.4	2	2	118	8.2	17:
38-b	SW	M	DEL	i2				6	ud	> 100	1.3	7	132	5.8	150
39	SW	M	i2	DEL				91	13.4	6.8	1	0.8	114	8	12:
40	SW	M	i2	i2	1029								116	8	
41	SW	M	i2	×3									116	12	_
42	SW	M	DEL	i2	2604		8.5	15	0.7	21	1	4	130	9.6	160
43	SW	M	i2*	conver	680	797			***		_		120	12	250
14	SW	4	i2	×3	300								117	10	_
15	SW	M	i2	DEL									111	7.6	175
16	SV	4	Q318ter	i2	387	957		22.5	28.3	0.8	3	7	137	4.1	143
17	SW	4	i2	i2	1936	,,,,		27	5.4	5	3	1	129	6	167
18-a	SW	4	i2	DEL	1730			6.1	74.9	0.1	7.7	36	127	5.5	140
19-b	SW	4	i2	DEL				8	5.5	1.5	1	30	124	6.6	160
50	SW	5	i2	i2				0	5.5	1.5	1	30	130	9	100
	SV		i2	i2									140	4	
51 52		3												8	
	SW	M	DEL	i2	1242	1202							113	8	
Mean					1242	1282									177
D					724	803									42
/ledian					1160	1062									175
Q3-Q1					1210	1040		-							40
						Mut	ation group	$\mathbf{B} (n=22)$						,,	
1	sv	M	CL6	I172N V281L									140	3.8	_
2	SV	M	i2	I172N				19	14	1.4	1.8	4	137	4.5	50
3	SV	4	conver	I172N	303								nl	nl	150
4	SW	3	R356W	I172N	440								135	6.8	175
5	SV	3	1172N	i2									nl	nl	
5	SW	4	I172N	DEL			40	37	12.3	3	0.6	0.3	132	7.9	167
7	SV	M	I172N	i2									nl	nl	
8-a	NC	1	I172N	Q318ter	64								nl	nl	75
)-b	SV	2	I172N	Q318ter	389	728							143	6.5	180
)-a	SV	M	DEL	1172N	269	1221							141		125
-b	SV	M	DEL	I172N	555	1534		8	61	0.1	0.1	1.4	141		100
2	SV	3	I172N CL6	I172N	605								139		150
}	SV	M	Q318ter	I172N				16	22.2	0.7	3.3	31	nl	nl	25
	SV	3	1172N	R356W									nl	nl	_
	SV	M	i2	I172N			74						nl	nl	_
	SV		i2*	I172N				9	55.3	0.2	2	2	nl	nl	60
	SV		I172N	conver									nl	nl	_
	SV		i2	I172N	151								nl		125
-a			i2	I172N	38								nl	nl	_
	NC							0.5		100					217
-b	NC SW	4	DEL.	I172N				0.7	ua	> 100	_	0.1	132	6.2	21/
-b	SW	4	DEL	I172N				0.7 94	ud 9	> 100 10.4	5		132	6.2	217
3-a 9-b			DEL I172N	I172N DEL	481	727		0.7 94 12	ua 9 ud	> 100 10.4 > 100	5 1	0.1 12 2	132 nl		150

Table V. (Continued)

						Mu	tation gro	up B (n = 22)							
Clinical		Mutation		Zona fasciculata function				Zona glomerulosa function							
ID	Dx	Prader stage	Maternal	Paternal	17OHP 0'	17OHP 60'	PΤ	PRA	Aldo	PRA/Aldo	Na balance	Age	Serum Na	Serum K	PH SCR
					nmol	/liter	μmol/d	ng/liter per s	nmol/m²/d		O:I	yr	nmo	l/liter	
Mean					329	1053									125
SD					199	397									56
Median					346	975									138
Q3-Q1					330	651									92
						Mı	utation gro	$\sup C (n = 16)$							
1-a	NC	0	DEL	V281L	35	363							nl	nl	80
2-b	NC	M	DEL	V281L	23	270							nl	nl	75
3	NC	M	i2	V281L	59	92		4.4	8	0.6	0.7	7	140	4	67
4-a	SV	M	P30L	R356W	323	1467							nl	nl	150
5-b	SV	4	P30L	R356W	0.20			3.3	178	0.2	1.5	16	140	4	80
6	NC	1	R356W	V281	27	283							nl	nl	80
7	NC	0	V281L	i2	12	187							nl	nl	60
8-a	NC	0	P30L	CL6									nl	nl	
9-b	NC	M	P30L	CL6									nl	nl	_
10	NC	1	I172N	P30L	61	425							nl	nl	100
11	SV	M	V281L	CL6	254	1564							nl	nl	12:
				F306+T											
				VAL281L											
12	SW	4	DEL	V281L	1289								na	na	_
13	SW	M	V281L	i2									138	5	_
14	NC	0	V281L	V281L	57	322							nl	nl	8
15	SV	3-4	R356W	P30L									nl	nl	15
16	NC	1	V281L	i2	61	651		7.2	1.6	4.5	1	0.6	135	4.8	10
Mean					229	563									9
SD					372	524									3
Median					60	343									8
Q3-Q1					258	381									3

In patient ID column, a and b indicate siblings. Under Dx, SW refers to salt-wasting patient, SV to simple virilizing, and NC to nonclassic patients. Prader stage for genital ambiguity is given where known for female patients; males are designated M. Description of the various mutations may be found in Table I: group A (including subgroups  $A_1$  and  $A_2$ ) are mutations and genotype combinations conferring no enzyme activity; genotype combinations in group B confer severely impaired enzyme activity; and genotype combinations group C confer moderately impaired enzyme activity. Mutation nomenclature in Table V: Deletion, DEL; large gene conversion detectable on Southern blot, conver; deletion of 8 bp in exon 3,  $\times$ 3; cluster of three mutations in exon 6, CL6; Phe306, insert T, F306+T; Gln318  $\rightarrow$  term, Q318ter; Arg356  $\rightarrow$  Trp, R356W; intro 2 bp656 A  $\rightarrow$  G, i2; Ile172  $\rightarrow$  Asn, I172N; Pro30  $\rightarrow$  Leu, P30L; Val281  $\rightarrow$  Leu, V281L. De novo mutations are indicated by asterisks. Serum 17-hydroxyprogesterone is given before and 60 min after corticotropin stimulation; PT indicates basal 24-h urinary pregnanetriol. PRA (plasma renin activity), Aldo (urinary aldosterone-18-glucuronide, ud signifies undetectable), and the ratio of PRA/aldosterone and sodium balance were measured under conditions described in Methods (> 100 indicates a ratio that was infinitely high). Age refers to the patients' age at the time of the PRA/aldo and sodium balance study. Serum electrolytes represent the nadir of Na and peak K at presentation. The method of calculating phenotype score is described in text. Descriptive statistics are given for 17-hydroxyprogesterone and phenotype score.

early life. Neither simple virilizing nor nonclassic patients, who were less frequently studied for aldosterone synthesis, had concomitant hyponatremia and hyperkalemia typical of salt-wasting patients. Two simple virilizing and one nonclassic patient had high PRA/aldosterone ratios in early childhood.

Genotype evaluation. Results of previously published in vitro mutagenesis and expression studies (18, 32, 35, 36) were used to predict the degree of enzyme deficiency for each mutation (Table I). The predicted enzymatic activity for conversion of 17-hydroxyprogesterone to 11-deoxycortisol of the most se-

vere mutation on each of the patient's two chromosomes was calculated (Table IV). For instance, homozygous deletion, or deletion in *trans* with a stop mutation, or the cluster of mutations at exon 6, all of which confer zero enzyme activity in vitro (35), would be predicted to result in 0% overall 21-hydroxylase activity in vivo. This type of patient would be categorized in group A, subgroup 1 ( $A_1$ ). A deletion conferring zero enzyme activity on the product of the maternal gene in combination with an intron 2 A  $\rightarrow$  G mutation, or homozygotes for the intron 2 mutation, were categorized in group A, subgroup 2

 $(A_2)$ . Homozygotes carrying the mutation Ile-172  $\rightarrow$  Asn, which confers  $\sim 2\%$  of normal activity on the gene product (35, 36), as well as compound heterozygotes for any of the group A mutations in *trans* with the Ile-172  $\rightarrow$  Asn mutation, were categorized as group B. Homozygotes for milder mutations, such as Val-281  $\rightarrow$  Leu and Pro-30  $\rightarrow$  Leu, which have  $\sim 50\%$  of wild-type enzymatic activity for conversion of 17-hydroxyprogesterone to 11-deoxycortisol in vitro (35, 36) (20% for progesterone to deoxycorticosterone), would be categorized in group C. Similarly, the combination of either of the latter mutations with Ile-172  $\rightarrow$  Asn (2% activity) or group A mutations (0% activity) would place the patient in group C with an average of  $\sim 25\%$  normal predicted enzymatic function for conversion of 17-hydroxyprogesterone to 11-deoxycortisol ( $\sim 10\%$  for progesterone to deoxycorticosterone).

Overall, mutations were detected on 95%, or 171/182 chromosomes examined (data not shown for patients in whom the mutation on one chromosome was not identified). When analyzed in this manner, ~ 74% of chromosomes were classified as bearing group A mutations. Large deletions comprised 19% of all chromosomes, large gene conversions (mutations in three or more contiguous exons) 5%, and intron 2 mutations 25%. Approximately 13% of mutations were found in group B, and 7% were found in group C. In 5% of chromosomes examined, no mutations were detected using this methodology.

Relationship of mutation group to clinical diagnosis. A total of 25 patients (23 families) were homozygous for mutations in group  $A_1$ , which result in no enzyme activity; all but one patient in this group had salt-wasting disease (Tables IV and V). 27 additional patients in group  $A_2$  (24 families) were homozygous for the intron 2 mutation or were compound heterozygotes for a group  $A_1$  mutation in trans with the intron 2 mutation. All but one of seven patients homozygous for the intron 2 A  $\rightarrow$  G mutation had salt-wasting disease; none of the seven has been studied beyond adolescence. 3 of 20 compound heterozygotes for the intron 2 mutation were simple virilizing patients with normal or mildly elevated PRA/aldosterone ratios (0.04–0.8) and no evidence of electrolyte derangements; the remainder were salt wasting.

22 patients (19 families) were categorized as Group B. Four were diagnosed as salt wasting, 16 were simple virilizing, and 2

MO PT FA

Arg356--Trp

NORMAL

MUTANT

Ile172--Asn

NORMAL

MUTANT

Figure 2. CYP21 allele-specific hybridization in mutation group B patient 14. CYP21 of this patient was amplified selectively in the PCR from the region of the 8-bp deletion in exon 3 through exon 8 as described. The patient has inherited a mutant allele for Arg-356 → Trp from her father and a mutant allele for Ile-172 → Asn from her mother. These hybridizations were performed by serial stripping and rehybridization.

were nonclassic patients. One female with the genotype combination deletion/Ile-172 → Asn had severe, unremitting saltwasting disease. A prototypic result of allele-specific hybridization in a patient who is a compound heterozygote for two different CYP21 mutations is illustrated in Fig. 2.

Group C mutations predicted to confer relatively mild impairment of enzyme activity consisted of 10 nonclassic patients, 4 simple virilizing, and 2 patients with salt-wasting disease. Among the nonclassic patients with intron  $2 A \rightarrow G/Val-281 \rightarrow Leu$ , two unrelated patients, one male and one female, had a sevenfold difference in 17-hydroxyprogesterone levels 60 min after corticotropin stimulation (92 vs. 651 nmol/liter).

Relationship of specific measures of in vivo 21-hydroxylase activity to mutation group. Mean basal serum 17-hydroxyprogesterone levels were significantly different among patients in mutation groups A, B, and C (mean $\pm$ SD: 1,241 $\pm$ 724 vs 330 $\pm$ 199 and 229 $\pm$ 372 nmol/liter, P=0.0001, Fig. 3). A significant, but less striking, difference was also observed among groups when 60-min ACTH-stimulated levels were compared (P=0.03). There were no significant differences between subgroups A<sub>1</sub> and A<sub>2</sub>.

Fig. 4 shows the relationship of total phenotype score and component scores other than 17-hydroxyprogesterone (serum electrolytes and sodium balance, PRA/aldosterone, and Prader stage) to mutation group. A significant difference in total phenotype scores was evident among the three mutation groups (P = 0.0001 for mean $\pm$ SD: 177 $\pm$ 42 vs. 125 $\pm$ 56 vs. 96 $\pm$ 31 in groups A, B, and C, respectively). Parallel differences were seen among mutation groups for each of the component scores, with the least significant difference observed in PRA/aldosterone.

Discriminant analysis on the 56 patients who were eligible for inclusion based on four or more data points in the phenotype score yielded cut-off points as follows: group A if score > 123; group B if score  $\geq$  89 and  $\leq$  123; and group C if score < 89. This classification rule resulted in group A, B, and C patients being correctly classified 90, 7, and 58% of the time, respectively. The overall correct classification rate was 63%. The rank transformed discriminant analysis yielded respective classification cut-off points of > 125, 91 to < 125, and < 91 for mutation groups A, B, and C, respectively. The correct classification rates were 87, 21, and 58%, respectively, with an overall correct classification rate of 64%. Therefore, the effect of any outliers was minimal in this set of data. In contrast to phenotype score, the sensitivity of the clinical diagnoses (salt wasting, simple virilizing, and nonclassic) in correctly classifying patients into mutation groups A, B, or C was 90, 71, and 59%, respectively. The overall correct classification rate for clinical diagnosis was 80%. The relatively poor sensitivity of the phenotype score in correctly classifying individuals in mutation group B could be ameliorated by broadening the range of scores in which patients could be classified in this group, at the expense of decreased accuracy of classification into groups A and C.

Data from sib pair studies. Four of five sib pairs classified as salt wasters in group A differed between sibs with respect to the ratio of PRA/aldosterone, sodium balance, or both. There were two additional families (Table V, group A, ID 43, group B, ID 16) in which the index case had acquired a de novo mutation, intron  $2 A \rightarrow G$ , which was clearly not inherited from either mother or father, whose parental status was confirmed by HLA serotyping. In one of these families, the molecu-

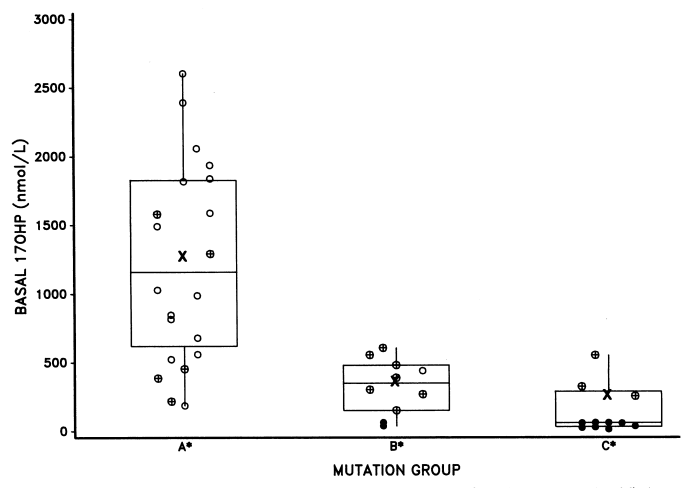


Figure 3. Relationship of serum 17-hydroxyprogesterone to mutation group. Basal serum levels of 17-hydroxyprogesterone (nmol/liter) were measured in the absence of glucocorticoid treatment at presentation in patients from each of the three mutation groups. Group A includes the most severely deleterious mutations, group B are intermediate mutations, and group C are mild mutations. The box encompasses the 25-75th percentiles, the bar indicates the median, the cross indicates the mean, and the vertical lines indicate the ranges. Open circles designate saltwasting patients, crossed circles designate simple virilizing patients, and filled circles designate nonclassic patients. The asterisk indicates statistically significant differences between groups (P = 0.0001).

lar genetic data explained the apparent paradox of HLA identity in phenotypically dissimilar siblings, one of whom was affected with ambiguous genitalia and elevated 17-hydroxyprogesterone levels, whereas the second-born child was normal.

In group B, the group with the widest range in phenotype scores, discordant clinical diagnoses were made in two of three genotypically identical sib pairs.

Finally, in one of three genotypically identical sib pairs in group C, the sister (Table V, group C, ID 1a) presented with normal genitalia and precocious adrenarche and subsequently developed severe hirsutism, whereas the brother (ID 2b) manifested no overt signs of his mild 21-hydroxylase deficiency through adolescence.

#### **Discussion**

Both methods of evaluating the clinical severity of disease, the clinical diagnosis and the phenotype score, were fairly well correlated with the degree of severity of the CYP21 mutation group. Thus, in vivo measures of 21-hydroxylase activity can to some extent be predicted from in vitro expression studies. Each

mutation group contained a number of patients whose phenotype was either more or less severe than predicted. Several explanations might be proposed for these observations. First, it is conceivable that individuals with phenotypes more severe than predicted or discordant from siblings have acquired additional, as yet unidentified, mutations within the CYP21 gene. Undoubtedly, other mutations that may not have arisen from gene conversion will be identified, but based on knowledge accumulated to date these are probably less frequent than those already described.

Second, with respect to discrepancies between predicted and actual phenotypes, it should be recognized that the least impressive intergroup differences between mutation group and phenotypic expression were those found for PRA/aldosterone ratios. Nor did separate examination of aldosterone alone allow improved distinction between mutation groups A and B, representing severely and moderately defective CYP21 genotype combinations, respectively. Based on data included in this and earlier reports (37), it is clear that patients with mutations conferring severe impairment of 21-hydroxylase activity have variable efficiency of aldosterone synthesis, even when the same patient is followed over time. In light of the variability in

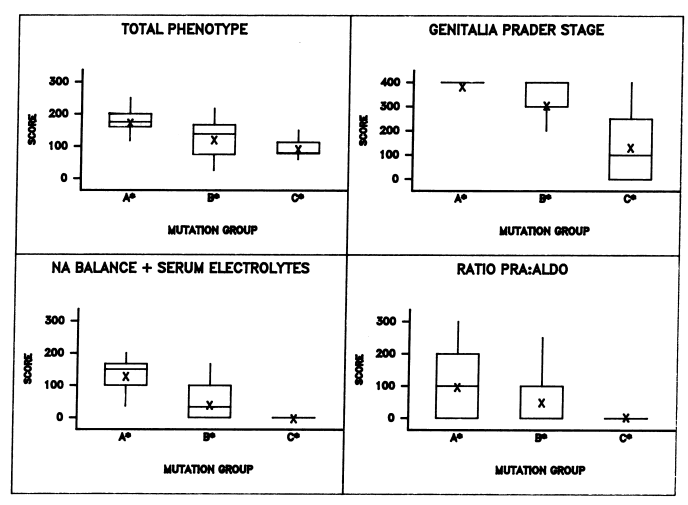


Figure 4. Relationship of phenotype score to mutation group. Adjusted phenotype scores (upper left) were calculated as described in the text. The other three panels show adjusted scores for the components of the total score, excluding 17-hydroxyprogesterone measurements. The box encompasses the 25-75th percentiles, the bar indicates the median, the cross indicates the mean, and the vertical lines indicate the ranges. Asterisks beside group letters show statistical differences between groups: P = 0.0001 among mutation groups A, B, and C for total phenotype score, Prader stage, and sodium balance + serum electrolytes; P = 0.04 among groups A, B, and C for the ratio of PRA/aldosterone (ALDO).

aldosterone production and sodium homeostasis in genotypically identical siblings in this and an earlier report (34), it is plausible that at least some differences in clinical disease expression are governed by factors remote from the CYP21 locus.

Differences between salt-wasting and simple virilizing patients with respect to 17-hydroxyprogesterone levels may actually be more significant than is evident from this sample, since in three cases 17-hydroxyprogesterone levels were recorded as extremely high and were excluded from analysis because the levels were not precisely measured. Data on patients with severe, salt-wasting 21-hydroxylase deficiency are sparse because of the difficulty encountered in obtaining hormonal measurements in the absence of treatment. Some patients included in this study (13 salt-wasting and 6 simple virilizing) were born and diagnosed before the availability of radioimmunoassay for 17-hydroxyprogesterone. It is not clear why one salt-wasting infant had unusually low levels of 17-hydroxyprogesterone.

Third, some variations in phenotype may represent "leakiness" of the intron 2 splicing mutation, with 4/27 patients in the  $A_2$  sub-group having been diagnosed as having simple virilizing disease. Splicing mutations have been associated with a variable degree of phenotypic severity when examined in thalas-

semia (38). In vitro transcription of  $\beta$ -thalassemia genes carrying intronic mutations that activate cryptic splice acceptor sites show that the cells' splicing machinery recognizes cryptic splice acceptors with different frequencies and that some normally processed mRNA can result from such genes (39). The in vitro activity of CYP21 bearing the intron 2 A  $\rightarrow$  G mutation has been examined in a single study, which indicated that the intronic mutation confers less severe enzymatic compromise than the Ile-172  $\rightarrow$  Asn mutation (36). It is not known whether this study, which involved transfection of cultured kidney epithelial cells, accurately describes in vivo activity. Precedent exists in at least one splicing mutation of the  $\beta$ -globin gene for finding an unexpectedly severe phenotype compared with the results of in vitro studies (40).

Finally, one might postulate that phenotypic severity is influenced by parental imprinting (41) or by negative allelic complementation, i.e., exaggerated gene dosage effect. Although there is no evidence of either of these phenomena from this sample, a larger number of families should be examined.

The molecular genetic epidemiology of 21-hydroxylase deficiency as described in this study is in good agreement with data from the single earlier study of point mutations in CYP21

in which the genotypes of both parents have been examined (20). Knowledge of the specific molecular genetic defects in a given family should enhance the accuracy of prenatal diagnosis when compared with allele-specific oligonucleotide hybridization using class II HLA probes, which carries an inherent error rate of  $\geq$  1% because of recombination between the HLA-DR locus and CYP21.

In conclusion, we have demonstrated with careful clinical and hormonal evaluations of patients that the severity of clinical disease generally correlates with what might be expected based on in vitro expression studies of discrete mutations. However, in each of the three mutation groups there were patients whose phenotype was different from that predicted, the reason for which is not immediately obvious and requires further investigation. In contrast to conclusions drawn by Okano et al. (42) in a recent elegant study of phenotype-genotype correlations in phenylketonuria, we are less certain that knowing the genotype will absolutely predict the course of the disease. Because of the observed discordance between phenotype and genotype in each mutation group, we believe that clinicians would be ill-advised to make therapeutic decisions based on genotype or to predict clinical outcome in a second-born affected child based on data gleaned from an older sibling. We advocate careful hormonal studies at presentation and periodically throughout childhood and adolescence to optimize therapy for each patient. Further studies of genetically identical sibling pairs may illuminate the source(s) of individual variability in phenotype. The relatively high rate of new mutations observed in this small sample ( $\sim 1\%$ ) should be further examined in a larger series of families.

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