Zurich, Zurich, Switzerland.
Male Pseudohermaphrodism due to 17,20-Desmolase Deficiency.
A $51 / 2$ year old $46, \mathrm{XY}$ pseudohermaphrodite presented with microphallus, third degree hypospadias, chordee and normal male internal genitalia. Serum LH and FSH levels were $<1.7 \mathrm{mIU} / \mathrm{ml}$. Results of adrenal and gonadal stimulation and Dexamethasone (DEX) suppression were:
Serum (ng/dl) BASAL ACTH DEX DEX and HCG
Progesterone
17-OH-Progesterone 17-OH-Pregnenolone Androstenedione
Testosterone
Cortisol ( $\mu \mathrm{g} / \mathrm{d}$ )
DHEA-S ( $\mu \mathrm{g} / \mathrm{d} 1$ )
Urine (mg/24hr)
Pregnanetriolone DHA

| BASAL | ACTH | DEX | DEX and HCG |
| ---: | ---: | ---: | :---: |
| 56 | 629 | 16 | 16 |
| 350 | 1350 | 34 | 34 |
| 173 | 225 | 37 | 42 |
| 61 | 48 | 10 | $<10$ |
| 40 | $<10$ | $<10$ | $<10$ |
| 35 | 46 | 1 | .9 |
| 31 | 46 | 9 | 10 |
|  |  |  |  |
| .2 | 3.93 | 0.01 | 0.04 |
| 4.01 | 4.01 | 4.01 | 0.01 |

In this patient the elevation of progesterone, 17-0H-Progesterone, 17-OH-Pregnenolone, and pregnanetriolone in conjunction with low DHEA-S, androstenedione, and DHA are consistent with gonadal and adrenal 17,20-desmolase deficiency which resulted in incomplete virilization.

Diagnosis and control of treatment of congenital adrenal hyperplasia (CAH) by semiautomated capillary gas-liquid-chromatography (OGIC) of steroid trimethylsilyl-enolethers (TMSEE).

CAH due to 21-hydraxylase (21-HD), 11-hydroxylase (11-HD) or $3 \beta$ hydroxysteroiddehydrogenase ( 3 B-HSD) deficiency can be diagnosed by detection of specific urinary steroid metabolites. Moreover, CAHtreatment can be controlled by quantitation of specific metabolites i.e.pregnane- $3 a, 17 a, 20 a \pm r i o l$ and pregnane- $3 a, 17 a, 20 a t r i o l-11$ on in 21-HD, tetrahydro-11-descxycortisol in 11-HD and A5pregnene-3 $\beta, 17 \alpha$ $20 a t r i o l$ and 16 ahydroxy- 45 pregnene $-3 \beta$ ol-20 on in $3 \beta-$-HSD. We have developed a sensitive, highly specific autamated method for the quantitative oGIC of TNSEE of these and other steroids. The method includes enzymatic hydrolysis, ether extraction, one-step derivatisation using N -methyl-N-trimethylsilyl-trifluoracetamid in presence of sodium acetate. The autamated oGic using solid injection separates about 10 samples/night. Because of the high resolution, no further purification step is needed. Mean coefficient of variation of the entire method is 128 . Unknown steroids can be detected by OGIC and identified by mass spectrometry. The TMSEE are convenient and stable enough for oGIC conditions. Ranges of steroid excretion in the different forms of CAH for four different age groups indicating optimal treatment have been established. In conclusion, oGIC with steroid TMSEE provides a precise, rapid and convenient tool for both diagnosis and treatment control of CAH.
 M. ZICBPMCN). Dopt.of Pediatrica, Soroke Med. Center, Foculty of Eealth Seiceese Ben-Garion Daiv.of the Hegev,
 Jeracalea. Congenital adrenal hyperplasia dee to 118 hydroxylaee deficioney. 25 casea of congenital edremal hyperplasia due to 118 hydroxylase doficioncy belonging to 17 fanilies have beon diagoosed in Iaral. These provided an unique apportanity to atudy the clinical apectrum of this disease. Pationts were of Nortb-ifrican Jewiah extraction.Diagnosia wae amopected by clinical evidonce of premature or abnormal virilisation acsociated in some cases with hypertonaion. The diagnosis confined biochenically by the presence of high urinary levols of tetra hydro 11 deexycortisol. In affected females clinical expression varied from onlarged elitoris to sevin -ly hypertrophied clitoris with penile urethra and fused labial acrotal folds. 10 out of 14 fanales whe were not diagnosed early in life wre reared as males, required corrective aurgery at puberty. Removed orarios aboved oystic changes. In these androgen oxcess was umpifective in suppressing gonedotropin secretion. Fopprtenaion was prosent in 15/25 cases and led to fatal vaceular accidents in 3 casea.Eppokalemia, obeerved in 9 patients was not correlated to hypertension. Kreept in infate, iow levels of ronin activity were found in all mitreated cases,indicating a state of volume oxpansiono№ correletion was found betwoen the degree of virilisation and biochemical ovidence of mimoralocerticoid excese.Preliminary data on an atteapt at antenatal diagnosis by meacuring THi in maternal urine and fatal aniotic fluid of affectad cases will be roported.

HLA genotyping and hormonal studies in 9 females with non-classical steroid 21-hydroxylase deficiency (AAH) indicate that this disorder is due to an autosomal recessive gene linked to HLA, similar to classical and cryptic 21-hydroxylase deficiency (21-OH def). They had normal genitalia at birth and presented between 9 mos to 16 yrs with varying degrees of virilization. Hormonal studies of the families revealed 2 fathers and their HLA identical sisters with 21-0H def. The remaining parents and the sibs sharing one HLA haplotype with the AAH patient responded to ACTH stimulation as heterozygotes for classical or cryptic 21-OH def. Five sibs who were HLA identical to their affected sib also had findings diagnostic of $21-0 \mathrm{H}$ def. The hormonal response to ACTH of the patients with AAH and their HLA identical sibs was similar to that of patients with cryptic 21OH def. Thus, individuals with these non-classical forms of $21-0 \mathrm{H}$ def and similar hormonal findings present with a clinical spectrum ranging from an asymptomatic deficiency to precocious pubic hair, acne, tall stature and advanced bone age, hirsutism, clitoromegaly and menstrual irregularities. The results of these studies support the concept that AAH, similar to classical and cryptic $21-0 \mathrm{H}$ def is due to an HLA linked autosomal recessive gene and that these disorders are due to allelic variants at the locus of steroid 21hydroxylase.
M.I.NEW, B.KOHN*, M.POLLACK*, S.PANG, D.LEVY*, G. RONDANINI*,F.LORENZEN*,A.LERNER*, B.DUPONT*, L.S.LEVINE, The New York Hosp-Cornell Med Ctr, SloanKettering Inst Cancer Research, NY USA and Univ Milan, Italy. Genotyping for 2l-hydroxylase deficiency: one or two genes?

We have devised nomograms relating the baseline and ACTH stimulable levels of $17-\mathrm{OHP}, \triangle 4$-androstenedione and testosterone for genotyping 21-hydroxylase deficiency. The nomograms provide a method for classifying the patient with congenital, late onset or cryptic 2l-hydroxylase deficiency as well as classifying the heterozygotes for each of these disorders. In addition, the subject predicted by HLA genotyping to be genetically unaffected can also be classified by these nomograms. Further the nomograms permit us to obtain evidence for genetic recombination between HLA and the 21-hydroxylase locus. For example a patient predicted by initial HILA genotyping to be unaffected was classified by the nomogram to be a heterozygote. When HLA-DR typing was performed an informative maternal HLA A:DR recombination was discovered. This recombination explained the heterozygote response of this subject. In another family a maternal DR:GLO recombination was found in an asymptomatic sister who was HLA identical to the patient with late onset 2l-hydroxylase deficiency. Although most recombinants have mapped the gene for 21 -hydroxylase between B and DR, this DR:GLO recombination presents evidence that there may also be a 21-hydroxylase locus between the DR-GLO loci. The nomograms thus provide a powerful tool to determine the 21-hydroxylase genotype by hormonal testing and assist in mapping the gene for 2l-hydroxylase deficiency.
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Middlesex, Great-Britain (Intr. by J.L. van den Brande).
Hypertension hypokalemia and retarded growth in a 17 month-old boy with 11 hydroxysteroid-dehydrogenase deficiency.

A 17 manth old boy, the first child of unrelated parents, presented with polydipsia, lack of appetite and failure to thrive. Blood pressure was $150 / 100 \mathrm{mmHg}$. He had hypokalemia ( $2.6 \mathrm{mEg} / \mathrm{l}$ ), suppressed renin activity ( $0.4 \mathrm{ng} / \mathrm{ml} / \mathrm{hr}$ ) and low plasma aldosterone level ( $2 \mathrm{ng} / 100 \mathrm{ml}$ ).

Urinary $11 / 3$-hydroxy-steroids were increased relative to 11 consteroids (THF: 110 , THE $8 \mu \mathrm{~g} /$ day; ratio 13.8 . Control ratio 0.4 ) indicating $11 \beta$-hydroxysteroid-dehydrogenase deficiency. Both parents were nomal. The hypertension and hypokalemda were unresponsive to spironolactone. Dexamethason treatment did not
influence the hypokalemia and even increased blood pressure. Triamterene ( $3 \times 25 \mathrm{mg}$ ) nomalised serum potassium, but addition of furosemide ( $2 \times 10 \mathrm{mg}$ ) was required for nomalisation of blood pressure. This treatment resulted in catch up growth (from -3.6 SD to -1.8 SD for height).
This is the youngest patient known with this syndrome. Unlike the other cases (Ulick et all. J.C.E.M., 49: 757, 1979) he did not respond to triamterene alone. Also the growth retardation and

