LARON TYPE DWARFISM IS NOT DUE TO A MUTATION IN OR 462 NEAR THE HUMAN GROWTH HORMONE GENE. John S. Parks, **402** Laura C. Sexton, Ruth Keret, Myrta Kalichman, Athalia Pertzelan and Zvi Laron. Emory University School of Medicine, Department of Pediatrics, Atlanta and Tel-Aviv University, Institute of Pediatric Endocrinology, Israel.

We have analyzed DNA restriction fragment length polymorphism to determine whether the mutation responsible for Laron Type Dwarfism (LTD) is linked to the human growth hormone (hGH) gene. An hGH haplotype represents the combination of presence absence of variable restriction sites for Bgl II, Hinc II and \underline{Msp} I at 5 locations in a single 55 kb hGH gene cluster. We studied 3 Israeli families with LTD. Different haplotypes are indicated by capital letters A-D:

	Famil	YI		1		II		III
Mother	LTD	LTD	Father	1	Mother	LTD	Father	LTD
AAA	AB	AIC	BIC	1	DE	DID	DA	DB
The affe	cted s	ibling	s in Fam	ily	I have	inher	ited dif	ferent hGH
genes from	n their	norma	l father	·. I	LTD pati	ents	in 3 fam	ilies show
a total	of 4	diffe	rent hG	H h	naplotyp	es.	Discord	ance for
haplotype	betwe	en af	fected s	ibl	ings and	l lack	of asso	ciation of
LTD with a	a parti	cular	hGH hapl	oty	pe indic	ate	that LI	D is not
linked to	the h	H gene	. This	resi	ult excl	udes	mutation	of hGH to
								stance and
								gene is
mutated :	in LTI), the	n this g	ene	is not	locat	ed in pr	oximity to
the hGH g	ene.							

ULTRASOUND EVALUATION OF NEWBORN THYROID STRUCTURE: <u>G. Carpenter</u>. Thomas Jefferson University, Depart 463 ments of Diagnostic Ultrasound and Radiologic Imaging and Pedi-atrics, Philadelphia, Pa.

Infants with suspect thyroid dysfunction need visualization of the structure of the thyroid gland in addition to the understood biochemical specific studies for diagnostic and follow up care. With the application of ultrasound real time (10 MHz high re-solution real time, prototype ATL, Bellevere, WA) 8 evaluations have revealed information aiding diagnosis and counseling poten-tial for parents. Infants with prematurity and low T4 studies compatible with the diagnosis of "sick euthroid" when seen to have a normal gland may be observed with less anxiety than those found to have no structure resembling thyroid. Gland size of infants whose mothers required antithyroid medication during pregnancy can be serially followed. Term newborns with screening values in the first weeks of life that indicate possible athyre-osis can have rapid confirmation when the gland is found absent; those with dysgentic thyroid structure may be followed with or without treatment depending on other individual balances of thyroid studies. Parents have indicated appreciation of this newer visualization technique utilizing ultrasound. The procedure of ultrasound evaluation of the newborn and pre-mature infant thyroid gland may supply a needed technique in the

mature infant thyroid gland may supply a needed technique in the optimum care of newborn and premature infant.

EVIDENCE FOR INDEPENDENT MUTATION OF THE 21-HYDROXY-LASE DEFICIENCY (210H DEF) GENE IN ALASKAN ESKIMOS WITH CONGENTRAL ADRENAL HYPERPLASIA (CAH). GM Petersen, JE MacCracken, JI Rotter, D Trotter, MI New, MS Park, PI Terasaki, RS Sparkes, JI Ward. Harbor-UCLA Med. Ctr., Torrance, CA; PHS Hosp. Bethel, Alaska; NY Hosp.-Cornell Med. Ctr.; UCLA Med. Ctr., Los Angeles, CA.

Worldwide, the salt-losing form of CAH (210H Def) has been reworldwide, the salt-losing form of CAR (210h ber) has been re-ported to have the highest known prevalence in Yupik speaking Alaskan Eskimos. The annual incidence is 1/490 with an estimated gene frequency of .045. We analyzed the HLA phenotypes and inferred genotypes in 11 Eskimo children with CAH, representing 7 families. All CAH children had at least one HLA-B27 allele (6 homozygotes), at least one HLA-C4 allele, and all were homozygous MLA-D20, WLA types chtsined on 118 healthy Refirm controls chtrod HIA-DR4. HLA types obtained on 118 healthy Eskimo controls showed significantly different frequencies for these alleles:

Allele Frequencies:	CAH Cases	Controls	Significance
HLA-B27	.93	.14	p<.001
HLA-C4	.50	.10	p<.001
HLA-DR4	1.00	.33	p<.001

The inferred haplotypes which carry the 210H Def gene include A24 B27 C4 DR4 and A2 B27 C2 DR4, plus A28 B61 C3 DR4, which occurs in only one family. The three haplotypes combined occur in .076 of the controls, a frequency only slightly greater than expected for CAH heterozygotes based on the observed CAH disease incidence, implying that most Yupik Eskimos with these haplotypes are heterozygous for 210H Def. These results differ markedly from other HLA population studies, suggesting that the 210H Def gene in Yupik Eskimos occurred by an independent mutational event.

DEFECTIVE TRANSFORMATION OF ANDROGEN-RECEPTOR COM-**† 465** PLEXES: A CLASS OF FAMILIAL, RECEPTOR-POSITIVE PARTIAL ANDROCEN RESISTANCE. Leonard Pinsky, Morris Kaufman. Lady Davis Institute and Centre for Human Genetics, McGill University, Montreal, Canada.

We have analyzed the androgen (A)-receptor (R) system in genital skin fibroblasts (GSF) of 2 subjects with external genital ambiguity. Subject 1 has 5 affected maternal relatives in 3 gene-(B_{max}) (receptor-positive), but an apparent equilibrium dissociation constant (K_d) of 1.2-1.4 nM for 5α -dihydrotestosterone (DHT) or the nonmetabolizable androgen, methyltrienolone (MT) (normal: Solution for the holinear and the second se regulation. In contrast, with MT his AR is normal GSF yield higher Kds (0.3-1.8 nM) for either ligand after 0.5compared to 2-h assays, and the GSF of both subjects catabolize DHT to a normal extent. Conclusions: (i) normal GSF transform initial, low-affinity A-R complexes to higher affinity states by a process that is codependent on time and initial ligand concentration; (ii) subject 1 forms DHT- and MT-complexes that remain in the initial, low-affinity state; (iii) subject 2 forms complexes that transform normally with MT but partially with DHT; and (iv) A-R complexes must attain the highest affinity state to act as the «signal» for up-regulation.

CORD SERUM THYROID STIMULATING HORMONE (TSH) AND THYROGLOBULIN (Tg) LEVELS DECLINE WITH INCREASING BIRTH 466 ROGLOBULIN (Tg) LEVELS DECLINE NIN AND A ROBERT PENNY WEIGHT IN NORMAL FEMALE NEWBORNS. Robert Penny

Carole A. Spencer, John T. Nicoloff. Depts. of Pediatrics and Medicine, Univ. of So. Calif., Los Angeles, California. Tg, TSH and T4 were determined in cord serum and related to birth weight and sex in normal newborns, 20 females and 19 males. Mean + SD birth weight of female infants (3299 + 282 grams) was significantly (p <0.005) less than that of male infants (3757 + 447 grams). Whereas, mean gestational age of female infants did not differ from that of male infants (40.1 + 0.5 vs 40.1 + 0.7 weeks). Mean Tg levels of female infants (31.1 + 8.9 ng/ml) was significantly (p <0.05) greater than that of male infants (26.1 + 7.7 ng/ml). TSH (14.9 + 11.3 vs 14.8 + 13.4 μ U/ml, p >0.1) and T4 (11.1 + 3.3 vs 11.7 + 3.5 μ g/dl, p >0.1) levels of female and male infants did not differ significantly. Further analysis indicated that Tg levels (r = -0.401, p <0.05) and the log of TSH levels (r = -0.576, p <0.005) correlated negatively with birth weight in female infants. Also, Tg levels correlated positively with the log of TSH levels (r = 0.401, p <0.05) in female infants. In contrast, none of these correlations were significant for male infants. We conclude that the sex differsignificantly (p <0.005) less than that of male infants (3757 significant for male infants. We conclude that the sex differ-ence in cord serum values and their correlations may be principally related to changes in body composition that accompany increasing birth weight.

467 CORD SERUM THYROID STIMULATING HORMONE (TSH) AND THY-ROGLOBULIN (Tg) LEVELS DECLINE WITH INCREASING WEIGHT IN LOW BIRTH WEIGHT NEWBORNS. Robert Penny, Carole A. Spencer, John T. Nicoloff. Depts. of Pediatrics and Medicine, Univ. of So. Calif., Los Angeles, California. Tg, TSH, free T4 index, and free T3 index were determined in cord serum and related to birth weight and sex in infants with weights of less than 2500 grams, 19 males and 19 females. Free index values are the product of the T3 resin uptake ratio and the appropriate thyroid function indice. The term of the second hypothesis that changes in body composition accompanying increasing birth weight are associated with an increase in thyroid gland responsiveness to TSH or a decrease in T4 clearance.