SLUGGISH RESPONSE TO ADRENOCORTICOTROPIN STIMULATION IN NEWBORNS

SLUGGISH RESPONSE TO ADRENOCONTICOTHOPIN'S IMULATION IN NEWBORNS WITH 21-HYDROXYLASE DEFICIENCY (CAH). <u>G. Kalaitzoqlou</u> and M.I. New, Dept of Pediatrics. The New York Hospital-Cornell Medical Center, New York, NY 10021, USA An unusually sluggish response to a single 0.25 mg dose of ACTH was noted in some newborns with congenita adrenal hyperplasia due to 21-hydroxylase deficiency (21-OHD) diagnosed by molecular genetic studies. Four of the 11 (36%) affected newborns that we studied, demonstrated baseline and ACTH slimulated 17-hydroxyprogesterone (17-OHP) levels clearly higher than the unaffected newborns, but only in the range of values established for heteroxygote rather than that for homozygote affected patients at an older ace. Therefore, we set out to establish the range of drepend sterior terponse an older age. Therefore, we set out to establish the range of adrenal steerol precisions of on ormat and affected infants in the newborn period. The serum concentrations of 17-OHP and Δ-4-Androstenedione (Δ-4-A), which have proven most useful in diagnosing CAH are summarized as follows:

	17-OHP (nmol/l)		Δ-4-A (nmol/l)	
	Baseline	Stimulated	Baseline	Stimulated
Affected	Mean: 514.8	Mean: 748.0	Mean: 275.9	Mean: 231.3
(M = 4;F = 7)	Range: 116.6-1075.4	Range 191.5-1721.4	Range: 22.6-970.1	Range: 94.1-432.1
Unaffected	Mean: 1.8	Mean: 5.6	Mean: 1.1	Mean: 2.8
(M = 6;F = 7)	Range: 0 0-4.0	Range: 0.8-31.6	Range: 0.0-2.9	Range: 0.0-6.8

We also measured baseline and stimulated values of 17-hydroxypregnenolone (17-OHPreg), 11-deoxycortisol (DOC), cortisol (F), deoxycorticosterone (B), dehydroepiandrosterone (DHEA), testosterone (T) and the calculated ratios of 17-OH Preg/17-OHP, DHEA/A-4 and 17-OHP/DOC. In conclusion. The response of 17-OHP to ACTH stimulation in newborns affected with CAH may be lower than that of older infants and children but higher than unaffected individuals. Physicians should be alerted to the possible significance of only moderately elevated 17-OHP levels in the early neonatal period.

93

MULLERIAN INHIBITING SUBSTANCE EIA IS YIELDING NEW INSIGHTS INTO GONADAL PHYSIOLOGY AND PATHOLOGY. ML Baker and IM Hutson, Surgical Research Laboratory, Royal Children's Hospital Research Foundation, Melbourne, Australia.

Mullerian inhibiting substance (MIS) is a testicular hormone made by fetal and postnatal Sertoli cells, and causes Mullerian duct regression during male sexual differentiation. We established an EIA to measure MIS serum levels in a number of physiological and pathological conditions. Significant levels of serum MIS are found in normal males during the first year of life (1). MIS levels decline throughout childhood and are undetectable after pubertal development (2). MIS levels were examined in patients previously diagnosed with Hodgkin's Lymphoma. Nine adults (>16 years) had low but measurable levels of MIS. Patients with Hodgkin's Lymphoma have azospermia, suggesting a possible link between MIS secretion and the spermatogonia. In very low birth weight babies (<1500g), MIS levels in cord sera were significantly higher than in full-term controls (79125µg/L vs 36±8µg/L; P<0.001). MIS levels also were examined previously in infants and boys with cryptorchidism, where the levels during the first year were significantly lower than normal (3). This suggested a possible link between MIS and germ cell maturation, which is occurring at that time. Recently we have confirmed experimentally that MIS does stimulate germ cell maturation in the neonatal mouse testis in vitro

(1) Baker ML, Metcalfe SM & Hutson JM. J Clin Endocrin Metab. 1990; 70: 11-15 (2) Baker ML & Hutson JM, J Clin Endocrin Metab (In Press)

(3) Yamanaka J, Baker ML, Metcalfe SM & Hutson JM. J Pediatr Surg. 1991; 26: 621-623

94

Baiyun Zhou MD and John M Hutson MD, FRACS Department of Surgical Research. Royal Children's Hospital Research Foundation, Melbourne, Australia. HUMAN CHORIONIC GONADOROPIN (hCG) FAILS TO STHULATE GONOCYTE DIFFERENTIATION IN NEWBORN MOUSE TESTES IN ORGAN CULTURE.

Postnatally, the germ cell differentiate through several steps to form primary spermatocytes, which are required for postpubertal spermatogenesis. This development is postulated to be controlled by the hypothalamic-pituitary-gonadal axis. To test the effect of hCG on germ cell development, newborn mouse testes (n=75) were cultured in vitro for 7 days. Exogenous hCG (0.102.0 IU/ml) or Mullerian inhibiting substance (0.5 μ /ml) (MIS) were added to serum-free medium containing transferring insulin and retinoic acid or fetal calf serum. Normal germ cell development was seen with added 10% fetal calf serum (P < 0.001) or exogenous MIS (P < 0.001), but was absent with medium ± growth factors or hCG. These results demonstrate that transformation of gonocytes to type- λ spermatogonia and other differentiated germ cells is regulated by MIS rather than hCG. As transformation of gonocytes to type- λ spermatogonia is deficient in boys with cryptorchidism, MIS may have a role in the clinical management of this common cause of infertility.

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RECEPTOR ACCESSORY FACTOR (RAF) ENHANCES SPECIFIC DNA BINDING OF ANDROGEN AND GLUCOCORTICOID RECEPTORS. <u>S.</u> <u>R. Kupfer</u>, K. B. Marschke, E. M. Wilson, and F. S. French, Laboratories for Reproductive Biology, University of North Carolina, Chapel Hill, NC 27599, USA

The biological responses to steroid hormones are mediated by a family of ligand-inducible intracellular receptors that activate or repress transcription of target genes. Protein-protein interactions are common among other transcriptional activators and may have important consequences for gene regulation by steroid hormone receptors. Using the mobility shift assay we have identified a factor that enhances specific DNA binding of truncated rat androgen (AR) and glucocorticoid (GR) receptors by 25-fold and 6-fold, respectively, through the formation of heteromeric complexes. This factor, designated receptor accessory factor, or RAF, also potentiates DNA binding of full-length human GR. RAF is temperature and trypsin sensitive and is present in a variety of cultured mammalian cells. By gel filtration RAF has a predicted molecular mass of 130 kDa. RAF enhancement of AR-DNA binding requires androgen response element DNA. RAF appears to interact directly with AR because deoxycholate, which interferes with proteinprotein but not protein-DNA interactions, prevents RAF-AR-DNA complex formation. Furthermore, RAF activity is recovered from an androgen response element DNA affinity column only in the presence of AR. Mutagenesis of truncated AR fragments indicates that a region in the NH2-terminal domain is required for RAF to enhance AR-DNA binding. The interaction of RAF with AR and GR suggests that RAF might influence the ability of these nuclear receptors to activate transcription.

97

NG-MONOMETHYL-L-ARGININE DIRECTLY STIMULATES IN VITRO LEYDIG CELL

N⁶-MONOMETHYL-L-ARGININE DIRECTLY STIMULATES <u>IN VITRO</u> LEYDIG CELL TESTOSTERONE PRODUCTION. <u>M. Watson</u>, M. Poth, and G. Francis. Department of Pediatrics, Walter Reed Army Medical Center, Washington, DC 20307, and Uniformed Services University of the Health Sciences, Bethesda, MD 20814, USA. Recent studies suggest nitric oxide (NO) may function as both an intercellular and intracellular signal (second messenger). Changes in NO generation are thought to affect neural and immunologic activity, vascular tone, platelet adhesion, and selected hormone production. Arginine analogues such as N⁶-monomethyl-L-arginine (L-NMMA) are thought to inhibit intracellular NO generation and have been used to study the effects of decreased NO in physiologic systems. A single <u>in vivo</u> study has suggested NO modulates testicular endocrine function but it is unclear if this was due to changes in vascular tone, neuronal activity, or Leydig cell steroidogenesis itself. The present study was performed to evaluate the <u>in vitro</u> effects of L-NMMA on basal and human chorionic gonadotropin (hCG)-stimulated production of adenosine 3':5'-cyclic monophosphate (cAMP) and T by Leydig cells. Rat Leydig cell-enriched cultures (ltN0⁶ cells/culture) were incubated 4 hr with L-NMMA, after which hCG (0.1 U/ml) was added. Levels of T in the media were determined at 20 hr and shown here as ng/10⁶ cells<u>t</u>SEM. (*pC0.02, *ppC0.001) <u>0.1 0.5 1.00</u>

		0.1	0.5	1.0
BASAL	1.9+0.03	2.94+0.09**	3.34+0.07**	3.06 <u>+</u> 0.11**
		10.0+0.5*		9.8 <u>+</u> 0.1*
L-NMMA	had no effec	ct on cAMP prod	uction (not sho	wn). We conclude
that L-	-NMMA increas	es T production	by both basal a	and hCG-stimulated
Leydia	cells withou	t increasing cA	MP. This sugge	sts that NO may be
		f Levdin cell T		