VASOPRESSIN (VP) RELEASE IN FETAL NEWBORN AND ADULT 449 SHEEP INDUCED BY HYPOXIA: CORRELATION WITH β -ENDORPHIN

IMMUNOACTIVITY (-EP) & PaO2. <u>Raymond I. Stark, Salha</u> <u>A. Daniel, Kasim H. Husain, Sharon L. Wardlaw & L.Stanley James, Div.</u> of Perin.Med., Depts. of Ped., Med., Coll. of P&S, Columbia Univ., NY.

To study the relationship of plasma VP(pg/ml), \$-EP(pg/ml) & PaO₂(mmHg)during hypoxic stress, we exposed pregnant ewes & lambs to an FIO₂ of 5 or 10% in N₂ for 30 min.In 11 fetuses of ewes exposed to 10% FIO2, VP increased from 1.7±0.5 to 56±17.8 (p <.001). In 6 newborn lambs with 10% FIO2, mean VP increased from 2.1±0.4 to 10.5±5.8 (p <.005) while with 5% FIO2, VP increased to 421±138 (p <.001). Exposure of 7 ewes to 10% FIO2 produced no change in mean VP, yet with 5% FIO2 mean VP increased from 1.7±5 to 38.9±15.7. There were significant negative correlations (p < .001) between Pa02 and log VP in the fetal & newborn lamb. Unlike VP control β -EP levels were higher (p < .01) in the fetus (134±31) & newborn (155±24) than in the ewe (50±11). Fetal β -EP in-

(13431) & newborn(15524)than in the ewe(50f11). Fetal β -EP increased during hypoxia to 544t81(p < .01) & values were positively correlated(r=.63,p < .005)with log VP. Lambs & ewes exposed to 10% FI02 had no increase in β -EP,while with 5% FI02 mean β -EP increased to 532±158 & 150±17(p < .01). β -EP levels correlated with VP(r=.91,p < .001). VP & β -EP values during hypoxia were higher(p < .02) in fetuses < 130 days gestation than those > 130 days.

We conclude that hypoxia strongly stimulates the release of VP & β -EP into the peripheral blood. With advancing gestation the fetus becomes more responsive to hypoxia. Data suggest parallel but independent release of VP & β -EP during hypoxia.

ZINC (Zn) AND BIRTH DEFECTS (BD). Constance Stewart, 450 Brad Katchan, Platon J. Collipp, Sanda Clejan, Scott Pudalov, and Shang Y. Chen. Nassau County Med Ctr., SUNY, Stony Brook Health Sciences Ctr., Department of Pediatrics, East Meadow, NY 11554.

Zn deficiency has been reported in infants with BD and in children born with achondroplasia. 304 families were studied comparing parental age, Zn nutritional status and infant weight at the time of birth.

Parental	H	air Zn (u	ıg/g)	Infant Wgt	Infant Hair			
Age(years)	Infant	Mother	Father	(g)	Zn			
14-20	192 [±] 23	174 [±] 33	142-28	∠ 2500	200 ± 30			
21-30	212±35	181±39	181 [±] 41	2500-3500	203±31			
31-40	199 ± 28	173 [±] 43	176±44	> 3500	192 ± 24			
There is a s	tatistica	lly signi	lficant co	orrelation (p 4	(.01) between			
infant hair	Zn and man	ternal ag	ge at time	e of birth. T	he decrease			
in infant hair Zn seen with teenage (14-20) and elder mothers								
(31-40) may	be relate	d to the	incidence	e of increased	BD which oc-			
curs in thes	e groups.	Infant	hair Zn w	vas lowest in	the heaviest			
infants and	in a group	p with BI) made up	of porencepha	ly, anence-			
phaly and mi	crocephal	y as the	presentin	ng sign (N=4;	Zn=134±10).			
The heavier children may represent the infants of potential dia-								
betic mother	s (who hav	ve more l	BD). Plac	ental Zn was	18-19 ug/g			
and did not	correlate	well wit	h materna	al or infant h	air Zn. Fin-			
ally, diet d	id not exp	plain par	cental Zn	status since	paired de-			
ficiencies d	id not oc	cur. The	erefore, a	n association	between Zn			
levels, high	-risk age	groups a	and those	children with	BD is ap-			
parent.								

INSULIN BINDING STUDIES IN THE ERYTHROCYTES OF 451 DIABETICS IN ONE KINSHIP. Larry D. Stonesifer, M. Joycelyn Elders, Victoria L. Herzberg, J. Mark
Boughter, Thomas J. Sziszak, and Donald E. Hill. University of Arkansas for Medical Sciences, Department of Pediatrics, Little Rock, AR.

Specific insulin binding to erythrocyte receptors has previously been reported to be unaltered in children and adolescents with insulin dependent diabetes mellitus (Ped. 66:385, 1980). I report here that five insulin dependent, fasting, non-ketotic We juvenile diabetics from the same close kinship have a significant Juvenile diabetics from the same close kinship have a significant reduction in the percent insulin bound to erythrocytes (Normal $B/T=8.32\pm2.71$, n=24±1SD vs 4.72%±0.84 for the diabetic child-ren). Three non-diabetic juveniles in the family have an inter-mediate value of 5.77%±1.08. The youngest family member has normal binding. Scatchard analysis or two-site analysis indicate that the reduced binding is due to a decrease in the number of receptor sites/cell. There is no correlation with circulating plasma insulin values nor with the severity of the diabetes, normed the insulin diabetics have an inordinate dehowever, the insulin dependent diabetics have an inordinate de-gree of ocular complications for the severity of the diabetes. Since diabetes mellitus is a heterogeneous disorder, there may be familial patterns to the abnormality seen with the erythrocyte receptor as in the above patients.

TREATMENT OF CUSHINGS DISEASE (C.D.) BY TRANSPHENOID-•451A AL MICROADENECTONY (Tr.M.) IN CHILDHOOD AND ADOLES-CENCE. D.M. Styne, J.B. Tyrell, F.A. Conte, S.L. Kaplan, M.H. Connors, G.P. August, C.B. Wilson, and M.M. Grum-bach. Dept. of Pediatrics, Univ. California San Francisco, CA. The efficacy of Tr.M. in the treatment of C.D. in children and adolescents has not been assessed. We treated 7 patients (7-6/12 to 18-9/12 years old) with C.D. before epiphyseal fusion with Tr.M. (followup 2-72 months postoperatively). Growth failure and weight gain were the first signs of C.D.; pubertal delay, viril-ization and fatigability were variable features. Preoperative evaluation showed elevated but highly variable urinary free cortisol and no diurnal variation in plasma cortisol or ACTH values; high dose dexamethasone therapy suppressed excretion of urinary free cortisol. Two of 7 patients lacked definite radiographic evidence of a pituitary adenoma. Transphenoidal exploration revealed 1-4 mm adenomas in 6 of 7 patients; no definite adenoma was noted in the 7th patient in spite of CT scan evidence of adenoma. Postoperative complications were limited to transient diabetes insipidus. The 6 treated patients had low cortisol and ACTH secretion postoperatively and required replacement glucocorticoid therapy for 6-12 months. Weight loss, growth and pubertal progression without recurrence of C.D. were noted in 5 patients with long term followup. Our experience indicates that ACTH secreting microadenomas are the principal cause of C.D. in childhood and adolescence and suggests that transphenoidal microadenectomy is the initial treatment of choice for Cushings disease in young patients.

DIRECT DETERMINATION OF 17a-HYDROXYLASE (170Hase) 452 DEFICIENCY IN A MALE PSEUDOHERMAPHRODITE BY IN VITRO STUDIES OF TESTICULAR STEROID BIOSYNTHESIS. A.Vargas, E.O.Reiter, K.Kula, L.J.Rodriguez-Rigau, E. Steinberger, A.W.Root. Univ. South Florida, Dept.Pediatrics, Tampa and Univ. Texas Health Sc.Ctr., Dept. Reprod. Med. and Biol., Houston. A 4 year old male (46XY) pseudohermaphrodite with 170Hase deficiency (Ped.Res.13:387A,1979) underwent bilateral orchi-ectomy 2 days after 3 days of HCG (2,000 u/day IM), during which dione, 170H-progesterone (170HP) and estradiol did not increase while progesterone increased from 52 to 69 ng/dl. Teased while progesterone increased from 52 to 59 mg/d1. Reased testicular tissue was incubated with louCi (3.3 nmol) of ³H-pregnenolone or ³H-170HP in 3 ml NADP fortified KRB buffer, pH 7.4, for 3 hr. at 37° C with and without 100 IU/ml HCG. <u>Metabolite</u> <u>pmol/20 mg tissue</u>

	3H-pr	³ H-pregnenolone				
	-HCG	+HCG	Normal Adult Male			
Progesterone	66.0	566.0	19.6			
170H-Pregnenolone	37.0	60.0	221.8			
170HP	0.9	66.6	115.0			
Testosterone	0.2	4.0	22.1			
	3	H-170HP	`			
Androstenedione	0.3	81.0				
Testosterone	59.0	96.0				
Thus the testes of th	is patient	were una	able to metabolize			
pregnenolone, but wer	e able to	metabolia	ze 170HP. The data ar			

consistent with deficiency of 170Hase activity.

SALT WASTING WITH POSTERIOR URETHRAL VALVES. John S. 453 Venglarcik, Dale Doerr, Virginia H. Peden, James A. Monteleone, St. Louis University School of Medicine, Cardinal Glemon Memorial Hospital for Children, Department of Pediatrics-Adolescent Medicine, St. Louis, MO. Clinical manifestations of posterior urethral valves (PUV) may

vary widely. Biochemical abnormalities have not been emphasized. Seven male infants (mean age 6 weeks, range 9 days to 6 months), seen over a ten year period, presented with laboratory evidence of salt wasting suggesting adrenal insufficiency. Admission sodium, potassium, BUN, creatinine, serum pH, and urine pH and specific gravity were recorded. When available simultaneous urine clific gravity were recorded. When available simultaneous unine and serum electrolytes were noted. In all cases PUV were con-firmed by voiding cystourethrogram. The mean sodium on admission was 117 \pm 11 mEq/L (range 95-127) while the mean potassium was 7.9 \pm 1.3 mEq/L (range 6.5-9.6). The BUN and creatinine were 76 \pm 42 mg/dl (range 2.2-136) and 3.7 mg/dl (range 2.2-5.8) respec-tively. The mean specific gravity of the urine was some and (range 1.005 \pm 1.017). In four patients simultaneous some and (range 1.005-1.017). In four patients simultaneous serum and urine electrolytes were determined. The mean serum sodium was 132 while the mean urine sodium was 9.25 mEq/Kg/L. All of the patients manifested an impaired ability to conserve water and so-dium. In addition, an impaired ability to excrete acids into the urine resulted in acidosis and hyperkalemia. The sodium, potassium, water and acid imbalance is not due to aldosterone defi-ciency but reflects a glomerulus-tubular dysfunction. Infants presenting with hyponatremia and hyperkalemia who have a very high BUN should have a urethral catheter implaced and a diagnosis of PUV ruled out.