1600 DEVELOPMENT OF NORMS FOR MEASUREMENT OF INTRACRANIAL STRUCTURES IN HIGH-RISK INFANTS: PRELIMINARY FINDINGS. Ronald L.Poland, Thomas L. Slovis, Seetha Shankaran. Departments of Pediatrics and Radiology, Wayne State University, Children's Hospital of Michigan, Detroit, Michigan.

25 real-time ultrasonic sector scans of the head were obtained through anterior fontanelles of high-risk infants who were judged graphic criteria. Thicknesses of frontal mantle (FR.M) and occi pital mantle (OC.M) were measured in sagittal planes. Distances between the inner tables of the calvaria were measured in coronal sections at the level of the caudate nuclei (BC) and of the atria of the lateral ventricles (BA). Distances between outer walls of lateral ventricles were measured at the same levels (VC and VA).

lateral ventricles were measured at the same levels (VL and VA). We recorded birthweight, present weight, length of gestation, age post-conception and head circumference at each examination. Infants in this study ranged from 29-48 weeks post-conception. They weighed 800 to 3140 g. All four measures of brain size or thickness (OC.M, FR.M, BA, BC) increased significantly with age and with body weight (P<0.01 in 6/8 and <0.05 in 2/8). VA also

increased significantly with age and weight but not so with VC.

All of the following ratios remained stable as weight and age increased:  $VC/BC=0.31\pm0.07$  (m±5.0.);  $VA/BA=0.54\pm0.09$ ; OC.M/FR.M=0.86±0.13. The ratios were normally distributed in the population.

We are expanding this series in search of quantitative meas-ures of normal brain structures that may be used to identify ventricular enlargement early in high-risk infants.

1601 NEUROPHYSIOLOGICAL, PSYCHOLOGICAL, AND NUTRITIONAL INVESTIGATIONS DURING DISCONTINUATION OF THE PHENYL-ALANINE-RESTRICTED DIET IN CHILDREN WITH CLASSICAL ALANINE-RESTRICTED DIET IN CHILDREN WITH CLASSICAL PHENYLKETONURIA (PKU). <u>Siegfried M. Pueschel, Laurie Fogelson-</u> <u>Doyle, Betsy Kammerer, and Yoichi Matsumiya</u>. (Spon. by Leo Stern) Brown University, Program in Medicine, Rhode Island Hospital, Department of Pediatrics, Providence, Seizure Unit, Clinical Neurophysiology, Department of Neurology, Children's Hospital Medical Center, Harvard Medical School, Boston, Massachusetts. Children with classical PKU had been maintained on a low-phenylalanine diet from the first few weeks of life until their Sth birthday when the phenylalanine restricted diet was citonwise

5th birthday when the phenylalanine-restricted diet was stepwise changed to a regular diet. Physical and neurological examinations, nutritional evaluations, anthropometric measurements, and psychological assessments were done at 4,  $4\frac{1}{2}$ , 5, and 6 years of age. In addition, neurophysiological investigations including at 4 years 10 months, 5 years, 5 years 2 months, and 5 years 6 months.

Analysis of relationships among the study variables revealed that children who had adequate dietary control and who were of average or above average intelligence displayed normal somatosensory evoked potentials, whereas those children with less adequate dietary control and lower IQ scores had atypical somato-sensory evoked potential waveforms. Yet, these multiple assess-ments did not uncover any significant changes when pre-diet termination data were compared with post-diet termination results.

ASSOCIATION OF HYPERTENSION AND FLUIDS WITH CLINICAL 1602 MANIFESTATION OF BELERE INTRACRANIAL HEMORRHAGE (ICH) IN VERY LOW BIRTH WEIGHT INFANTS: TOnse N.K.Raju, G. Cybulski, G.J. Ramirez-Lavin, D. Vidyasagar, University of Illinois Department of Pediatrics, Chicago, Illinois

Fluid overload in VLBW infants and induced hypertension in animals have been implicated in the etiology of ICH. We compared 51 clinical and biochemical variables from 34 VLBW infants with autopsy proved severe ICH, and 38 randomly selected control infants (C). No differences were found between the groups in:4 prenatal stress factors, mode of presentation and delivery, admission blood gases and diagnosis. ICH group had 2/3 symptoms of severe ICH by 29.8+20 (SD) hours. C infants had no symptoms, normal CSF and are develop-ing normally. B.Wt and G.A. were, ICH:0.93kg and 28Wks: C group 1.2kg and 30 Wks (p<0.05).0ther results, (x+) SE values) are in the Table. ICH infants were smaller 1.5 Wks here return and

LI	e labie.	ICH infa	nts were	smaller,	1.5 Wks less	mature more
	Apgar		Adm	BP	BP(Torr)	Fluidsm1/kg/d
	1'	5'	Temp.	Adm.	Day 1 Day	2 Day 1 Day 2
H	*	*	*	[	1	*   *
H	3.7±0.4	-			36.5±5.2 52	±5 115±10 149±10
U	5.2±0.3	7.1±3.5	96.6±1.4	40±2.2	37.9±1.6 40	±2 100±6 120±7

asphyxiated and colder on admission. (\*p<.05) They received slightby more fluid (N.S) on day 1. On day 2 however, they became more symptomatic, had elevated mean BP, and had received about 30ml/Kg. more fluids(\*pc.05). We feel that in VLBW infants, asphyxia sets up a stage for ICH.Vigorous resucitation, and rapid volume expansion trigger a hypertensive response precipitating the final episode of massive ICH.

1603 CEREBRAL PERFUSION PRESSURE(CPP) IN BIRTH ASPHYXIA(BA) AND ITS RELATION TO THE QUALITY OF OUTCOME.<u>Tonse N.K.</u> <u>Raju</u>, <u>Dharmapuri</u> Vidyasagar, <u>Usha</u> Doshi,University of Illinois, Department of Pediatrics, Chicago, Illinois.

In 10 normal infants and 14 BA infants ICP changes were studied by a noninvasive technique (Ladd Monitor). Systemic BP,mean ICP and CPP(Mean BP-ICP) were calculated during acute phase(3 days) and recovery (6 days) in BA infants. 4 infants died and had evidence of brain damage and 2 of the 10 survivors had cerebral palsy (CP) at 3 years. 8 remaining survivors were normal(26-46 mos) Clinical, ICP and CPP values from normals. (Gr.I) BA with normal long term out-come(Gr.II), BA with brain damage (Gr.III) are shown below: (values mean + SE) \*p<0.05, \*\*p<0.025, + p<0.001. CPP during acute phase

	B.Wt.	GA.	Apgar 5 Min.	Ad.pH	ICP	Admin. BP Torr	Least CPP Torr
Gr.I n=2	3.3±0.5	39±0.1	9 <sub>±</sub> 1		8.2±0.8		
Gr.II n=8	2.9 <sub>±</sub> 0.3	38±.3	4.4±0.8	7.18±0.05	13.5±1*5	44±2.5	31.3±2**2
Gr.III n=6	3.4±0.5	39±1.5	3 <sub>±</sub> 1	7.06±0.04	22.8±4.7	38.3 <u>4</u> 2.2	13.5 <u>+</u> 1.5

was low in Gr.II and improved during recovery. In Gr.III profound falls of CPP were due to elevation of ICP (cerebral edema) and fall of BP. In 12 infants who had accurate CPP measurements, 6 had CPP <30 Torr (mean-3 SD). 5 of these had brain damage and the other has amblyopia. None of 6 with >30 Torr CPP had brain damage. We conclude: A)Very low CPP in BA infants is associated with brain damage.(p=0.015,Fishers Ex.Test),B) CPP should be utilized in the management and to prognosticate the outcome.

PROSTAGLANDIN LEVELS IN THE CEREBROSPINAL FLUID OF 1604 INFANTS AND CHLOREN WITH FEVER. Frank Ritter, Richard Siegler, William Jubiz. Dept. of Ped. and Internal Medicine, University and V.A. Hospitals, Univ. of Utah Col. of Med. Salt Lake City, Utah. It has been proposed that central nervous system (CNS) prosta-

glandins (PG) may be mediators of fever. In some animal models, cerebrospinal fluid (CSF) PGE is elevated in pyrogen-induced fever; and the intraventricular injection of PGE results in a febrile response. To investigate the role of CNS Prostaglandins in fever, PGE and PGF2 $\sigma$  concentrations were measured by radio-immunoassay in the CSF of 16 pediatric patients who were febrile (average temp. 39.0°C). They were free of CNS disease and were on no prostaglandin synthetase inhibiting drugs. Their values were compared to those of 19 patients who were neither febrile or on prostaglandin synthetase inhibitors. The following results (mean + SEM) were obtained:

	Febrile	Afebrile
PGE (pg/ml)	47 + 15	39 <u>+</u> 19
PGF20 (pg/ml)	169 <u>+</u> 26	133 <u>+</u> 37

There were no significant differences in the CSF levels of either PGE or PGF20, between the febrile and the afebrile groups. These observations do not support the hypothesis that QNS prostaglandins mediate fever.

PROSTAGLANDIN LEVELS IN THE CEREBROSPINAL FLUID (CSF) 1605 OF CHILDREN WITH MENINGITIS. Frank Ritter, Richard Siegler, William Jubiz. Dept. of Ped. and Internal Medicine, University and V.A. Hospitals, Univ. of Utah Col. of Medicine, Salt Lake City, Utah.

Prostaglandins (PG) are synthesized in the Central Nervous System (CNS) during inflammatory processes. In order to study the relationship between meningitis and CNS PG production, PGE and PGF20 concentrations were measured by radioimmunoassay in the initial CSF sample (day 1) of 7 children with bacterial and 4 with aseptic meningitis. Follow-up measurements were also ob-tained at 3 and 12 days in the bacterial meningitis group. The following results (mean  $\pm$  SEM) were obtained.

TOTTOMING LCOUT	co (incert _ bri	i) were obcarin		
	Aseptic Mening	itis Bacte	erial Mening	itis
	(Day 1)	(Day 1)	(Day 3)	(Day 12)
PGE (pg/ml)	173 + 41	1683 + 229	257 + 116	80 + 37
PGF2 (pg/ml)	473 + 145	1333 + 232	530 <del>+</del> 160	125 + 33
The childre	n with bacteri	al meningitis	had signifi	cantly high-
er levels of bo	th PGE (p < 0.	.001) and F20	(p< 0.03)	than did
those with asep				
ed the clinical				
patient (not in				
the initially h	igh levels (p	(ml) of PGE(1	950) and PGF	20 (2100)
persisted at 3	days (PGE 225	), PGF20 2200)	and 12 days	(PGE 2300,
PGF2 A 2480).			•	

Thus, CSF PG measurements may be helpful in distinguishing between bacterial and aseptic meningitis, and may be useful in mon-itoring the clinical course of bacterial meningitis.