SIMULTANEOUS DETERMINATIONS OF BREATH CONCENTRATION 614 AND PULMONARY EXCRETION RATE OF HYDROGEN IN NEWBORNS.

**614** AND PULMONARY EXCRETION RATE OF HYDROGEN IN NEWBORNS. <u>Clinton R. Ostrander, Barrett E. Cowan, John A.</u> <u>Kerner, David K. Stevenson and John D. Johnson, Depts. of Peds.</u> <u>Stanford Univ. Sch. of Med., Stanford, CA, and Univ. of New</u> <u>Mexico School of Medicine, Albuquerque, New Mexico.</u> <u>Production of hydrogen (H2) was simultaneously estimated by</u> <u>both end-tidal breath sampling (ETH2) and by direct excretion</u> <u>rate determinations (VeH2), using a flow-through system with gas-</u> <u>tight head hood and reduction gas detector (res. = .010 ppm/2.5</u> <u>ml sample). Studies were performed on 8 normal formula-fed</u> <u>infants of various postnatal and gestational ages immediately</u> <u>after feeding with sampling at 30-minute intervals until the next</u> after feeding with sampling at 30-minute intervals until the next feeding (2-3 hrs). Linear regression analysis of the normalized data showed no significant elevation in  $H_2$  production over the first 1, 2, or 3 hrs after feeding. ETH<sub>2</sub> and VeH<sub>2</sub> for each infant were thus taken to be the respective means of the multiple separate determinations during the sampling period. For the infant studies with complete data (n=10), mean ETH<sub>2</sub> was  $31\pm24.9$  (S.D.) ppm (range 6.85 to 90.6 ppm); mean VeH<sub>2</sub> was  $1.47\pm.89$  (S.D.)ml/hr (range .18 to 2.96 ml/hr). The failure of feeding to elicit an increase in either ETH<sub>2</sub> or VeH<sub>2</sub> and the persistence of high values relative to adult levels may be contributed to by the continuous presence of carbohydrate substrate in the gut because of frequent feeding. This hypothesis is supported by data from one extended study (5 hrs) during which no increase in ETH<sub>2</sub> was noted after feeding, but a drop to 50% of the initial mean value occurred after 3 hours (r = .87).

## $615 \stackrel{\text{ACCELERATED DEGRADATION OF ESSENTIAL FATTY ACLOS}{\text{DEF}_{1257}, 10} \\ \frac{1}{1257} \\ \frac{$

State Univ. Sch. of Med., Hutzel Hospital, Depts. of Pediatrics, Detroit, MI.' We have previously demonstrated, in vitro (Ped. Res. 10:429, 1976) by measuring 02 consumption and the products of lipid peroxidation (thiobarbituric acid reactants or TBA and diene conjugates) that the oxidation of EFA (linoleic and linolenic a.) is accelerated by exposure to light (460 nm), 02 and photosensiti-zing agents (bilitubin or methylene blue). This report describes the phenomenon to also occur, in vivo, in premature infants on PT. MTHODS: Seven premature infants while on PT for treatment of jaundice were serially tested during the first 3 days of PT for the concentration of linoleic acid (218:2) and triene/tetra-ene ratio in the phospholipid and sterol ester fractions of the serum by as liquid chromatography. Serum TBA and diene conju-gates, were measured spectrophotometrically. RESULTS: During PT, there was a conthnuous fall in Cl8:2 concentration both in the phospholipid (-0.19010.252 mg/dl/h) and sterol (-0.37210.580 mg/dl/h) fractions of their serum and the decay was greatest during the first 24 h of PT (-0.259 mg/dl/h) in phospholipid and -0.543 mg/dl/h for sterol. The rate of decay of Cl8:2 was also a simultaneous rise in triene/tetratene ratio (14 to 0.066, but none achieved critical EFA deficiency of a products of lind peroxidation in The was decised for the serum of the phospholipid and the decay before was also a simultaneous rise in triene/tetratene ratio furing PT, there was also a simultaneous rise in triene/tetratene ratio furing PT, there was also a first of 0.4 to 0.366, but none achieved critical EFA deficiency of di-ene conjugates + 0.013 00 mg Cl3 act, tone of the first of the cond there was also for the peroxidation at The theology of the infants for event atio level of 0.4 to 0.366, but none achieved critical EFA deficiency for the decay. Furthermore, the decay be the ending of the infants of EFA decay. Furthermore, the decay be the ending of the infants meyer tor the ending of the ending of the

616 SERUM LEVELS OF ANTIOXIDANTS IN BREASTFED VS BOTTLE-FED INFANTS: A POSSIBLE ROLE OF BREAST MILK IN PRO-TECTING INFANTS ACAINST OXYGEN TOXICITY. Enrique M. Ostrea. Jr., James E. Balun, and Ruth Minkler. Wayne State Univ. Sch. of Med., Hutzel Rospital, Depts. of Pediatrics, Detroit, MI.

Pediatrics, Bette univ. Sch. of Med., Mutzer Mospital, Depts. of Pediatrics, Detroit, M.
Vitamin E (E) and beta-carotene (C) are 2 naturally occurring antioxidants and we have previously shown that C is 100x more potent an antioxidant than E (Ped. Res. 14;1013; 1980). In this study, we measured spectrophotometrically, the levels of E (mgs/dl) and C (ug/dl) in paired cord blood samples and maternal sera, in human breast milk and formula, and in 4 breast and 4 bottle-fed infants. RESULTS: There is a direct correlation (re0.44, p<0.01) between maternal serum and cord blood side (b, 940.01) we measured spectrophotometrically, the levels of C, but hot of E. Maternal serum is signif. (p:0000) higher than Cord blood in levels of C (12).2446.6 vs 05, 02940.030). Migher than Cord blood in levels of C (12).2446.6 vs 05, 02940.030). Migher than Cord blood in levels of C (12).2446.6 vs 05, 02940.030). Migher than compared to formula breast milk (particularly first day colostrum, has very high levels of C (25)4463 vs 4.240.5) and E (2.0411.1) vs 1.0340.060). Thus, breastfed infants show increasing serum conc. of C (cord=15.944, 4, pay 3=30.0111.1, pay 6=66.2429.00 and E (Cord=0.3010.06, hay 3=0.0610.29, pay 6=1.0610.471 to levels that almost approximate those of the adult. In bottle fed infants, there was a rise in the serum level of E (Cord=0.3010.06, Day 3=0.0610.29, pay 3=13.345.3). COM-GLUSION: We have demonstrated that (1) breastmilk, particularly the premature, is very rich in its content of antioxidants, particularly the premature, is very low compared to the adult. Meeveer, (3) breastfed infant satin levels of C and E that almost approach adult levels of C and E that almost approach adult levels of the adult. We therefore speculate the vent of a the serum level of the dault. The bottle fed infants, attricularly the premature, is very low compared to the adult. However, (3) breastfed infant satia in levels of C and E that almost approach adult levels of C and E that almost approach adult levels within 6 day

HUMAN MILK TRACE METALS - APPLICATION OF X-617 RAY FLUORESCENCE SPECTROMETRY TO QUANTITA-TION AND SCREENING. <u>Paul A. Palma, Richard M.</u> Caprioli, R. Rodney Howell. Univ. of Texas Medical School at Houston,

Departments of Pediatrics and Analytical Chemistry, Houston. X-ray fluorescence spectrometry (XRF) was examined as a tool for quantitative analysis of trace metals of biological importance in human milk. The validity of XRF was demonstrated by study of within day, day to day, and operator to operator variability with an accuracy of >97%. Standard curves were constructed for Ca, V, Cr, Mn, Fe, Co, Ni, Cu and Zn and were highly significant ( $r \ge 0.99$ ). Correlation of results to atomic absorption spectrophotometry was excellent.

Forty women provided 250 samples of human milk obtained from 2-559 days of lactation. Trace metals were segregated almost exclusively in the aqueous fraction of milk. Ca, the only major metal analyzed, was found in fairly constant concentration throughout lactation, 25-40 mg/dl. Fe, Cu, and Zn concentrations declined during lactation; this decline was most marked during the first 30 days. There was a 10-fold decrease in milk zinc concentrations by 6 months of lactation. The physiologic significance of these findings is unclear. It does not appear to be a dilutional effect but may reflect a change in maternal nutritional status, or, teleologically, a change in infant nutritional requirements based on improved assimilation of trace metals.

The advantages of XRF for quantitation and screening of trace elements in human milk are 1) accurate, simultaneous analysis of multiple elements from a single sample 2) ease of sample preparation which limits contamination risk 3) non-destructive analysis which permits repeated analysis of a single sample.

**618** EFFECTS OF DIETARY THERAPY ON ETHANOL ELIMINATION IN **618** TYPE I GLYCOGEN STORAGE DISEASE. P.H. Parker, A. <u>Hoyumpa, H.L. Greene</u>, Departments of Pediatrics and Medicine, Vanderbilt University, Nashville, TN. Type I Glycogen Storage Disease (GSD-1) is associated with a wide variety of metabolic abnormalities including an extremely rapid elimination of ethanol. As dietary therapy aimed at main-taining normoglycemia corrects many of the metabolic abnormali-ties, the effect of this therapy on ethanol elimination was studied. We measured the half-life (T+min) and Clearance (C1 ml/ min) of 16ml/m<sup>2</sup> of intravenous ethanol in 3 groups of patients with GSD-1. Group I consisted of 3 patients who were treated; Group III con-sisted of 4 patients who were partially treated and were develop-Group II consisted of 3 patients who were treated; Group III consisted of 4 patients who were partially treated and were developing metabolic imbalances. Results: Group I showed T½ 11.79± 0.8 and Cl 1059±240; Group II showed T½ 28.3±1.5 and Cl 489±105; Group III showed T½ 20.6±2.3 and Cl 770±170; and the Control showed T½ 28.5±6.4. The T½ and Cl of ethanol was significantly shortened in Group I when compared with Group II patients (p < 0.05). The T½ and Cl in Group II patients was intermediate between group L and Group II patients.

tween group I and Group II patients. Conclusion: (1) The elimination of ethanol is increased in untreated GSD-I. (2) Treatment aimed at correcting metabolic imbalances results in a normal ethanol elimination in GSD-I, and cessation of therapy results in a return toward pretreatment values. (3) This suggests the rapid elimination of ethanol in GSD-I is not intrinsic to the disease but associated with the secondary metabolic derangements. (4) Evaluation of the elimi-nation of other drugs is indicated in patients with GSD-I.

SALICYLATES AND REYE'S SYNDROME. Jacqueline S. Partin, 619 John C. Partin, William K. Schubert and Jeanne Hammond School of Medicine, SUNY at Stony Brook, Department of Pediatrics and Children's Hospital Medical Center, Cincinnati. Serum salicylate levels ([ASA]) were obtained at admission from 172 cases of Reye's Syndrome (RS), 130 of whom were biopsy proven. Shown are [ASA] for biopsy proven cases, by clinical grade, and age matched, community matched, sick and well control children. 
 Mean
 12
 + SD
 7.7 mg/d1
 Range
 1-33

 13
 + SD
 15.7
 0-48

 11
 + SD
 10.2
 0-35
 Grade I (n=42)Grade II (n=13) Grade III (n=32) Grade IV (n=34) 13 + SD 10.7 Grade V (n=9) 13 + SD 11.7 Age matched controls collected 1978-80: 0-46 0-33  $\begin{array}{c} 1.6 + SD & 2.0 \\ 1.0 + SD & 0.5 \\ 0.9 + SD & 0.8 \end{array}$ 0.5-9.1 Varicella (n=17) URI (n=37) 0.3 - 4.6(n=79) 0.0-1.1 Well Liver biopsy ultrastructure of RS pts with high [ASA] (>20mg/d1) was not different from that of comparably ill pts with low [ASA] ( $\leq 2mg/dl$ ); both differed greatly from that of a non-RS pt with acute ASA intoxication (ASA 66 mg/dl). There is no correlation between clinical grade, mean [ASA] or outcome; but RS pts had

10x higher [ASA] than controls. We propose the higher [ASA] in RS pts are due to liver injury and reduced metabolism rather than the cause of the injury.