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METABOLISM OF GLUCOSE IN LOW BIRTH WEIGHT NEWBORN INFANTS IN THE FIRST HOURS OF LIFE. Van Asselt WA, Baarsma R, Chapman T, Reyngoud D-J, Okken A, Berger R. Div. Neonatology, Dept. Pediatrics, University of Groningen, Groningen, The Netherlands. Glucose metabolism was studied in low birth weight (LBW) infants before any oral feeding was given. We

have measured glucose utilization using a stable isotope dilution technique and calculated glucose oxidation from oxygen consumption and carbondioxide production measured with open circuit indirect calorimetry. Birth weight of the infants ranged from 1.29-2.26 kg, gestational age from 32-39 weeks. Results are shown in the table. serum glucose glucose ovidation glucose utilization

Intant	(hrs)	(mmo1/1)	(mg/kg/min)	(mg/kg/min)	
1	5	2.5	5.8	5.3	
2	3	3.8	5.8	5.1	
3	5	2.9	5.6	5.5	
4	14	3.4	3.9	5.9	
5	5	4.6	3.4	5.8	
6	8	4.6	5.7	8.1	
7	6	5.1	7.5	8.1	

The discrepancies between glucose utilization and glucose oxidation suggest that in some LBW infants substrates other than glucose are oxidized, and in some infants not all utilized glucose is oxidized.

> ENDOGENOUS GLUCOSE PRODUCTION IN SMALL FOR DATES NEWBORN INFANTS IN THE FIRST HOURS OF LIFE.

R Baarsma, T Chapman, WA van Asselt, D-J Reyngoud, A Okken. Div. Neonatology and Research Lab, Dept. Pediatrics, University of Groningen, Groningen, The Netherlands.

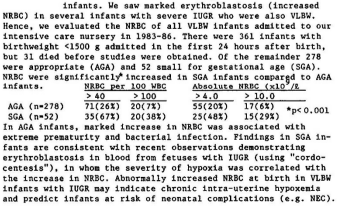
Most studies on endogenous glucose production (GPR) in Most studies on endogenous glucose production (GPR) in newborn infants, reported so far, have been carried out beyond the first day of life. Moreover, most infants had been fed orally. This study was performed to measure GPR in small for dates newborn infants in the first hours of life, using the prime dose-constant rate infusion technique with 6.6-dideuteroglucose. During the study the glucose levels remained within the normal range. Results:

Birthweight	Gest.Age	Postnatal	Glucose i.v.	GPR
(gm)	(wk)	Age(hours)	(mg/kg per	min)
1780	36	5	3.3	2.0
2255	38	3	3.0	2.1
2170	39	5	3.1	2.4
1870	36.3	14	2.7	3.2
1835	37.6	5	3.2	2.7

In conclusion: Near term and term small for dates newborn infants do not seem to have a limited endogenous GPR.

> ERYTHROBLASTOSIS AND FETAL GROWTH RETARDATION (IUGR): AGS Philip, AM Tito, Department of Pediatrics, Maine Medical Center, Portland, Maine, USA.

Little has been published about circulating nucleated red blood cells (NRBC) in very low birth weight (VLBW)



THE EFFECT OF SEVERE HYPOGLYCAEMIA AND DIA-BETES ON THE EEG IN DIABETIC CHILDREN Acsádi G., and Soltész G. University Department of Paediatrics

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Insulin hypoglycaemia(H)is the commonest form of H in childhood and severe and recurrent H can cause Institutional hypogrycaemia (h) is the commonest form of H in childhood and severe and recurrent H can cause brain damage. Serial EEG recordings were made in 70 diabetic children (DC) (age 11,2+0,5 yr, duration of diabetes (DD) 5,1+0,4 yr, mean+SE) and EEG findings were related to age (at EEG and at diagnosis), DD, daily insu-lin dose, long-term metabolic control assessed by HbAI and severe H episodes. EEG abnormalities (EEG-A) were found in 26 p.d.of DC. There was no relationship between EEG-A and DD, daily insulin dose or HbA1. DC with EEG-A were younger (9,6+0,7 vs. 12,8+0,7 yr, p 0,01), had an earlier onset of diabetes (4,8+0,6 vs. 7,6+0,6 yr) and 78 p.c.of them had severe antecedent H, whereas EEG-A were found in only 22 p.c.of DC with no H(p<0,001). All DC with H and convulsions had perma-nent EEG-A.The degree of metabolic control has no ef-fect on the EEG during the early years of D, but se-vere antecedent H, young age and early onset are important risk factors.

fatty acid oxidation in artificially fed infants during continuous infusion of  $^{13}\mathrm{C}\text{-trioctanoin}$  and  $^{13}\mathrm{C}\text{-}$ 87 TRIOLEIN, H. Paust, G. Knoblach, H. Brösicke, W. Park, T. Keles, H. Helge. Children's Hospital Freie Universität Berlin, FRG.

Previous investigations of MCT in infants showed a high energetic utilization of the given substrate after single injection of  $^{13}$ C-MCT. To what extent these patients can oxidize MCT during continuous application, especially in comparison to LCT, is still uncertain. Therefore we measured with the  $^{13}\mathrm{CO}_2-\mathrm{breath}$  test the oxidation of MCT and LCT in infants during continuous infusion of  $^{12}\mathrm{c}$ triglycerides. 9 infants, fed with supplementary parenteral nutrition, were investigated. The carbohydrate intake was 7.4-19.7 g/kg \*24h. For investigation the parenterally applied fat emulsion was substituted by an isocaloric amount of the testemulsion either enriched with  $^{13}\mathrm{C-trioctanoin}$  or  $^{13}\mathrm{C-triolein}$ . The  $^{13}\mathrm{C-content}$ , measured by mass-spectrometry, represents fatty acid oxidation. We found a higher MCT-oxidation compared with LCT: x=35.6±15.0% vs.  $x=18.6\pm8.3$ %. Both oxidation rates revealed a negative correlation to concomitant carbohydrate supply. The results show, that all patients can utilize parenterally applied MCT to a greater amount than LCT. MCT oxidation is also reduced by concomitant carbohydrate intake like LCT. On account of their high energetic utilization MCT in general are an adequate substrate for parenteral nutrition in infancy.

DETERMINANTS OF THE GROWTH OF BREAST-FED INFANTS IN NORTHERN THAILAND. DA Jackson, MW Woolridge, SM Imong, 88 A Silprasert, L Wongsawat, K Rungruengthanakit,

RF Drewett, K Amatayakul, & JD Baum. Res. Inst. for Health Sciences, Chiang Mai Univ, Chiang Mai, Thailand, & Inst. of Child Health, Univ of Bristol, Bristol, U.K. Predictors of growth were examined cross-sectionally in a random

sample of 65 infants under 1 year of age in a rural Thai community. Infants were studied for a continuous 48hr period in their own home; variables measured included: nutrient intake from breast milk (b.m.) and supplementary food (s.f.); morbidity (over the previous month); and total bacterial intake from s.f. and water

Overall age, sex & <u>birthweight</u> accounted for most variance (87%) in attained weight. Total <u>protein</u> intake (g/24hr, summed for b.m. & s.f.) predicted current weight, accounting for a further 1.3% of the variance in weight (p=.02), while total <u>calorie</u> intake (kcal/24hr) showed no association (p=.10). Morbidity did not predict attained weight at any stage of analysis (p=.61), but total bacterial intake was inversely related to weight (p=.01).

These findings differ prior to 6 mos of age when either protein or calories from b.m. predict infant weight (p=.004). After 6 mos total calorie intake no longer predicts weight, and the association with protein is now predominantly due to s.f. intake. This change is related to the observed increase in s.f. at around 6 mos of age, with a concomitant reduction in the number of breast feeds.