EFFECT OF DIETARY COPPER DEFICIENCY ON BRAIN CATECHOLAMINES C NELBOON DATE

Mercedes Lázaro, Ignacio Monreal, Ignacio Villa Elízaga Pediatric Research Laboratory. Faculty of Medi University of Navarra. Pamplona. Spain. 131 Medicine.

To investigate the effect of dietary copper deficiency in brain catecholamines of newborn rats an experimental model was developed. Three groups of 10 female Sprague - Davley rats were used. Group 1, copper deficient groups was fed with a copper-free diet 5 weeks pre-gestation and during gestation until delivery. Group 11, the control, received a copper addecuste diet (6pm). Group 11, the pairfed, received a copper addecuste diet, but restringed according to the ca-loric input of the copper deficient group. Inmediatily after birth, the newborn rats from a total of 7; 4 and 7 litters of the groups 1, 11 and 111, respecti-vely were decapited. Eleven assays of each group were performed. For each assay we used 1, 2, 3, or 4 whole brains mirus cerebella. Whole brain dopamine and noradrenaline levels were quantified by high performance liquid chromato-graphy with electrochemical detector. Copper deficient group showed a highly significant decrease in brain moradrenaline levels (50% lesser than others two groups: p = 0.0341), whereas dopamine concentrations remained largely unalte-red. The levels of brain copper in the newborn rats in group 1 were signifi-cantively reduced (p = 0.0001) as well as brain weight (p = 0.0012). The CA sector neuronal counting was unaltered, however the girus dentatus counting showed a significant decrease in neuronal cellularity in Group 1. We conclude that the materno-fetal copper deficiency altered the synthesis of noradrenali-ne. This could be a consequence of a functional impairment (possibly Dopamine - hidroxilase enzyme disminished activity), however the possible role of the hypocellularity as the one observed in the <u>girus dentatus</u> sector can not be excluded.

METHYLMALONATE AND PROPIONATE EFFECTS ON CO2 PRODUCTION AND LIPID BIOSYNTHESIS IN BRAIN OF SUCKLING RATS

132 Clóvis M. D. Wannmacher, Janice C. Dutra, Moacir Wajner, Silvia E. C. Cardoso and Eduardo R. Motta - Departamen-to de Bioquímica, Universidade Federal do Rio Grande do Sul, Porto Alegre, Brasil.

Methylmalonic acidaemia and propionic acidaemia are inherited metabolic disorders characterized respectively by methylmalonate (MMA) and propionate (PA) accumulation in tissues of affected patients. Both diseases have severe clinical expression and many undiagnosed patients have a fatal outcome. Those who survive presente neurological dysfunction and a variable degree of neuromotor delay/mental retardation. We investigated the in vitro effects of MMA and PA on CO<sub>2</sub> production and lipid biosynthesis in rat brain. Brain prisms of suckling animals were incubated in Krebs Ringer bicarbonate buffer in the presence of  $[2^{-14}C]$ -glucose and 5.0 mM MA or 5.0 mM PA. Controls did not contain the metabolites. After incubation,  $I^4CO_2$ was trapped into Whatman No. 3 paper filter, lipids were extracted from brain prisms and the radioactivity determined in a scintillation counter. We found that MMA but not PA inhibits signicantly CO<sub>2</sub> production from glucose. Lipid biosynthesis was also diminished Methylmalonic acidaemia and propionic acidaemia are inherited production from glucose. Lipid biosynthesis was also diminished by MMA but not by PA. These results indicate that MMA may impair brain energy metabolism and normal brain development.

133 CEREBELLAR INVOLVEMENT IN BILIRUBIN ENCEPHALOPATHY DUE TO CRIGLER-NAJJAR TYPE I DISEASE. Philippe Labruna\*, Catherine Narcy, Anne Myara\*\*, Jeanne Francoual\*\*, François Trivin\*\*, Michel Odièvre\*, \*Service de Pédiatrie, Hôpital Antoine Béclère and \*\*Laboratoire de Biochimie, 92141 Clamart, and \*\*Laboratoire de Biochimie, Hôpital St Joseph, 75014 Parie.

Patients suffering from Crigler-Najjar (CN) type I disease, a rare inborn error of bilirubin metabolism due to a deficiency in hepatic bilirubin uridine diphosphate glucuronosyltransferase (B UDPG-T) activity, are at permanent risk of developing neurotoxic effects of bilirubin (kernicterus). While cerebellar hypoplasia occurs in new born Gunn rats, the animal model of CN type I, cerebellar symptoms are not prominent in kernicterus observed in children.

kernicterus observed in children. <u>Cases reports</u>: 3 infants (2 girls and 1 boy) became jaundiced at 2 days. Despite treatment all of them underwent exchange-transfusions at the age of 9 days. Then, phototherapy had to be continued and phenobarchial was not efficient. At the age of 4 months, hepatic B UDPG-T was found to be nit in the 3 patients. At the age of 3 years, 4 years and 6 years, respectively, these children presented with cereballar symptoms a week after a reduction of phototherapy. Serum bilirubin concentrations at the time of admission were 581, 514 and 600µmoVL, respectively. Under correct treatment cerebellar symptoms slowly and partially decreased in the 3 children, while moderate mental retardation became obvious; none of them has any evidence of dealness and brain CT scan is corrul in 2 of them. scan is normal in 2 of them.

<u>Conclusions:</u> 1) These 3 cases illustrate a new clinical aspect of kernicterus, 2) kernicterus may have different clinical manifestations in infants and in older children, 3) the cerebellum may be the preferential target of bilirubin in children while it is not so in infants.



135

PROGRESSIVE ATAXIA AND MENTAL RETARDATION ASSOCIATED WITH LEQCODYSTROPHY IN 4 PATIENTS WITH L-2-HYDROXY-GLUTARIC ACIDURIA Hoffmann, G.F., Barth, P., Lehner, W., Duran, M., Hunneman, D.H., Jacken, J., Rating, D., Schutgens, R.B., Vos, W. and Hanefeld, F. Depts. Pediatrics Heidelberg, Freiburg, Göttingen, FRG, Amsterdam, Urrecht, The Netherlands, and Louvain, Belgium. We present the clinical, neuroradiological and biochemical data of 4 patients with L-2-hydroxy-glutaric aciduria, which appears to be a new neurometabolic disorder. Two male siblings of Maroceco ancestry presented with psychomotor retardation and dystrophy in early childhood. Consequently they developed a progressive ataxia, slight extrapyramidal signs and mental retardation. (Do 48 at the age of 18 y and 10=4 k at the age of 12 y). One of them suffered from convulsions. Two additional female patients (To, A., age 15y, and Fi., A., age 19) of Turkish and Greek ancestry followed a similar course with short staure, progressive ataxia and mental retardation. In addition, To, A., developed an oligoepilepsy with generalized scizures in early infancy and showed pyramidal signs with the age of 10 y. Neuroradiological investigations revealed distinct extensive subcortical hypotensities, resembling spongy degeneration and a (sub)cortical and cerebellar attophy in all four patients. Neuroradiological investigations revealed distinct extensive subcortical hypodensities, resembling spongy degeneration and a (sub)cortical and cerebellar attrophy in all four patients. Urinary exerction of L-2-hydroxyglutaric acid was in the order of 1-3 not/mot of crea. in all patients as determined by GCMS and HI-NMR spectroscopy (controls < 0.002 mol/not of crea.). Plasma levels of L-2-hydroxyglutaric acid ware 20-30 µmol/l and in the CSF 40-50 µmol/l (controls n. d.). The very similar clinical findings as well as the distinct neuroradiological picture in 4 patients from 3 families pinpoint a hitherto unrecognized neurometabolic disorder. The relation of L-2-hydroxyglutaric acid ware a small decrease in the urinary exerction of L-2-hydroxyglutaric acid under a carbohydrate rich dict and an increase after an oral load with plant oil. In To, A., a 24h fast produced a 4-fold increase in L-2-hydroxyglutaric aciduria. A number of other <u>in vivo</u> loading experiments and <u>in vitro</u> investigations in fibrohasts using radiobabelled precursors were unrewarding. investigations in fibroblasts using radiolabelled precursors were unrewarding.

> LONGTERM FOLLOW UP OF 16 CHILDREN AFFECTED WITH THE "LATE ONSET" VARIANT OF ARGININOSUCCINATE LYASE DEFICIENCY: NO IMPAIRMENT OF THE NEUROLOGICAL AND PSYCHOMOTORIC DEVELOPMENT.

Sonja Koch, Susanne Scheibenreiter, Elisabeth Knoll, Kurt Widhalm. Department of Pediatrics, University of Vienna, A-1090 Vienna, Währinger Gürtel 18-20, Austria.

At present, two variants of ASL-deficiency, the second most common enzyme defect of the urea cycle, have been described according to the mode of onset and the clinical course. Nearly all previous studies reported on an impaired neurological and intellectual development in affected children. Our study demonstrates for the first time, that these children can develop physically and mentally appropriate for age if treated with low protein diet and/or arginine supplementation.

Since 1975 16 Austrian children have been detected to be suspicious for this enzyme detect. After having confirmed the diagnosis, a daily arginine supplementation (3-4 mmoles/kg body weight/day) was initiated in all but three children in addition to normal diet respectively to recommended protein restriction of 1,2 - 1,5 g/kg body weight/day. Except for slight elevations, blood ammonia levels remained within the normal range over the time. EEG patterns showed paroxysmal abnormalities in 5 out of 16 children, whereas physical and mental development were within the normal range.

physical and mental development were within the normal range. It can be concluded, that early diagnosis of ASL-deliciency within a routine screening program followed by dietetic treatment (low protein and/or arginine substitution) seems to be able to prevent intercurrent hyperammonemic episodes, that cause severe neurological and psychomotoric abnormalities. Furthermore, an accumulation of argininosuccinic acid and its anhydrids does not seem to have a neurotoxic effect.

SEIZURES ASSOCIATED WITH TOXIC LEVELS OF CYCLOSPORINE A IN LIVER TRANSPLANTED CHILDREN.

Maria Roberta Cilio, Olivier Danhaive, Etienne Sokal, Jean-François Gadisseux, Jean-Bernard Otte, spn by 136 Jean-Paul Buts

Jean-Paul Buts Department of Pediatrics, Catholic University of Louvain St-Luc Hospital, Brussels In our series of 210 children having undergone 256 liver trans-plantations from 4/1984 to 12/1990, four patients presented with convulsions unrelated to a common etiology. Clinically they dis-played tonico-clonic seizures either generalised or lateralised with secondary generalisation. EEG showed either generalised or lateralised encephalopathy pattern with irritativity. None showed hypertensive encephalopathy on onhtalmic examination. None showed hypertensive encephalopathy on ophtalmic examination. None showed electrolyte, glucose or hepatic function abnormality. Magnesium electrolyte, glucose or hepatic function abnormality. Magnesium was normal. In two children cholesterol was normal (no data for others). LCR analysis showed no infection. In all four children Cyclosporine (CSA) through serum level reached supra-therapeutic levels, the "toxic" period lasting from 1 to 2 days and occuring from 2 to 10 days before scizures. CSA levels on specific RIA were normal or moderately perturbated. CT-scan in one patient showed subcortical and cortical diffuse hypodense lesions 3 days after scizures resolving aftervarde. Saizures recorded well to after seizures, resolving afterwards. Seizures responded well to antiepited rugs. No child showed permanent impairment. We conclude that seizures under cyclosporine therapy may be related with toxic levels of CSA and that discordance between specific and non specific assay suggests the role of metabolites.