SYMBIOTIC INTERACTIONS BETWEEN THE COLONIC MICROFLORA AND THE PROTEIN TURNOVER IN IN-89

MICROFLORA AND THE PROTEIN TURNOVER IN INFANTS

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The utilization of ¹⁵N-nitrogen from ¹⁵N labeled Bifidobacteria for whole body protein synthesis was studied in a total of 9 infants by means of oral or colonic pulse labelings. The microbes were harvested from synthetic culture mediums containing ¹⁵N enmionium chlorid as the only source of nitrogen. ¹⁵N enrichment of the cells amounted to 95 %. A tracer dose of 3 mg ¹⁵N/kg was chosen. The pulse labeling was followed by a 48 hours collection of urine and feces. Following oral single pulse labeling of the native and heat-treated yeast cells 77 % on an average of the heavy nitrogen was retained in the protein pool.

Comparatively high utilization rates of microbial ¹⁵N were found after instillation of ¹⁵N labeled Bifidobacteria into colonic segments in 3 infants with colostomies. The retention was confirmed by elevated ¹⁵N concentrations of the plasma proteins.

The results show that microbial nitrogen is intensively used for whole body protein synthesis in infants. The colonic microflora contributes to improvement of non essential endogenous and exogenous nitrogen. non essential endogenous and exogenous nitrogen.

THE IMMUNOLOGICAL RESPONSE TO GLIADIN IN COELIAC DISEASE SEEMS NOT TO BE OF PRIMARY PATHOGENETIC 90 IMPORTANCE.

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Patients with untreated coeliac disease have circulating antibodies against gliadin. We have previously demonstrated that the gliadin-antibody pattern is dominated by reactivity against a few polypeptides, tentatively identified as gamma-gliadins by their migration in electrophoresis. A varying degree of reactivity ty to alfa- and beta-gliadin was detected, but the sera did not react with glutenins. It was accordingly hypothesised that the reaction pattern was due to a specific primary reactivity against some gliadin polypeptides, with a secondary immunization with other polypeptides, initiated by epithelial damage and increased intestinal permeability.

Gliadin, digested with pepsin and trypsin (PT-gliadin), has been shown to be toxic. In order to further clucidate the connection between gliadin antigenecity and toxicity, the antibody reactivity in coeliac sera against PT-gliadin was investigated by immunoblotting. The enzymatic digestion of gliadin accomplished low molecular weight polypeptides, which notably did not display antibody reactivity. antibody reactivity.

It is therefore suggested that characteristic gliadin antibody pattern in patients with coeliac disease is not of primary pathogenetic importance, but instead correlated to common HLA-antigens of most of the patients.

CHANGES IN THE LIVER HBV-DNA PATTERN EXPRESSION IN CHILDREN WITH HBV CHRONIC INFECTION

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The antiviral effect of r-iFNa on HBV replication has been proven "in vivo". However there is no information about the changes induced by the r-iFNa in the patterns of HBV-DNA, specially in children. The aim of this work was to study the effect of the r-iFNa over the liver HBV-DNA patterns in children with chronic hepatitis B.

30 children with CAH histologically proved were included. All of them had HBeAg and serum HBV-DNA for at least 6 months prior to the beginning of the study. Two liver biopsies were obtained from all the patients, one just before the treatment and the second at the 15th month.

HBV-DNA was tested in serum by dot blot and in liver by southern blot hybridization in the first liver sample all children had replicative intermediates of the viral DNA and in one of them the HBV-DNA was also integrated in the host genome. In the second liver biopsy, in the children who did not respond to the therapy (HBV-DNA+ in serum) the replicative forms of the HBV-DNA remained in the liver and in 3 children inegrated HBV-DNA was detected simultaneously. Among the 9 patients who detectable in their liver, in the other child integrated and episomal forms non detectable in their liver, in the other child integrated and episomal forms non replicatives of the HBV-DNA were detected. None of the responder children lost the HBSAg in serum. To investigate this fact we looked for the presence of HBV-DNA in peripheral blood cells by dot-blot. None of these children showed viral DNA in these cells.

In conclusion, HBV-DNA can be integrated in the host genome early in the natural history of the HBV-chronic infection. Our results suggest the antiviral effect of the r-IFNa of hepatic level. The persistence of HBSAg in the responder children is not maintained at expenses of HBV-DNA expression in mononuclear blood cells.

PREDICTIVE FACTORS OF THE RESPONSE TO RECOMBINANT INTERFERON IN CHRONIC HEPATITIS B IN CHILDHOOD M.Ruiz-Moreno, V.Carreno, J.Jimenez, J.Bartolome, A.Morena and J.C.Porres

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The antiviral effect of recombinant interferon (rIFN) therapy of chronic hepatitis type B in children has been demonstrated. In order to determine the predictive factors of the response to rIFN, the basal features of responder and non-responder patients (age, sex, ALT levels, HBV-DNAp activity, time of HBsAg carriers, Knodell's index and percentage of infected cells of HBcAg in the liver biopsy) have been evaluated. Twenty eight children day age 2-14 years) with viral replication markers (HBV-DNA positive) have been included. Eight children had received 10 MU of rIFN-a2X (Roberon)/m body surface, I.N., twice a week during six months; 8 children were treated with 7.5 MU under the same conditions and 12 received 10 MU of rIFN-a2A (Roberon)/m body surface, I.N., thrice weekly during 3 months. All of them had an histologically proven CAH. All the end of the follow-up period (15 months) 8 children became HBV-DNA and HBeAg negative and were considered as rIFN-responders and 20 remained positive for these markers (non-responders).

The responders patients had a significantly lower activity of HBV-DNAp (265.7± 323.6 vs 719.3± 480.3. p(0.05) and less percentage of HBcAg infected liver cells than the non-responders (14.17 ± 6.83 vs 52.8± ± 32.73 p(0.05). In addition, the ALT (210.5± 75.2 vs 128.8± 76.1, p(0.05) and the liver Knodell's index of histological activity in the liver biopsies (12.25± 2,41 vs 7.64 ± 3.23 p(0.05) were higher in the responders than in the non-responders.

In conclusion, the children who responded to rIFN therapy had a more active liver disease (ALT, Knodell's index). Furthermore these patients had a relatively low level of viral replication (HBV-DNAp, HBcAg in liver cells).

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PANCREATIC FUNCTION IN ESSENTIAL FATTY ACID DEFICIENT (EFAD) RATS 93 Ahren B; Andrén-Sandberg Å; Böttcher G*; Hjeite L, An Strandvik B.

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Patients with cystic fibrosis (CF) have EFAD which might contribute to their symptoms (Scand J Gastroenterol 1988; 23 (Suppl 143):1-4). About 90% of the patients have pancreatic insufficiency and 5-10% develop diabetes mellitus. Pancrease has been shown to be highly develop diabetes mightures. Particles as been shown to be nighty sensitive to EFAD as revealed by the triene-tetraene ratio. To study the influence of EFAD on pancreatic endocrine function the insuin response was studied after administration of glucose intravenously in infusion or bolus and in addition the glucagon response after arginine infusion in EFAD female rats (120 days old) and agematched controls. The exocrine function was studied in isolated pancreatic acinar cells after stimulation with alcohol and carbacholine chloride. The amount of secreted amylase was assessed colorimetrically in computarized spectrophetrator (carbichaterial) rimetrically in computerized spectrophotometer. Gastrointestinal peptides were studied by immunochemical methods.

The EFAD rats showed a higher insulin secretion than control rats

with all stimulations whereas no differences in glucose and gluca-gon levels were seen. The isolated pancreatic acinar cells showed a normal amylase secretion in EFAD rats. Morphologically no changes were seen in the pancrease and the immunohistochemistry pattern of insulin-, glucagon-, somastotatin- and pancreatic polypeptide cells was not different from the controls. The results of this study indicate that the endocrine pancrease seems more sensitive to EFAD than the avercine pant than the exocrine part.

UNCOUPLING OF BILIARY LIPID SECRETION FROM BILE ACID SECRETION BY ORGANIC ANIONS, DUE TO INTRACANALICULAR INTERACTION WITH BILE ACIDS.
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Biliary secretion of phospholipids (PL) and cholesterol (CH) is regulated by bile acids (BA). However, a number of organic anions (OA) has been shown to inhibit PL and CH secretion without affecting that of BA. The mechanism of this OA-effect is unclear. We studied this uncoupling phenomenon with 3 different OA in normal Wistar (NW) rats and Croningen Yellow (CY) Wistar rats. The CY strain has a genetic defect in biliary secretions of various OA (JCI 81: 1593-9,1988). NW and CY rats with 8-day bile diversion were injected intravenously with ampicillin (18 mol.100 g BW), sufated taurolithocholic acid (STLC, 1.0 umol/100gBW) or idocyanine green (ICG, 0.6 umol/100gBW). At 1 hr after injection recoveries in bile were ampicillin, 4.1% and 0.5%; STLC 98% and 32%; ICG, 39% and 9%, in NW and CY rats, respectively. Ampicillin and STLC caused a strong uncoupling in NW rats (maximal BA.(PH-tH) -ratiof-70% (ampicillin) and +147%(STLC), but no (ampicillin) or a much smaller (STLC +65%) uncoupling in CY rats. ICG injections did not induce an uncoupling, either in NW rats or in CY rats. The hetatic uptake of the used OA appeared to be unaffected in GY rats. Cel filtration chromatography (Sepharose CL-4B) showed that ampicillin and STLC coeluted with BA, while ICC coeluted with the PL and CH fraction. We conclude that the uncoupling of biliary PL and CH from BA secretion by ampicillin and STLC is not due to disturbance of processes at intracellular or bile canalicular membrane level, but to interactions with BA inside the canalicular lumen which disturb PL and CH solubilization.

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