Cerebroprotective effect of dexamethasone (DXM) in newborn piglets with experimental pneumothorax (EPT). P. TEMESVARI, F. JOO, M.KOLTAI, E.ECK, G.ADAM and 29 P. These with experimental preumothoray (BPT). P. These Natt, F. JOO, M.KOLTAT, E.EXK, G.ADAM and D. BODA. Pediatric Dett., thiv. Med. School, Szeged, Hungary. The effect of DMM treatment in meshorn piglets with ET was studied on 30 piglets. 20 animals were subjected to ET (group A and B) and 10 piglets served as controls without ET (group C). In the ET animals 10 piglets were injected with 5 mg/bake go DMM 4 hours prior to ET (A). Each animal was paired with untreated sibling having the same ETT (B). In the ET piglets 4 hours after the critical phase (as apnea appeared, MMEP fell, the ET was terminated) brain tissue water and Evans blue dye were determined. There was a highly significant difference between A and B piglets regarding the time necessary to the development of critical phase (A X-69, GaDe27.5 vs B 28, 7²10.1 minutes; p < 0.001). In the untreated ET animals (B) the Evans blue content of the parietal ortex and cerebellum exceeded significantly the values obtained in DMM treated (A) and control piglets (C) -parietal cortex A 0.18²0.12, B.088²0.57, C 0.17²0.07; cerebellum A0.14²0.02, B 0.78²0.36, C 0.17²0.07 µg/dye/g wet tissue. The same phenomenon was observed regarding the water content of above train regions, as well. In conclusion, DMM treatment increases the tolerance to hypoda and prevents brain cedema in mestorn piglets with ET. with EPT.

30 T-CELL DEFICIENCY IN AN AFRICAN FAMILY. T. Jonckheer¹, P. Van de Perre¹, P. Lepage², <u>H. Taelman², C. Muller¹, N. Clumeck¹, T. Dab¹</u>. Brussels. 2. Hôpital Kigali Rwanda. 3. Tropical Insti-

tute Antwerp. We report the case of a 19-month old African boy born from parents with clinical and immunological features suggesting prodromal acquired immunodeficiency syndrome suggesting prodromal acquired immunodeficiency syndrome (AIDS). He showed early failure to thrive, generalized lymphadenopathy, bilateral parotitis. Later he develop-ped persistent interstitial pneumonia, persistent oral thrush and mild hepatosplenomegaly. Immunological stu-dies showed hyper IgG level (3.2 g/d1), reversed T-hel-per/T-suppressor (T4/T8) ratio (19/68) and T-cell defect (cutaneous anergy for different antigens, depressed in vitro response for concanavalin, phytohaemagglutinin and Pokeweed mitogens). Open lung biopsy showed inters-titial nodular lymphocytic infiltrates on optic micro-scopy and cytoplasmic particles on electron microscopy which are of uncertain origin. Recently the two older which are of uncertain origin. Recently the two older brothers were investigated. The oldest (6 years) sho-wed chronic parotitis, hyper IgG level (4.7 g/d1), re-versed T4/T8 ratio (18/49) and depressed in vitro res-ponse to mitogens. The second (5 years) showed slight parotitis, reversed T4/T8 ratio (29/32) and depressed in vitro response to mitogens. Both had normal chest X-rays The clinical and immunological presentation of the three children resemble that of infants with AIDS-like syn-drome described by others in the USA. We suggest speci-fic modes of transmission could be involved in familial acquiring of AIDS (transplacental-route, breast feeding, or other routine close-contact).

INFLUENCE OF GROWTH HORMONE (GH) TREATMENT ON 31 INFLIEXCE LE GAWH HONDRE (GH) THEATMANT ON INSULIN SINGITIVITY AND INSULIN RECEPTOR BIN-DING IN PATIENTS WITH GH-DEFICIENCY. H.Frisch⁺, R.Frager⁺, E.Schober⁺, G.Schernthaner⁺ (In-trod. by W.SNGBOD). Pediatric Dept. of Medi-cine II, Univ.Vienna and L.Boltzmann Institute for Endocrinology, Vienna, Austria.

In patients with GH-deficiency various disorders of carbohydrate metabolism have been reported. The aim of the study was to examine the influence of GH on inof the study was to examine the influence of GH on in-sulin sensitivity and insulin receptor binding by means of euglycemic CLAMP technique and radioreceptor assay using mononuclear leucocytes as target cells. 5 pa-tients with hypopituitarism were studied during GH therapy (phase A) and after withdrawal of treatment for 4 weeks (phase B). Informed consent from the pa-tients and their parents to partizipate in the study was obtained. Results: 1.) Basal insulin levels were not different in phase A and B. 2.) Peripheral glucose utilisation was significantly elevated in phase B com-pared to phase A (6.3 vs. 3.9mg/kg BM·min, p<0.05). 3.) Insulin receptor binding to monocytes remained unchanged after withdrawal of GH-therapy (specific binding fraction 3.18% vs. 3.66%, n.s.). These data indicate that the influence of GH substitution on pe-ripheral glucose metabolism is most likely due to postreceptor mechanisms. postreceptor mechanisms.

Surfactant and sudden infant death (SIDS) MORLEY CJ, HILL CM*, BROWN BD*, BARSON AJ*, SOUTHALL D*, DAVIS JA. Dept of Paediatrics Addenbrooke's Hospital, Cambridge. Dept Pathology St Marys Hospital, Manchester, Cardiothoracic Unit, Brompton Hospital. London.

Hospital, London. To detect abnormalities of pulmonary surfactant in SIDS surfactant was lavaged at autopsy from the lungs of 59 infants who died from SIDS, analysed for its phospho-lipid composition and compared with the composition of surfactant from 39 babies dying from HMD,8 newborn with normal lungs, 9 infants with normal lungs, 8 tracheal aspirates from living adults and 15 from living infants. The percentage phosphatidylcholine (PC) was reduced in SIDS and sphingomyelin(SM) increased as shown.mean+sem. SPC (p)

| | | \$PC (p) | | | | SM | (1 | p) | | | | |
|---|---------------------|--|-------------------------|----------------|-------------------------------------|------------------------------------|----------------|-------------------------|-----------------|-------------------|---------------------------------|---------------|
| SIDS | 59 | 60 + 1 | | | | 12 | + | 1 | | | | |
| HMD | 39 | 63 + 2 | ns | 3 | | 9 | + | 1 | P | < | .05 | |
| Newborn | 8 | 78 + 2 | P | < | .01 | 2 | ± | 1 | P | < | .01 | |
| Dead infants | 9 | 70 + 2 | P | < | .01 | 6 | + | 1 | P | < | .01 | |
| Live infants | 15 | 70 + 3 | P | < | .01 | 3 | + | 1 | P | < | .01 | |
| Live adults | 8 | 76 + 4 | P | < | .01 | 1 | + | 1 | P | < | .01 | |
| For the indiv parameters. differentiato | idua Howa | al group ever the The 90th | ra | at. | ere is io of tile F | PC:SM PC:SM | o i: rat | s a tio | lap re fe | eat | sonabl SIDS | se e is |
| 10. At this 1 infants, 20% In conclusion whether patho | evel live the | l there e infant e change ical or | are s, s i det | 1 In rei | 22% ne 3% liv surfa lopmer | wborn re adu actant atal, | is | 22% s a omp su | nd os ch | f 6 it a | dead 7\$ HMI ion const |). ant |
| feature of SI component of | DS the | that it mechani | is sm | 1 t | ikely hat co | to be ntrib | auto | fu | to | t | ental he | |

NUTRITIONAL IRON DEFICIENCY (NID) DELAYS THE 33 MATURATION OF INTESTINAL ABSORTIUE FUNCTIONS BY ALTERATION OF SURFACE MEMBRANE GLYCOPROTEIN SYNTHESIS. J.P. Buts, D. Delacroix, F. De Craeker, N. De Keyseë and R. De Meyer. Dept. of Ped., Univ. of Louvain, Brussels, Belgium.

De Keysef and R. De Meyef. Dept. of Ped., Univ. of Louvain, Brussels, Belgium. In children and in growing rats, NID produces intestinal malabsorption of carbohydrates by decreasing the activi-ty of jejunal disaccharidases. Crypt cell surface membra-ne glycoproteins such as the secretory component (SC) of immunoglobulins are also depressed. To clarify, the mechanism(s) by which NID affects these intestinal func-tions, suckling rats made iron deficient in utero were studied at day 12 after birth. SDS-polyacrylamide gel electrophoresis of purified jejunal brush border showed that the protein band corresponding to lactase was virtu-ally absent as was the incorporation of D-[1Cl⁴]-gluco-samine into lactase protein (3315 + 302 controls. vs 140 + 10 cpm.mg prot.⁻¹, NID, $p \downarrow 001$). In a second experiment, rats with NID were studied after weaning (day 28). Measuring SC concentration into subcellular fractions of isolated jejunal cells, we found similar. differences of SC content in the cytosol fraction (1.2 + 0.3 vs 5.65 + 1.2 µg.g.tissue⁻¹, p(0.05) and in the brush border (0.7 + 0.1 vs 3.58 + 0.6 µg.g.tissue⁻¹, p(0.05) between iron deficient rats and controls. synthesis of intestinal brush border glycoproteins rather than the intracellular transport or the final membrane assembly of these proteins.

34 Rotavirus infection in hospitalized newborn infants. U.B. SCHAAD* and R. ZBINDEN*. Dept. of Pediatrics, Univ. of Berne, Switzerland. Rotavirus (RV) gastroenteritis is found in 40% of our hospitalized pediatric pat. with acute diarrhea. This 3-year experience favorably compares to that of other cen-tres in temperate climates. Nosocomial spread (fecal-oral, resp. droplets) of RV infection among pediatric pat. and medical personnel is well documented and is re-sponsible for 7 % of our RV gastroenteritis cases. Recen-tions in the community have been reported from Sidney, Melbourne, London and Washington: fecal RV excretion was found in 30-50% and remained asymptomatic in 70-92%. Since April 1983 we conduct a prospective surveillance study for RV infection in our referral 6-bed intensive and 8-bed special care nurseries. From each pat. fecal and 8-bed special care nurseries. From each pat. fecal specimens on admission plus 3x weekly are examined for RV by ELISA technique (Rotazyme). From April to Sept. 1983 we observed RV in the stools of 31 (17.1%) of 181 1983 we observed RV in the stools of 31 (17.1%) of 181 neonates. In the majority of cases RV was detected be-tween the 1st and 5th day of life and RV excretion lasted from 1 to 3 days only. Careful analysis of clinical and laboratory data revealed that all 31 neonatal RV infec-tions were totally asymptomatic. During the first 6 study months recovery rates remained constant, but there was no community outbreak. Preliminary data during our RV season (winter months) indicate increased incidence also in recovers the bit of the sumptome on given season (winter months) indicate increased incidence aix in neonates, but clearly RV related symptoms or signs were never detected. In our experience neonatal RV in-fection is rather common but of only short duration and extremely benign. It is suggested that both noscomial spread and immunologic protection might explain these observations.