GLUTATHIONE RECOVERY STUDIES DURING OXIDATIVE STRESS IN NEONATAL RED BLOOD CELLS.



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Red blood cell (RBC) peroxide catabolism, via the synergistic (glutathione peroxidase and the glutathione recycling system (glutathione peroxidase and reductase), helps protect the lung against oxygen toxicity (Am Rev Resp Dis 1989;140:531). Using serial changes in reduced (CSH) and oxidized (CSSG) glutathione as a marker, the ability of RBCs to deal with a hydrogen peroxide (H_2O_2) load was compared in vitro in preterm (n=8) and term (n=9)the set of the set of depletion produced a slower GSH recovery time (r=-0.79, $\rho{<}0.001$). Neonatal RBCs showed significantly less depletion and quicker recovery of GSH than those of adults (p<0.001). Partial inhibition of H₂O₂ catabolism by catalase inactivation produced 50% loss of intracellular glutathione and slower GSH recovery (p<0.005) in all subjects, but recovery remained quicker in the babies (p<0.01). There was a positive correlation between gestational age and recovery time (r=0.68, p<0.02). The effective peroxide catabolism in neonatal RBCs may partly compensate for deficiencies in antioxidant defenses of the immature lung.

PLATELET ACTIVATION IN THALASSEMIC CHILDREN.



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Thromboembolic events, which are associated with significant morbi dity and mortality, occur in eta thalassemia major patients. Eldor et al. (Am. J. Hemat. 32:94, 1989) reported findings of increased circulating platelet aggregates and short platelet lifespan, suggesting platelet activation. We studied the expression of the platelet selectin CMP-140 on intact cells from thalassemic patients, as a mar ker of platelet activation.Blood was collected in glutaraldehyde solution from 10 children and from 10 health adult donors. Platelets were isolated and the expression of GMP-140 was measured by flow-cytometry, using the monoclonal antibody CLB-Thromb./6. The mean of positive cells was 18 ± 6 vs 5 ± 2 (Wilcoxon test:p <0.01) Our study indicates that in fact platelets are activated in vivo in children affected with thalassemia major.



1073 TUMOR NECROSIS FACTOR alpha(INF-alpha) (NTH1UVIENILE RILEUMATOID ARTINKITIS (JRA) Jaces J. Pietrzy, Wojciech Uracz, Barbara Hajto, Teresa Marek-Szydlowska, Marek Zembala Ist Dept. of Pediatrics, Dept. of Clinical Immunology, Institute of Pediatrics, Medical Academy, Kraków, Poland of Istanov, Marking Marking Marking Marking Marking Children and Io Infection-free matched controls were entolled to the study. The TNF level in the sera was determined by ELISA test. Spontaneous (NIL) and induced production of TNF was assessed. To induce TNF production, patients Mø were stimulated with LPS and fibroblasts (from healthy donor and a selected child with JRA). The analyses were performed at AS and LR. For statistical evaluation non parametric test was used. Results: Lower TNF levels in the sera of JRA patients at AS in comparison to LR was observed (z=-1.491 p = .07). Mø of AS patients revealed significantly lower (p= 0.01), and LR children significantly higher (p= 0.0002) NL production in comparison to the controls. A similar patiern was observed for Mø after LPS simulation. The production of 'TNF by Mø of patients (AS) stimulated by JRA fibroblasts was significantly higher (233 U/ml vs 50 U/ml) (z=-2.273 p=_001) in comparison to the situation when the fibroblasts from a healthy donor were used as stimulations. No such the fibroblasts from a healthy donor were used as stimulation with fibroblasts from a healthy donor were used as timulation with fibroblasts from ARA patient indicate the possibility that Mø-fibroblasts interaction may participate in the pathomechanism of JRA.

INFECTIOUS DISEASES

E.COLI ENDOTOXIN (LPS) GIVEN IN INTRAVENOUS INFUSION RESULTS IN BLOOD-BRAIN BARRIER (BBB) OPENING FOR NA-FLUORESCEIN (NaF) IN NEWBORN PIGLETS 110

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Cerebral complications like brain edema, bleeding, thrombosis, etc. are very frequent in the course of neonatal bacterial infections. We investigated in vivo the reactions of pial vessels by fluorescence macroscope giving LPS (E.coli O 111 B 4) to newborn piglets in pial vessels by fluorescence macroscope giving LPS (E.coli O 111 B 4) to newborn piglets in intravenous infusion in doses 0.1 µg/kg bw./h (<u>Group I</u>], n=6) and 1.0 µg/kg bw./h (<u>Group I</u>], n=6) through 4 hours. 6 animals were given 0.9% NaCl, and served as contools (<u>Group II</u>). The physiological parameters (HR, MABP, CVP, body- temperature, blood gases and acid-base state) were monitorized continuously. Using 1% NaF as BBB permeability tracer extravastion was observed in Group I (128.3±27.7 min after the start of infusion), and in Group II (177.3±28.3 min, N.S. vs. Group I). Brain NaF uptake was higher in Group I (2.1±0.4 µgNaFxmg³ protein/hgNaFxµl³ serum), than in Group II (1.4±0.5 µgNaFxmg³ protein/hgNaFxl³ serum, N.S. vs. Group I) and was non-detectable in Group III. Continuous administration of both doses of LPS produced hyperdynamic state with increased cardiac output (180% of baseline) and significantly decreased lung-thorax compliance (80% of initial value) at the time of BBB-opening. During endotoxin infusion WBC-counts in sera of treated groups elevated gradually: they reached an approximately 2-fold increase when NaF extravasation occurred, and an almost 7-fold increase 2 hours after the end of the infusions. At this time a moderate metabolic acidosis and pleocylosis was found in Group I (1800±431 cells/µl CSF) and in Group II (345±79 cells/µl CSF, pc:0.01). All the parameters studied were within normal range in Group III. It is concluded, that LPS given in similar doses as found frequently in plasma from septic newborns opens the BBB for NaF in piglets. These brain microcirculatory disturbances were accompanied with significant leucocytosis and pleocytosis augmented them. (All values are meantSE.)

> NEUTROPHIL ELASTASE IN DIAGNOSIS OF NEONATAL INFECTION. Alistair G.S. Philip, Christian P. Speer and Leon Sann.

Alistair G.S. Philip, Christian P. Speer and Leon Sann. Pediatrics Depts, Maine Medical Center, Portland, USA, Göttingen Univ, FRC and Höpital Debrousse, Lyon, France. Elastase (5) released from neutrophile during phago-cytosis is rapidly bound and inactivated by gi-protein-ase inhibitor. As previously shown, the complex is a sensitive and rapidly responsive indicator of neonatal aepsis using a time-consuming ELISA method (J. Pediatr. 1986, 108:987). In this 3 center prospective study we measured E with a rapid concer (1940) Consuming ELISA method (J. Pediatr. 1986, 108:987). In this 3 center prospective study we measured E with a rapid assay (IMAC-Elastade, Merck; 15 min) and compared it with immature/total neutrophils (I/T) and C-reactive protein (CRP) in infants with suspected inflection. Normal IMAC-E values (n=319) were obtained from 125 controls (upper limits: day 0-2, 130 µg/1; day 3-5, 95 µg/1; day 6-28, 65 µg/1). An additional 252 neonates of di-verse birth weights and gestational ages were evaluated for in-fection. Service upper dis 10 gestational ages were evaluated for infection. Sepsis was proved in 10 and pneumonia (positive tracheal aspirate culture and x-ray) in 23.

				E+	E+	I/T+	E+ I/T
	Е	I/T	CRP	I/T	CRP	CRP	+CRP
Sensitivity#	75	52	52	43	44	23	23
Specificity	62	82	92	93	95	98	100
Pos. Pred. Value	19	25	44	41	52	54	88
Neg. Pred. Value	96	94	94	94	94	92	92
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Values derived from infected (sepsis/pneumonia) vs. non-infected IMAC-E is a useful adjunct in diagnosing meonatal infection, but combining E with I/T and/or CRP markedly increases PPV.

ELEVATION OF PROSTAGLANDIN LEVELS IN PREGNANCIES COM-PLICATED BY PREMATURE RUPTURE OF THE MEMBRANES (PROM)

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Paired maternal and fetal prostaglandin levels were estimated in patients with pregnancies complicated by premature rupture of the membranes (PROM). Their results were compared to those of controls who had pregnancies with intact membranes. Fetal samples were obtained by cordocentesis, no patient was in labour at the time of cordocentesis. No control pregnancy was complicated by oligohydramnios or infection and none of the fetuses had renal disease. Prostaglandin levels were assessed by estimation of PGEM levels. Nine patients with PROM were recruited, cordocentesis was performed at a median of 4 days following PROM and at a median of 28 weeks gestational age. 12 controls were recruited, median ges-To were selectional age. 12 controls were recruited, median ges-tational age at cordocentesis 27 weeks. Maternal PGEM levels were higher in the PROM patients (mean 348 pg/ml) than the controls (mean 262 pg/ml), (95% confidence intervals 2.0 to 172), p<0.05. Fetal PGEM levels were also higher in the PROM patients (mean 349 pg/ml) than the controls (mean 216 pg/ml), (95% confidence interv-pale 41 to 224). vals 41 to 224), p<0.01. We conclude prostaglandin levels are elevated in pregnancies complicated by premature rupture of the membranes.