BONE MARROW TRANSPLANTATION IN CONGENITAL IMMUNODEFICENCY STATES - ULM EXPERIENCE

101

Wolfgang Hartmann and Wilhelm Friedrich spn. by Enno Kleihauer - Department of Pediatrics, University of Ulm, D7900 Ulm, Germany

Between 1983 and 1990 we treated 53 patients for congenital combined immunodeficency by bone marrow transplantation. 50 patients had SCID including 5 with ADA deficency and 3 with Omenn syndrome, 2 were treated for bare lymphocyte syndrome, two siblings suffered from a functional T-cell defect. The overall survival was 67% with a mean follow up of 4.2 years. After HLA - identical transplantation 7/9 (77.7%) and after haploidentical transplantation 29/45 (64.4%) survived. After haploidentical transplantation with T - cell depletion and without pretransplant conditioning, GvHD occured in 1/23, after pretreatment with ATG in 1/5 and after conditioning with busulfan and cyclophosphamide in 7/17 pts.. There was a slightly higher early mortality in patients with conditioning, but no difference in long time survival. However conditioned patients had a benefit in respect to B - cell reconstitution. The occurence of maternally derived lymphocytes (40% of pts.) had no influence on survival. The outcome of all these patients seems mainly influenced by their clinical state prior to BMT.

102

BONE MARROW TRANSPLANTATION FOR CHRONIC GRANULOMATOUS DISEASE X/bj. M.M. Manzionna, P. Di Bartolomeo +, B. Di Girolamo +, B. De Mattia, G. Torlontano +, F. Schettini.

Dipartimento di Biomedicina dell'Età Evolutiva - Clinica Pediatrica I Università di Bari and + Divisione di Emalologia, Cattedra di Ematologia, Ospedale Civile di Pescara, Università di Chieti - Italy.

An B year old boy affected by a rare form of X-linked Chronic Granulomatous Disease (CGD) and present cytochrome b underwent successful bone marrow transplantation on August 1989 in Personal 3.4 x10° nucleated cells/Kg) from his 6 year old histocompatible sister [HLA-identical and HLC negative) after a preconditioning regimen of Bussifan (3.25 mg/kg/day for 4 days) followed by Cyclophosphamide (50 mg/kg/day for 4 days). The patient ag/Ag/day for a days followed by Lyclophosphaside 150 ag/Ag/day for a days. The patient was protected by Left isolator and by our decontaination protocol. Cyclosporine madministered for 12 months and oral Acyclovir for 270 days. The clinical course was uncomplicated and the boy was engrafted promptly with PNH cours mormalizing on day +23 and the complete reversal of neutrophil function defect. Cytogenetic showed complete engraftent of donor origin at 4 months after BNT. Eighteen months after BNT the boy still continues to be free of infections. In fact NBT was already positive at the nineth month for 2 was an (2521) and for DNA (2821). for Zymosan (92%) and for PMA (91%). The $Q_{\overline{g}}$ generation with PMA was 8.88 nmol/min/10° cells and with fMLA was 13.01 nmol/min/10° cells. Cytochrome b spectra showed peaks at 428, 530, 558 nm (7.81 pmol/10* cells) and all his neutrophils stained for cytochrome bon the cell surface by using monoclonal antibodies 705 (98%). Earlier attempts of BMT for CGD by bone marrow from related or unrelated donors have failed either because of slow loss of graft and gradual deterioration in neutrophil function or because of lethal infections and/or GvH disease. We conclude that as well as in CGB $\chi/b-$, previously described by us (Bone Marrow Transplant 4, 695, 1989), a complete neutrophil engraftment can be achieved also in this rare form CGD X/b+. Supported by a grant from M.B.R.S.I.,

CLINICAL AND GENETIC HETEROGENEITY OF RECESSIVE FORM 103 OF HEREDITARY SPHEROCYTOSIS.

A.lolascon,S.Perrotta,E.Miraglia del Giudice,L.Pinto and S.Cutillo Department of Pediatrics - University of Naples (Italy)

α-spectrin synthesis exceeds β by a factor four (Hanspal 1987). This statement suggests that \alpha-spectrin deficiency would be evident only in the homozygous state. Agre (1982,85) described some cases of recessively inherited HS characterized by severe spectrin deficiency and life-threatening hemolysis. Marchesi (1989) founded in 9 out of 2O of these kindreds an acidic shift of all domain. More recently Forget (1990) analyzing the genomic DNA of these subjects demonstrated an Ala-Asp (GAT-GCT) substitution at position 972 of α-sp. In order to assess clinical findings of rHS and verify the linkage with a-spectrin gene we studied 11 unrelated kindreds of italian extraction, characterized by affected propositus with hematologically and biochemically normal parents. Two of these had severe HS, whereas the remaining have a clinical picture of typical HS. We examined 3 polymorphic sites (Xbal,Pvull,Mspl) founded in linkage disequilibrium with α-sp gene (Hoffman,1987)(3O21-E1 cDNA probe was given by courtesy of B.G.Forget). In the informative families rHS is associated with the following haplotype X(+),P(-),M(+) in three chromosomes and with ++-, -+- and --- in three others. The haplotype +-+ is the most frequent in normal population of our country. This finding suggests the heterogeneity of molecular defect at the α -gene level. Furthermore in two cases the haplotype assignement allows to the exclusion of α -sp gene involvement. This suggests that the \alpha-spectrin gene abnormalities couldn't cause all rHS.

NEWBORN ERYTHROCYTES ARE MORE RESISTENT. THAN ADULT TO FREE RADICAL-MEDIATED DAMAGE CAUSED BY LIPID EMULSION.

S Kljuchnikov, O Pitkänen, K Raivio and S Andersson - 4th Children's Hospital, Univ. of Moscow, USSR; and Children's Hospital, Univ. of Helslnki, Helsinki, Finland.

Intravenous lipid emulsions containing polyunsaturated fat are susceptibe to lipid peroxidation, and may trigger cell damage. We incubated adult (A) or term neonatal (N) erythrocytes with 0.44 % Intrallpid® (IL) in phosphate-buffered saline for 17 h. Hemolysis was 40.5±2.1% in A and 22.4±1.3% in N (p<0.001). MCV increased by 69.7% in A and 48.5% in N (p<0.001). Glutathione (GSH) decreased in A from 63.6±4.7 to 21.6±4.5 (66.0 %) and in N from 61.1±5.9 to 25.7±5.0 mg/dl (57.9%)(changes p<0.001). However, malondialdehyde and conjugated dienes were higher in N (both p<0.001). Desferoxiamine or sodium etidronate Inhibited the effects of IL in both A and N. We conclude that despite higher lipid peroxidation in the membrane, the newborn red cell is more resistant to free radical damage than that of the adult.

> HIGH DOSES OF HUMAN RECOMBINANT ERYTHROPOLETIN (CHUEPO) ARE EFFECTIVE IN THE TREATMENT OF AMERICA OF PREMATURITY (AP)

105

Virgilio Carnielli, Rosalia Da Riol, Graziella Zacchello, Felice Cantarutti, Giovanni Montini Department of Pediatrics, University of Padova, Via Giustiniani 3, Padova, Italy,

rHuEPO has been preaviously used for the treatment of AP with uncertain results; dosages batween 40 and 300 IU/Kg/week have generally failed to reduce the need for blood transfusions in premature infants, although no major side-effects were shown. We performed a pilot study to demonstrate the safety and the efficacy of higher doses (120-300-600-1200 IU/kg/week) of rHuEPD administered to premature infants. Only doses of 1200 1U/Kg/week were shown effective to increase the reticulocyte count; we describe these data. Patients: Inclusion criteria were: gestational age (GAGE) less than 33 weeks and birth weight (BW) less than 1.75 Kg; admission to our meonatal intensive care unit. Group A: 8 premature infants (sean BW 1.3810,26; GAGE 30.211.6) were given 1200 IU/Kg/week in 3 doses and 20 ag/Kg/week of iron. Group B: 11 premature infants (BW 1.3410.26; GAGE 29.612.4) were used as controls. Results: no side-effects of the treatment, especially for blood pressure, white blood cell and platelet count, were shown. Main results are shown in the table,

	Number	RC transfused	Blood sampled	Hean Hct	Mean retics
	transfusions	(ml/Kg)	(m1/Kg)	(X)	(X)
EPO	1.0±1.7	16129	20±23	48.1±7.0	4.6±0.3
CONTR.	3.1±2.2	48134	27±19	41.913.6	1.610.4
p	0.019	0.023	0.21	0.048	0.0001
CONCLUS	ONS: From these	preliminary dat	a high doses of	rHuEPO and	iron supplementation are
effectiv	e in reducing t	he number of blo	od transfusions	in prematur	re infants.

106

PREMATORE NEONATES: PRELIMINARY RESULTS OF A DOUBLE BLIND CONTROL TRIAL Vasiliki Soubasi, George Kremenopoulos, Chaido Tsantali, Sotiria Mastrogianni, George Kyriakides, Dimitris Tsakiris, (spn. by Dimitris Anagnostakis) — Department of Neonatology, University of Thessaloniki, 546 22 Thessaloniki, Greece

EFFECT OF RECOMBINANT HUMAN ERYTHROPOIETIN (EPO) IN PREMATURE NEONATES: PRELIMINARY RESULTS OF A DOUBLE

We investigated whether preventative treatment with higher doses of EPO We investigated whether preventative treatment with higher doses of EPO in the premature neonates is safe and reduces the need for transfusion. 20 infants were randomized to an EPO (n:12,8W:1176±234g,GA:28,2±2,29) and a control group (n: 8, BW: 1232±239g, GA: 29,3±1,6w). EPO 150U/kg was given s.c. every 3rd day for 6w early from the 1st w of life and iron 3mg/kg/d from day 14. The 2 groups were similar (BW,GA, clinical status). During the 6 w of therapy with EPO, there were no differences between control and EPO groups in Ht, Hb, total leukocytes, neutrofils, HbF, transfusion requirements, mortality or morbitidy. Reticulocytes and thrombocytes were increased from the 3rd w and serum ferritin decreased from the 6th w of EPO treatment compared to control group (table).

pre-EPO 2nd w 4th w 6th w

pre-EPO 4th w 3.3 ± 1.7 Reticulocytes EPO 5,6 ± 1,6 p<0,001 1,25 ± 1,1 1.37 ± 1 12 ± 4.8 Thrombocytes EPO control control 269±69 262±166 213±116 290±143 Ferritin EPO 196±221 273±80 284±147 200±193 pco.05 control 210±67 181±85 335±122 547±279 In conclusion EPO administration of 300 U/kg/w appears to be safe. However, although the need for transfusion was not reduced the reticulocyte production was significantly increased.