RETROLENTAL FIBROPLASIA (RLF) II - ASSOCIATION OF

1414 RETROLENTAL FIBROPLASIA (RLF) II - ASSOCIATION OF OXYGEN THERAPY. <u>Malini Satish, Gerald Katzman,</u> <u>Venkatesan Krishnan, Daniel Marcus, Jerald Bovino,</u> Jose Urrutia, Irwin Weinfeld, P.L.S. Amma, Sidney Kripke. (Spon. by <u>M. G. Robinson</u>) Medical College of Ohio, The Toledo Hospital, Dept. of Ped., Toledo, Ohio. 50 neonates with RLF identified between 1/75 and 12/79 were

compared with a group of matched controls without RLF. There was a significantly greater number of hours of oxygen exposure in the RLF group over the control group. Analysis of duration of exposure to FiO\_ above room air is presented. No significant association was found with duration of exposure to FiO\_>.5. RLF neonates were exposed to significantly longer periods of F102 less than .5 than controls. Increased duration of oxygen exposure at any FiO<sub>2</sub> seems to influence RLF development more than the actual magnitude of the FiO<sub>2</sub>.

|                | RLF   |       | CONT  | p*   |       |
|----------------|-------|-------|-------|------|-------|
| ,              | Thrs. | S.D.  | x hrs | S.D. |       |
| Total Duration |       |       |       |      |       |
| Oxygen Therapy | 974   | 755   | 386   | 510  | .001  |
| 100% Oxygen    | 5.11  | 10.9  | 8.13  | 44.3 | NS    |
| 71-100% Oxygen | 16.66 | 38.7  | 8.8   | 29   | NS    |
| 51- 70% Oxygen | 50.9  | 127.2 | 22.7  | 75.2 | NS    |
| 41- 50% Oxygen | 97.3  | 212.2 | 25.8  | 74.2 | 0.05  |
| 31- 40% Oxygen | 198   | 231   | 72    | 124  | 0.01  |
| 21- 30% Oxygen | 589   | 404   | 279   | 334  | 0.001 |

\*t test also confirmed with Cochrane test

STUDIES IN RETROLENTAL FIBROPLASIA (RLF) I - ASSOCIA-1415 TION OF ARTERIAL PaO2. Malini Satish, Gerald Katzman, **1415** <u>Venkatesan Krishnan, Daniel Marcus, Jerald Bovino,</u> <u>Jose Urrutia, P.L.S. Amma, Irwin Weinfeld, Sidney Kripke.</u> (Spon by <u>M. G. Robinson</u>) Medical College of Ohio, The Toledo Hospital, Dept. of Ped., Toledo, Ohio. (Spon.

Of 569 oxygen-treated neonates examined by two retinologists between 1/75 and 12/79, 50 had RLF. The severest grade in either eye is indicated in Table I. A com- TABLE #1 NUMBER % #Examined #RLF parison of the duration of exposure to 569 50 8.8 several PaO<sub>2</sub> levels was made between RLF neonates and 50 weight-matched Grades: 29 5.0 1.75 controls without RLF. Linear trend-II 10 0.9 ing of the PaO<sub>2</sub> between individual III IV PaO2 determinations was assumed in order to estimate the time each V 0.9 5 patient was exposed to a given PaO2 range. Only the period when arterial blood gases were TABLE Duration S.D. Value #2 obtainable from indwell-Hrs Exposure Pa02 ing catheters was ana-RLF RLF/C lyzed. Short periods of Range Control 1.56 exposure to high PaO2's do 150 1.67 1.26 NS not seem to have a signi-1.74 100-150 10.84 6.71 NS 9.18 ficant relationship to the 4.94 genesis of RLF. However, even modestly elevated 80-99 24.02 10.62 21.61 .01 Pa02's (80-99) for signi-7.37 ficant durations may influence RLF development. -Peak Pa02 x NS 192.9 174.3 60

RETROLENTAL FIBROPLASIA (RLF) III - ASSOCIATION OF 1416 PHYSIOLOGIC STATE AND THERAPEUTIC MODALITIES OTHER THAN OXYGEN. <u>Malini Satish, Cerald Katzman</u>, <u>Venkatesan Krishnan, Daniel Marcus, Jerald Bovino</u>. (Spon. by <u>M. G. Robinson</u>) Medical College of Ohio, The Toledo Hospital,

Dept. of Ped., Toledo, Ohio.

50 neonates with RLF and their matched controls were studied. 16 of 34 parameters analyzed had significance with RLF and are presented. These suggest that compromised neonates are susceptible to RLF.

| TABLE #1              | RI    | .F     | CONTH   | ROL   | P      |
|-----------------------|-------|--------|---------|-------|--------|
|                       | x     | S.D.   | x       | S.D.  | Value  |
| IMV - hours           | 526.6 | 506.31 | 185.9   | 324.5 | < .001 |
| CPAP - hours          | 136.3 | 129.34 | 55.1    | 84.4  | < .01  |
| Pa02 50 (hours)       | 6.55  | 7      | 3.7     | 4.5   | < .02  |
| *Bradycardic spells   | 6.2   | 10.2   | 1.42    | 2.82  | < .001 |
| *Apneic spells        | 2.1   | 2.8    | 1.06    | 1.94  | < .005 |
| Peak PCO <sub>2</sub> | 58    | 11.25  | 49.4    | 8.8   | < .005 |
| *requiring bagging    |       |        |         |       |        |
| TABLE #2              | RLF   | (50)   | Control | (50)  |        |

| ·                    | Yes | No |   | Yes | No | X <sup>2</sup> | . p   |
|----------------------|-----|----|---|-----|----|----------------|-------|
| Exchange transfusion | 22  | 28 |   | 8   | 42 | 8              | <.005 |
| FF Plasma Given      | 11  | 39 |   | 0   | 50 | 8.6            | <.005 |
| Abd. Distention      | 35  | 15 |   | 22  | 28 | 5.8            | <.025 |
| Ileus & NPO >10 days | 18  | 32 |   | 7   | 43 | 5.3            | <.025 |
| PDA-LA/A0 >1.3       | 15  | 35 | 1 | 5   | 45 | 5              | <.005 |

Analysis of therapeutic modalities also seem to reflect the need for greater support in these patients. Therapies to improve tissue perfusion and oxygenation may parodoxically add to RLF risk.

HYPOXANTHINE (HX) CONCENTRATIONS AS INDICATOR OF HY-•1417 POXIA. O.D. Saugstad, Bruce Kessel, Brian Saunders, Louis Gluck. Univ. of Calif. San Diego, Depts. OB/GYN 8 Pediatrics, La Jolla, CA; Kaiser Foundation Hosp. Dept. Ped. The HX concentration is a specific indicator of tissue hypox

ia. The plasma level of this purine metabolite can be used to assess intracellular energy status. Extensive animal studies. but less clinical data have evaluated monitoring this metabolite in routine clinical work. In the present study plasma HX has In routine Clinical work. In the present study plasma HX has been determined according to the micromethod previously descri-bed (Saugstad, 1975). Umbilical cord blood from normal deliv-ery & after asphyxia (by Apgar score & cardiotocographic trac-ings) were studied. The mean HX concentration in non-asphyxia-ted babies was 9  $\mu$ mol/1 ( $\frac{1}{2}2SD=0-22 \ \mu$ mol/1). In babies with mod-erate asphyxia, the HX level was elevated (26-40  $\mu$ mol/1). Art-erial plasma HX from neonates with & without tissue hypoxia (by clinical signs & acid-base status) in 18 samples showed linear correlation between plasma HX & base deficit (BD): (BD=0.44HX correlation between plasma HX & base deficit (BD): (BD=0.44HX -1.8 r=0.66 p<0.01). BD ranged between -6 & 21 µmol/1. The relation between HX & pH was: (pH= -0.006HX + 735 (r=0.46, p=0.05). No baby received sodium bicarbonate. Conclusion: The present results demonstrating a good corre-lation between HX & BD is in agreement with animal studies.

(Saugstad et al, 1978, Thiringer et al, 1980). Hypoxanthine determination is rapid  $\varepsilon$  simple. HX more specifically than BD or lactate reflects tissue hypoxia. We therefore suggest that HX be measured routinely in clinical neonatology for assessment of hypoxia.

INCREASED INCIDENCE OF EARLY ONSET HYPERBILIRUBINEMIA 1418 IN BREAST FED VERSUS BOTTLE FED INFANTS. Kenneth L. **1410** IN DECASI FED VARUE DUTING (Spon, by Alan B. Lewis). University of Southern California School of Medicine, Childrens Hospital of Los Angeles, Department of Pediatrics, Los Angeles.

We performed a 9 month retrospective study on 1351 full term healthy neonates (874 breast fed; 477 bottle fed) to compare the incidence of early onset hyperbilirubinemia (EOH) (defined as serum bilirubin >8 mg/dl within 48 hours after birth and/or >12 mg/dl after 49 hours but before 7 days). Excluded from the study were prematures (<2500 gms), infants with direct hyperbilirubin-emia and/or Rh or ABO blood group incompatibility. 379 (43%) breast fed and 182 (38%) bottle fed infants had serum bilirubin levels measured because of visible levels induces (p > 0.5). EOH was found in 293 (34%) breast fed and 109 (23%) bottle fed infants, (p < 0.001). In addition, marked EOH (>15 mg/dl) was found in 40 (5%) breast fed and 4 (1%) bottle fed infants (p < 0.001).

We have found a significantly greater incidence of EOH in breast fed as compared with bottle fed infants. Possible eti-ologies include dehydration and/or inadequate nutrition while the breast milk supply is coming in, or contributory factors in the breast milk itself. Breast fed infants should be carefully monitored for EOH in the first seven days of life.

QUICK EVALUATION OF CARDIOPULMONARY ADAPTATION OF

QUICK EVALUATION OF CARDIOPULMONARY ADAPTATION OF **1419** EWBORNS BY OXYGEN-CARDIORESPIROGRAPHY. <u>Harald</u> <u>Schachinger</u>, Univ. Children's Hospital, Free Univ. of Berlin, Germany (FRG). (Intr. by John C. Sinclair) Blood gas and acid base values in cord blood are often not predictive of neonatal cardiopulmonary adaptation. We employed oxygen-cardiorespirography to measure beat-to-beat heart rate, respiratory rate, thoracic impedence and transcutaneous Po2 in 337 newborns. 16 infants had severe cardiorespiratory problems in the first hours (II respiratory distress, 5 congenital heart disease); two of these had a cord blood pH below 7.15, and one showed a low Po2. In all 16 infants oxygen-cardiorespirography was abnormal in at least one of the parameters. In clinically abnormal infants the tcPo2 was low, and frequently a decrease of long-term variability of the heart rate was seen. In 10 of 16 infants, a pathological pattern was recognized by oxygeninfants, a pathological pattern was recognized by oxygen-

cardiorespirography before clinical symptoms appeared. A hyperoxia test adds additional information. Within 2 minutes following hyperoxia, a distinction can be made between healthy newborns, respiratory problems and congenital heart dis-ease. Shunting through an open ductus arteriosus can be shown ease. Shunting through an open ductus arteriosus can be shown by a difference in tcPo2 measured simultaneously from the thorax  $% \left( {\left[ {{\left( {{{\left( {{{\left( {{\left( {{\left( {{{\left( {{{\left( {{{\left( {{{\left( {{{\left( {{{\left( {{{\left( {{{\left( {{{}}}} \right)}}}} \right.}$ and abdomen.

Oxygen-cardiorespirography allows continuous multiparametric data collection and permits the early recognition of important trends in neonatal cardiopulmonary adaptation.

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