POSTNATAL STEROID TREATMENT AND OCCURRENCE OF CEREBRAL PALSY AT THE AGE OF TWO YEARS AMONG NEONATES BELOW 32 WEEKS. THE EPIPAGE COHORT

UNIXI J. Mourdie<sup>1</sup>, P. Truffert<sup>2</sup> <sup>1</sup>Hôpital Port-Royal and INSERM U149, Médecine néonatale, Paris, France; <sup>2</sup>Hôpital Jeanne de Flandre, CHRU Lille and INSERM U149, Médecine néonatale, Lille, France Background: The increasing survival of very low birthweight babies and the improvement of their management are not

Background: The increasing survival of very low birthweight babies and the improvement of their management are not associated with a decreased incidence of cerebral palsy (CP). Among many risk factors, postnatal systemic steroids have been pointed out by clinical and experimental studies. The aim of this work is to explore the relationship between these treatments and CP at the age of two years in a population based cohort. Methods: We conducted a population based study on the Epipage cohort, including 2358 premature infants of less than 32 weeks of gestation, and surviving at discharge, born in 1997 in 9 french areas. From this population 408 were lost to follow-up (17 %). The analyses have been performed on 1950 neonates followed up to the age of two years. We choosed the definition of cerebral palsy used by the SCPE. There was no data about the type of steroid used. All infants were evaluated from a medical form filled by a physician. We made an univariate analysis to identify risk factors of CP in this cohort, then a multivariate analysis by logistic regression. We also studied two subgroups of preterm neonates: below 28 weeks, and those free of cerebral lesions. weeks, and those free of cerebral lesions.

Weeks, and nose ree of cerebral lesions. **Results:** mean gestational age was 30 weeks ( $\pm$  0.1) and the average weight 1380 g ( $\pm$  9.1). The incidence of cerebral palsy was 7.6 % and 14 % of infants treated by postnatal steroids (OR = 1.8 [1.1 - 2.8]). The other risk factors after multivariat analysis were cerebral hypoxia (OR = 5.5 [1.8 - 8.9]), and severe intraventricular haemorrhages (stage 3 or 4) (OR = 6.1 [1.6 - 6.3]). The decreasing gestational age was nearly significative (OR = 1.2 [1.0 - 1.3]). Protective factors were maternal hypertension (OR = 0.6 [0.3 - 1.0]), and fetal growth restriction (OR = 1.2 [1.0 - 1.3]). Although not significantly, the same trend was observed in the studied subgroups (neonates below 28 weeks (OR = 2.1 [0.9 - 4.8]) and neonates free of neonatal cerebral lesions (OR = 1.3 [0.6 - 2.4])).

Since includes nee of neonate certorial restors (OK = 1.5 (0.6 ± 2.4))). Conclusion: These results are consistent with the data from randomized trials, and are supported by many physiopatho-logical arguments. Analysis concerned a broader population than ventilated newborns included in these trials. Prescription of postnatal systemic steroids should be restricted, and only used in rescue for babies with severe respiratory distress.

## 194

SUPERIORITY OF A NOVEL SURFACTANT, SURFAXIN (LUCINACTANT), OVER EXO-SURF (COLFOSCERIL PALMITATE) IN PREVNTING RESPIRATORY DISTRESS SYN-DROME IN VERY PRETERM INFANTS: A PIVOTAL , MULTINATIONAL, RANDOM-IZED TRIAL

IZED TRIAL
F.R. Moyul, J. G. Gadzinowski<sup>2</sup>, E. B. Bancalari<sup>3</sup>, V. S. Salinas<sup>4</sup>, B. K. Kopelman<sup>5</sup>, A. B. Bancalari<sup>6</sup>, M. K. Kornacka<sup>7</sup>, R. S. Segal<sup>8</sup>, C. S. Schube<sup>2</sup>, H.T. Tad<sup>3</sup> University of Tesus, Pediatrics, Houston, T.K. United States: "Karol Marcinkowski University of Mean Pediatrics, Manual Chemory, and Chemory, an

	Surfactant				
	Surfaxin	Exosurf	Survanta	P-value	
Outcome	N=527	N=509	N=258		
RDS at 24 hours	206(39%)	240(47%)	86(33%)	0.005	
RDS -related death by Day 14	25(4.7%)	49(9.6%)	27(10.5%)	0.001	

\*Surfaxin vs. Exosurf, based on logistic regression adjusting for center and BW strata

Conclusion: Surfaxin is significantly more efficacious than Exosurf in the prevention of RDS and in reducing RDS-related mortality through 14 days after birth.

# 195

# COMPARISON OF INCIDENCES OF ALL-CAUSE MORTALITY BETWEEN THE NOVEL

COMPARISON OF INCIDENCES OF ALL-CAOSE MORIALITY BETWEEN THE NOVEL SURFACTANT, SURFAXIN (LUCINACTANT) AND THE ANIMAL DERIVED SURFAC-TANTS SURVANTA (BERACTANT) AND CUROSURF (PORACTANT ALFA) <u>F.M. movy</u>, SS Sinha<sup>2</sup>, R S Segal<sup>3</sup>, C S Schaber<sup>4</sup>, H T Sua<sup>3</sup> <sup>1</sup>University of Texas, Pediatrics, Houston, TX 01548, United States; "The James Cook Hospital, Pediatrics, Middlesbrough, United Kingdon; "Discovery Laboratories, Clinical Research, Doylestown, PA 18901, United States; <sup>4</sup>Discovery Laboratories, Clinic of Neonatology, Doylestown, PA 18901, United States

United States Background: Animal-derived, protein-containing surfactants perform better vs. non-protein-containing surfactants for prevention and treatment of Respiratory Distress Syndrome (RDS). A new generation, non-animal-derived surfactant, Surfaxin, containing a peptide that mimics the action of human SP-B, has been shown to be effective in animals, in studies of inflants with RDS, I and in large, controlled studies in the prevention of RDS in preterm inflants. Our objective was to compare the incidence of all-cause mortality at Days I 4 and Qs, and at 36 weeks postconceptional age of Surfaxin vs. Survanta and Surfaxin vs. Curosurf across two clinical trials measuring their respective safety and efficacy in the prevention of RDS. Surfaxin (S A mL/S or 175 mg/kg) vs. Survand at (40 mL/kg or 100 mg/kg) and Curosurf (22 mL/kg or 175 mg/kg). MI andonized patients were included. Endpoint definitions were the same for both trials. Adjustments were performed for birth weight (BW 0) strata and study centers. Inclusion eriteria: BW 600-1250 ge gestational age -32 weeks, and BW strata.

Results: The pooled analysis of the two trials comparing Surfaxin with animal-derived surfactants is presented below: See table.

		Animal-derived	Surfaxin vs.	
All-cause mortality	Surfaxin	Surfactant	Animal-derived surfactant	
	(N=651)	(N=386)	p value	
By Day 14	99(15.2%)	66(17.1%)	0.245	
By Day 28	116(17.8%)	83(21.5%)	0.044	
By 26 weeks PCA*	132(20.3%)	93(24.1%)	0.045	

\*Conceptual age as per last known menstruation. Analyses of further endpoints are in process and will be presented

Conclusion: Combined data from these trials show lower all-cause mortality with Surfaxin than with the animal-derived surfactants (combined results of Survanta and Curosurf). Since Surfaxin has a higher concentration of surfactant apporteni-equivalents—in the form of sinapultide—than any other commercially available surfactant, these results are consistent with the hypothesis that both the presence of, and a higher concentration of, surfactant protein amplify the benefits of surfactant therapy.

#### 196

RESUSCITATION WITH 100% O2 INCREASES PULMONARY MMPS IN HYPOXIC PIG-LETS

LETS <u>B H Munkeby</u><sup>1</sup>, W B Borke<sup>1</sup>, K Bjørnland<sup>2</sup>, L I B Sikkeland<sup>3</sup>, G Borge<sup>4</sup>, J Lomo<sup>5</sup>, S Rivera<sup>6</sup>, B Halvorsen<sup>7</sup>, O D Saugstal<sup>8</sup> <u>7</u> Riskhospitalet University Hospital, Dept of Pediatric Research/Inst for Surgical Research, Oslo, Norway; <sup>2</sup> Riskhospitalet University Hospital, Inst for Surgical Research, Oslo, Norway; <sup>5</sup> Riskhospitalet University Hospital, Centre for Occupa-tional and environmental medicine. Oslo, Norway; <sup>4</sup> HER Jean Roche. Faculté de Médecine Nord, Marseille, France; <sup>7</sup> Riskhospitalet University Hospital, Research Institute for Internal Medicine, Oslo, Norway; <sup>8</sup> Riskhospitalet University Hospital, Dept of Pediatric Research, Oslo, Norway; <sup>9</sup> Background: Research Stat, Norway Background: Research Noslo, Norway

**Background:** Respiration recovers faster in neonates resuscitated with ambient compared to 100% O<sub>2</sub>. Matrix Metalloproteinase(MMPs) -9 and 2 play a role in pulmonary extracellular matrix remodelling and degradation after ischemia-repertision. **The objective:** Resuscitation of piglets with 100% oxygen is detrimental to pulmonary tissue compared to ambient air. We assessed MMP-9 and 2 in piglets after global hypoxia and subsequent resuscitation with ambient air or 100% O<sub>2</sub> after a hypoxic insult, and thereafter observed for 150min. In pulmonary tissue extracts and tracheal aspirate, MMPs were analysed by gelatine zymography and in situ zymography. **Results:** In pulmonary tissue gelatin zymography pro-MMP-9 was increased in the resuscitated or properties to baseline (p<0.005). In tracheal aspirate pro-MMP-9 and 2 were doubled in the ones resuscitated with 100% oxygen compared to ambient air (p<0.005). In tracheal aspirate pro-MMP-9 and 2 were doubled in the ones resuscitated with 100% oxygen compared to ambient air (p<0.005). In tracheal aspirate pro-MMP-9 and 2 were doubled in the ones resuscitated with 100% oxygen compared to ambient air (p<0.005). In tracheal aspirate pro-MMP-9 and 2 were doubled in the ones resuscitated with 100% oxygen compared to ambient air. In situ zymography gave the same results. **Conclusion:** 100% oversus 21% oxygen for resuscitation of newborn piglets gives increased MMP-9 and 2 in pulmonary tissue and tracheal aspirate. Our data suggests that ambient air is less toxic to the lungs than 100% oxygen.

### 197

IMPAIRED IL-4-ASSOCIATED GENERATION OF CCR4-EXPRESSING T CELLS IN NE-

ONATES AT HEREDITARY ALLERGY RISK U Haddeland<sup>1</sup>, G B Sletter<sup>2</sup>, P Brandtzaeg<sup>1</sup>, <u>B Nakstad<sup>3</sup></u> <sup>1</sup>The Laboratory for Immunohistochemistry and Immunopa-thology (LIIPAT), Institute of Pathology, Rikshospitaler University Hospital, Oslo, Norway; <sup>5</sup>Department of Chemistry, National Veterinary Institute, Oslo, Norway; <sup>3</sup>The Department of Pediatrics, Akershus University Hospital, Oslo, Norway Background: Reduced microbial exposure in early life may be partly responsible for an increase in atopic diseases in 'westernised' societies but the underlying mechanisms remain elusive. Objective To examine how exposure to bacterial Increducesholid (J BS) dynamic the first entire accounter micht influence the methodic of learned homehoid elle and

lipopolysaccharide (LPS) during the first antigen encounter might influence the maturation of neonatal lymphoid cells, and to analyse possible differences in this respect between neonates with a high risk of allergy due to family history (FH+) and controls with no apparent risk (FH-). Methods: Cord blood monouclear cells from the FH+ or FH- group were stimulated with pure LPS or â-lactoglobulin

(a-LG) in its inherent LPS mileu. T-cell expression of chemokine receptors CCR4 and CXCR3 was determined by flow cytometry and RT-PCR. Cellular expression of IL-4 was analysed by quantitative RT-PCR, whereas IFN-å was analysed by both quantitative RT-PCR and ELISA.

by both quantitative R1+PCR and ELISA. **Results:** Stimulation with LPS, or à-LG together with LPS, induced up-regulation of CCR4 (P<0.05) and CXCR3 (P<0.05). For CCR4 such up-regulation was related to the level of IL-4 produced by the same T cells (rs=0.49, P=0.03), whereas CXCR3 expression was negatively correlated with the IL-4 levels (rS=-0.56, P=0.02). Compared with the FH-group, the FH+ group showed a significantly lower capacity for induction of CCR4+ T cells (mean % of total T cells: FH+, 242% vs. FH, 5.74%, P<0.01), and tended to express less IL-4 mRNA. Conversely, induction of CXCR3 and IFN-à was not significantly different between the two groups. C **Conclusion:** When the immune system in early life encounters antigen together with LPS, T-cell localization for further immune induction within barboici discone is doubleted by CCR4 and CYCR4 and expression. In accounter, take of

immune induction within lymphoid tissue is facilitated by CCR4 and CXCR3 expression. In neonates at hereditary risk of allergy this homeostatic mechanism is jeopardised due to poorly up-regulated CCR4. Conversely, Th1 responses to antigen in the presence of LPS is not reduced compared with controls.

## 198

IMPAIRMENT OF PUTATIVE REGULATORY T CELLS IN CORD BLOOD FROM NEO-

NATES WITH HEREDITARY ALLERGY RISK U Haddeland<sup>1</sup>, P Brandtzaeg<sup>1</sup>, <u>B Nakstad<sup>2</sup></u> <sup>1</sup> Akershus University Hospital, The Laboratory for Immunohistochemistry and Immunopathology (LIIPAT), Akershus, Norway; <sup>2</sup> Akershus University Hospital, The Department of Pediatrics, Akershus, Norway

Background: The hygiene hypothesis suggests that the increasing prevalence of allergy in "westernised" countries is due to reduced bacterial exposure in early life, but the underlying mechanism remains elusive. Objective: To analyse the

the to reduce doctrial product lipoplysaccharide (LPS) on the generation of regulatory T ( $R_{\rm g}$ ) (loc) (loc) is an any set of the doctrial product lipoplysaccharide (LPS) on the generation of regulatory T ( $R_{\rm g}$ ) (loc) in neonates with allergy risk due to a family history of atopy (FH<sup>+</sup>) and in controls without such hereditary risk (FH). Methods: Cord blood mononuclear cells from the FH<sup>+</sup> and FH<sup>-</sup> groups were stimulated with bovine å-lacoglobulin in an inherent LPS milieu. T-cell phenotypes indicative of T<sub>R</sub> cells (CD25<sup>+</sup>, CD25<sup>ligh</sup>) and integrin  $\frac{1}{R_{\rm g}}^{-1}$  and the proliferation antigen Ki-67 were quantified by flow cytometry. Release of TGF-â1 from its inactive complex was determined by U.S.

proliferation antigen Ki-67 were quantified by flow cytometry. Release of TGF-â1 from its inactive complex was determined by ELISA. **Results:** The FH<sup>+</sup> group showed reduced capacity for generation of CD25<sup>+</sup> T cells (FH<sup>+</sup>, 16.2% vs. FH<sup>-</sup>, 34.9%; p < 0.01) and  $a_E^{-+}$  T cells (FH<sup>+</sup>, 2.1% vs. FH<sup>-</sup>, 3.9%; p < 0.05). Moreover, the CD25<sup>high</sup> p cleal subset themded to be impaired in the FH<sup>+</sup> group (FH<sup>+</sup>, 5.1% vs. FH<sup>-</sup>, 3.9%; p < 0.05). Moreover, the CD25<sup>high</sup> phenotype (r = -0.54, p < 0.05), and also to the activation-induced release of TGF-â1 (r = -0.80, p < 0.001). **Conclusions:** This study suggested that early-life exposure to a dietary antigen in physiological LPS milieu may generate T<sub>R</sub> cells, including the CD25<sup>high</sup> phenotype that was shown to be related to TGF-â activity and suppressed proliferation. This capacity was impaired in neonates with hereditary allergy risk, but clinical follow-up will be required to determine the effect on allergy emergence.