TYPE 1 DIABETES IN NEW ZEALAND CHILDREN AND ADOLESCENTS: INCIDENCE,

PREVALENCE AND MORTALITY <u>B A Darlow¹</u>, J A Willis², R § Scott², C M Florkowski² ¹Christchurch School of Medicine, Department of Paediatrics, Christchurch, New Zealand; ²Christchurch School of Medicine, Lipid and Diabetes Research Group, Christchurch, New Zealand

Background: Type 1 diabetes results from the autoimmune destruction of pancreatic beta cells. Epidemiological studies from many populations and geographical contexts reveal increases in the incidence of type 1 diabetes with time. This study in Canterbury (New Zealand) determined the incidence of type 1 diabetes from 1970 to the present, the prevalence mid-way through the study period and mortality rates at 9 and 15 years follow-up. Methods: Prospective ascertainment of incident cases aged 0–19 years commenced in 1982. Cases presenting 1970–88

were ascertained from hospital records. The prevalence of type 1 diabetes in the population was determined in January 1994 via a community survey based in retail pharmacies. The prevalence cohort was followed up at 9 and 15 years to establish vital status.

Results: Incidence rates ranged from 2.40 to 26.59 cases/100 000 person years over the three decades, increasing by 0.59 Results: Incidence rates ranged from 2.40 to 26.59 cases/100 000 person years over the three decades, increasing by 0.59 cases/100 000 per year or 5% annually. The prevalence of type 1 diabetes mid-way through the observation period was 0.3 and 1.8 per 1000 for the 0–9 and 10–19 year age groups respectively. At follow-up after 15 years, the standardised mortality ratio for individuals diagnosed with diabetes before age 30 was 2.7 for females and 3.0 for males. That is, a diagnosis of diabetes prior to 30 years of age is associated with a three-fold increase in mortality compared with non-diabetic children in the background population. While cardiovascular disease accounted for the highest absolute mortality rate, the standardised mortality rates for hypoglycaemia, and for renal failure were 7.52 and 35.74 respectively. Conduction: This incidence are of pume 1 diabetes pin increasing and for renal failure were and 3.574 respectively. Conclusion: The incidence rate of type 1 diabetes is increasing significantly over time. A diagnosis of diabetes is ciated with considerable excess mortality

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NF-KAPPA B ACTIVATION AND APOPTOSIS IN LUNG NEUTROPHILS FROM INFANTS

WITH HYALINE MEMBRANE DISEASE F C Cheah¹, <u>B A Darlow¹</u>, C C Winterbourn², M B Hampton², M C M Vissers² ¹Christchurch School of Medicine, Department of Paediatrics, Christchurch, New Zealand; ²Christchurch School of Medicine, Department of Pathology, Christchurch, New Zealand

Christchurch, New Zealand Background: Sequestration and the persistence of neutrophils in the lungs of infants with hyaline membrane disease (HMD) have been associated with pulmonary injury and the subsequent development of chronic lung disease (CLD). While activation of the transcription factor, NF-kappa B, in these cells could upregulate the expression of various mediators that enhance the pulmonary inflammatory process, apoptosis of neutrophils may promote the resolution of lung inflammation. We aimed to determine the proportion of tracheal aspirate samples from infants with HMD that had neutrophils showing evidence of NF-kappa B activation and/or apoptosis.

Methods: Tracheal aspirates were collected from mechanically ventilated infants who were diagnosed to have HMD. The cells in the samples were pelleted after centrifugation, fixed and stained for immunofluorescence microscopy. The neutrophils from each sample were stained using anti-p65 antibody to assess for NF-kappa B activation and cleaved

neutrophils from each sample were stained using anti-p05 antibody to assess tor Nt-kappa B activation and cieavea caspase-3 antibody for the detection of apoptosis. **Results:** A total of 172 tracheal aspirates from 59 premature infants were studied. The median gestational age and birthweight were 27 weeks (range, 23 to 36 weeks) and 855 g (range, 500 to 3200 g) respectively. The median age when aspirates were obtained was day four of life and the median number of samples per infant was two. Fifty-four (31%) samples from 33 (56%) infants had neutrophils with evidence of Nt-kappa B activation. A total of 56 tracheal aspirates from 27 infants were available for concomitant assessment for neutrophil apoptosis. Eleven (16%) samples from seven (26%) infants showed evidence of neutrophil apoptosis. With the exception of one sample, tracheal aspirates with *neutrophile beuing* Nt-kappa B activation or anontosis were multially exploiting requires the state of the sample of the state of the sample of the sample for the sample samples in the sample for the sample samples in the sample for a sample in the sample sample for the sample sample for an anontosis were multially exploiting perturbations the sample sample sample for the sample for the sample sample for the sample for the sample sample for the sample sample for a nontonic same multially exploiting perturbations and the sample sample for the sample sample sample for the sample for the sample sample for the sample for the sample sample sample for the sample sample sample sample for the sample sample

(corr) manual advances of the end of the conditions that are anti-apoptotic, and thus be as sociated with persistent neutrophilic inflammation implicated in the process of lung injury and the development of CLD.

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VITAMIN K DEFICIENCY BLEEDING (VKDB) IN NEW ZEALAND INFANTS: RESULTS

OF SURVEILLANCE OVER FIVE YEARS (1998 TO 2002) <u>B A Darlow</u> Christchurch School of Medicine, Department of Paediatrics, Christchurch, New Zealand Background: There have been several changes in the recommended prophylaxis of VKDB in New Zealand over the

past decade, Precedent the introduction of a new preparation, Konakion MM, in 2001, a consensus statement from professional organisations representing paediatricians, midwives, nurses, GPs and obstetricians was published. This recommended that all babies should receive vitamin K prophylaxis and that the preferred route is intramuscular (i.m.). Tecominenced that an obtions should receive vitamin K prophytaxis and that the predicted role is minimuscular (Linf). Where parents did not agree to i.m. vitamin K, a regime of three repeat oral doses was recommended. We aimed to determine the annual incidence of, morbidity from and circumstances surrounding, cases of VKDB, defined as "sponta-neous bruising or bleeding associated with prolonged cloting time, but not due to an inherited coagulopathy or disseminated intravascular coagulation, in an infant <6 mths old". Methods: The standard methods of the NZPSU were used. Each month all paediatricians were sent either a reply-paid card or email to indicate if they had seen a child with VKDB in the previous month. Following a "yes" response, a two page questionmaire was sent to ascertain details of the case. VKDB was included on the NZPSU card from its inception is between VOB or db near survivore vibror as they may the sentence of the transmitter or the sentence of the transmitter o

in January 1998 and has remained under surveillance subsequently

in January 1998 and has remained under surveillance subsequently. Results: 19 cases were reported in the five years 1998 to 2002; 1 report was not valid, 3 were double reports. Of the 15 remaining cases, 6 were in the first week of life ("Early") and 9 were from day 8 to 6 months ("Late-onset"). Amongst early, 4 were confirmed of Classical type (day 2 to 7), all were breast fed and none received vitamin K; 2 cases were in the first day of life and unusual (pulmonary haemorrhage, sub-dural haemorrhage) and were only possible cases. Amongst late-onset, 1 had i.m. vitamin K at birth, had severe liver disease of unknown cause and required repeat doses of vitamin K; 8 were fully breast fed and had not received vitamin K, 4 had liver disease, 3 had intracerbal haemorrhage (all mathematics) and the liver back to be and the core unit is the new to be under the intracerbal haemorrhage (all were fully breast fed and had not received vitamin K, 4 had liver disease, 3 had intracerbal haemorrhage (all mathematics) and the core unit is the new to be under the intracerbal haemorrhage (all the new to be under the liver the core unit is the new to be under the intracerbal haemorrhage (all the new to be under the liver the wriving. 2 wriving a wriving a set of the formation for the set of the set of

prophylactic vitamin K at birth either i.m. or orally

SIZE MATTERS: THE RISK OF CHRONIC LUNG DISEASE AND SIGNIFICANT RETI-NOPATHY OF PREMATURITY

<u>B A Darlow¹</u>, J L Hutchinson², J M Simpson², D A Donoghue², D J Henderson-Smart², N Evans on behalf of the Australian and New Zealand Neonatal Network (ANZNN)^{2 1}Christchurch School of Medicine, Department of Paediatrics,

Australian and New Zealand Neonatal Network (AIX2N)⁶⁻¹ Christchurch School of Medicine, Department of Paediatrics, Christchurch, New Zealand, ² University of Sydney, Centre for Perinatal Health Services Research, Sydney, Australia Aims: To identify prenatal risk factors for hospital–based outcomes in very preterm infants. Methods: ANZNN prospectively audits all high-risk infants (gestation <32 completed wks or birthweight (BW) <1500g) admitted to each level III newborn intensive care unit in Australia and New Zealand. Two outcomes were selected for individual analysis; chronic lung disease (CLD-any respiratory support at 36wks post-menstrala lage-PMA), significant retinopathy of prematurity (ROP-stage III or greater). As local criteria for ROP screening is most oflen <31 wks gestation or <1250g BW, a subset of babies born at <29 wks was used to prevent bias towards small for gestational age infints. Data were excluded if no eye examination was recorded (n=176, 7.7%). Univariate analysis of 23 variables was undertaken and variables not significant at P <0.05 eliminated; the remainder being entered stepwise into a multivariate logistic model and reiected when they lost significant at P <0.01. rejected when they lost significance at P <0.01.

rejected when they lost significance at P < 0.01. **Results:** The study group consisted of 5599 babies born during 1998–99 who survived to 36wks PMA. Of these, 1235 (22%) had CLD. On univariate analysis having a mother who had a previous perinatal death, antepartum haemorrhage, male sex, 1 min Apgar <4, lower gestational age, lower BW for gestation, ethnicity, pregnancy induced hypertension, and both the method and presentation of birth were associated with CLD. Pre-term pre-labour rupture of the membranes was protective. ROP exams were recorded for 2111 babies born at 22 to 28 weeks gestation, BW for gestation, 1 min Apgar <4, or and produced purture of the method with ROP on univariate analysis were gestation, BW for gestation, 1 min Apgar <4, were and produced purture of the method with ROP on univariate analysis were gestation, BW for gestation, 1 min Apgar <4, were and produced purture of the method with ROP on univariate analysis were gestation, BW for gestation and the function in the function of the second produced in the second produced produced in the second produced purture of the method in the second produced purture of the produced in the second produced produced produced produced produced in the second produced pr in 205 (90%). Significantly associated with KOP on univariate analysis were gestation, BW for gestation, Imm Apgar <4, sex and prolonged rupture of membrane. For both models, after simultaneous adjustment, factors that remained in the multivariate model were gestation, sex and BW for gestation. A dose response was demonstrated for gestation (trend P <0.0001) and BW for gestation (trend P <0.0001), with the risk of poor outcome increasing with decreasing gestation and increasing growth restriction. Being female was protective. The models were validated on comparable groups born in 2000–2001 with all Receiver Operating Characteristic curve statistics >0.80. **Conclusion:** The association of CLD with reduced fetal growth is biologically plausible and may be causal, as it has been demonstrated in other populations. The relationship is less clear for ROP, where historically populations have been

based on BW.

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GENDER-RELATED DIFFERENCES IN PLASMA LEPTIN LEVELS AMONG OBESE

CHILDREN WITH IMPAIRED GLUCOSE TOLERANCE OR TYPE 2 DIABETES *B B Garany-Bogacka¹, <u>G D Dawid²</u>, <i>A J Horodnicka-Jozwa², M S Syrenicz¹, A G Gebala¹, J G Goral¹ Pomeranian Medical University, Independent Laboratory of Propedeutics of Childrens Diseases, Szezein, Poland, ²Pomeranian Medical University, Second Department of Pediatrics, Szczein, Poland* Plasma leptin has been shown to correlate positively with mary indices of obssity, as well as insulin resistance. Previous studies have reported that leptin levels are similar in diabetic and nondiabetic individuals. However, these studies were not performed in newly diagnosed diabetics, and other write/log (univ new conformed the carryle). and other variables (such as gender) could have confounded the results.

and other variables (such as gender) could have confounded the results. Aim: The aim of the study was to explore the influence of the glucose tolerance status on plasma leptin in obese children, separately girls and boys. Material and methods: The study group consisted of 67 obese subjects (28 boys and 39 girls), aged 7–17 years. Among them 61 were glucose intolerant (IGT) and 6 had diabetes mellitus type 2, according to ADA criteria. The control group comprised 64 obese children (30 boys and 34 girls). Both groups were well matched for age, % overweight, and body mass index (BMI). Measurements of weight, height, waist and hip circumferences were performed with the using of standardized techniques. Fat mass was measured by bioimpedance method. Plasma insulin was measured

With the Using of standardized techniques, rat mass was measured by poloimpeance method. Plasma limit was measured using a commercialised enzyme immunoassay kit. Fasting plasma leptin was determined by RIA (Linco Research). **Results:** Mean plasma leptin level was significantly lower in entire study group than in control group (17.0±7.5 ig/L, vs. 21.0±7.4 ig/L; p<0.05). Leptin levels were, on average, 36% lower in grifs with diabetes or IGT versus control girl (17.1±6.5 ig/L vs. 23.1±7.7 ig/L; p<0.001). No such between-group difference was observed in boys. In a multiple regression model adjusting for age and BMI in the female subgroup, plasma insulin and glucose concentrations 2-h after glucose load were the best predictors of fasting plasma leptin (r=0.38, p<0.05 and r=0.06); p<0.001, respectively). **Conclusion:** Leptin synthesis by adipose tissue is more susceptible to regulation by insulin and glucose in girls than in

boys. Lower plasma leptin levels in obese girls with IGT or type 2 diabetes than in control girls suggest that hyperglycemia may interfere with the stimulatory effect of insulin on leptin synthesis.

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ABNORMAL ORAL MUCOSAL LIGHT REFLECTANCE IN BRONCHOPULMONARY DYSPLASIA

DYSPLASIA C <u>De Felice</u>'s S Parrin², A Barducci³, G Chitano⁴, F Bagnoli⁵, G Latini⁶ ¹ Azienda Ospedaliera Universitaria Senese, Neonatal Intensive Care Unit, Siena, Italy; ²University of Siena, Odontostomatological Sciences, Siena, Italy; ³Institute of Applied Physics "Nello Carrara", National Research Council of Italy, Information Engineering, Firmer, Italy; ⁴University of Pise and ISBEMErron Mediterramen Scientific Biomedical Institute, Cardiothoracci, Brindisi, Italy; ⁴Neonatal Intensive Care Unit, Pediatrics, Obstetrics and Reproductive Medicine, Siena, Italy; ⁶Perrino Hospital, Neonatology, IEC CON Large Science Device, Italy IFC-CNR, Lecce Section, Brindisi, Italy

Background:Bronchopulmonary dysplasia(BPD)remains an important cause of mortality and morbidity in preterm infants. A disordered vascular development and a decreased production of angiogenic factors in BPD have been recently reported. Extracellular matrix (ECM) is known to play an important role on angiogenesis and blood vessel geometry and changes in ECM components have been previously reported in experimental models and patients with BPD. In the present study, we tested the hypothesis of abnormal reflectance of the oral mucosa in BPD. Methods:Fifteen preterm infants with BPD(M:9,F:6; gestational age:27.5±2.0 wk, birth weight:850±125 gr) and 15

Methods:Fifteen preterm infants with BPD(M:9,76; gestational age:27.5 \pm 2.0 wk, birth weight:S05 t=125 gr) and 15 gender and gestational age-matched control infants without BPD (M:9, 76; gestational age: 72.6 \pm 2.0 wk, birth weight: 865 \pm 135 gr) were examined. Mean blood concentrations of major chromophores, including hemoglobin and bilirubin (P= 0.95) and total bilirubin (P= 0.88) were comparable between the groups. Reflectance was measured on high-resolution photographs of the lower lower gingival and vestibular oral mucosa, using an imaging spectrophotometer. Spatially averaged spectra were used in order to estimate the oral mucosal color in the 400–700 nm wavelength electromagnetic spectral range. Derived spectral data were reproducible [intra- and inter-observer coefficients of variation (mean \pm SD),1.46 \pm 0.899% and 3.72 \pm 1.75%, respectively], and qualitatively comparable (\pm 2%shift) to those obtained by direct measurements (absolute spectral range of the deflectance values from 50–100 different artifact-free and vessel-free areas were measured, and mean values were used for data analysis. The predictive accuracy of oral spectrophotemytry as calculated using receiver operating characteristic curve analysis.

calculated using receiver operating characteristic curve analysis. **Results:**BPD patients showed significantly lower light reflectance values in the red (610–700 nm; t values range:4.552– 6.775, df=28,P<0.0001), with higher values in the violet (400nm, P=-0056;430nm, P=-014), and blue-green (480–500 nm, P≤.024) sections of the spectrum. A low reflectance value in the 640–700 nm wavelengths interval was found to identify BPD patients with 100% sensitivity and 100% specificity (640nm: cutoff=44.91%,650nm:≤4 45,64%;660nm:≤45.95%;070nm:≤47,56%;690nm:≤48,59%;700nm:≤58,508,1%). **Conclusion:**These findings indicate the presence of previously unrecognised, early abnormalities of the average optical properties of the oral mucosa in BPD infants.