circulatory rate and oxygenation after birth. The role of SI on major neonatal outcomes remains controversial.

**Methods** We conducted a systematic review and meta-analysis of randomised clinical trials that evaluated the effects of SI and IPPV on mortality and bronchopulmonary dysplasia (BPD). Descriptive and quantitative information was extracted; relative risk (RR) and risk difference (RD) estimates were synthesised under a random-effects model. Heterogeneity was assessed using the Q statistic and I^2.

**Results** Pooled analysis of 4 trials (n = 611) showed significant reduction in the need of mechanical ventilation within 72 h after birth (RR=0.87 [0.77–0.99], RD=-0.10 [-0.17, -0.03]), number-needed-to-treat=10) in preterm infants treated with an initial SI compared to IPPV. However, significantly more infants treated with SI received treatment for patent ductus arteriosus (RR=1.27 [1.05–1.54], RD=0.09 [0.02, 0.16], number-needed-to-harm=11). There were no differences in BPD, death at latest follow-up, the combined outcome for BPD or death, and other major neonatal outcomes between the two approaches.

**Conclusions** Compared to IPPV, preterm infants initially treated with SI at birth required less mechanical ventilation within 72 h after birth with no improvement in the rate of BPD and/or death. SI should currently only be used in randomised trials until future studies demonstrate the efficacy and safety of this lung aeration manoeuvre.

**Poster abstracts**

**PO-0673** IMPROVING QUALITY OF CARE FOR ELBW IN OUR NEONATAL UNIT: AN INTERDISCIPLINARY PROJECT

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**Background** Our hospital attends 90 VLBW and 40 ELBW infants/year. Vermont Oxford Network is used as benchmarking tool.

**Aims** To improve quality of care delivered to ELBW’s during the first week of life. Primary outcome is IVH reduction. Secondary outcomes are mortality, late onset sepsis (LOS) and ROP reduction.

**Methods** An interdisciplinarian group leaded by 4 nurses and 3 neonatologists was created in January 2012. Management of ELBW infants was reviewed to identify weaknesses and strengths. The original team was then divided into subgroups that worked together for problem resolution. 2 gynaecologists, 1 surgeon and 60 members of the neonatal unit were involved (70% of nurses and 90% of neonatologists). Specific actions were developed for 1. Delivery Room: plastic wrapping, delayed cord clamping and blood sampling; 2. Admission and first hours: temperature and humidity targets, blood sampling, calostrum administration, enamas and positional care. Results were transmitted in oral sessions and written guidelines to the rest of the Unit.

**Results** No changes were found in 2012 compared to the last five years. However, in 2013, a decrease was seen in inborns severe IVH (from 11.6 to 10.9%), LOS (15.8% to 9.8%) and severe ROP (10.1 to 5.9%).

**Conclusions** A high percentage of the neonatal unit has been involved in this project. Short term outcomes have improved for the first time last year. A qualitative change difficult to quantify has also been produced. Caution must be taken when interpreting numerical results as they reflect improvement of only one year.

**PO-0674** INCIDENCE OF INDIRECT HYPERBILIRUBINEMIA AT NEWBORN, ASSOCIATED PATHOLOGY AND THE ROLE OF PHOTOTHERAPY

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**Background** Indirect hyperbilirubinemia at newborn is a distinct and various epidemiological described entity, having multiple causes and mechanisms, in which bilirubin level rises to 15–20 mg/dL.

**Aim of the study** The aim of the study was to evaluate the incidence of jaundice with indirect bilirubin at newborns admitted at the Newborn Department of the Childrens Clinic Hospital from Brasov, Romania. As secondary objectives we considered the associated pathology along with the usefulness of phototherapy.

**Patients and methods** We undergone a retrospecitive study on 1020 patients admitted at our hospital during one year period (January 2012–December 2012). We have evaluated...
demographic data, clinical and laboratory data along with the prescribed therapy.

Results From the 1020 studied cases in 260 there was a diagnosis of indirect hyperbilirubinemia, associated pathology consisted of urinary tract infection in 15 cases, piodermatitis in 12, otitis media in 7, acute diarrhoea in 14 cases and severe dehydration in 9 cases.

Only one case complicated with kernicterus. From the 260 cases only 60 had phototherapy in the treatment schedule.

Conclusion we have shown that one quarter of the jaundice at newborn is due to indirect hyperbilirubinemia and that phototherapy is still a useful treatment.

**PO-0675 PREVALENCE OF MUTATION C.11864G >A (P. TRP3955X) IN THE USH2A GENE IN PATIENTS WITH Usher II SYNDROME FROM VOLGA-Ural REGION OF RUSSIA**

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Usher Syndrome (US) is an autosomal recessive condition characterised by a combination of congenital hearing impairment and retinitis pigmentosa. To date, ten genes have been associated with US, representing up to 90% of cases. Three types of US are known and differ by onset of the symptoms, severity and progression of deafness and additional vestibular dysfunction. Patients with type II US have congenital bilateral sensor neural hearing loss that is mild to moderate in the low frequencies and severe to profound in the higher frequencies, intact vestibular responses, and bilateral retinitis pigmentosa.

40 unrelated Usher II type families (60 patients) from Volga-Ural Region of Russia were studied using genotypeing micro array (Usher, Asper-Biotech) for screening 614 mutations in genes CDH23, MYO7A, PCDH15, USH1C, USH1G, USH2A, GPR98, CLRN1, DFN3B1 and automatic sequencing of Usher’s genes. Diagnosis was based on pedigree data, ophthalmologic, audio logical and vestibular examination.

We revealed homozygous and heterozygous genotypes for the c.11864G >A (p. Thr3955X) mutation (USH2A) in six unrelated families among Russian, Tatar and Chuvash patients with Usher II syndrome. We found four pathogenic mutations in coding region of 8 patients (p. Glu4459fs, p. Thr3955X, p. Glu4078fs, and p. Gly1392X), confirming their clinical diagnosis. The most frequent USH2A gene mutation was c.11864G >A (9/80 alleles; 11.25%). Mutation c.11864G >A in heterozygous state was also found in one Russian subject out of 1066 examined individuals from 16 various populations of Eurasia: Bashkirs, Tatars, Chuvashes, Udmurts, Komi-Permyakhs, and Mordvins, Russians, Belarusians, Ukrainians, Veps, and Karelians, Abkhazians, Kazakhs, Uzbeks, Yakuts, Altaians. Study was supported by grants (No12–04–00342_a, No12–04–98520_r_vostok_a, 14–04–97002_r_povolgie_a, 14–04–97007_r_povolgie_a,14–04–01741_A).

**PO-0676 PREDICTIVE VALUE OF POSTNATAL NASOGASTRIC TUBE AND CHEST/ABDOMINAL XRAY FOR ANTENATAL POLYHYDRAMNIOIS AND/ OR SMALL STOMACH IN THE DIAGNOSIS OF OESOPHAGEAL ATRESIA-TRACHEO- OESOPHAGEAL FISTULA**

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Introduction Oesophageal atresia (OA) ± tracheo-oesophageal fistula (TOF) is a rare congenital anomaly which occurs in ~1/3500 total births.

Prenatal diagnosis of OA-TOF is desirable as it may improve patient outcome by optimising the pre- and postnatal management. Therefore, a high prenatal detection rate is desirable.

Prenatal ultrasound is one of the modalities used in improving the diagnostic rate. Indirect or non-specific signs of suspected OA-TOF include polyhydramnios (10–20% of which are associated with fetal abnormalities, one of it being OA-TOF), absent/small stomach bubble, a blind ending upper oesophageal pouch and observation of fetal swallowing.

In our institution, all foetuses with Isolated polyhydramnios ± small/absent stomach have postnatal NGT insertion followed by xray if follow up scans remained the same.

Objectives To evaluate the clinical relevance of postnatal NGT insertion followed by xray for prenatal finding of isolated polyhydramnios ± absent/small stomach bubble in a cohort of newborn babies to assess for the diagnosis of OA-TOF.

Methods

- Retrospective cohort review January2011-March2014 of all foetuses/neonates with suspected EA-TOF on prenatal ultrasound, isolated polyhydramnios [deepest pool >8 cm, Amniotic Fluid Index >95th percentile for gestational age and/or small/absent stomach bubble] or perinatally identified to have >1500 mls amniotic fluid and therefore needing postnatal NGT insertion+ Xrays pre-feed were identified.
- Course and tip of NGT, stomach-bubble, vertebral anomalies or any other abnormalities detectable on Xray.
- Time taken from birth to when the postnatal Xrays were taken was calculated (time to first feed).
- Babies were also examined for clinical signs of OA-TOF or other abnormalities.
- Clinical data on all OA-TOF cases that were diagnosed postnatally over the same 3 yr period.
- Exclusion – above two features with other any other abnormalities on antenatal scans.

Results

- Total deliveries over period = 17,200.
- Total no. fulfilling criteria of isolated polyhydramnios ± small/absent stomach only = 48.
- No. that resolved antenatally and therefore did not have NGT +Xray assessment postnatally = 12.
- Total No. who had NGT +Xray assessment = 36.
- No. without OA-TOF = 35.
- No. diagnosed with OA-TOF = 1.
- 7 cases of OA/TEF (1 had above 2 features); of which 3 did not have polyhydramnios ± absent/small stomach; 3 had multiple abnormalities including 2 with polyhydramnios.
- Above two antenatal features had a Sensitivity = 25%; Specificity =99.8%; positive predictive value=2.78% for diagnosis of OA-TOF.
- Mean time for NGT+Xray (away from mothers and before their first) = 156 mins (range 81–332).