as it happens in IUGR. To assess changes in neural connectivity in the hippocampus in IUGR animals, the hippocampus synaptic network has been analysed through three different synaptic proteins, Postsynaptic Density Protein 95 (PSD95), Synaptophysin and Synaptosome-associated Protein of 25 KDa (SNAP25).

**Methods** IUGR was induced by meso-ovarian vessels’ cautery in pregnant rats. Sham surgery was performed in control animals. The pups were divided into: Control, Ischemic and IUGR (birth weight < 2 SD). 25 days after birth, animals were subjected to an aquatic learning test. At day 35, they were sacrificed. Synaptic protein levels were analysed by immunohistochemistry staining and Western blotting.

**Results** There were differences in the learning outcomes between Control, Ischemic and IUGR animals. The analysis of PSD95, showed a gradual staining reduction from Controls to Ischemic to IUGR. There were no differences between groups in Synaptophysin immunostaining. The intensity of SNAP25 staining was lower in Ischemic and IUGR than in Controls. These results were corroborated by western blot analysis.

**Conclusions** IUGR animals displayed reduced protein levels of PSD95 and SNAP25 in the hippocampus with respect to Control animals, suggesting a decrease in functional synapses.

**1256 QUALITY OF GENERAL MOVEMENTS AFTER TREATMENT WITH LOW-DOSE DEXAMETHASONE IN PRETERM INFANTS AT RISK FOR BRONCHOPULMONARY DYSPLASIA**

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**Background** Postnatal dexamethasone (DXM) is widely used to treat preterm infants at risk for bronchopulmonary dysplasia. Previously, it was reported that high-dose DXM leads to deteriorated quality of general movements (GMs). We determined neurological functioning in infants after low-dose DXM treatment, assessed by the GM-quality until three months post term.

**Methods** We included preterm infants, admitted to our NICU between 2010−2012 and treated with DXM (starting dose 0.25 mg/kg/d). GM-quality was assessed before (day 0), during and after treatment until three months post term. We determined the change in GM-quality by comparing the GM-quality of day 0 with the GM-quality of the last video recording. Additionally, we calculated a motor optimism score (MOS), ranging from 0 (low optimality) to 18 (high optimality).

**Results** Sixteen infants were included [median GA 26.9 wks (25.0−29.7); BW 800 g (620−1665)]. Before treatment, 4 infants had normal GMs which remained normal after starting treatment. GM-quality improved in 8 of 12 initially abnormal infants (Mc Nemar: P=0.008), whilst MOS slightly increased: median 10.5, 12.0 and 12.5 on days 0, 1 and 7, respectively (NS). Cumulative DXM doses, treatment duration and postnatal ages at starting DXM were not associated with change in GM-quality. Infants whose GMs improved were ventilated for a shorter period than infants whose GMs remained the same quality (P=0.065).

**Conclusions** GM-quality did not deteriorate after DXM treatment but rather improved in infants with initial abnormal GMs. Our findings suggest that neurological functioning until three months post term is not adversely affected after low-dose DXM.

**1257 THE NEURODEVELOPMENTAL ASSESSMENT OF VERY LOW BIRTH WEIGHT INFANTS AT 4–6 YEARS OF AGE**

doi:10.1136/archdischild-2012-302724.1257

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The major and minor neurodevelopmental morbidities among premature infants become an important issue because of the increase in the number of surviving premature newborns.

The Aim of this study was to examine the cognitive, neuromotor, emotional and behavioral outcomes of the premature newborns at 4−6 years of age born with very low birth weight and to investigate the relationship between neuromotor and neurocognitive development.

The neuromotor status of 68 children were evaluated according to Touwen neurological examination, 64 children were assessed using Stanford-Binet and Peabody Picture Vocabulary Test, 65 children using Strength and Difficulties Questionnaire and Vineland Adaptive Behavior Scale.

Three cases were already diagnosed and followed as CP. According to Touwen examination 28 (%41.2) children were normal, 35 (%51.5) had simple minor neurological dysfunction (MND), 2 (%2.9) had complex MND. The mean IQ score was 90±10.9. The rate of hyperactivity, behavioral problems and emotional problems were in order %60, %33.8 and %53.8. The children were diagnosed as having a delay of 14.9±10.6 month for conducting, 10.6±8.6 month for daily activities, 10.7±11.5 month for social competence and a delay of 6.3±10.2 month for motor behavior. The cognitive and neuropsychological results of the 35 children with MND and 28 children with normal neuromotor status were compared. The children with MND had significantly lower mean IQ score, more delay for motor behavior and they were more hyperactive.

Majority of the children who were considered as normal had cognitive impairment, language, behavior/emotional and neuromotor problems in various degrees.

**1258 HEARING IN PRETERM INFANTS WITH POSTNATALLY ACQUIRED CYTOMEGALOVIRUS INFECTION**

doi:10.1136/archdischild-2012-302724.1258

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**Background and Aims** Long-term sequelae of breast milk-associated cytomegalovirus (CMV) infection in preterm infants are insufficiently evaluated. We studied the hearing of preterm infants with postnatal CMV infection within the first and second year of life.

**Methods** Preterm infants (GA<32wks) admitted to our NICU between 2005 and 2011, and diagnosed with CMV infection using CMV PCR of urine at 40wks were included. Congenital infection was excluded in all. Hearing was tested using auditory brainstem response (ABR) in the neonatal period and during the first and second year of life. Neurodevelopmental outcome was estimated using the Griffiths mental developmental scale (GMDS) at 18 months.

**Results** Eighty-eight preterm infants were diagnosed with postnatal CMV infection of whom four were lost to follow-up. All infants had normal hearing in the neonatal period. ABR-tests were performed in 64/84 (76%) infants during the first year of life (median corrected age 7 months, range 2−11) and in 18/84 (21%) infants during the second year (median corrected age 33 months, range 12−50). None of the infants developed SNHL. Mean GMDS score evaluated so far in 58/84 (69%) infants at 15.8 months corrected age (range 13.0−21.0) was 104.4 (SD 9.9) and mean score of the language sub-scale was 16.7 months (SD 2.1). There were no differences in clinical data, cerebral ultrasonography results, viral load and GMDS scores between infants with hearing tests and non-tested infants.
Conclusion Postnatally acquired CMV infection among preterm infants is not related with SNHL during the first and second year of life.

Abstracts

1259 GESTATIONAL AGE AND PREGNANCY DISORDERS AS RISK FACTORS FOR CEREBRAL PALSY
doi:10.1136/archdischild-2012-302724.1259
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Background Both pregnancy disorders and low gestational age (GA) are associated with an elevated risk of cerebral palsy (CP). As many pregnancy disorders are associated with preterm birth, it is unclear whether the increased risk of CP in preterm children can be attributed to the pregnancy disorder or the low GA.

Aim To disentangle the risk of CP according to GA and pregnancy disorders in children born preterm.

Method Norwegian, national cohort study with linkage of the Medical Birth Registry of Norway to the Statistics Norway and the National Insurance Scheme. All births from 1967 to 2001 were followed through 2005, and 1,499,705 individuals were included in the cohort.

Results Placental abruption, chorioamnionitis, placenta previa, multiple birth, prolonged rupture of membranes, cervical conisation, unspecified bleeding and pre-eclampsia were associated with an increased risk of CP. The risk of CP increased with decreasing GA (23–27 wks RR 83.3 [95% CI, 69.8–99.4], 28–30 wks RR 49.5 [95% CI, 43.2–56.6], 31–33 wks RR 17.4 [95% CI, 15.3–19.7], 34–36 wks RR 3.3 [95% CI, 2.9–3.8]). The association between GA and CP was not substantially weakened after adjustment for pregnancy disorders.

Conclusion Although several pregnancy disorders were associated with an increased risk of CP, low GA appeared to be a more important risk factor for CP in preterm children.

1260 NEUROSENSORY OUTCOMES AND HEALTH CARE NEEDS IN SCHOOL-AGE CHILDREN BORN AT < 26 WEEKS' GESTATION AFTER ACTIVE PERINATAL CARE
doi:10.1136/archdischild-2012-302724.1260
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Background and Aims To examine neurosensory outcomes and special health care needs in 10 to 16-year-old extremely preterm (EPT) children born at 2 tertiary care centers in Sweden adhering to a policy of universal resuscitation of all infants born alive.

Design and methods The outcomes of 122 surviving EPT children were compared to a matched control of children born at term. Neurosensory impairments (NSI) were assessed by review of pediatric case records, intelligence by WISC-III, and functional limitations and special health care needs by validated parental interviews (QUICC).

Results Of 213 consecutive EPT live births, 140 (66%) survived to discharge home and 6 infants died in the first year of life. Of survivors, 122 children (91%) were recruited for the study. Table I shows rates of disabilities. EPT children had significantly higher rates of functional limitations and special health care needs than controls but the vast majority was free from severe disability that curtail them from activities of daily life.

Abstract 1260 Table 1 Rates of Disability

<table>
<thead>
<tr>
<th>Disability</th>
<th>EPT (N = 121)</th>
<th>Control (N = 100)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>45.4% (55)</td>
<td>90%**</td>
</tr>
<tr>
<td>Mild</td>
<td>28.9% (35)</td>
<td>7%**</td>
</tr>
<tr>
<td>Moderate</td>
<td>19.8% (24)</td>
<td>3%**</td>
</tr>
<tr>
<td>Severe</td>
<td>6.6% (8)</td>
<td>0%*</td>
</tr>
</tbody>
</table>

Severe Disability: Severe CP (inability to walk), Blindness (visual acuity < 20/200 in better eye), or IQ < 3 SD; Moderate disability, moderate CP (considerable difficulty in walking with or without appliances), deafness requiring bilateral aids or worse or IQ – 3 SD to < 25 SD; Mild disability; mild CP (walking with minimal limitation), or an IQ – 25 SD to < 1 SD; *P<0.05; **P<0.005

Conclusions Disability rates and special health care needs remain high in EPT children, but have not increased since 1990s despite the marked increase in their survival.

1261 NEURODEVELOPMENTAL OUTCOME IN EXTREMELY PRETERM INFANTS BORN IN SWITZERLAND BETWEEN 2000–2008, PRELIMINARY DATA OF THE SWISS NEONATAL NETWORK
doi:10.1136/archdischild-2012-302724.1261
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Introduction So far, national outcome data on extremely premature infants in Switzerland were not available, and discussions on the care of these patients were based on earlier studies from other countries. This national study assessed neurodevelopment in Swiss infants born between 2000 and 2008 at 24 0/7 to 27 6/7 weeks gestational age.

Methods Neurodevelopment was assessed at 2 years using the BSID II. Moderate neurodevelopmental disability (ND) was defined as a mental (MDI) or psychomotor (PDI) development index of 55–70, or mild cerebral palsy (GMFCS level 2). Severe ND was defined as a MDI or PDI < 55, cerebral palsy (GMFCS level 2S), deafness or blindness. Multivariate logistic regression was performed.

Results Among the 1147 extremely preterm born during the study period 303 (26%) died. Follow-up information was available in 684 (81%) survivors. 440 (64%) showed normal development, 166 (24%) moderate ND, and 72 (12%) severe ND. Severe ND was significantly (p<0.05) associated with earlier year of birth, major intra-cerebral lesions, bronchopulmonary dysplasia, grade 3 retinopathy of prematurity, and lower socioeconomic status. In contrast, birth weight, gestational age and sex showed only trendwise associations with severe ND.

Conclusion Based on these preliminary analyses, we now are ready to establish representative Swiss national data on the outcome of extremely premature infants. These will offer guidance to obstetricians, neonatologists, neurologists and parents based on Swiss data.

1262 PREVALENCE OF HEARING IMPAIRMENTS AT AGE FOUR YEARS IN EXTREMELY PRETERM INFANTS
doi:10.1136/archdischild-2012-302724.1262
1A Granh, 1S Horsch, 1B Sköld, 1M Blennow, 1I Uhlén, 1B Hallberg. 1Neonatology; 2Audiology, Karolinska Institutet and University Hospital, Stockholm, Sweden

Table 1: Prevalence of hearing impairments at age four years in extremely preterm infants

<table>
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Conclusions Disability rates and special health care needs remain high in EPT children, but have not increased since 1990s despite the marked increase in their survival.