Abstracts

**Background** Screening for trisomy 21 (T21) in the West Midlands has evolved from double screening in 1995 to combined screening in 2011, with the perception of improved detection amongst the public and providers.

**Methods** Cases of T21 (n = 2,608) were identified using a regional, population-based, multiple source anomaly register covering a birth cohort of 1,140,866 between 1995 and 2011. Regional data on invasive testing were available for the same period.

**Results** The total prevalence of T21 increased by approximately 50% from 18.2 to 27.5 per 10,000 births over the study period. The proportion of cases with a prenatal karyotype was unchanged (annual 50% from 18.2 to 27.5 per 10,000 births over the study period). The rate of amniocentesis and CVS combined for T21 indications decreased by 76% (4.7% of births in 1995 to 1.1% in 2011).

**Conclusion** The time and effort to develop the T21 screening programme has resulted in safer pregnancies for unaffected cases. Mothers of affected pregnancies have seen no improvement in prenatal diagnosis. The combination of an increasing prevalence of T21 with no change in prenatal diagnosis nor TOP rates means the live birth prevalence continues to rise at the same pace.

**PF06** NEONATAL OUTCOMES FOLLOWING EXTREMELY PRETERM PRELABOUR RUPTURE OF MEMBRANES (EPPROM) IN MULTIPLE PREGNANCY

doi:10.1136/archdischild-2013-303966.018

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**Aims and Objectives** To assess the neonatal outcome of multifetal pregnancies that are complicated by Extremely Preterm Premature Rupture of Membranes (EPPROM) under 24 weeks of gestation.

**Study design** The following is a retrospective observational study of twins and triplets who were referred to the Centre for Fetal Care (CFC) at Queen Charlotte’s and Chelsea Hospital (QCCR) who had an antenatal diagnosis of spontaneous EPPROM under completed 24 weeks of gestation. The population were subdivided in to the following three groups: monochorionic, dichorionic, and trichorionic. The scan data base, the obstetrics and the neonatal records of 52 women and their 108 fetuses, were the sources of the information. Our study covered a 10 year period (2002–2012), and then result was analysed.

**Results** The median gestation at PPROM was 22 + 5 weeks (range 13–23 + 6). The latency period Median was 10 days (range 1–91 days), while the mean gestational age at delivery was 25 + 2 weeks (range 18–37) and the median fetal weight was 650 g (range 290–3500 g), pregnancy loss of 33.3% and neonatal of 29.6%. The overall survival rate was 36.11% (39 fetuses).

**Conclusion** Neonatal survival in EPPROM is very poor (36.11%), with 60% of neonatal deaths being due to prematurity complications and perinatal/neonatal survival. MCDA twins had worse outcome than DCDA, in terms of antenatal complications and survival (36.36% vs 72.97%).

**PF07** EXPRESSION OF 2, 3-BISPHOSPHOGLYCERATE MUTASE (BPGM) IN HUMAN PLACENTA

doi:10.1136/archdischild-2013-303966.019

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**Introduction** BPGM is an enzyme in erythrocytes and trophoblasts, which synthesises 2, 3-bisphosphoglycerate (2, 3-BPG), a facilitator of oxygen liberation from haemoglobin. It has insulin-like growth factor II knockout mouse model of intrauterine growth restriction (IUGR), placental BPGM expression is lower than in wild type animals, implicating BPGM in the pathophysiology of IUGR and suggesting a role for 2, 3-BPG in oxygen delivery to the fetus.

**Methods** Human placental messenger RNA encoding BPGM was quantified by TaqMan RT-PCR. The relative expression of BPGM was assessed a) over the course of pregnancy at 7–11, 12–20, 24–34 weeks of gestation (w) and term (total n = 68), b) in IUGR placenta at early (24–34 w, n = 15) and late (37–39 w, n = 5) gestations and compared with appropriately grown fetus (AGA) controls (n = 8 early, n = 26 late).

**Results** BPGM mRNA expression significantly increased with advancing gestation (ANOVA < p = 0.001). There was a 6 and 7-fold increase from 7–11 w to 24–34 w and term respectively, and 3-fold between 12–20 w and term (p < 0.05 for all). There were no statistically significant differences in BPGM mRNA expression between IUGR and AGA placenta in either gestational age group.

**Discussion** Levels of BPGM increased in a time-dependant manner to term. This may indicate a protective mechanism to avoid oxidative stress damage during the early stages of fetal development, with BPGM expression increasing over time in response to greater oxygen demand from the growing fetus. Placental BPGM expression does not appear to be implicated in the pathogenesis of IUGR in human.

**PF08** QUANTITATIVE FIBRONECTIN CAN BE USED FOR EARLIER PREDICTION OF PRETERM BIRTH FROM 18 WEEKS, BUT THE POSITIVE THRESHOLD NEEDS REDEFINING

doi:10.1136/archdischild-2013-303966.020

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**Introduction** Fetal fibronectin (fFN) is an excellent predictor of spontaneous preterm birth (sPTB) and is used qualitatively (<50 ng/ml negative threshold) from 22 weeks gestation. Its value at earlier gestations using a quantitative test (qfFN) is unknown.

**Methods** A prospective secondary analysis of 431 asymptomatic women at high risk of sPTB, who underwent qfFN testing at 18–21 + 6 weeks. 327 women underwent later testing at 22–26 + 6 weeks (acting as their own controls). The end-points were sPTB/ preterm premature rupture of membranes and delivery before 30, 34 and 37 weeks gestation and within 8 weeks of testing.

**Results** Early qfFN predicted delivery within 8 weeks of testing, <30, <34 and <37 weeks with receiver operating characteristics (ROC) areas of 0.66 (0.54–0.80, p < 0.05), 0.68 (0.56–0.79, p < 0.01), 0.68 (0.58–0.78 p < 0.001) and 0.64 (0.57–0.72, p < 0.001). 22-week test prediction was ROC areas of 0.77 (0.63–0.91 p < 0.001), 0.78 (0.61–0.95, p < 0.001) and 0.79 (0.70–0.89, p < 0.001) respectively. A qfFN result of <10 ng/ml at earlier gestations had only 1%, 2% and 4.3% of women delivered within 8 weeks, <30 and <34 weeks gestation, rising to 6.7%, 8.1% and 14.1% with values between 10–49.9 ng/ml (all differences statistically significant, p = 0.03, 0.02, 0.004 by Fishers-Exact). The 22-week test had 1.0%, 1.0% and 2% respectively, rising to 6.7%, 8.1% and 14% with values between 10–49.9 ng/ml (p = 0.02, 0.23, 0.004).

**Conclusion** qfFN is valid for screening for sPTB at 18 weeks, but has inferior predictive value to 22 weeks. Early identification might enable earlier targeted management. A threshold of <10 ng/ml is more appropriate than current practise to define low risk at 18 weeks.

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*Abstracts are reprinted from Arch Dis Child Fetal Neonatal Ed. First published as 10.1136/archdischild-2013-303966.018 on 26 April 2013. Copyright © BMJ Publishing Group Ltd 2013*