Conclusion  CRL discordance is of poor predictive value for adverse perinatal outcome after 14 weeks of gestation in either MC or DC twin pregnancies.

PP.11  ONE THIRD OF NON-ANOMALOUS TERM STILLBIRTHS ARE ASSOCIATED WITH ABNORMAL FETAL GROWTH
doi:10.1136/archdischild-2013-303966.292
AA Moraitis, GC Smith. University of Cambridge, Cambridge, UK

Background  Approximately one third of all stillbirths occur at term. These losses are potentially preventable by early delivery at 37 weeks. One means of screening to detect babies at risk of term stillbirth would be to use ultrasound to assess abnormalities of fetal growth. However, there is limited information on the importance of fetal growth in determining the risk of stillbirth at term.

Methods/Results  We studied registries of Scottish maternity and stillbirth data and identified 668,887 eligible records of term births since 1992. Birth weight was classified on the basis of sex and gestational age specific percentiles. The relationship between birth weight percentile and the risk of stillbirth (all causes except congenital anomaly) was assessed referent to birth weight percentiles in the range 20–80th. The risk (odds ratio [95% CI] P) of stillbirth was increased with birth weight between the 1st–3rd percentile (OR = 8.2 [CI:6.8–9.9] P < 0.001), 4th–10th percentile (OR = 3.5 [CI:2.9–4.0] P < 0.001) and the 11th–20th percentile (OR = 1.8 [CI:1.4–2.2] P < 0.001). The risk of stillbirth was also increasing among the largest infants: 98th–100th percentile (OR = 2.3 [CI:1.7–3.1] P < 0.001). The attributable fraction for stillbirth associated with these birth weight categories was 33% (95% CI: 28 to 37%). The associations were minimally affected by adjusting for maternal characteristics, including smoking.

Conclusion  Approximately one third of all term stillbirths are associated with growth abnormality. These data indicate that population screening for growth abnormality could identify babies at increased risk of term stillbirth.

PP.12  WEIGHT DISCORDANCE AND PERINATAL MORTALITY IN TWINS: THE STORKE MULTIPLE PREGNANCY COHORT
doi:10.1136/archdischild-2013-303966.293
F D’Antonio, A Khalil. T Dias, A Bhide, B Thilaganathan. Fetal Medicine Unit, Division of Developmental Sciences, ST. George’s University of London, London, UK

Objectives  The main aim of this study was to ascertain the performance of BW and ultrasound EFW discordance in the prediction of perinatal loss in twin pregnancies.

Methods  A retrospective study of all twin pregnancy births from a large regional cohort of 9 hospitals over a ten year period. Intertwin BW and ultrasound EFW discordance were related to stillbirth and neonatal death of one or both twins obtained from a mandatory national register. Receiver operating characteristic (ROC), survival and logistic regression analyses were performed to evaluate the contribution of weight discordance in determining perinatal loss.

Results  A total of 2,161 twin pregnancies were included in the analysis. A BW discordance of ≥25% was associated with the highest AUC for the prediction of stillbirth and neonatal death (BW AUC = 0.72, 95% CI: 0.65–0.80). The perinatal loss in twins with a BW discordance of ≥25% was significantly greater (60.9 per 1,000 fetuses) than that in twins with an EFW discordance <25% (8.6 per 1,000 fetuses) (p < 0.0001); the predictive accuracy was similar using either BW or ultrasound EFW discordance (EFW AUC = 0.69; 95% CI: 0.62–0.77, p = 0.62). Logistic regression analysis demonstrated that BW discordance and gestational age, but not chorionicity or individual fetal size percentile, were independently associated with perinatal mortality.

Conclusions  An EFW discordance of ≥25% represents the optimal cut-off for the prediction of stillbirth and neonatal mortality irrespective of chorionicity or individual fetal size. A policy of increased fetal surveillance commencing from 26 weeks’ gestation and elective delivery by 38 weeks might be reasonable.

doi:10.1136/archdischild-2013-303966.294
AA Moraitis, GC Smith. University of Cambridge, Cambridge, UK

Background  A previous study (Lancet 2003; 362:1779–84) reported an increased risk of unexplained stillbirth in women with previous caesarean delivery among women having second births in Scotland, 1992–1998. Subsequent studies have yielded heterogeneous results but have employed data, analytic approaches and interpretation of variable quality.

Methods/Results  We replicated our previous methods and analysed 128,585 eligible singleton second births between 1999 and 2008. There were 88 stillbirths among 23,683 women with a previous caesarean (2.33 per 10,000 women per week) and 288 stillbirths in 104,897 women who previously delivered vaginally (1.67 per 10,000 women per week, P = 0.002). When analysed by cause, women with a previous caesarean had an increased risk (hazard ratio [95%CI] P) of unexplained stillbirth (1.47 [1.12–1.94] P = 0.006) and the excess risk was apparent from 34 weeks onwards (1.75 [1.23–2.49] P = 0.002). When the analysis was confined to 96,983 women with linked records from the first and second pregnancy (confirming exact mode of previous delivery) the association was stronger (2.12 [1.55–2.88] P < 0.001). Adjustment for maternal characteristics and first pregnancy complications had a minimal effect (1.97 [1.43–2.72] P < 0.001). The association was similar whether the previous caesarean was performed prior to labour (2.1 [1.24–3.80] P = 0.007) or during labour (2.50 [1.53–4.08] p < 0.001) and when the analysis was confined to previous term births (2.55 [1.50–3.58] p < 0.001).

Conclusion  We confirm that previous finding that previous caesarean delivery is a risk factor for unexplained stillbirth. The association is independent of maternal characteristics, obstetric outcome or the indication for the caesarean delivery.

PP.14  PROGESTERONE MODULATES CERVICAL ANTIMICROBIAL IMMUNITY
doi:10.1136/archdischild-2013-303966.295
1CP James, N Klein, *M Bajaj-Elliott, D Peebles. 1UCL Institute for Women’s Health, London, UK; *UCL Institute of Child Health, London, UK

Introduction  Human beta defensins (HBD1, 2.3) have antimicrobial and immunomodulatory properties and are components of the cervical mucus plug. Vaginal progesterone delays delivery in women with cervical shortening, but the mechanism(s) underlying this effect remain undetermined. This study describes the expression of HBDs by cervical epithelia in response to stimulation with classical infective and inflammatory agonists and progesterone.

Methods  The human endo-cervical cell-line End1/E6E7 and ectocervical cell-line Ect1/E6E7 were stimulated with bacterial (Lipo polysaccharide, LPS, Peptidoglycan, PGN) and inflammatory (Interleukin 1 beta, IL-1B; Interferon gamma, IFNγ) agonists and progesterone for up to 24 hours. HBD secretion was assessed by ELISA.
**Introduction** Sickle cell disease (SCD) in pregnancy has been associated with adverse pregnancy outcomes. However, with a multidisciplinary approach and surveillance, there is a suggestion of improvement in the overall outcome. The aim of this study was to evaluate the pregnancy outcomes in women with known SCD in a singleton pregnancy seen in our clinic, and to assess whether the presence of SCD alters the maternal serum analytes (Free B-human chorionic gonadotrophin and Pregnancy-associated plasma protein -A) at combined screening.

**Methods** Case-control study of all pregnant women with SCD undergoing combined screening from 01/01/2008 to 31/12/2011. Each case was matched with 3 non-SCD controls. Pregnancy outcomes in the two groups were compared.

**Abstract PP16 Table**

<table>
<thead>
<tr>
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<th>SCD (n = 54)</th>
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<td>Gestation at delivery, wks, median (IQR)</td>
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<td>Birth weight (grammes), median (IQR)</td>
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<td>B-HCG MoM, median (IQR)</td>
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*P < 0.05

**Conclusion** Women with SCD, when followed closely from the first trimester, have successful pregnancy outcomes. However, our data suggests an increased incidence of pre-eclampsia and lower birth weights compared to non-SCD controls. There was also a significant difference noted in the maternal serum analytes. Larger prospective studies are therefore required to assess the impact of SCD on combined screening.

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**PP17 PRE-GESTATIONAL DIABETES AND THE RISKS OF FETAL AND INFANT DEATH IN NORMALLY-FORMED OFFSPRING**

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