

295 INTERNAL ANAL SPHINCTER INJURY PREDICTS CONTINENCE OUTCOME FOLLOWING OBSTETRIC SPHINCTER TRAUMA RHONA MAHONY¹, LESLIE DALY², MICHAEL BEHAN³, CATRIONA KIRWAN³, COLM O'HERLIHY¹, RONAN O'CONNELL⁴, ¹University College Dublin, Obstetrics and Gynaecology, Dublin, Ireland, ²University College Dublin, Public Health Medicine and Epidemiology, Dublin, Ireland, ³University College Dublin, Radiology, Dublin, Ireland, ⁴University College Dublin, Surgery, Dublin, Ireland, Ireland

OBJECTIVE: The correlation between obstetric anal sphincter injury, as defined by endonal ultrasound, and altered postpartum fecal continence is weak. To improve the predictability of fecal incontinence symptoms postpartum, we assessed a cohort of 500 women following third degree perineal tear, with a view to identifying a risk profile predictive of debilitating postnatal fecal incontinence.

STUDY DESIGN: Five hundred consecutive patients, who attended our Perineal Clinic at three months postpartum following primary repair of a recognized third degree tear, constituted our study group. Their evaluation included continence assessment, anal manometry and endoanal ultrasound. Severe incontinence was defined as a modified Wexner score of >9/20. The relationship between various potential risk factors and the presence of severe incontinence symptoms was examined using univariate and multivariate analysis.

RESULTS: On univariate analysis, increasing maternal age and parity, instrumental delivery and a manometric resting pressure <35 mm Hg significantly correlated with the presence of severe symptoms (*P* < .05). On multiple regression analysis, a sonographic internal anal sphincter (IAS) defect was significantly predictive of severe incontinence (*P* < .05, OR7.1), while neither external anal sphincter defects nor manometric squeeze pressures correlated significantly.

CONCLUSION: IAS defects identified endosonographically at three months postpartum were highly predictive of severe fecal incontinence symptoms, likely to significantly impair quality of life.

296 D-DIMER LEVELS DURING DELIVERY AND THE POSTPARTUM PERIOD MANUELLA EPINEY¹, FRANCOISE BOEHLER², MICHEL BOULVAIN¹, GUIDO REBER², ERIC ANTONELLI¹, MICHEL-ANGE MORALES¹, PHILIPPE DE MOERLOOSE², OLIVIER IRION¹, ¹University of Geneva, Gynecology and Obstetrics, Geneva, Switzerland, ²University of Geneva, Angiology and Hemostasis, Geneva, Switzerland

OBJECTIVE: To define reference intervals for D-dimer in the postpartum period. To determine when D-dimer levels return to values below 500 ng/ml, which have a high predictive value to rule out venous thromboembolism.

STUDY DESIGN: We included 150 women delivering at term, either vaginally (n = 100) or by cesarean section (n = 50) after an uncomplicated pregnancy. D-dimer levels were measured at the end of pregnancy, delivery, day 1, 3, 10, 30 and 45 postpartum (VIDAS DD NEW assay).

RESULTS: D-dimer levels are significantly higher in women with cesarean section or instrumental vaginal delivery than with spontaneous delivery. All measurements were above 500 ng/ml at term, delivery, day 1 and day 3 postpartum. There was a marked elevation at delivery, especially when instrumental. An increase in D-dimer levels ten days after delivery was observed, followed by a progressive decrease at day 30 and 45. The number of women with D-dimer levels below 500 ng/ml is very low up to 30 days after delivery; at day 30 and day 45, respectively, 79% and 93% of women in the vaginal delivery group and 70% and 83% in the cesarean group had levels below 500 ng/ml. Bleeding, breastfeeding and heparin prophylaxis did not significantly modify D-dimer levels.

CONCLUSION: We provide reference intervals for D-dimer for the postpartum period. Cesarean section and instrumental delivery were associated with significantly higher levels. Using a cut-off at 500 ng/mL, D-dimer measurements cannot be used for excluding venous thromboembolism for about four weeks postpartum.

Median D-dimer levels (ng/mL; 5th-95th percentiles) at selected sampling days

	At term	Day 0	Day 3	Day 10	Day 45
SVD(n = 80)	1385 (689-2450)	3641 (1698-8501)	1203 (707-2102)	1214 (536-3079)	241 (123-554)
IVD(n = 20)	1489 (993-3542)	4986 (2006-9130)	1521 (993-2706)	1668 (1106-3199)	223 (144-680)
CS(n = 50)	1544 (735-2455)	3432 (1167-10168)	1542 (678-2602)	2061 (1028-3863)	279 (159-636)

SVD, Spontaneous vaginal delivery; IVD, instrumental; CS, cesarean section.

297 THYROID STIMULATING HORMONE AND FREE THYROXINE IN PREGNANCY-ARE LEVELS REALLY UNCHANGED? JOSEPH BIGGIO JR¹, JOHN MAHAN¹, ¹University of Alabama at Birmingham, Obstetrics/Gynecology, Birmingham, Alabama

OBJECTIVE: Based on data derived from older generation assays, the levels of thyroid stimulating hormone (TSH) and free thyroxine (fT4) are believed to be unchanged during pregnancy. With the advent of more precise assays, we sought to determine whether this postulate is accurate.

STUDY DESIGN: Excess clinical samples were obtained from four groups of asymptomatic patients: Non-pregnant, 1st (1 TRI), 2nd (2 TRI), and 3rd trimester (3 TRI). TSH (3rd generation assay, mIU/mL) and fT4 (ng/dL) measurements were performed on a Beckman Coulter Access (Brea, CA) using commercially available chemiluminescent assays. Data were analyzed for each group and for all pregnant women in composite. The 95% central interval of distribution (CID) was used to derive reference intervals (RI). The utility of the non-pregnant RI as a normal range for pregnant women was examined.

RESULTS: Values are displayed in the tables below (mean ± SD). Application of the non-pregnant RI for TSH (0.09-5.32) resulted in acceptable test performance: 4% of results were abnormal. However, had the non-pregnant RI for fT4 (0.64-1.34) been applied to the interpretation of levels from pregnant women, 44 % would have been deemed abnormal.

CONCLUSION: The levels of TSH and fT4 are affected by pregnancy and vary throughout gestation. Interpretation of these results using non-pregnant ranges may lead to misdiagnosis; consideration should be given to establishing pregnancy-specific reference ranges, especially for fT4.

TSH and fT4: Pregnant versus non-pregnant

	Pregnant(n = 667)	Non-pregnant(n = 100)	<i>P</i> value
TSH	1.29 ± 0.9	1.69 ± 1.1	< .001
95% CID	0.06 - 3.78	0.09 - 5.32	
fT4	0.67 ± 0.2	0.83 ± 0.2	< .001
95% CID	0.46 - 0.95	0.64 - 1.34	

TSH and fT4: By trimester

	1st TRI(n = 208)	2nd TRI(n=355)	3rd TRI(n=104)	<i>P</i> value
TSH	1.06 ± 0.8	1.15 ± 0.8	1.87 ± 1.2	< .001
95% CID	0.04 - 2.80	0.18 - 4.02	0.15 - 4.17	
fT4	0.74 ± 0.2	0.64 ± 0.1	0.63 ± 0.1	< .001
95% CID	0.55 - 1.25	0.45 - 0.85	0.44 - 0.90	

298 ACUTE MYOCARDIAL INFARCTION DURING PREGNANCY AND POSTPARTUM ANDRA JAMES¹, MARGARET JAMISON¹, GEETA SWAMY¹, EVAN MYERS¹, ¹Duke University, Obstetrics and Gynecology, Durham, North Carolina

OBJECTIVE: The purpose of this study was to determine the incidence, mortality and risk factors for acute myocardial infarction (MI) during pregnancy and postpartum.

STUDY DESIGN: The Nationwide Inpatient Sample (NIS), from the Healthcare Cost and Utilization Project of the Agency for Healthcare Research and Quality (AHRQ), for 2000-2001 was queried for all pregnancy-related discharges. The records were classified as antepartum, delivery-related or postpartum. The ICD9 code 410 was used for acute MI. Counts, rates and standard errors were calculated using methods accounting for the survey design of the NIS.

RESULTS: There were 525 records with a diagnosis of acute MI or 6.3 events per 100,000 deliveries. 46% were antepartum, 28% were delivery-related and 26% were postpartum. There were 25 deaths from MI for a case fatality rate of 4.8%. The risk of MI increased with age. Women younger than 20 had 0.4 events per 100,000 deliveries while women 40 and older had 27.6. Due to the limited number of cases for analysis, no statistically significant difference existed between racial groups, but black women, with 9.9 events per 100,000 deliveries, appeared to be at greater risk than those from other groups. Statistically significant risk factors are listed in the table.

CONCLUSION: Acute MI is a rare event in pregnancy. Risk factors include not only recognized risk factors for coronary artery disease, but also complications of labor and delivery such as postpartum infection, postpartum hemorrhage and transfusion.

Risk factors for myocardial infarction

Risk factor	Odds ratio	Confidence interval
Hypertension	8.9	5.0, 16.0
Thrombophilia	11.7	2.9, 48.0
Smoking	5.9	4.2, 11.8
Anemia	1.9	1.1, 3.3
Postpartum infection	7.9	2.4, 25.4
Postpartum hemorrhage	3.6	1.9, 6.8
Transfusion	11.5	4.5, 29.7