Background and aims Congenital malformations (CM) are defined as abnormal structure of the organism resulting from disrupted embryogenesis. Many factors influence the appearance of CM. Regarding different criteria and authors, the incidence of CM at newborns is between 2−7%. The aim of this paper was to determine the incidence of CM at liveborns delivered at SHGO “Mother Teresa” – Cair, Skopje, during the five years period (2000–2004) and five years period (2005–2009). Also, the distribution among organ system had been analysed.

Methods A retrospective analysis of 19097 liveborns delivered at SHGO Cair, during 2000–2004, has been performed. Database (Access 2000) from Neonatal Unit has been used. The incidence and percentage of CM among different systems have been determined and a retrospective analysis of 15293 liveborns delivered at SHGO Cair, during 2005–2009, has been performed. Database (Access 2005) from Neonatal Unit has been used. The incidence and percentage of CM among different systems have been determined.

Results Period 2000–2004, among a total of 19097 liveborns, 736 or 3.85% have had CM. The incidence between different years was:4.28% in 2000, 3.92%−2001, 3.79% 2002, 3.20%−2003 and 3.98% in 2004. Regarding different organ systems the distribution was: 58.1% of all CM were the anomalies of musculoskeletal system, 14.6% - cardiopathies, 8.1% - anomalies of urogenital system, 7.2% -CM of gastro intestinal system, etc. Perid 2005–2009, among a total of 15293 liveborns, 573 or 3.75% have had CM. The incidence between different years was:4,02% in 2005, 3.42%−2006, 3.22%− 2007, 4.23%−2008 and 3.85% in 2009. Regarding different organ systems the distribution was: 40,31% of all CM were the anomalies of musculoskeletal system, 21,81% - cardiopathies, 20,07% - anomalies of urogenital system, 12,04% -CM of gastro -intestinal system, 5,76% of CNS, etc.

Conclusions During the five years period (2000–2004), the incidence of CM 3.85%, which is in accordance with data from literature. Among years, the incidence varies from 3.2 to 4.3%. During the five years period 2005–2009, the incidence of CM 3.75%, which is in accordance with data from literature. Among years, the incidence varies from 3.2 to 4.3%. The anomalies of musculoskeletal system are the most frequent, followed by those from cardiovascular, uro-genital, gastrointestinal and central nervous system. Comparing the five-year periods 2000–2004 and 2005–2009 shows almost identical incidence. Congenital malformations still remain an important medical and social problem requesting more serious nationwide engagement, as in medical aspect, socioeconomic, ecological, etc.

Background and aims The American Academy of Paediatrics recommends the hour-specific evaluation of either skin (SB) or total serum bilirubin as screening tool for neonatal jaundice. However, serum bilirubin is not a good predictor of kernicterus, while SB represents a molecular species already extravasated and passed into a tissue. Circulating unbound bilirubin (UB) is actually related to the neurotoxicity, being the portion of bilirubin capable to freely pass from circulation to the tissues. We aimed at exploring the relationship between SB and UB. Thus, simultaneous measurements of SB (using 2nd generation transcutaneous devices) and UB were performed in 35 term jaundiced neonates.

Results In term neonates (mean GA: 38 weeks; BW 3095 g; postnatal age 74,8 h; TSB 250.8 µmol/L (SD 63); UB 0.48 (SD 0.2) µg/dL), a positive correlation was found between SB and UB (r = 0.70; p < 0.001; Figure 1). This correlation remained significant after adjustment for birth weight, gestational or postnatal age (partial correlation r = 0.68, p < 0.001; r = 0.69, p < 0.001; r = 0.64, p < 0.001).

Abstract PO-0693 Figure 1

Background and aims Congenital malformations (CM) are defined as abnormal structure of the organism resulting from disrupted embryogenesis. Many factors influence the appearance of CM. Regarding different criteria and authors, the incidence of CM at newborns is between 2−7%. The aim of this paper was to determine the incidence of CM at liveborns delivered at SHGO “Mother Teresa” −Cair, Skopje, during the five years period (2000−2004) and five years period (2005−2009). Also, the distribution among organ system had been analysed.

Methods A retrospective analysis of 19097 liveborns delivered at SHGO Cair, during 2000−2004, has been performed. Database (Access 2000) from Neonatal Unit has been used. The incidence and percentage of CM among different systems have been determined and a retrospective analysis of 15293 liveborns delivered at SHGO Cair, during 2005−2009, has been performed. Database (Access 2005) from Neonatal Unit has been used. The incidence and percentage of CM among different systems have been determined.

Results Period 2000−2004, among a total of 19097 liveborns, 736 or 3.85% have had CM. The incidence between different years was: 4.28% in 2000, 3.92%−2001, 3.79% 2002, 3.20%−2003 and 3.98% in 2004. Regarding different organ systems the distribution was: 58.1% of all CM were the anomalies of musculoskeletal system, 14.6% - cardiopathies, 8.1% - anomalies of urogenital system, 7.2% -CM of gastrointestinal system, etc. Period 2005−2009, among a total of 15293 liveborns, 573 or 3.75% have had CM. The incidence between different years was: 4.02% in 2005, 3.42%−2006, 3.22%−2007, 4.23%−2008 and 3.85% in 2009. Regarding different organ systems the distribution was: 40.31% of all CM were the anomalies of musculoskeletal system, 21.81% - cardiopathies, 20.07% - anomalies of urogenital system, 12.04% -CM of gastrointestinal system, 5.76% of CNS, etc.

Conclusions During the five years period (2000−2004), the incidence of CM 3.85%, which is in accordance with data from literature. Among years, the incidence varies from 3.2 to 4.3%. During the five years period 2005−2009, the incidence of CM 3.75%, which is in accordance with data from literature. Among years, the incidence varies from 3.2 to 4.3%. The anomalies of musculoskeletal system are the most frequent, followed by those from cardiovascular, uro-genital, gastrointestinal and central nervous system. Comparing the five-year periods 2000−2004 and 2005−2009 shows almost identical incidence. Congenital malformations still remain an important medical and social problem requesting more serious nationwide engagement, as in medical aspect, socioeconomic, ecological, etc.

**THE VALUE OF RISK FACTORS IN SCREENING FOR DEVELOPMENTAL DYSPLASIA OF THE HIP (DDH). CAN SELECTIVE SCREENING REPLACE UNIVERSAL SCREENING?**

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**Background** Ultrasonography is accepted as a reliable tool in detecting DDH. Universal screening is criticised to be financially demanding, time consuming and leads to overtreatment. Selective ultrasonography is proposed as the alternative.

**Methods** A retrospective study of all newborn in a period of 2 years. All newborn with risk for DDH or positive clinical signs were screened with ultrasonography (selective screening). Data of babies who presented after the age of 6 months with DDH.

**Results** Between May 2008 and April 2010, 33768 live births were registered. 966 newborn had risk for DDH, 532 with breech presentation, 81 had positive family history, 440 were premature, 191 product of multiple pregnancy, 67 had foot abnormality, 3 with torticollis, and 7 with other anomalies.

12 babies who were born in the same period and were not screened because they had neither risk for DDH or positive clinical finding presented after the age of 6 months with dislocated hips. In screened group 15 newborn could be discovered early with DDH, whereas 12 were missed and presented late.

**Conclusions** Only 56% of newborn with DDH could be detected with the selective screening program, which is statistically not acceptable, and selective screening program in our hand is not an adequate tool to detect DDH. Risk for DDH can detect only 1 out 9 with DDH only, and cannot be considered alone as tool for screening.

**PO-0693 UNBOUND BILIRUBIN CORRELATES WITH SKIN BILIRUBIN MEASURED IN JAUNDICED NEONATES**

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**Background and aims** The American Academy of Pediatrics recommends the hour-specific evaluation of either skin (SB) or total serum bilirubin as screening tool for neonatal jaundice. However, serum bilirubin is not a good predictor of kernicterus, while SB represents a molecular species already extravasated and passed into a tissue. Circulating unbound bilirubin (UB) is actually related to the neurotoxicity, being the portion of bilirubin capable to freely pass from circulation to the tissues. We aimed at exploring the relationship between SB and UB.

**Methods** Simultaneous measurements of SB (using 2nd generation transcutaneous devices) and UB were performed in 35 term jaundiced neonates.

**Results** In term neonates (mean GA: 38 weeks; BW 3095 g; postnatal age 74.8 h; TSB 250.8 µmol/L (SD 63); UB 0.48 (SD 0.2) µg/dL), a positive correlation was found between SB and UB (r = 0.70; p < 0.001; Figure 1). This correlation remained significant after adjustment for birth weight, gestational or postnatal age (partial correlation r = 0.68, p < 0.001; r = 0.69, p < 0.001; r = 0.64, p < 0.001).
Conclusions These preliminary data suggest that SB is strongly correlated to UB. SB could be related to neurotoxicity as it may be formed by UB passed from circulation to a tissue. Further investigations are needed to clarify this relationship and possible influencing factors.

Abstract PO-0694 Figure 1

Conclusions The use of LR is better than GA and BW in predicting M of VLBWs. RAM can be used as a tool for quality improvement.

Abstract PO-0695 Figure 1

**Background and aims** Preterm babies have higher mortality than terms. Risk-adjusted mortality (RAM) is useful for making comparisons among different NICUs. GA, BW, sex, singleton birth and antenatal steroid have been used to estimate mortality (M) of preterm. The aims of this study are 1. To compare the performance of GA, BW, and Logistic Regression (LR) in predicting M of VLBW infants. 2. To compare the RAM in different areas and periods.

**Methods** Cohort data from 2000 to 2011 were used. M is defined as death prior to discharge. Exclusion criteria included 1) Transferred after 24 h of age; 2) Death within 24 h of admission and 3) Lethal malformation. We developed a LR model to predict M [expected probability (Pro)]. ROC curves were used for assessing performance of predicting M. To compare the RAM, we calculated (O-E) Pro (observed Pro – expected Pro) values in each patient and used these values for comparisons.

**Results** 9207 VLBWs were enrolled. The calculated probability of death by LR model was: $P = \frac{1}{1+e^{-z}}$, where $e$= natural logarithms and $z = (-0.62*\text{[prenatal steroid]} - (0.219*\text{GA}) - (0.004*\text{BBW}) - (0.327*\text{singleton}) + (0.286*\text{[male]}) + 8.438$. Area under ROC were 0.858 for LR (95% CI: 0.847–0.869), 0.841 for BBW (95% CI: 0.829–0.853) and 0.827 for GA (95% CI: 0.815–0.839).

There were significant differences of RAM in different locations and years (Figure 1).