**ORIGINAL ARTICLE**

**Neonatal bilirubin production-conjugation imbalance: effect of glucose-6-phosphate dehydrogenase deficiency and borderline prematurity**

M Kaplan, M Muraca*, H J Vreman, C Hammerman, M T Vilei, F F Rubaltelli, D K Stevenson


**Objective:** To evaluate relations between production and conjugation of bilirubin in the pathophysiology of jaundice in glucose-6-phosphate dehydrogenase (G6PD) deficient neonates.

**Methods:** Term and borderline premature (35–37 weeks gestational age), healthy, male, G6PD deficient neonates were studied close to the beginning of the 3rd day. Blood carboxyhaemoglobin corrected for inspired CO (COHbc; an index of bilirubin production) and serum total conjugated bilirubin (TCB; a reflection of bilirubin conjugation) were measured in simultaneously drawn blood samples by gas chromatography and reverse phase high performance liquid chromatography respectively. A bilirubin production-conjugation index comprising COHbc/TCB was determined; a high index reflects imbalance between the bilirubin production and conjugation processes. COHbc and TCB individually and the production-conjugation index were studied in relation to serum total bilirubin (STB) concentration.

**Results:** Fifty one G6PD deficient neonates were sampled at 51 (8) hours. COHbc values did not correlate with STB \( (r = 0.22, p = 0.15) \). TCB did correlate inversely with STB \( (r = 0.42, p = 0.004) \), and there was a positive correlation between the production-conjugation index and STB \( (r = 0.45, p = 0.002) \). The production-conjugation index (median [interquartile range]) was higher in the premature \( (n = 8) \) than term neonates \( (2.31 [2.12–3.08] v 1.05 [0.53–1.81], p = 0.003) \). This difference was the result of changes in TCB.

**Conclusions:** The data show that jaundice in G6PD deficient neonates is the result of an imbalance between production and conjugation of bilirubin with a tendency for inefficient bilirubin conjugation over increased haemolysis in its pathogenesis. Borderline premature infants are at special risk of bilirubin production-conjugation imbalance.

Both increased production of bilirubin and diminished elimination from the body, primarily by conjugation, contribute to the mechanism of neonatal jaundice. Equilibrium between these processes should result in a degree of jaundice that does not exceed the physiological range. However, should the rate of bilirubin production exceed the body’s capacity to conjugate it, serum total bilirubin (STB) concentrations are expected to increase, and hyperbilirubinaemia may ensue.1

The bilirubin production and conjugation processes can be evaluated using blood carboxyhaemoglobin measurements, corrected for ambient carbon monoxide (CO) to give COHbc, and serum total conjugated bilirubin (TCB) respectively. The principle behind the former test is that, for each molecule of biliverdin, and subsequently bilirubin, produced from haem catabolism by the enzyme haem oxygenase, equimolar quantities of CO are produced, and COHbc values will reflect the rate of haem catabolism.2 As a small fraction of conjugated bilirubin effluxes from the hepatocyte into the plasma, the amounts of total conjugated bilirubin (TCB) measurable in the plasma or serum are regarded as a reflection of conjunctive capacity.3,4 Thus, in infants with efficient bilirubin conjugating capacity, greater quantities of conjugated bilirubin, relative to the total bilirubin load, should be formed than in those with less effective conjugation. The combined effect of bilirubin production and its conjugation has been represented mathematically using a “production-conjugation index” (unitless) comprising COHbc divided by TCB.4 Accordingly, low bilirubin production along with efficient bilirubin conjugation can be expected to result in low COHbc and high TCB and therefore a low index. Conversely, high bilirubin production relative to conjugation should result in a raised index.

Two risk factors highly associated with neonatal hyper-bilirubinaemia with the potential of bilirubin encephalopathy or kernicterus include G6PD deficiency and borderline prematurity (35–37 weeks gestation). The former is a common X linked enzymatic deficiency, which may nowadays be encountered world wide.5 The enzyme plays an important role in antioxidant mechanisms. In some instances, overwhelming haemolysis akin to favism may result in hyperbilirubinaemia.6 However, in many G6PD deficient neonates, only moderately raised haemolysis has been detected by COHbc techniques, and no correlation been found between COHbc and STB.7 In contrast, there appears to be a predilection for diminished bilirubin conjugation in the mechanism of jaundice in the G6PD deficient state.8,9

With regard to borderline prematurity,10 activity of the bilirubin conjugating enzyme, UDP-glucuronosyltransferase 1A1 (UGT), has been shown to increase in concert with gestational age.11 Incomplete maturation of this enzyme in premature infants may therefore result in diminished bilirubin conjugation, placing these neonates at especial risk of hyperbilirubinaemia.

We quantified COHbc and unconjugated and conjugated serum bilirubin fractions in blood and serum sampled simultaneously from G6PD deficient neonates, to enable

**Abbreviations:** COHb, carboxyhaemoglobin; COHbc, COHb corrected for inspired (room air) CO; G6PD, glucose-6-phosphate dehydrogenase; HPLC, high performance liquid chromatography; STB, serum total bilirubin; TCB, total conjugated bilirubin

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being cared, representing inspired air, was collected and a sample of room air from the nursery in which the baby was subsequently shipped on dry ice to the University of Padua, The study cohort comprised healthy consecutive neonates of this cohort.56 The study cohort comprised healthy consecutive neonates of this cohort.56 The study protocol, approved by the institutional review board of the Shaare Zedek Medical Center, was identical with that used previously for G6PD normal neonates of this cohort.14 The study cohort comprised healthy consecutive neonates delivered at or near term at the Shaare Zedek Medical Center all of whom were tested for G6PD deficiency. In this study, we analysed those found to be G6PD deficient. Only male infants were included, to avoid the problem of designation of G6PD status may be difficult.7 Infants with female G6PD deficient heterozygotes for whom accurate allele specific oligonucleotide hybridisation, as described elsewhere.13 was determined at The Scripps Research Institute, La Jolla, California, USA (by Ernest Beutler) by polymerase chain reaction followed by polymerase chain reaction followed by allele specific oligonucleotide hybridisation.

DNA was prepared from peripheral leucocytes by a high salt extraction method.22 The G6PD Mediterranean C563T mutation, which is encountered in Sephardic Jews,13 23 was determined at The Scripps Research Institute, La Jolla, California, USA (by Ernest Beutler) by polymerase chain reaction followed by allele specific oligonucleotide hybridisation, as described elsewhere.11

The relative roles of increased haem catabolism and diminished bilirubin conjugation in the pathogenesis of neonatal bilirubinaemia were compared by correlating COHbc and TCB respectively with STB, using multivariate analysis. The effects of these processes in combination were evaluated by calculating the bilirubin production-conjugation index, COHbc/TCB, and testing these values for correlation with STB using linear regression analysis. To confirm these findings, the value for the 90th centile for COHbc, TCB, and STB of neonates with values of each of the three variables >90th centile were compared with those <90th centile using Student’s t test. The effect of borderline prematurity on the production-conjugation index and its constituents was studied by comparing values for neonates <37 weeks gestational age with those >37 weeks, using Student’s t test or the Mann-Whitney rank sum test, as appropriate, for stored in a special container designed for this purpose pending determination of its CO concentration (Bistable Gas Sampler; Chemical Projects Limited, Toronto, Ontario, Canada).

COHb was determined by a sensitive and accurate gas chromatographic method, and its concentrations expressed as a percentage of total haemoglobin, which was quantified by the cyanmethaemoglobin method, both as previously described.10 17 The within day and between day coefficients of variation for reference blood samples with this method are 3% and 8% respectively.16 The CO concentration of the specimens of room air was determined at the Shaare Zedek Medical Center using a sensitive electrochemical CO analyser supplied by Stanford University.17 These CO concentrations were used to correct measured COHb for the effect of inspired CO (COHbc) by a previously derived formula (COHbc = measured COHb − 0.17 µl CO/litre room air).20

Unconjugated and conjugated bilirubin fractions were quantified using alkaline methanolysis followed by reverse phase high performance liquid chromatography (HPLC), by the method of Muraca and Blanckaert.21 For this method, the within day coefficient of variation is 5–8%, and the between day coefficient of variation is 6–13%. The sum of the measured concentrations of serum unconjugated, monoconjugated, and diconjugated bilirubin was defined as serum total bilirubin (STB), and that of the monoconjugated and diconjugated bilirubin concentrations comprised the total conjugated bilirubin (TCB) value. The latter was expressed as a percentage of the STB concentration, as explained elsewhere.2

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A sample of blood was drawn near the beginning of the third day of life. This time was chosen to coincide with routine metabolic screening, avoiding the need to take blood expressly for this study. The study blood sample was divided into aliquots: whole blood for COHb determination (150 µl) was collected into custom made capillary tubes containing heparin and saponin, as previously described16 17 and stored in a special container designed for this purpose pending determination of its CO concentration (Bistable Gas Sampler; Chemical Projects Limited, Toronto, Ontario, Canada).

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Figure 2 Regression analysis between production-conjugation index and serum total bilirubin (STB).

Figure 1 (A) Regression analysis between blood carboxyhaemoglobin corrected for inspired (room air) carbon monoxide (COHbc) and serum total bilirubin (STB). (B) Regression analysis between total conjugated bilirubin (TCB) and STB. Hb, Haemoglobin.
values with a parametric or non-parametric distribution respectively. Parametric data are presented as mean (SD), and non-parametric data as median (interquartile range). Significance is defined as p<0.05.

RESULTS
A total of 51 G6PD deficient neonates was sampled at 51 (8) hours of age. Gestational age was 39 (1) weeks, and birth weight was 3118 (462) g. Eight were born at ≥37 weeks gestation. Ten (20%) were delivered by caesarean section, 26 (51%) were exclusively breast fed, and a further nine (18%) received a combination of breast feeding and formula. Sixteen (31%) developed hyperbilirubinaemia, defined as a plasma total bilirubin ≥256 μmol/l (15.0 mg/dl).

Measured values for the entire G6PD deficient neonatal population included mean (HPLC) STB concentration 144 (67) μmol/l, COHbc level 0.83 (0.19)%, and median (interquartile range) TCB value 0.65% (0.39–1.27%). Multivariate analysis with STB as the dependent variable and COHbc and TCB as independent variables showed no correlation with COHbc, but did show correlation with TCB (r = 0.50, p = 0.09, and p = 0.003 for COHbc and TCB respectively). Figure 1 shows the correlation analysis between STB, and COHbc, but did show correlation with TCB (r = 0.50, p = 0.09, and p = 0.003 for COHbc and TCB respectively). Table 1 gives values for the 90th centile of COHbc, TCB, and STB for neonates >37 weeks gestation and those ≤37 weeks gestation.

Table 1: Comparison of STB concentrations for respective COHbc, TCB, and production conjugation index values >90th centile versus <90th centile

<table>
<thead>
<tr>
<th>Variable</th>
<th>≥90th centile</th>
<th>&lt;90th centile</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>COHbc (90th centile 1.09)</td>
<td>134 (49)</td>
<td>146 (70)</td>
<td>0.7</td>
</tr>
<tr>
<td>TCB (90th centile 2.65)</td>
<td>67 (68)</td>
<td>154 (62)</td>
<td>0.005</td>
</tr>
<tr>
<td>Production-conjugation index (90th centile 3.0)</td>
<td>205 (94)</td>
<td>137 (61)</td>
<td>0.03</td>
</tr>
</tbody>
</table>

Values are mean (SD). STB, Serum total bilirubin; COHbc, carboxyhaemoglobin corrected for inspired carbon monoxide; TCB, total conjugated bilirubin.

DISCUSSION
This study differs from previous ones of this nature in that COHbc and TCB were analysed in blood and serum sampled simultaneously from the same patients, allowing direct comparison between the bilirubin production and conjugation processes at the same point in time, and thereby providing insight into the physiological relations between production and conjugation in these infants. Furthermore, whereas in previous studies COHbc in G6PD deficient neonates was analysed in relation to STB concentrations determined by the diazo method, we used a highly accurate reverse phase HPLC method, which allowed determination of both unconjugated and conjugated bilirubin fractions.

The G6PD deficient neonates of this study differ from the previously reported G6PD normal counterparts drawn from the same cohort in that STB concentrations were higher, and the risk of hyperbilirubinaemia greater. For reference, STB concentrations were 144 (67) μmol/l v 114 (67) μmol/l (p = 0.003), and the incidence of clinically determined hyperbilirubinaemia (plasma bilirubin ≥ 256 μmol/l by the diazo method) was 31% v 6% (p<0.001). The study design, by which measurements reflecting haem catalolysis and bilirubin conjugation were quantified from simultaneously drawn samples from identical babies, facilitated comparison of the roles of these processes in the pathophysiology of jaundice. As previously shown using serum bilirubin determinations obtained by the diazo method, COHbc did not correlate with HPLC determined STB concentrations in the G6PD deficient neonates. Correspondingly, COHbc values ≥90th centile were not associated with STB concentrations greater than those with COHbc values <90th centile, as would be expected in a G6PD normal population. In contrast, TCB did diminish progressively along with increasing STB concentration, whereas STB concentrations in neonates with TCB bilirubin concentrations ≥90th centile, representing those with the most efficient bilirubin conjugation, were significantly lower in this group. Although STB concentrations were not significantly different between the two groups, there was a trend to a weak correlation between diminishing gestational age and increasing STB concentration (r = 0.28, p = 0.06).

Table 2: Comparison of values for the production-conjugation index, COHbc, TCB, and STB for neonates >37 weeks gestation and those ≤37 weeks gestation

<table>
<thead>
<tr>
<th>Variable</th>
<th>≤37 weeks gestation</th>
<th>&gt;37 weeks gestation</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Production-conjugation index (#unitless)</td>
<td>2.31 (2.12–3.08)</td>
<td>1.05 (0.53–1.81)</td>
<td>0.003</td>
</tr>
<tr>
<td>COHbc (% Hb)</td>
<td>0.88 (0.21)</td>
<td>0.82 (0.20)</td>
<td>0.46</td>
</tr>
<tr>
<td>TCB (% STB)</td>
<td>0.39 (0.31–0.42)</td>
<td>0.74 (0.44–1.69)</td>
<td>0.009</td>
</tr>
<tr>
<td>STB (μmol/l)</td>
<td>160 (35)</td>
<td>141 (72)</td>
<td>0.48</td>
</tr>
</tbody>
</table>

Values are mean (SD) or median (interquartile range), as appropriate. STB, Serum total bilirubin; COHbc, carboxyhaemoglobin corrected for inspired carbon monoxide; TCB, total conjugated bilirubin; Hb, total haemoglobin.
Neonatal Hyperbilirubinaemia. Furthermore, in a recent study by the American Academy of Pediatrics Subcommittee on hyperbilirubinaemia in G6PD deficient neonates. The production-conjugation index, reflecting imbalance between bilirubin production and its conjugation, increased in concert with increasing STB concentration, confirming the concept of imbalance between production and conjugation of bilirubin in the pathogenesis of jaundice in a neonatal subgroup with a higher than average incidence of hyperbilirubinaemia. Even when possible outliers with the two highest indices were removed, the correlation between STB and the index maintained significance (fig 2).

Borderline prematurity is an important contributing factor to hyperbilirubinaemia and kernicterus. The high risk status of neonates of 35–37 weeks gestation has been emphasised by the American Academy of Pediatrics Subcommittee on Neonatal Hyperbilirubinaemia. Furthermore, in a recent report of 61 neonates, originally discharged as healthy, but readmitted for kernicterus within the first seven days, 23% were of 35–36 weeks gestational age. In our study, although there was only a trend to a weak correlation between diminishing gestational age and STB concentration, production-conjugation indices, in contrast, were significantly higher in the near term subgroup. Further analysis pinpointed this difference to the TCB component of the index, implying diminished bilirubin conjugation, but not increased haem catabolism, in the near term neonates. Although the effect of borderline prematurity on STB concentration was minimal, that on the production-conjugation index was highly significant, and may contribute to our understanding of the high risk of near term neonates. The degree of imbalance between bilirubin production and conjugation at the time the neonates were sampled was insufficient to result in significantly higher STB concentrations. However, in a situation with either diminished bilirubin conjugation or increased haem catabolism, however slight, any additional or superimposed ichterogenic factors may have a much greater effect, with the potential of severe hyperbilirubinaemia, than if the bilirubin production and conjugation processes were working at normal capacity.

The COHbc measurements we used were obtained by state of the art technology. Red blood cell indices and haaptoglobin or haemopexin determinations are unreliable indicators of haemolysis in the newborn because of overlap between the normal and haemolytic states. As the predominant source of endogenous CO in the human body is from the degradation of haem by haem oxygenase, with little originating from non-haem sources, measurement of blood COHbc reflects primarily the rate of bilirubin formation from haemoglobin. The serum conjugated bilirubin is believed to reflect the intrahepatic bilirubin profile, as the liver is the only organ that can esterify bilirubin to any significant extent, and conjugated bilirubin compounds are not absorbed from the bowel. In the absence of hepatocellular disease or cholestasis, this method provides a minimally invasive assessment of bilirubin conjugation. The assay has been used for this purpose in neonates, children, and adults. In individual cases, variation in the balance between the rate limiting steps and mild disparities in bile flow may affect interpretation of the results; however, use of large numbers of subjects should offset this potential inaccuracy. The pathogenesis of neonatal jaundice is clearly multifactorial. COHbc and TCB studies evaluate some, but not all, of the contributing factors. Some of the latter include hepatic uptake, bilirubin excretion, and reabsorption of bilirubin via the enterohepatic circulation.

Arithmetic deficiencies of this index or its components include the facts that COHb is measured as a percentage of total Hb, but that catabolism of haem is what the COHbc is expected to reflect. TCB is expressed as a percentage of the STB, but it is the STB that is subsequently correlated with the production-conjugation index. Notwithstanding these mathematical dependencies and the already mentioned difficulties in interpretation, the index does contribute to our understanding of the mechanism of neonatal hyperbilirubinaemia. An increase in the index in line with rising STB concentration confirms the importance of the concept that imbalance between bilirubin production and its conjugation is a crucial factor in the mechanism of neonatal jaundice. Diminishing gestational age has a further effect on increasing the degree of imbalance. Additional genetic or environmental factors that either increase haemolysis or diminish bilirubin conjugation may exacerbate any imbalance between these processes, potentiating the risk of severe neonatal hyperbilirubinaemia.

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