Evaluating the beneficial and detrimental effects of bile pigments in early and later life

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INTRODUCTION

The generation of bile pigments occurs through a unique pathway for the degradation of heme, limited by the enzyme hemeoxygenase (HO). This enzymatic reaction requires molecular oxygen (O2) and NADPH as a reducing equivalent and results in the formation of biliverdin and the release of carbon monoxide (CO) in an equimolar ratio. In addition, reduction of heme iron from Fe2+–Fe3+ occurs with the transfer of electrons from oxygen with NADPH as a source of reducing equivalents. Also, oxidative cleavage of the α-methene carbon bridge in the heme molecule forms biliverdin, releases carbon monoxide and iron (Figure 1). In most circumstances, heme does not accumulate freely but rather, it is bound to hemo proteins that are essential for cellular metabolism. An intricate enzymatic cascade regulates the production of heme (Ryter and Tyrrell, 2000). In pathological conditions, free heme is released from hemoglobin and can deposit in tissues because it is lipophilic and can lead to the formation of oxygen radicals therefore, the enzymatic reaction of HO is essential to preventing this since HO-1 is highly inducible by the substrate heme or by oxidative stress (Chang et al., 2005). The constitutive isoenzyme HO-2 can also catalyze the degradation of heme and is found in high abundance in the brain and testes (Maines et al., 1986; Maines, 1988). In the last steps of the reaction, biliverdin reductase (BVR), a microsomal enzyme, converts biliverdin into bilirubin in a non-rate-limiting fashion (Figure 1). Bilirubin, unlike biliverdin is not water soluble, but rather lipophilic, and can penetrate cellular membranes. To be made water soluble and therefore excretable in the gastrointestinal tract, it must be conjugated. The latter is regulated by UDP-glucuronosyl transferase 1A1 (UGT1A1), an enzyme that adds two glucuronic acid residues to bilirubin to render it water soluble (Figure 1).

The heme degradation pathway has been conserved throughout phylogeny and allows for the removal of a pro-oxidant and the generation of unique molecules including bile pigments with important cellular functions. The impact of bile pigments on health and disease are reviewed, as is the special circumstance of neonatal hyperbilirubinemia. In addition, the importance of promoter polymorphisms in the UDP-glucuronosyl transferase gene (UGT1A1), which is key to the elimination of excess bilirubin and to the prevention of its toxicity, are discussed. Overall, the duality of bile pigments as either cytoprotective or toxic molecules is highlighted.

Keywords: neonatal jaundice, kernicterus, UDP-glucuronosyltransferase, antioxidant, polymorphisms

BILIRUBIN

EFFECTS OF BILIRUBIN IN THE NEONATE

Newborns have elevated numbers of red blood cells with a shortened life-span. When these cells lyse, heme is released from hemoglobin. In addition, due to a reduced ability to conjugate bilirubin formed during the degradation of heme, this pigment can accumulate in the serum in the first days of life leading to a transient hyperbilirubinemia, which typically resolves within the first weeks of life. In fact, the average full-term newborn infant has a peak serum bilirubin concentration of 5–6 mg/dL (86–103 µmol/L). This level is referred to as physiologic jaundice. However, in some circumstances, such as increased accumulation of heme (i.e., birth trauma, bruising, hemolysis), serum bilirubin levels can increase beyond the physiologic range. If serum bilirubin values are between 7 and 17 mg/dL (104–291 µmol/L), this is then referred to as exaggerated physiologic jaundice. However, this must be judged according to the infant’s age in hours on the bilirubin nomogram, as serum bilirubin levels change rapidly during the first weeks of life. Serum bilirubin concentrations higher than 17 mg/dL in full-term infants are considered pathologic and can be associated with adverse sequelae. Fortunately, most infants will not be affected until their bilirubin levels are significantly higher than 17 mg/dL because the toxicity of bilirubin is dictated by many factors including age (the younger, the more vulnerable), maturity (prematures are more vulnerable), and associated illnesses (hemolysis, sepsis, acidosis worsen bilirubin toxicity) amongst other factors. The most severe manifestation of bilirubin toxicity is kernicterus, a rare but devastating condition with acute
Investigations do confirm that bilirubin has significant antioxidant benefit to this reaction (McDonagh, 1990) and laboratory life and what is its value. Physiologic hyperbilirubinemia is a guide to preventing this problem by using age-based serum bilirubin thresholds (Bhutani et al., 2007; Bhutani et al., 1999). In the present day, we have adopted a more thoughtful approach to the preventable problem of hyperbilirubinemia (Maisels and Newman, 1983). In addition, reports of the beneficial, antioxidant effects of bilirubin (Stocker et al., 1987, 1990; McDonagh, 1990) made this aggressive approach seem even more unwarranted. Alas, with the risk of kernicterus increases because free bilirubin enters tissues and causes its toxic effects (Ahlfors et al., 2009). Conditions that alter the blood–brain barrier, such as infection, acidosis, hyperoxia, sepsis, prematurity, and hyperosmolarity, may also affect the entry of bilirubin into the brain (Dennery et al., 2001).

**MECHANISMS OF TOXICITY OF BILIRUBIN**

Bilirubin has high affinity for membrane phospholipids thereby entering cells and inhibiting mitochondrial enzymes (Chuniaud et al., 1996; Rodrigues et al., 2002a), interfering with DNA synthesis, inducing DNA-strand breaks (Rosenstein et al., 1983), and inhibiting protein synthesis and phosphorylation. In the brain, bilirubin inhibits the uptake of tyrosine, reduces synaptic transmission, and inhibits N-methyl-D-aspartate-receptor ion channels. Overall, bilirubin can interfere with neuro excitatory signals and impair nerve conduction particularly in the auditory nerve (Bratilid, 1990). Unconjugated bilirubin directly interacts with mitochondria influencing membrane lipid and protein properties, redox status, and cytochrome c content (Rodrigues et al., 2002b). It can work in concert with amyloid β peptide to activate apoptosis in neural cells (Rodrigues et al., 2000). Interestingly, younger animals are less susceptible to bilirubin-related mitochondrial injury (Rodrigues et al., 2002c) and the toxicity of bilirubin is not restricted to neonates. In patients with Crigler–Najjar I syndrome and absent activity of the UGTA1, therefore mitochondrial injury (Rodrigues et al., 2000) and the auditory nerve preferentially (Shapiro and Nakamura, 2001). This occurs if bilirubin is not bound to albumin or is unconjugated or if there has been damage to the blood–brain barrier. As an example, a newborn infant with a serum albumin concentration of 3 g/dL can bind 25 mg/dL of bilirubin. In current clinical practice, the bilirubin albumin ratio is taken into consideration to determine levels at which physicians should proceed to more aggressive management of hyperbilirubinemia (i.e., exchange transfusion). If the serum albumin concentration is low, the risk of kernicterus increases because free bilirubin enters tissues and causes its toxic effects (Ahlfors et al., 2009). Conditions that alter the blood–brain barrier, such as infection, acidosis, hyperoxia, sepsis, prematurity, and hyperosmolarity, may also affect the entry of bilirubin into the brain (Dennery et al., 2001).

**FIGURE 1 | Heme degradation pathway.** Heme from hemoglobin and cellular hemo proteins is metabolized in a rate-limiting step by HO. The reaction utilizes molecular oxygen and NADPH as a reducing equivalent. This leads to the release of iron, water, and CO as well as biliverdin. The latter is converted in a non-rate-limiting fashion to bilirubin by BVRI. Since bilirubin is not water soluble, it is conjugated to monoglucuronide and diglucuronide forms, which can then be excreted. These glucuronide residues can be removed by intestinal bacteria and allow unconjugated bilirubin to re-enter the circulation.
Unconjugated bilirubin induced protein oxidation and lipid peroxidation and reduced antioxidant defenses in neuronal cells in culture (Brito et al., 2008). Although toxic to cultured neuroblastoma cells, exposure to unconjugated bilirubin induced genes involved in the endoplasmic reticulum stress response in surviving cells thereby enhancing cellular homeostasis (Calligaris et al., 2009). In astrocytes, unconjugated bilirubin up-regulated the multidrug resistance-associated protein-1 and increased its trafficking to the plasma membrane, thus reducing its cytotoxicity by preventing its intracellular accumulation (Gennuso et al., 2004).

The toxic effects of bilirubin are not limited to the brain. Unconjugated bilirubin can mediate apoptosis in cultured hepatocytes by increasing oxidative stress and enhancing caspase-9 activity (Oakes and Bend, 2005). In erythrocytes, high bilirubin concentrations can induce hemolysis and lead to membrane disruption, which could theoretically worsen hemolytic anemia (Brites et al., 1997). We observed that in erythrocytes derived from cord blood, concentrations of bilirubin equal to or exceeding 30 mg/dL were associated with increased protein oxidation, decreased erythrocyte glucose-6 phosphate dehydrogenase and adenosine triphosphatase activity as well as altered cell membrane integrity (Mireles et al., 1999). There was a correlation between the release of unconjugated bilirubin and hepatotoxicity after TNF-α administration, in mice and this was resolved with HO inhibitors (Van Molle and Libert, 2003). Another important cytotoxicity of bilirubin involves its effects on complement activation, a key element of immune defense. Unconjugated bilirubin interferes with the interaction between C1q and immunoglobulins, which results in decreased complement activation via the classical pathway (Basiglio et al., 2010).

**ANTIOXIDANT BENEFITS OF BILIRUBIN**

Despite the fact that bilirubin may be toxic at higher concentrations, there is still significant evidence that it is a potent antioxidant at micromolar concentrations *in vitro* and *in vivo*. In fact, bilirubin is the most abundant cellular antioxidant. *In vitro*, bilirubin is a chain-breaking molecule that can scavenge the hydroxyl radical better than α-tocopherol, a well-known antioxidant against lipid peroxidation (Stocker et al., 1987; Mireles et al., 1999). Although incubation with bilirubin and albumin at concentrations greater than 30 mg/dL was associated with dose-dependent injury in erythrocytes derived from cord blood, protection against lipid peroxidation was seen at lower concentrations (Mireles et al., 1999), indicating the duality of bilirubin as a cytotoxic and cytoprotective molecule. Similarly, hemeoxygenase is both detrimental and beneficial based on levels of activity. Nevertheless, in that study, no change in bilirubin levels could be detected to explain the toxicity of hemeoxygenase (Suttner and Denney, 1999).

Similarly, *in vivo*, there are several examples of the beneficial effects of hyperbilirubinemia. In 50 patients older than 40 years with Gilbert syndrome, a relatively benign condition leading to mild to moderate unconjugated hyperbilirubinemia because of impaired glucuronidation, occurrence of ischemic heart disease was compared to that of a large cohort of patients without the disease. Ischemic heart disease occurred in only 2% of the Gilbert patients compared to 12.1% of the controls and, interestingly, hyperbilirubinemia rather than elevation of HDL cholesterol levels seemed to be more important in protection from ischemic heart disease (Vitek et al., 2002). In a case report, resolution of corticosteroid- and cyclophosphamide-resistant pulmonary fibrosis occurred with onset of hyperbilirubinemia due to biliary obstruction in a patient who developed elevated conjugated bilirubin levels (Ohrui et al., 2001), suggesting that higher serum bilirubin levels could reverse pulmonary fibrosis. The mechanisms by which this could occur are not yet explored. In organ transplantation, bilirubin can be protective against graft rejection (Ollinger et al., 2007). Injection of bilirubin in mouse organ recipients prolonged islet allograft survival and induced tolerance induction and graft acceptance via a regulatory T cell-dependent mechanism involving CD4(+) and CD25(+) cells. In fact, bilirubin enhanced *de novo* generation of regulatory T cells in the recipients thereby preventing rejection (Rocuts et al., 2010). Another novel mechanisms by which bilirubin may be protective is by the regulation of rapid eye movement sleep and by mediating some of the antidepressant effects of ambient light (Oren, 1997). Whether the antioxidant effects mediate these benefits is not yet clear.

Overall, the beneficial effects of bilirubin have been demonstrated in various models but beyond a certain threshold, bilirubin is clearly toxic.

**EPIDEMIOLOGIC EVIDENCE OF BILIRUBIN AS A CYTOPROTECTIVE MOLECULE**

To further understand whether bilirubin is cytoprotective in humans, epidemiologic studies can provide a clue. Regulation of bilirubin conjugation is key in the accumulation of bilirubin and its potential benefits or toxicity, therefore, studies comparing patients with differences in the ability to conjugate bilirubin may provide clues. The promoter of the *UGT1A1* gene has regions of TA repeats, which regulate its transcriptional efficiency. In Caucasian populations, an additional TA repeat (*TA_2* vs. *TA_0*) is necessary but not sufficient to cause Gilbert syndrome (Bartlett and Gourley, 2011). Strong associations between polymorphisms in the *UGT1A1* gene and human disease have been shown. In particular, there have been associations with altered bilirubin conjugation and the occurrence of various cancers. The common *UGT1A1*∗28 allele results in elevated plasma bilirubin levels and is strongly associated with Gilbert syndrome in Caucasians. Low serum bilirubin levels observed in a Caucasian cohort with predicted high activity of *UGT1A1* were associated with an increased risk of esophageal cancer. Interestingly, the *UGT1A8* and *UGT2B4* genotypes, associated with decreased UGT enzyme activity and increased unconjugated bilirubin levels, were also significantly associated with increased risk of esophageal cancer (Dura et al., 2012). In another study, the *UGT1A1* gene cluster on chromosome 2q37.1 was identified in a cohort of patients with bladder cancer suggesting that enhanced UGT1A may protect from bladder cancer by increasing the removal of carcinogens from bladder epithelium (Tang et al., 2012). In a meta-analysis of 21 case-control studies cancer risk was associated with intermediate, and low activity of *UGT1A7* genotypes, found predominantly in Asians (Lu et al., 2011). In contrast to the other studies, the TA repeat polymorphism of *UGT1A1* gene did not alter prostate cancer risk susceptibility in Caucasian men (Karatzas et al., 2010).
Overall, these studies suggest that the concentration of bilirubin in the serum determines whether it is beneficial or detrimental. Not only does the UGT1 gene play a role in cancer, it appears to have important effects in other diseases. For example, the homozygous state associated with higher serum bilirubin levels appeared to be protective against Crohn’s disease (de Vries et al., 2012).

Serum bilirubin, independent of variation in UGT promoter activity, is also associated with diseases in large populations, in particular in cardiovascular disease. In a Swedish cohort, plasma bilirubin was lower in 231 cases of ischemic stroke than in 462 matched controls but the difference reached significance only in women (Ekbloom et al., 2010). In males with coronary artery disease, there was inverse association between serum total bilirubin and coronary artery calcification score. Additionally, bilirubin was associated with reduced c-reactive protein levels, which could explain the lower calcification scores (Zhang et al., 2012). In another study, bilirubin levels were also inversely associated with the presence of coronary heart disease. Interestingly, bilirubin levels were significantly raised after treatment with 80 mg simvastatin independent of changes in liver enzymes (Nolting et al., 2011).

Despite the beneficial effects of bilirubin, the biggest challenge remains determining a specific threshold at which bilirubin is toxic vs. beneficial. It seems paradoxical that early events in bilirubin toxicity may involve increased oxidative stress and changes in redox status (Tell and Gustincich, 2009) yet conversely, bilirubin alleviates oxidative stress.

**ANTIOXIDANT PROPERTIES OF BILIVERDIN**

Although biliverdin does not accumulate in mammals, since it is rapidly converted to bilirubin through the action of biliverdin reductase (BVR), it may have important signaling effects. In macrophages, bilirubin activates endothelial nitric oxide, resulting in NO-dependent S-nitrosylation of BVR. The mechanisms by which biliverdin mediates this effect was via the repression of Toll-like receptor-4 (Wegiel et al., 2011). In another study, rats injected intraperitoneally with biliverdin before undergoing lung transplantation had less evidence of inflammation, oxidative injury, and apoptosis suggesting that biliverdin has anti-inflammatory and anti-apoptotic effects (Wang et al., 2010). Despite these data, the most plausible effect of biliverdin is to serve as a signaling molecule that may regulate BVR (Lerner-Marmarosh et al., 2008). The properties of this enzyme have been reviewed at length in a previous issue of this journal (Gibbs et al., 2012).

**OTHER POSSIBLE MEDIATORS OF HO-RELATED CYTOPROTECTIVE EFFECTS**

Although the byproducts of the HO-1 reaction are important cytoprotective molecules, a likely factor that influences the beneficial effects of the HO reaction is the degradation of a potent oxidant, heme. In one example, induction of HO-1 prevented photodynamic therapy-induced tumor necrosis, but neither bilirubin, biliverdin nor CO was responsible for this cytoprotection. In fact, the iron chelator desferrioxamine enhanced the cytotoxic effects of photodynamic therapy suggesting that heme was key important to enhancing the tumor killing effects of this therapy (Nowis et al., 2006).

**SUMMARY**

Overall, the byproducts of the HO reaction are important cytoprotective molecules that have clinically significant effects in various diseases. Nevertheless, in most cases, these molecules can also be cytotoxic under specific circumstances and/or at high concentrations. Potential therapeutic interventions will need to balance the potential benefits with the risk of toxicity to be most effective.

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