Modulation of Mrp1 (ABCC1) and Pgp (ABCB1) by Bilirubin at the Blood-CSF and Blood-Brain Barriers in the Gunn Rat

Silvia Gazzin1*, Andrea Lorena Berengeno1, Nathalie Strazielle2, Francesco Fazzari1, Alan Raseni3, J. Donald Ostrow4, Richard Wennberg5, Jean-François Ghersi-Egea6, Claudio Tiribelli1

1 Centro Studi Fegato, Basovizza Trieste, Italy, 2Brain-i at INSERM U842, Université de Lyon, Lyon1, Faculté de Médecine Laennec, Lyon, France, 3 S.C. Laboratorio Analisi Cliniche IRCCS Burlo Garofolo, Trieste, Italy, 4 Gastroenterology/Hepatology Division, University of Washington School of Medicine and VA Puget Sound Health Care System/Seattle Division, Seattle, Washington, United States of America, 5 Division of Neonatology, University of Washington School of Medicine, Seattle, Washington, United States of America, 6 INSERM U842, Université de Lyon, Lyon1, Faculté de Médecine Laennec, Lyon, France

Abstract

Accumulation of unconjugated bilirubin (UCB) in the brain causes bilirubin encephalopathy. Pgp (ABCB1) and Mrp1 (ABCC1), highly expressed in the blood-brain barrier (BBB) and blood-cerebrospinal fluid barrier (BCSFB) respectively, may modulate the accumulation of UCB in brain. We examined the effect of prolonged exposure to elevated concentrations of UCB on expression of the two transporters in homozygous, jaundiced (jj) Gunn rats compared to heterozygous (Jj) littermates at different developmental stages (2, 9, 17 and 60 days after birth). BBB Pgp protein expression was low in both jj and Jj pups at 9 days (about 16–27% of adult values), despite the up-regulation in Jj animals (2 and 1.3 fold higher than age matched Jj animals at P9 and P17–P60, respectively); Mrp1 protein expression was barely detectable. Conversely, at the BCSFB Mrp1 protein expression was rather high (60–70% of the adult values) in both jj and Jj at P2, but was markedly (50%) down-regulated in jj pups starting at P9, particularly in the 4th ventricle choroid plexuses: Pgp was almost undetectable. The Mrp1 protein down regulation was accompanied by a modest up-regulation of mRNA, suggesting a translational rather than a transcriptional inhibition. In vitro exposure of choroid plexus epithelial cells obtained from normal rats to UCB, also resulted in a down-regulation of Mrp1 protein. These data suggest that down-regulation of Mrp1 protein at the BCSFB, resulting from a direct effect of UCB on epithelial cells, may impact the Mrp1-mediated neuroprotective functions of the blood-cerebrospinal fluid barrier and actually potentiate UCB neurotoxicity.

Introduction

Severe jaundice from unconjugated bilirubin (UCB) can occur transiently in newborn infants with immature hepatic conjugating capacity [1,2,3,4,5] and lifelong in patients with Crigler-Najjar type I disease [6,7]. Over 99.9% of UCB in blood is bound to plasma proteins (primarily albumin) that do not enter the brain [8,9,10]. Only the small fraction of unbound bilirubin may diffuse into the brain and cerebrospinal fluid (CSF). With severe jaundice, the serum binding sites approach saturation and unbound UCB (free bilirubin, Bf) will rise dramatically even at lower bilirubin/albumin molar ratios [2,11]. Under these conditions, the accumulation of UCB in brain (kernicterus) can produce toxicity and result in permanent brain injury.

The exchange of unbound UCB between the blood and the brain may be modulated by two blood-brain interfaces that control cerebral homeostasis [12]. Directly contacting the neuroglia, the blood-brain barrier consists of tightly bound, specialized endothelial cells lining the brain microvessels [13]. A second interface, the blood-cerebrospinal fluid barrier is provided by the epithelial cells of the choroid plexuses, and controls exchanges between plasma and CSF. Both barriers display a large surface area for exchanges [14]. Tight junctions [15,16] preclude the paracellular passage of hydrophilic compounds [17], while transporters [18] and metabolizing enzymes [19,20,21,22] control the neural access of lipid-soluble substrates.

Two trans-membrane proteins, belonging to the ATP binding cassette (ABC) family, have been identified as potential UCB transporters, which export the pigment from the cells. MDR1 (ABCB1: Pgp, Mdr1a/1b in rodents) displays a low affinity for UCB [23] and MRPI (ABCC1: Mrp1 in rodents) that possess a very high affinity for UCB [Kn for Bf = 10 nM] [24]. In rat as well as in human brain, Pgp is specifically expressed in microvessels, localized at the luminal (blood) side of the endothelium, while Mrp1 is mostly localized at the basolateral membrane of the choroidal epithelium, facing the stromal/blood space. Pgp protein increases during post-natal development, whereas Mrp1 is highly expressed in choroid plexuses, even at birth [14,18,25,26]. Both ABC transporters may participate in limiting the entry of UCB by increasing its export from the central nervous system [27,28].
In this study we investigated the effect of sustained unconjugated hyperbilirubinemia on the developmental protein expression of Mrp1 in the lateral and 4th ventricle choroid plexuses, and of Pgp in brain microvessels. We used immature and adult Gunn rats [29], a well established animal model for chronic unconjugated hyperbilirubinemia and kernicterus [30]. Homozygous, recessive (jj) Gunn rats, like patients with Grigler-Najjar syndrome type I (CNS I), develop severe, lifelong, non-hemolytic, unconjugated hyperbilirubinemia, due to the congenital absence of UDP-glucuronosyl transferase (UDPGT: EC 2.4.1.17) 1A1, the enzyme that synthesizes excretable conjugates of UCB. The same enzyme has transient low activity in jaundiced human newborns [31,32]. Heterozygous Jj animals, used as controls, have only a minimally reduced activity of UGT1A1 compared with Sprague-Dawley animals [33].

Materials and Methods

Animals

Gunn rats (Hds Blue-Gunn-UDPGTj) [29], originally purchased from Harlan (Harlan, IL, USA) in 2006, were maintained in the animal facility of the University of Trieste. Animal care and procedures were conducted according to the guidelines approved by Italian Law (decrees 116-92) and by European Community directive 86-609-EEC. Sacrifice was performed after anesthesia (isofluorane) in order to avoid animal pain or stress, within the law 116-92. Because animals spontaneously develop the pathology, and no treatments have been applied, additional ethical approval was not required. All efforts were made to avoid suffering of animals.

Littermates were obtained by breeding Jj females with jj males. Parturitions were synchronized to obtain a sufficient number of littersmate pups of each genotype and post-natal age (P±24 hrs). Jj pups from the same litter were used as controls for jj animals. Post-natal ages to study were selected based on developmental vulnerability to bilirubin neurotoxicity established by others [34,35,36,37,38]. The number of rats pooled for each batch and the number of batches analyzed for each transporter are listed in Table 1.

Plasma bilirubin determination

A heparinized sample of blood was obtained by jugular puncture from each anesthetized animal. After centrifugation (1500 g for 20 min at room temperature), plasma was collected under dim light to minimize bilirubin photo-oxidation and immediately frozen at -20°C until assayed. The diazo-reaction [39] (Boehringer-Mannheim Kit 1552414, Monza, Italy) was used immediately frozen at 20°C until assayed. The diazo-reaction was performed after anesthesia (isofluorane) in order to avoid animal pain or stress, within the law 116-92. Because animals spontaneously develop the pathology, and no treatments have been applied, additional ethical approval was not required. All efforts were made to avoid suffering of animals.

Sample preparation and Western blot quantification

Freshly isolated microvessels, and lateral and 4th ventricle choroid plexuses were homogenized in buffer (0.25 M Sucrose; 50 mM K Phosphate; 1 mM EDTA; 0.1 mM DTT; pH 7.4), using a Dounce-type glass-glass homogenizer. Total protein content was determined using bicinchoninic acid (BCA), following the manufacturer’s protocol (Procedure # TPRO 562, Sigma, St Louis, MO, USA).

Proteins were separated on 10% SDS-polyacrylamide gels. Before transfer, the gel was cut longitudinally; the lower part (containing the actins) was blotted on PVDF membrane (0.2 μm; BioRad Laboratories, Hercules, CA, USA) at 100 V in transfer buffer (25 mM Tris-base; 192 M glycine, 0.1% SDS; 20% methanol) for 1h, while the upper part of the gel (Mrp1 and Pgp) was blotted for 2 h, to achieve maximal transfer. Completeness of transfer was assessed by lack of Coomassie blue coloration of the gels after blotting.

Table 1. Animals and batches of samples used for the ABC transporter analysis.

| Post-natal age | P2 | P9 | P17 | P60 | P120 |
|---------------|----|----|-----|-----|------|
| MVs Animals* | N.P. | 8 | 6 | 4 | N.P. |
| Batches | N.P. | 3 | 4 | 5 | N.P. |
| CPs Animals* | 4 | 4 | 3 | 4 | 3 |
| Batches | 4 | 4 | 4 | 4 | 3 |

MV= microvessels; CP= choroid plexuses; * Animals pooled together to generate one batch. N.P.: not performed.

doi:10.1371/journal.pone.0016165.t001

Tissue dissection and preparation

Animals were euthanized under isofluorane anesthesia by decapitation (within the law 116-92). Two lateral ventricle choroid plexuses, one from each hemisphere, 4th ventricle choroid plexuses, cerebral cortices and cerebella were dissected individually from four P2, P9, P17 and P60 animals of each phenotype (see Table 1) in Krebs-Ringer buffer (in mM: 135 NaCl, 4 KCl, 2.2 CaCl2, 1.2 MgCl2, 6 NaHCO3, 10 HEPES, 5 glucose, pH 7.4), at 4°C under stereomicroscope vision. Spleens from Jj and jj animals were also dissected for MRP1 analysis. Brain microvessels were obtained as described previously [18]. Briefly, pooled cerebral cortices were cleaned from all apparent meninges under a stereomicroscope. For mature animals (P60), tissues were homogenized in a Dounce-type glass-glass homogenizer after the addition of 5 vol/g tissue of 1% bovine serum albumin-supplemented Krebs-Ringer. The microvessels were separated from larger vessels and brain parenchyma material by filtering through decreasing pore diameter mesh sieves and the myelin was removed by centrifugation on a 17.5% 70 KDa-Dextran gradient in Krebs-Ringer buffer at 3000 g for 30 min. The microvessel fraction retained on the 40 μm sieve was recovered in 0.1% albumin in Krebs-Ringer buffer, centrifuged, and suspended in a small volume of the same buffer. Because of the fragility of brain tissue, cerebral cortices of P17 animals were homogenized by eight loose-type pestle strokes, while five loose-type pestle strokes were used for P9 pups. Preparations contaminated with larger vessels or tissue remnants were discarded. All steps were performed at 4°C within 4 hours of decapitation. Samples were stored at -80°C until use for Western blot.

UCB Induced Mrp1/Pgp Modulation at the BBI

Table of contents

1. Materials and Methods
   a. Animals
   b. Plasma bilirubin determination
   c. Sample preparation and Western blot quantification
   d. Tissue dissection and preparation
2. Results
   a. Organ distribution of Mrp1
   b. Mrp1/Pgp expression in the choroid plexus
   c. Mrp1/Pgp expression in the brain cortex
3. Discussion
4. Conclusion

References

[1] DOI 10.1371/journal.pone.0016165

ImageQuant software (GE Healthcare Europe GmbH, France).
Reference controls to quantify the relative contents of Pgp and Mrp1 were obtained by serial diluted of pooled microvessels and 4th ventricle choroid plexuses obtained from P60 Jj Gunn rats and run for each Western blot procedure (Fig. 1). A standard curve was generated by non-linear regression (MMF model, each r>0.99) of plots of optical density values versus the amount of reference sample protein. Relative expression of the ABC transporters, normalized for the actin signal, was calculated by CurveExpert 1.38 software (Hixon, TN, USA). The results are expressed as mean ± S.D., n = 3 to 5 sample batches for each post-natal age (see Table 1).

Immunohistochemical detection of Mrp1 in rat brain sections

Several rat brains of each genotype and developmental stage were immediately snap-frozen after decapitation, and 10 µm-thick slices were cryo-sectioned, using a Micron HM550 cryostat (Bio-Optica, Milan, Italy). Sections were fixed at room temperature in 4% paraformaldehyde in phosphate buffer (PBS; in mM: NaCl 150; Na2HPO4 12; KH2PO4 2; pH 7.4) for 10 min, then blocked 4% paraformaldehyde in phosphate buffer (PBS). After overnight incubation at 4°C with the anti-Mrp1 antibody (final concentration 1 µg/mL) in blocking solution, secondary anti-rabbit Alexa-conjugated antibodies (MolecularProbes, Carlsbad, CA, USA, 2 µg/mL, in 0.3% Triton-PBS) were added and incubated for 2 hr at room temperature.

Nuclei were stained with Hoechst 33342 (0.1 µg/mL) in PBS for 10 min at room temperature and sections were mounted with cytometry fluorescence mounting medium (S3023, Dako, Glostrup, Denmark). Negative controls were performed by omitting the first antibody. Fluorescent immunoreactions were observed using a Leica DM2000 fluorescence microscope, equipped with the Digital Camera Leica DFC490 and the software Leica LAS Image Overlay (Leica, Ernst-Leitz-Strasse 17–37, Wetzlar, D-35578, Germany).

Real Time-PCR analysis of Mrp1 mRNA levels in choroid plexuses of P9 animals

Lateral and 4th ventricle choroid plexuses of Jj and jj Gunn rats were collected in TRIReagent (Sigma-Aldrich, St Louis, MO, USA). Total RNA was isolated according to the manufacturer’s instructions. Briefly, tissues were lysed with the reagent, chloroform was added and cellular RNA was precipitated by isopropyl alcohol. After washing with 75% ethanol, the RNA pellet was dissolved in nuclease-free water and stored at −80°C until further analysis. RNA was quantified spectrophotometrically at 260 nm, and the RNA purity was evaluated by measuring the ratio A260/A280, considering RNAs with appropriate purity those showing values between 1.8 and 2.0. RNA integrity was evaluated by gel electrophoresis.

Total RNA (1 µg) in 15 µl of nuclease-free water was added to 4 µL reaction mix, and 1 µL reverse transcriptase using the iScript™cDNA Synthesis kit (Bio-Rad Laboratories, Hercules, CA, USA) according to the manufacturer’s instructions. The reaction was performed for 5 min at 25°C (annealing), 30 min at 42°C (cDNA synthesis), and 5 min at 85°C (enzyme denaturation).

Three house-keeping genes (β-actin, 18S and GAPDH), each corresponding to a different function in the cell, were employed to normalize the results. The PCR reaction was performed on 25 ng cDNA with gene-specific sense and anti-sense primers for Mrp1, β-actin, GAPDH (all Mrp1; β-actin and GAPDH 250 nM) and 18S (100 nM) (Table 2) with iQ SYBR Green Supermix (Bio-Rad Laboratories, Hercules, CA, USA) in an i-Cycler IQ thermocycler (Bio-Rad Laboratories, Hercules, CA, USA). The thermal cycler conditions consisted of 3 min at 95°C (cDNA synthesis) and 80 cycles of 0.5°C increments (10 s each) beginning at 55°C for 20 s, 60°C for 20 s and 72°C for 30 s. Specificity of the amplification was verified by a melting-curve analysis, performed immediately after the amplification protocol, under the following conditions: 1 min denaturation at 95°C, 1 min annealing at 55°C, and 80 cycles of 0.5°C increments (10 s each) beginning at 55°C. Non-specific products of PCR were not found in any case. A standard curve was generated using a “calibrator” cDNA (chosen among the cDNA samples), which was serially diluted and analyzed for all genes to determine the amplification efficiencies of individual genes. The relative quantification was made using Genex software (Bio-Rad Laboratories, Hercules, CA, USA) based on ΔΔCt method, taking into account the efficiencies of individual genes and normalizing the results to the three house-keeping genes. The levels of Mrp1 mRNA were expressed relative to a selected 4th ventricle choroid plexus Jj reference sample. The results are expressed as mean ± S.D., n = 3 pools of choroid plexuses dissected from 4 animals.

Effect of bilirubin on Mrp1 protein in an in vitro model of the blood-CSF barrier

Choroid plexus epithelial cells were isolated and cultured on Costar Transwell Clear inserts (0.4 µm pore size, 6.5 mm diameter) as
previously described [41,42]. Cells were treated in the basolateral compartment of the bicameral system and exposed for 6 consecutive days to daily renewed 7.5 mM UCB, starting on day 4 after plating. UCB was dissolved in DMSO (0.16 and 0.4% final volume, for low and high dose, respectively) and diluted in culture medium supplemented with 11.5% fetal bovine serum containing 30 mM albumin. UCB concentration was adjusted to yield bilirubin/albumin ratios calculated to have free bilirubin concentrations of approximately 40 and 140 nM [43]. Control monolayers were treated in parallel with the same concentration of DMSO. On day 10, filters were cut and transferred into 1/6 Lysis Buffer (Cell Signalling Technology, Saint-Quentin Yvelines, France) and kept at 280°C, until processed for Western blot analysis of both Mrp1 and actin as previously described in detail [18]. Duplicate monolayers were included for each condition. Actin was used to assess protein load of each sample.

Statistical analysis

Data are reported as mean ± SD. Statistical differences between hyperbilirubinemic animals and controls (jj) were analyzed using paired Student’s t-test for Western blot ABC transporter quantification. Anova following by a Tukey-Kramer test was used to assess variation in total plasma bilirubin concentration over time. One tailed Student’s test for unequal variances was applied to assess differences in cerebellar growth and plasma bilirubin concentration and in mRNA levels between Jj and jj animals. Differences were considered statistically significant at a p value lower than 0.05.

Results

Plasma bilirubin and cerebellar growth

A transient postnatal increase in total plasma bilirubin concentration was observed in Jj animals, decreasing to the low adult values by P17 (Fig. 2A). The total bilirubin in plasma of jj rats was significantly higher (p<0.005) than in Jj rats at all post-natal ages and reached the peak level (250 μM) at P9. A hallmark of bilirubin toxicity in jj rats is cerebellar hypoplasia [37,44,45]. Cerebellar weights in jj and Jj animals were identical at P2, but cerebellar growth was impaired thereafter in jj animals and almost completely arrested by P17 (Fig. 2B). Thus, the jj cerebellar weight was 25% lower at P9 (p<0.01), 40% lower at P17 (p<0.005) and 50% lower at P60 (p<0.005) than in Jj animals.

Characterization of isolated microvessels and choroid plexus tissues

It was technically impossible to obtain microvessel preparations of adequate purity from rat brain tissue at ages earlier than 9 days [18]. On phase contrast microscopic examination, microvessel preparations consisted mostly of long, branched capillaries, with the thin structure of the microvessels better appreciated at higher magnifications (Fig. 3, B-
No contaminating larger vessels, cellular debris, or meninges were observed (Fig. 3, B), and red blood cells were rarely found. Despite a softer consistency of the cortices dissected from jaundiced jj rats, vascular morphology was similar to Jj rats and no cellular alteration (granulations, bubbles, etc.) was detected (Fig. 3, panel D, F for Jj and panel C, E, G for jj). Similarly, no difference was noted in the dimensions or morphology of either lateral or 4th ventricle choroid plexuses dissected from jaundiced or control Gunn rats.

**Pgp protein expression in microvessels**

In agreement with previous data on rats and mice, Pgp expression in microvessels strongly increased from the early post-natal age in days. jj: hyperbilirubinemic rats; Jj: controls.

doi:10.1371/journal.pone.0016165.g003
Mrp1 expression in the spleen did not differ between jji and JJJ rats at any age (P9: 100.8±2.6 vs. 101.8±2.5; P17: 101.2±9.0 vs. 90.4±8.1, jji vs. JJJ animals). When Mrp1 mRNA was analyzed by real time PCR in P9 choroid plexus preparations, the expression was comparable between JJJ and jji rats, despite a 50% drop in Mrp1 protein (Fig. 6).

The absence of either molecular weight shift or lower molecular weight bands for Pgp in micro vessels and Mrp1 in choroid plexuses in jji preparations suggests that hyperbilirubinemia does not alter the post-translational maturation nor produce degradation of these transporters. In both JJJ and JJJ Gunn rats, Pgp expression in choroid plexuses and Mrp1 expression in microvessels were barely detectable (data not shown) as previously shown in Sprague-Dawley rats [18].

Immunofluorescence analysis of Mrp1 cellular localization in rat brain

Mrp1 was localized at the basolateral membrane of choroid plexus epithelial cells, in both JJJ and JJJ Gunn rat brain slices, even at P2 (Fig. 7, A, D and E), in agreement with previous studies in Sprague-Dawley rats [18]. No cytoplasmic staining was detected in jji rats (compare D with E, and F with G), suggesting that UCB is unable to reallocate the transporter in vivo. When the nuclei were counterstained by Hoechst 33422, no apoptotic bodies were detected, in keeping with the absence of macroscopic cellular damage due to UCB.

Effect of UCB on Mrp1 protein expression in cultured choroidal epithelium

Monolayers of choroidal epithelial cells were cultured in bicameral devices, reconstituting in vitro the barrier properties of the blood-CSF interface. Cells were exposed to 2 different concentrations of unbound bilirubin (40 and 140 nM, which mimic levels measured in physiological and pathological jaundice, respectively) at the basolateral membrane site for 6 consecutive days to mimic the in vivo prolonged postnatal exposure. This treatment resulted in a decrease in Mrp1 protein at the higher bilirubin concentration (Fig. 8). This effect was not accompanied by any alteration in the paracellular permeability of the monolayers to sucrose (data not shown), and therefore does not reflect a pleiotropic toxicity. These data are in agreement with our in vivo results and suggest that the reduction of Mrp1 protein observed in jji animals in the blood-CSF barrier results from a direct action of UCB on barrier cells.

Discussion

UCB can enter brain parenchyma either directly via the blood-brain barrier, or by transfer across the choroid plexuses. The ABC transporters, Pgp and Mrp1, have been proposed to modulate brain entry of UCB [14]. In vitro, Pgp, mainly expressed in microvessels, is a UCB transporter weaker than Mrp1 [23,24], the latter being dominant in the choroid plexuses [18]. The effects of prolonged exposure to high UCB levels in vivo as seen in severely jaundiced newborns or Crigler-Najjar type I patients, on the post-natal expression of Pgp and Mrp1 at the blood-brain interfaces have not been reported.

We observed an up-regulation of Pgp protein in microvessels, and a down-regulation of Mrp1 in choroid plexuses isolated from hyperbilirubinemic jji Gunn rat pups as compared with their non-jaundiced JJJ littermate controls. Mechanisms mediating these opposite responses are still uncertain. UCB has been show to affect the cellular redox [48,49,50,51] state and Pgp expression is increased by oxidative stress [52,53], suggesting that oxidative...
stress may be a mediator of UCB-induced Pgp protein level at the blood-brain barrier.

UDPGT activity is present in rat choroid plexuses [19,54] and glucuronosyl conjugates are one of the substrates for Mrp1. In jj Gunn rats, UGT1A1 activity is completely abolished by a genetic mutation, suggesting that the resulting limited availability of Mrp1 substrates might lead to an Mrp1 down regulation in jj animals. However, the normal protein expression of Mrp1 in choroid plexuses of P2 pups (Fig 5) and in spleen and in liver (data not shown) indicates that the Mrp1 protein reduction is not due to the decrease in UGT activity but rather to a direct effect of the prolonged exposure to high levels of UCB. This is further supported by the in vitro data showing that UCB is per se able to reduce the protein content of Mrp1 in choroid plexuses.

Lipophilic compounds such as UCB cross cellular membranes mainly by passive diffusion [17,57]. The protein up-regulation of Pgp we observed is consistent with a defensive role proposed for this transporter against bilirubin entry into brain. However, despite this up-regulation, the expression of Pgp in response to the elevated levels of UCB in jj Gunn rats in the first 2.5 weeks after birth (P2–P17) remained very low compared to adult values (16 and 27% of adult Jj level, respectively). This suggests that Pgp offers, at best, a marginal defense against bilirubin entry into brain during the critical period (P2–P17) when the UCB-related neurological damage occurs, as the diffuse yellow coloration of the brains demonstrates (not shown). At the same time, the Mrp1 protein level in the choroid plexuses of jj animals is significantly reduced, which may further impair the ability of the brain to protect itself from UCB accumulation and toxicity. This has been shown in vitro where silencing of Mrp1, but not Pgp, is associated with a greater cellular toxicity induced by UCB exposure [58].

Figure 5. Mrp1 relative expression in the lateral and 4th ventricle choroid plexuses dissected from Jj and jj Gunn rats. The Mrp1 protein amount present in each sample is expressed as % of the Mrp1 amount in the Reference Sample (pooled P60 Jj 4th V CP preparations). White bars: heterozygous (Jj) rats, black bars: hyperbilirubinemic homozygous (jj) animals. 4th V CP: Forth Ventricle Choroid Plexuses; LV CP: Lateral Ventrices Choroid Plexuses. Statistical significance is indicated.
doi:10.1371/journal.pone.0016165.g005

Figure 6. Comparison of Mrp1 mRNA and protein relative expression in choroid plexuses. White bars: heterozygous (Jj) P9 rats, black bars: hyperbilirubinemic homozygous (jj) P9 animals. The values are expressed as mean ± SD. Statistical significance is reported. NS: not significant difference.
doi:10.1371/journal.pone.0016165.g006
Mrp1 expression in choroid plexuses matures early during development and is involved in regulating cerebral availability of endogenous biologically active compounds (leukotriene [59], steroid hormones [60], glutathione, glucuronide and sulfate conjugates [61], reduced glutathione [62]) and therapeutic drugs such as etoposide [63,64]. In line with the functional relevance of these transport activities, the involvement of the transporter in inflammation [65] and oxidative stress [66] has been demonstrated in vivo in Mrp1 KO mice.

We speculate that Mrp1 protein down-regulation observed in jaundiced jj Gunn rat pups and maintained into adulthood, may increase the central nervous system accumulation of toxins or drugs, impair immune and endocrine response, and lead to increased oxidative stress impairing neuronal development independent of bilirubin itself. In summary, we provide evidence that major alterations in Pgp and Mrp1 expression are induced in the BBB and BCSFB, respectively, by prolonged exposure to elevated levels of UCB, as seen in jj Gunn rats.

Although the mechanisms of these alterations in expression are currently undetermined, we propose that the modulation of UCB...
transporters in vivo results from the direct exposure of barrier cells to bilirubin, possibly mediated in part by an alteration in the cellular redox state.

Acknowledgments

We thank Sylvie Cavagna for her skillful technical help.

References

1. Gouley GR (1997) Bilirubin metabolism and kernicterus. Adv Pediatr 44: 173–229.
2. Ostrow JD, Pascoli I, Shapiro SM, Tiribelli C (2003) New concepts in bilirubin encephalopathy. Eur J Clin Invest 33: 988–997.
3. Ostrow JD (1987) Bile pigments and jaundice Marcel Dekker, Inc.
4. Kaplan M, Muraca M, Hammerness C, Rahalbelti F, Yedzi MT, et al. (2002) Imbalance between production and conjugation of bilirubin: a fundamental concept in the mechanism of neonatal jaundice. Available: http://pediatrics.org/cgi/content/full/110/4/e47.
5. Shapiro SM, Bhutani VK, Johnson I, (2006) Hyperbilirubinemia and kernicterus. Clin Perinat 33: 387–410.
6. Bosma PJ, Goldhoorn B, Ouide Elfenink RP, Sinaasappel M, Oostra BA, et al. (1993) A mutation in bilirubin uridine 5’-diphosphate-glucuronosyltransferase isoform 1 causing Crigler-Najjar syndrome type II. Gastroenterology 105: 216–220.
7. Kadakol A, Ghosh SS, Sappal BS, Sharma G, Chowdhury JR, et al. (2000) Genetic lesions of bilirubin uridine-diphosphoglucuronate glucuronosyltransferase (UGT1A1) causing Crigler-Najjar and Gilbert syndromes: correlation of genotype to phenotype. Hum Mutat 16: 297–306.
8. Ostrow JD, Mekerjee P, Tiribelli C (1994) Structure and binding of unconjugated bilirubin: relevance for physiological and pathophysiological function. J Lipid Res 35: 1713–1737.
9. Diamond I, Schmid R (1966) Experimental bilirubin encephalopathy. The mode of entry of bilirubin-14C into the central nervous system. J Clin Invest 45: 678–689.
10. Takahashi M, Sugiyama K, Shumiya S, Nagase S (1984) Penetration of bilirubin into the brain in albumin-deficient and jaundiced rats (AJR) and Nagase albuminemic rats (NAR). J Biochem 96: 1705–1712.
11. Wennberg RP, Ahlfors CE, Bhutani VK, Johnson LH, Shapiro SM (2006) Toward understanding kernicterus: a challenge to improve the management of jaundiced newborns. Pediatrics 117: 474–485.
12. Wennberg RP (2009) The blood-brain barrier and bilirubin encephalopathy. Cell Mol Neurobiol 29: 97–109.
13. Rodriguez-Baeza A, Reina-de la TF, Poca A, Marti M, Garnacho A (2003) Morphological features in human cortical brain microvessels after head injury: a three-dimensional and immunocytochemical study. Anat Rec A Discov Mol Cell Morphol 275: 583–593.
14. Gheresi-Egea JF, Gazzin S, Strazielle N (2009) Blood-brain interfaces and bilirubin-induced neurological diseases. Curr Pharm Des 15: 2903–2907.
15. Liebner S, Kniesel U, Kalbacher H, Wolburg H (2000) Correlation of tight microvessels: differential expression of gamma-glutamyl transpeptidase and alkaline phosphatase. Anat Embryol (Berl) 199: 29–34.
16. Lippoldt A, Liebner S, Andbjer B, Kalbacher H, Wolburg H, et al. (2000) Localization of drug-metabolizing enzyme activities to blood-brain interfaces. J Comp Neurol 510: 497–507.
17. Cannon C, Daood MJ, O’Day TL, Watchko JF (2006) Sex-specific regional brain bilirubin content in hyperbilirubinemic Gunn rats pups. Biol Neonate 90: 40–46.
18. Johnson JA, Hayward JJ, Kornguth SE, Siegel FL (1993) Effects of hyperbilirubinaemia on glutathione S-transferase isoenzymes in cerebral cortex of the Gunn rat. Biochem J 291 (Pt 2): 433–461.
19. Conlee JW, Shapiro SM (1997) Development of cerebral hypoplasia in jaundiced Gunn rats: a quantitative light microscopic analysis. Acta Neuropathol 93: 450–460.
20. Ostrow SM (2002) Somatosensory and brainstem auditory evoked potentials in the Gunn rat model of acute bilirubin neurotoxicity. Pediatr Res 52: 844–849.
21. Ehrlich P (1988) Sulfadoxine, ein Reagens auf Bilirubin. Cenr Krank 4: 721–723.
22. Ferranti N, Pascoli L, Podica E, Brennaro R, Stelbel M, et al. (2001) Preparation of an antibody recognizing both human and rodent MRPs. Biochem Pharmacol 63: 1064–1069.
23. Strazielle N, Belin MF, Gheresi-Egea JF (2003) Choroidal plexus regulates brain availability of anti-HIV nucleoside analogs via pharmacologically inhibitable organic anion transporters. AIDS 17: 1473–1485.
24. Strazielle N, Gheresi-Egea JF (1999) Demonstration of a coupled metabolism-efflux process at the choroid plexus as a mechanism of brain protection toward xenobiotics. J Neurosci 19: 6273–6289.
25. Tsai CE, Daood MJ, Lane RH, Hansen TW, Gruetzacher EM, et al. (2002) P-glycoprotein expression in mouse brain increases with maturation. Biol Neonate 81: 58–64.
26. Matsuoka Y, Okazaki M, Kitamura Y, Taniguchi T (1999) Developmental expression of P-glycoprotein (multidrug resistance gene product) in the rat brain. J Neurobiol 39: 383–392.
27. Cesarrato L, Calligaris SD, Vasconcello G, Cesarrato L, Calligaris SD (2007) Bilirubin-induced cell toxicity involves P-gp inhibition via NR1H3 activation through an APE1/Ref-1-dependent pathway. J Mol Med 85: 1099–1112.
28. Tell G, Gustincich S (2009) Redox state, oxidative stress, and molecular mechanisms of protective and toxic effects of bilirubin on cells. Curr Pharm Des 15: 2904–2914.
29. Brito MA, Lima S, Fernandes A, Falcao AS, Silva RF, et al. (2008) Bilirubin injury to neurons: contribution of oxidative stress and rescue by glycodeoxycholic acid. Neurotoxicology 29: 259–269.
51. Brito MA, Rosa AI, Falcao AS, Fernandes A, Silva RF, et al. (2008) Unconjugated bilirubin differentially affects the redox status of neuronal and astroglial cells. Neurobiol Dis 29: 30–40.

52. Felix RA, Barraud MA (2002) P-glycoprotein expression in rat brain endothelial cells: evidence for regulation by transient oxidative stress. J Neurochem 80: 64–72.

53. Hong H, Lu Y, Ji ZN, Liu GQ (2006) Up-regulation of P-glycoprotein expression by glutathione depletion-induced oxidative stress in rat brain microvessel endothelial cells. J Neurochem 90: 1463–1473.

54. Strazielle N, Ghezzi-Egea JP (1999) Demonstration of a coupled metabolit- efflux process at the choroid plexus as a mechanism of brain protection toward xenobiotics. J Neurosci 19: 6275–6289.

55. Elferink MG, Olinga P, Draaisma AL, Merema MT, Faber KN, et al. (2004) LPS-induced downregulation of Mrp2 and Bsep in human liver is due to a posttranscriptional process. Am J Physiol Gastrointest Liver Physiol 287: G1008–G1016.

56. Jones BR, Li W, Cao J, Hoffman TA, Gerk PM, et al. (2005) The role of protein synthesis and degradation in the post-transcriptional regulation of rat multidrug resistance-associated protein 2 (Mrp2, Abcc2). Mol Pharmacol 68: 701–710.

57. Mediavilla MG, Pascolo L, Rodriguez JV, Guibert EE, Ostrow JD, et al. (1999) Uptake of [3H]bilirubin in freshly isolated rat hepatocytes: role of free bilirubin concentration. FEBS Lett 463: 143–145.

58. Corich L, Ariand A, Carrassa L, Bellarosa C, Ostrow JD, et al. (2009) The cytotoxic effect of unconjugated bilirubin in human neuroblastoma SH-SY5Y cells is modulated by the expression level of Mrp1 but not Mdr1. Biochem J 417: 305–312.

59. Leier I, Jellitschky G, Buchholz U, Cole SP, Dedley RG, et al. (1994) The MRP gene encodes an ATP-dependent export pump for leukotriene C4 and structurally related conjugates. J Biol Chem 269: 27807–27810.

60. Sugiyama D, Kusuhara H, Lee YJ, Sugiyama Y (2003) Involvement of multidrug resistance-associated protein 1 (Mrp1) in the efflux transport of 17beta estradiol-D-17beta-glucuronide (E217betaG) across the blood-brain barrier. Pharm Res 20: 1394–1400.

61. Jellitschky G, Leier I, Buchholz U, Barnouin K, Kurz G, et al. (1996) Transport of glutathione, glucuronate, and sulfate conjugates by the MRP gene-encoded conjugate export pump. Cancer Res 56: 988–994.

62. Hirrlinger J, Konig J, Keppler D, Lindenau J, Schulz JB, et al. (2001) The multidrug resistance protein MRP1 mediates the release of glutathione disulfide from rat astrocytes during oxidative stress. J Neurochem 76: 627–636.

63. Cole SP, Sparks KE, Fraser K, Loe DW, Grant CE, et al. (1994) Pharmacological characterization of multidrug resistant MRP-transfected human tumor cells. Cancer Res 54: 5902–5910.

64. Wijnholds J, deLange EC, Schiefere GL, van den Berg DJ, Mol CA, et al. (2000) Multidrug resistance protein 1 protects the choroid plexus epithelium and contributes to the blood-cerebrospinal fluid barrier. J Clin Invest 105: 279–285.

65. Wijnholds J, Evers R, van Leusden MR, Mol CA, Zaman GJ, et al. (1997) Increased sensitivity to anticancer drugs and decreased inflammatory response in mice lacking the multidrug resistance-associated protein. Nat Med 3: 1273–1279.

66. Lorico A, Rappa G, Finch RA, Yang D, Flavell RA, et al. (1997) Disruption of the murine Mrp (multidrug resistance protein) gene leads to increased sensitivity to etoposide (VP-16) and increased levels of glutathione. Cancer Res 57: 5238–5242.