Treatment of jaundice in low birthweight infants

M J Maisels, J F Watchko

Exchange transfusion and phototherapy remain the staples of intervention for the jaundiced newborn. Clinical management of the jaundiced low birthweight infant is discussed.

Although it is generally believed that pre-term and low birthweight (LBW) infants are at greater risk of developing bilirubin associated brain damage than term infants, quantification of the magnitude of this risk has proven elusive, as has a consensus among experts on the concentration of total serum bilirubin (TSB) at which treatment should be initiated.

Nevertheless, there is little doubt that kernicterus is currently a very rare event in premature infants in neonatal intensive care units (NICUs), although a recent report in two infants at 31 and 34 weeks of gestation, neither of whom were acutely ill and whose TSB levels were 13.1 mg/dl (224 µmol/l) and 14.7 mg/dl (251 µmol/l), has raised renewed concerns about low bilirubin kernicterus in the premature infant. We do not know why kernicterus has more or less disappeared from the NICU population. It could be the result of overall improvements in the care of LBW infants and/or the aggressive use of phototherapy.

EXCHANGE TRANSFUSION AND PHOTOTHERAPY

Exchange transfusions and phototherapy remain the staples of intervention for the jaundiced newborn. Phototherapy was introduced to the medical world at the Rochford (Essex) General Hospital in 1958 by Cremer et al. In the closing paragraph of their paper the authors, with typical reserve, had the following to say about their new discovery: "No prospect can be entertained that this light treatment will prove a substitute for exchange transfusion in the erythroblastotic infant with active haemolysis, but the method may be turned to clinical advantage in controlling the level of serum-bilirubin in cases of jaundice of prematurity." It would be no exaggeration to say that phototherapy has succeeded beyond their wildest dreams. Just how effective it has been can be gauged from the impact it has had on the number of exchange transfusions performed for hyperbilirubinaemia (fig 1), and nowhere has this difference been more dramatic than in infants weighing < 1500 g.

In a cohort of 833 infants with birth weights of 500–1500 g born in North Carolina between 1985 and 1989, only two infants (0.24%) underwent exchange transfusion and at William Beaumont Hospital in Royal Oak, Michigan, between 1988 and 1997 no exchange transfusions were performed in 1213 live births of infants weighing < 1500 g (fig 1). Certainly phototherapy, if used appropriately, is capable of controlling the bilirubin levels in almost all LBW infants, with the possible exception of the occasional infant with severe erythroblastosis fetalis or pronounced bruising. Furthermore, exchange transfusion, when performed at the low bilirubin levels used for treatment in LBW infants, is very inefficient and is less effective than phototherapy in achieving prolonged reduction of TSB in infants with non-haemolytic jaundice.

EXCHANGE TRANSFUSION COMPLICATIONS

Sick, preterm infants are much more likely than term infants to experience a serious complication of exchange transfusion, such as an arrhythmia, thrombosis, thrombocytopenia, necrotising enterocolitis, and infection among others, or to die during or soon after the procedure. In a US study of 331 exchange transfusions, there was one death (a rate of 0.3/100 procedures, 95% confidence interval (CI) 0 to 0.9) and significant morbidity in 5%. In 1472 exchange transfusions, Hove and Siimes identified four deaths “possibly related” to the procedure, a rate of 0.3/100 procedures (95% CI 0 to 0.5). Jackson reviewed the records of the Children’s Hospital Medical Center and the University of Washington Medical Center in Seattle, Washington, for 1980–1995, during which 106 infants had an exchange transfusion. Eighty one were healthy, and there were no deaths in these infants, although one child developed severe necrotising enterocolitis requiring surgery. There were 25 sick newborns; three had serious complications from the exchange transfusion and two (8%) of these infants died (95% CI 0% to 19%). In three additional infants, deaths were considered “possibly due” to the exchange transfusion, so that the mortality could have been as high as 20%. Four surviving infants had permanent serious sequelae: one of the 80 surviving well infants and three of 20 (15%) sick infants (95% CI 0% to 31%).

These rates, however, may not be generalisable to the current era. Experience with exchange transfusion is decreasing (fig 1), and it is
reasonable to assume that, like most procedures, frequency of performance is an important determinant of risk. It is certainly very difficult, if not impossible, to teach pediatric residents in the United States to perform exchange transfusions, and there is legitimate concern about our ability to train even neonatal fellows to perform this once common procedure. Indeed, with newer phototherapy technologies emerging, as well as the potential for pharmacological inhibition of bilirubin production, exchange transfusion in the NICU is in danger of becoming extinct. As discussed previously, the validity of traditional criteria for exchange transfusion in the LBW population has been questioned, and the introduction of more effective phototherapy for these infants has rendered much of the debate on exchange transfusion in this population moot.

**Table 1** Guidelines for the use of phototherapy and exchange transfusion in low birthweight infants based on birth weight

<table>
<thead>
<tr>
<th>Birth weight (g)</th>
<th>Phototherapy†</th>
<th>Exchange transfusion‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;1500</td>
<td>5–8 (85–140)</td>
<td>13–16 (220–275)</td>
</tr>
<tr>
<td>1500–1999</td>
<td>8–12 (140–200)</td>
<td>16–18 (275–300)</td>
</tr>
</tbody>
</table>

Note that these guidelines reflect ranges used in neonatal intensive care units. They cannot take into account all possible situations. Lower bilirubin concentrations should be used for infants who are sick—for example, presence of sepsis, acidosis, hypoprolactinemia—or have haemolytic disease.

*Consider initiating treatment at these levels. Range allows discretion based on clinical conditions or other circumstances. Note that bilirubin levels refer to total serum bilirubin concentrations. Direct reacting or conjugated bilirubin levels should not be subtracted from the total.

†Used at these levels and in therapeutic doses, phototherapy should, with few exceptions, eliminate the need for exchange transfusions.

‡Levels for exchange transfusion assume that bilirubin continues to rise or remains at these levels despite intensive phototherapy.

**Table 2** Guidelines for use of phototherapy and exchange transfusion in preterm infants based on gestational age

<table>
<thead>
<tr>
<th>Gestational age (weeks)</th>
<th>Total bilirubin level (mg/dl (µmol/l))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Phototherapy</td>
</tr>
<tr>
<td>36</td>
<td>14.6 (250)</td>
</tr>
<tr>
<td>32</td>
<td>8.8 (150)</td>
</tr>
<tr>
<td>28</td>
<td>5.8 (100)</td>
</tr>
<tr>
<td>24</td>
<td>4.7 (80)</td>
</tr>
</tbody>
</table>

*Rhesus disease, perinatal asphyxia, hypoxia, acidosis, hypercapnia

**Table 3** Guidelines according to birth weight for exchange transfusion in low birthweight infants based on total serum bilirubin (mg/dl) and bilirubin/albumin ratio (mg/g) (whichever comes first)

<table>
<thead>
<tr>
<th>Birth weight (g)</th>
<th>Total bilirubin 13</th>
<th>B/A ratio 5.2</th>
<th>Total bilirubin 10</th>
<th>B/A ratio 4.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;1250</td>
<td>15</td>
<td>6.0</td>
<td>15</td>
<td>6.0</td>
</tr>
<tr>
<td>1250–1499</td>
<td>17</td>
<td>6.8</td>
<td>17</td>
<td>6.8</td>
</tr>
<tr>
<td>1500–1999 g</td>
<td>18</td>
<td>7.2</td>
<td>17</td>
<td>6.8</td>
</tr>
<tr>
<td>2000–2499 g</td>
<td>18</td>
<td>7.2</td>
<td>17</td>
<td>6.8</td>
</tr>
</tbody>
</table>

*Risk factors: Aggar <3 at five minutes; PaO₂ <40 mm Hg; pH <7.15; PaCO₂ >2 h; birth weight <1000 g; haemolysis. Clinical or central nervous system deterioration; total protein <4 g/dl or albumin <2.5 g/dl. B/A ratio, Bilirubin/albumin ratio.

Bilirubin is a powerful antioxidant and may have a physiological role as an antioxidant in the human neonate. It has also been suggested that keeping TSB levels low with phototherapy (and thus reducing the antioxidant level) could facilitate the development of retinopathy of prematurity. In a study of 157 23–26 week gestation infants, however, no relation was found between TSB levels and retinopathy of prematurity. More than 90% of extremely LBW infants (birth weights <1000 g) receive phototherapy, and concern for the possible negative consequences of aggressive phototherapy has led the NICHD Neonatal Research Network to initiate a prospective randomised trial...
to compare aggressive with conservative phototherapy in these infants (B Morris, personal communication, 2002). Table 4 shows the TSB levels for intervention with phototherapy and exchange transfusion in this continuing trial. The primary outcome will be death or neurodevelopmental impairment at 18–24 months corrected age. The results of this study should provide important information about the risks and benefits of phototherapy in this vulnerable population.

Although the “aggressive” arm of the NICHD study (table 4) calls for phototherapy to be used prophylactically, in the absence of a documented increase in TSB, this is a practice that currently should be discouraged. In a randomised trial in LBW infants, prophylactic phototherapy (treatment initiated directly after birth) was compared with phototherapy when the TSB reached 5 mg/dl (85 μmol/l). Prophylactic phototherapy significantly lengthened the duration of treatment but had no beneficial effect on the bilirubin levels. This is hardly surprising as phototherapy probably acts on bilirubin in the capillaries of the skin or interstitial space, and until there is some increase in the serum bilirubin, phototherapy is ineffective.

Tables 1–3 provide suggested guidelines for the management of hyperbilirubinaemia in LBW infants. Phototherapy is generally used according to a sliding scale: the lower the birth weight or gestation, the lower the TSB level at which phototherapy is instituted. However, as discussed above, there is little good evidence to support this practice except perhaps for the influence of gestation on bilirubin-albumin binding. These guidelines reflect ranges used in different NICUs and cannot take into account all possible situations.

### ALBUMIN BINDING AND THE BILIRUBIN/ALBUMIN RATIO

Bilirubin is transported in the plasma tightly bound to albumin, and the portion that is unbound or loosely bound can more readily leave the intravascular space and cross the intact blood-brain barrier. Rises in unbound bilirubin have been associated with kernicterus in sick, preterm newborns. In addition, raised unbound bilirubin concentrations are more closely associated than TSB levels with transient abnormalities in the audiometric brain stem response in both term and preterm infants. There are, however, no contemporary long term studies relating unbound bilirubin concentrations in LBW infants to developmental outcome. In one follow up of 224 infants born in 1974–1976 with birth weights below 2000 g and evaluated at age 6 years, no relation was found between measures of bilirubin-albumin binding and intelligence quotient scores. In addition, clinical laboratory measurement of unbound bilirubin is not generally available.

The ratio of bilirubin (mg/dl) to albumin (g/dl) does correlate with measured unbound bilirubin in newborns and has been used as an approximate surrogate for the measurement of unbound bilirubin. It must be recognised, however, that albumin binding capacity varies significantly between newborns, is impaired in sick infants, and increases with increasing gestational age and postnatal age. Furthermore, the risk of bilirubin encephalopathy is not simply a function of the TSB level or the concentration of unbound bilirubin, but is a combination of both—that is, the total amount of bilirubin available (the miscible pool of bilirubin) as well as the tendency of bilirubin to enter the tissue (the unbound bilirubin concentration). An additional factor is the susceptibility of the cells of the central nervous system to damage by bilirubin. The bilirubin/albumin ratio can therefore be used together with, but not in lieu of, the TSB level as an additional factor in determining the need for exchange transfusion (table 3).

### USING PHOTOThERAPY EFFECTIVELY IN THE LBW INFANT

Given that the mainstay of treatment for neonatal jaundice in preterm infants is phototherapy, we review aspects of this treatment that influence its effectiveness.

#### Light source and dose

We use phototherapy in the NICU at quite low TSB levels, to prevent the TSB from rising to a level where exchange transfusion may be necessary. Like any therapeutic agent, phototherapy should be administered in an adequate dose, and this can be estimated quite easily using a radiometer obtained from the manufacturer of the phototherapy device. Figure 2 illustrates the dose-response relation between the irradiance used and the rate at which the TSB level declines in term infants under phototherapy.

Most commonly used phototherapy units deliver enough output in the blue-green region of the visible spectrum to be effective for conventional phototherapy in LBW infants. On the other hand, if bilirubin levels approach a range at which exchange transfusion is indicated, then more intensive therapeutic forms of phototherapy should be used. This is achieved by increasing the irradiance and the surface area exposed (see below). The most effective light source currently commercially available is that provided by special blue fluorescent tubes. These tubes are labelled F20 T12/BB or PL52/20W, and they provide much greater irradiance than regular blue tubes (labelled F20T12/B) (fig 3). Special blue tubes are more effective because they provide light with wavelengths predominantly in the blue-green spectrum,
where light penetrates the skin well and is absorbed maximally by bilirubin. Fibre optic phototherapy systems have several advantages over conventional phototherapy. Non-distressed neonates can be swaddled or dressed with the fibre optic blanket placed inside the blankets or clothes. These infants do not need to have their eyes covered (a major benefit to the family), but if the infant is lying (undressed) on a fibre optic pad, eye patching is advisable. In some studies of LBW infants, fibre optic phototherapy has been as effective as conventional phototherapy but it is less effective when compared with special blue phototherapy lamps.

**Distance from the light**

Spectral irradiance increases dramatically as the distance between the light source and infant decreases, and this effect is greatest when special blue tubes are used (fig 3). If bilirubin levels are rising in spite of phototherapy—for example, severely bruised infants or those with haemolytic disease—a free standing bank of special blue fluorescent tubes should be placed about 10 cm from the infant. This proximity is accomplished by placing the fluorescent bank of lights between the radiant warmer and the incubator. If halogen spot phototherapy lamps are used, they must not be positioned closer to the infant than recommended by the manufacturer because of the risk of a burn. In addition, if two or more halogen lamps are used, they should never be focused on the same area of the infant’s skin, as this may also cause a burn.

**Surface area**

The efficacy of phototherapy is closely related to the surface area of the infant exposed to the phototherapy lights, and the simplest way of increasing the surface area exposed is to place fibre optic pads below the infant with phototherapy lamps above. In preterm infants, this type of “double phototherapy” is approximately twice as effective as single phototherapy. If this intervention does not produce the desired decrease in the serum bilirubin level, additional exposure can be achieved by lining the sides of the radiant warmer or the incubator with a reflecting material such as white linen or aluminium foil.

**Intermittent versus continuous phototherapy**

Clinical studies comparing intermittent with continuous phototherapy have produced conflicting results, but in most circumstances phototherapy does not need to be continuous. As long as the serum bilirubin level is being controlled, phototherapy can certainly be interrupted during feeding or parental visits. It is particularly important during parental visits to switch off the phototherapy lights and remove the eye patches. Not being able to see their baby’s eyes is extremely disconcerting for parents.

**Hydration**

Some studies suggest that phototherapy significantly increases insensible water loss in preterm infants, but more recently it has been shown that as long as skin temperature is kept constant (by servo-control) and infants are not subjected to heat stress, phototherapy does not lead to an increase in oxygen consumption or insensible water loss through the skin or the respiratory tract. As we have simple and effective ways of monitoring newborn hydration (such as a daily weight and measurements of electrolytes), there is no indication for providing additional fluids routinely to infants who are receiving phototherapy. They should, of course, be kept adequately hydrated, not because phototherapy produces dehydration but because adequate urine output is important for effective phototherapy. The bilirubin isomer, lumirubin, is excreted in the bile and the urine, and lumirubin excretion appears to be an important element in the bilirubin lowering function of phototherapy.

**Complications**

Major complications associated with phototherapy are exceptionally rare. Perhaps the most important clinical complication encountered during the use of phototherapy in the LBW infant is that associated with the presence of direct hyperbilirubinaemia or cholestatic jaundice (usually after prolonged parental nutrition). When infants with direct hyperbilirubinaemia are exposed to phototherapy, they may develop a dark, greyish-brown discolouration of the skin, serum, and urine (the “bronze baby syndrome”). The pathogenesis of this syndrome is unknown, but it is possibly related to the accumulation of porphyrins or other pigments in the plasma in the presence of cholestasis. Although few deleterious consequences of the bronze baby syndrome have been described, two infants with this syndrome who died were shown to have kernicterus at autopsy. Except for these case reports, there are no other reports of significant complications in infants who develop the bronze baby syndrome, although impaired binding of bilirubin to albumin has been detected in three infants with this syndrome and in infants with cholestasis. If there is a need for phototherapy, the presence of direct hyperbilirubinaemia should not be considered a contraindication to its use, particularly in the sick newborn, and, as a rule, the direct serum bilirubin level should not be subtracted from the total bilirubin concentration when decisions are made about initiating phototherapy or an exchange transfusion. In infants who develop the bronze baby syndrome, exchange transfusion should be considered if phototherapy does not lower TSB. Because of the paucity of data, however, firm recommendations cannot be made. Rarely, purpuric bullous eruptions have also been described in infants with severe cholestatic jaundice receiving phototherapy.
REFERENCES


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