Albumin Administration in Exchange Transfusion for Hyperbilirubinaemia

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Exchange transfusion for neonatal hyperbilirubinaemia is usually carried out using twice the infant's blood volume of donor blood over a period of about 100 minutes (Forfar et al., 1958; Valaes, 1959). The importance of the albumin fraction in the plasma has been emphasized by Odell (1959a, b), who showed that the diffusion of bilirubin between plasma and the extravascular tissues is influenced by the capacity of albumin to bind the pigment. During an exchange transfusion the albumin in the donor blood replaces the infant's bilirubin-laden protein, and thus promotes movement of bilirubin from extravascular tissues into the circulation.

Kitchen, Krieger, and Smith (1960) claimed that by substituting 12.5 g. salt-poor human albumin in 50 ml. water for an equal volume of whole blood at the beginning and again half-way through an exchange transfusion the efficiency of bilirubin removal was increased by about 50%. Odell, Cohen, and Gordes (1962) 'primed' the infant with an intravenous injection of human albumin given 1 to 4 hours before exchange and in a dose of 1 g./kg. They obtained an apparent increase of 41% in bilirubin removal. Using one dose of 12.5 g. salt-poor albumin in 50 ml. water in place of donor plasma, Waters and Porter (1964) obtained an increase in efficiency of 26%.

The object of the present investigation was to evaluate the effect of human albumin administered either by substitution or by priming on bilirubin removal, albumin concentration, and bilirubin shift in the plasma of infants undergoing exchange transfusion.

**Material and Methods**

Infants requiring exchange transfusions for hyperbilirubinaemia were treated by three different methods. All those over 10 hours old (at which time plasma shift would be complete (Gairdner et al., 1958)) and whose pre-exchange bilirubin was over 12 mg./100 ml. were included. Infants were allotted to one of three groups by random selection, and their composition as regards age, maturity, birthweight, haematocrit, and pre-exchange bilirubin level is shown in Table I.

*Group I (controls):* 19 infants who were treated by simple exchange transfusion.

*Group II (substitution exchanges):* 17 infants who at every 100-ml interval during the exchange had 10 ml. (2.5 g.) salt-poor human albumin substituted for the blood, similar to the method of Kitchen, and using a total of about 20 g. albumin.

*Group III (primed exchanges):* 17 infants whose exchange transfusion was preceded by the intravenous injection of 2.5–10 g. albumin (approximately 1.75 g./kg.) 1–6 hours before the procedure. This was a larger dose of albumin than Odell's 1 g./kg.

Infants in all three groups were given two-volume exchange transfusions using the umbilical vein. Samples were taken into heparinized containers before and after exchange. Acid-citrate-dextrose blood with a haematocrit of approximately 50% was exchanged in 10 ml. aliquots over 60–146 minutes. All blood removed from the infant was collected in a dark-glass container with 2500 units heparin added. Care was taken to prevent contamination of waste blood by either saline washings, donor blood, or albumin. After measuring the volume removed, samples were taken for the various investigations. Haematocrit, bilirubin, and albumin concentrations were estimated within a few hours of collection when this occurred during laboratory working hours; otherwise haematocrit estimations were done by the authors, and plasma from centrifuged specimens was stored in the dark at 4°C.

Plasma bilirubin was estimated spectrophotometrically (White, Haidar, and Reinhold, 1958) and plasma albumin concentration by a modification of the method described by Bartholomew and Delaney (1966).

Calculations of blood volume, plasma volume, total mass of intravascular bilirubin, and total mass of bilirubin removed were based on the work of Valaes (1963).

1. **Blood volume (BV).** Mollison, Veall, and Cutbush (1950) found the mean blood volume in newborn infants to be 85 ml./kg., but that it differed according to age, early or late clamping of the cord, maturity, and venous haematocrit. By matching the three groups these factors should largely cancel out; blood volume was therefore calculated in ml. as: birthweight in kg. × 85.
TABLE I
Comparative Data of Infants in Groups I, II, and III

<table>
<thead>
<tr>
<th>Data</th>
<th>Group I</th>
<th>Group II</th>
<th>Group III</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of infants</td>
<td>19</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td>Maturity (wk.)</td>
<td>38-1</td>
<td>37-8</td>
<td>37-5</td>
</tr>
<tr>
<td>Birthweight (kg.)</td>
<td>35-41</td>
<td>36-40</td>
<td>34-40</td>
</tr>
<tr>
<td>Age (hr.)</td>
<td>2-96</td>
<td>3-03</td>
<td>2-92</td>
</tr>
<tr>
<td>Haematocrit (%)</td>
<td>2-06-3-75</td>
<td>2-47-3-75</td>
<td>2-05-4-12</td>
</tr>
<tr>
<td>Pre-exchange bilirubin (mg./100 ml.)</td>
<td>10-5-112</td>
<td>20-78</td>
<td>15-104</td>
</tr>
<tr>
<td>Time taken for exchange (min.)</td>
<td>46-6</td>
<td>44-1</td>
<td>41-9</td>
</tr>
<tr>
<td></td>
<td>39-66</td>
<td>31-53</td>
<td>33-51</td>
</tr>
<tr>
<td></td>
<td>20-1</td>
<td>20-0</td>
<td>21-0</td>
</tr>
<tr>
<td></td>
<td>12-4-32-8</td>
<td>13-7-33-5</td>
<td>15-1-25-8</td>
</tr>
<tr>
<td></td>
<td>82-9</td>
<td>85-4</td>
<td>78</td>
</tr>
<tr>
<td></td>
<td>60-145</td>
<td>61-115</td>
<td>60-114</td>
</tr>
</tbody>
</table>

*No significant difference between Groups I and III or Groups II and III.

(2) Plasma volume (PV) in ml. = BV x 100 — haematocrit.

(3) Total mass of intravascular bilirubin in mg. = PV x pre-exchange bilirubin concentration in mg./ml.

(4) Total mass of bilirubin removed by exchange in mg. = volume of blood removed x 1 — haematocrit of blood removed

100

x plasma bilirubin concentration of blood removed.

(5) Bilirubin clearance % =

total mass of bilirubin removed x 100.

total mass of intravascular bilirubin (3) x 100.

Although approximately twice the blood volume was exchanged in each case, variations in haematocrit inevitably led to variations in the size of the plasma volume exchanged, the actual range being 1-6-2-8. Therefore, as it is the plasma and not the red cell fraction which is concerned with bilirubin removal, bilirubin clearance has been expressed per 2 volumes of plasma exchanged.

Results

Bilirubin removal. In Table II the figures show that the amount of bilirubin removed was increased significantly in Group II and marginally in Group III as compared with the control group. This increase applied whichever method was used to express the amounts of bilirubin removed from the three groups.

Albumin concentration. Table III shows the mean differences in serum albumin concentrations before and after exchange transfusion in the three groups. In the control group there was a mean fall of 0-5 g./100 ml. In Group II the concentration was raised to 5-03 g./100 ml., an increase of 1-42 g./100 ml. over the pre-exchange level. Group III showed an over-all mean decrease of 0-04 g./100 ml., the post exchange concentration being 3-59 g./100 ml.

Effect of albumin priming in Group III (primed) cases. Table IV sets out the effects of administration of albumin on the haematocrit, bilirubin concentration, total bilirubin mass, and albumin concentration, in the time interval between priming with albumin and before starting the exchange. In each case there was a rise in bilirubin and albumin concentrations, the mean increases
As the albumin concentration increased as the exchange was made the bilirubin estimation becomes less accurate.

**Discussion**

The administration of large amounts (20 g.) of human albumin as an adjunct to exchange transfusion resulted in a significant increase in bilirubin removal as compared with simple exchange transfusion. The probable explanation of this effect is that bilirubin diffusion from the extravascular into the intravascular space is accelerated when large numbers of binding sites are made available by the addition of albumin (Sproul and Smith, 1964). In terms of bilirubin removal the results reported here are less impressive than Kitchen's. His cases varied widely as regards age, weight, and especially in pre-exchange serum bilirubin concentration, the mean in the control group being 19⋅3 mg./100 ml. compared with 10⋅7 mg./100 ml. in the albumin-treated group. We found that in our cases with pre-exchange bilirubin levels of less than 10 mg./100 ml. (not included in this series) the mean clearance was 154% as compared with 123% in our control cases with pre-exchange bilirubin levels above 10 mg./100 ml. In addition, these lower levels are of slight clinical interest only, while the method of bilirubin estimation becomes less accurate. Our Group III (primed) cases showed a statistically insignificant increase in bilirubin removal. Expressing our results as mg. bilirubin removed per kg. birthweight, our control group was nearly as effective as Odell's albumin-primed group (see Table V). This difference is difficult to explain unless our control cases were given more plasma relative to the total blood volumes exchanged.

In Group III cases the effect of albumin priming was similar to that in Odell's series, in that there was a marked rise in the serum bilirubin concentration of the order of 3 mg./100 ml. As the mean time interval was 2½ hours, this implies extraction of bilirubin from the tissues and its discharge into the blood. Had no such shift occurred, the bilirubin concentration should have decreased since the plasma volume increased as evidenced by the fall in haematocrit.
TABLE V
Comparison of Bilirubin Removal (expressed as mg. bilirubin removed per kg. infant's birthweight)

<table>
<thead>
<tr>
<th>Author</th>
<th>Simple Exchange</th>
<th>Albumin Substituted</th>
<th>Albumin Primed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Odell et al. (1962)</td>
<td>9-3</td>
<td>12-33</td>
<td></td>
</tr>
<tr>
<td>Present study</td>
<td>5-7-13-3</td>
<td>7-7-17-2</td>
<td>14-0</td>
</tr>
<tr>
<td></td>
<td>11-9</td>
<td>4-4-20-3</td>
<td>10-8-18-2</td>
</tr>
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</table>

Bilirubin removal by exchange transfusion is only one aspect of the protection of infants against bilirubin toxicity, as this procedure may only temporarily reduce the level of serum bilirubin. Studies by Diamond and Schmid (1966) suggest that brain damage occurs only when the sites for binding bilirubin to serum albumin are saturated either with bilirubin or competing substances. It follows that by increasing the serum albumin concentration, and therefore the number of available binding sites, the infant should be able to tolerate a level of serum bilirubin which would otherwise lead to brain damage. Having shown in Group II infants that large amounts of albumin administered during the exchange resulted in an increase of serum albumin from a mean of 3.61 g./100 ml. to 5.03 g./100 ml., and that this was well tolerated, the infants in Group III who had a slight fall in total serum albumin at the end of the exchange were then given a booster dose of 2.5 to 3.75 g. albumin to provide further binding sites. The effect of this procedure on the bilirubin-binding capacity of the serum will be discussed in a later publication. Preliminary investigations on these lines, based on the method of Porter and Waters (1966) and Waters (1967), are encouraging, but follow-up periods are still short.

Possible disadvantages in the administration of albumin suggested by Ruys and van Gelderen (1962) were not encountered, care being taken not to use albumin in severely anaemic infants, the lowest haematocrit being 35. The number requiring subsequent 'top up' transfusion did not exceed expectation.

Conclusion

Human albumin administered either during, or before and after, exchange transfusion in the amounts quoted is safe in the non-anaemic infant. Within these limits the more albumin given the more bilirubin will be removed. Odell's method of priming before an exchange may provide some immediate protection for the brain while awaiting collection and cross-matching of suitable donor blood. A further boost afterwards should provide further cover during the rebound phase by providing additional binding sites.

Summary

A total of 53 infants undergoing exchange transfusion for hyperbilirubinaemia were divided into two albumin-treated groups, and a control group. One group received a total of about 20 g. human albumin intravenously during the course of exchange; a significant increase in bilirubin removal resulted. The other group received a 'priming' dose of about 1.75 g./kg. albumin intravenously 1–6 hours before exchange; no significant increase in bilirubin removal resulted.

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REFERENCES


