Peritoneal Dialysis in the Reduction of Blood Ammonia Levels in a Case of Hyperammonaemia

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Four disease entities associated with hyperammonaemia and corresponding to lack of each of the specific enzymes required in the Krebs Henseleit urea cycle have now been reported (Russell et al., 1962; Freeman et al., 1964). A closely allied condition, lysine intolerance, where lysine competes with arginine for arginase has also recently been described (Colombo et al., 1964). Patients with these conditions have extremely high blood and CSF levels of ammonia, frequently much higher than in hepatic coma. The acute neurological sequence of spasticity, confusion, and coma during hyperammonaemic crises in these patients is analogous to that in hepatic coma precipitated by ingestion of protein in cirrhotic patients with an Eck fistula. Chronic ammonia intoxication, on the other hand, is characterized by periodic vomiting, deterioration of intelligence, progressive spasticity, and cerebral atrophy. Both haemodialysis and peritoneal dialysis have been shown to be of limited use in the treatment of ammonia intoxication in hepatic failure (Kiley et al., 1958; Krebs and Flynn, 1967; Maxwell et al., 1959; Nienhuis, 1966). Increase in blood ammonia is commonly associated with hepatic coma but is frequently not the precipitating cause. This may explain why reduction in ammonia levels by dialysis has not been more successful (Brown, 1967).

This paper reports a patient with hyperammonaemia due to deficiency of ornithine transcarbamylase in whom peritoneal dialysis was of considerable value in reducing blood ammonia levels.

Case Report

Full clinical and biochemical details of the patient are reported elsewhere in this issue (Hopkins et al., 1969) and are here given in summary.

The patient, a female aged 18 months, had had two previous hospital admissions because of vomiting, irritability, and failure to maintain normal physical and mental development. At 13 months hyperammonaemia was suspected and blood ammonia was found to be 428 μg./100 ml. and 734 μg./100 ml. on 2 estimations (normal range for adults < 60 μg./100 ml.). A low protein diet of high caloric and essential amino acid content (1·5 g. protein/kg. per day) resulted in marked clinical improvement. Blood ammonia fell to 100 μg./100 ml. over the following 2 weeks and subsequently to normal levels (26 and 56 μg./100 ml.) over the next 2 months.

At 18 months an open liver biopsy was performed under nitrous oxide and halothane anaesthesia. Her condition remained satisfactory for 12 hours after operation, but rapidly deteriorated over the ensuing 12 hours. During this time she took only small quantities of oral glucose and water, and blood ammonia rose from 72 μg./100 ml. immediately after operation to 1000 μg./100 ml. after 24 hours. This was associated with dehydration, acidosis, intermittent convulsions, and profound coma. Intravenous therapy was started with 5% dextrose in ½ isotonic saline and added bicarbonate. After rehydration blood ammonia fell to 670 μg./100 ml., but by the next day this had risen to 1970 μg./100 ml. despite a protein sparing regimen with intravenous fructose and anabolic steroids.

Peritoneal dialysis was started 54 hours after operation, using standard (1·5% dextrose) dialysis fluid containing KCl 6 mEq/l. and exchanges of 400 ml. (i.e. 40 ml./kg.) in 1-hourly cycles. The effect on blood ammonia is shown in the Fig. Blood ammonia had fallen to 800 μg./100 ml. after 4 hours, at which time the child was responding to painful stimuli. 12 hours later the level had fallen to 640 μg./100 ml., with further improvement in conscious state. She now made spontaneous purposeful movements and was able to tolerate a short period without assisted respiration. However, deterioration occurred, she became cyanosed and acidic, and once again required assisted respiration. Dialysis was continued for a total of 36 exchanges over 36 hours, by which time blood ammonia had fallen to 330 μg./100 ml., a level previously unassociated with coma in this patient. Dialysis was suspended with the catheter left in situ, but blood ammonia rapidly rose to 780 μg./100 ml.
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μg./100 ml. and dialysis was restarted. No clinical improvement was observed after her second deterioration and she died 10 hours later. 5% dextrose in ½ isotonic saline was continued at approximately maintenance rates during the dialysis.

![Graph](https://via.placeholder.com/150)

**Fig.** Effect of peritoneal dialysis on blood ammonia levels. The abscissa represents time after open liver biopsy.

**Discussion**

Ammonia intoxication in the body occurs most commonly in hepatic failure when ammonia absorbed from the gut bypasses the liver due to porto-systemic shunting. Russell *et al.* (1962) reported 2 cases of hyperammonaemia due to ornithine transcarbamylase deficiency, where blood ammonia and CSF ammonia levels of 980 μg./100 ml. and 360 μg./100 ml., respectively, were found. These are much higher than the levels usually considered toxic in hepatic coma.

Both haemodialysis and peritoneal dialysis have been used to treat ammonia intoxication in hepatic failure. Kiley *et al.* (1958) showed that haemodialysis with a Kolff system removed blood ammonia, and improvement in conscious state followed some hours later. Nienhuis (1966) showed clinical improvement which paralleled fall in blood ammonia levels in a patient who recovered from hepatic coma after peritoneal dialysis, and Maxwell *et al.* (1959) showed temporary improvement in their patient with hepatic coma though blood ammonia levels were not estimated. Krebs and Flynn (1967) used peritoneal dialysis to prevent ammonia intoxication during exchange transfusion of a patient with acute hepatic failure. They were able to maintain blood ammonia levels at 200 μg./100 ml. even though dialysate levels were very low and ammonia levels of stored blood were high.

In our case peritoneal dialysis was chosen in view of the patient’s grave condition and the problems of cannulation in the very young. In addition, the relatively greater peritoneal membrane area makes peritoneal dialysis more efficient in a child (Esperanca and Collins, 1966). It was possible to produce an initial fall in blood ammonia level and with continuing dialysis to maintain a steady state (see Fig.).

Return to consciousness lagged behind the fall in blood level as in those cases reported by Kiley *et al.* (1958). After an initial rapid fall, blood ammonia fell more slowly to 300 μg./100 ml. This demonstrated the possibility of maintaining such a level of blood ammonia in the face of a persisting high rate of ammonia production, as evidenced by the rapid rise in blood ammonia from 300 to 780 μg./100 ml. when dialysis was suspended.

In previously reported cases of hepatic coma treated by dialysis ammonia levels have been much lower and dialysis less efficient. The higher levels present in our case would lead to more efficient removal of ammonia initially, owing to the high concentration gradient. Though direct measurement of ammonia in dialysis fluid was not performed, a Berthelot (phenolic hypochlorite) reaction was performed on dialysis fluid, and showed removal of large amounts of ammonia and glutamine nitrogen in the early stages, with a fall to much lower levels as the dialysis continued.

The rapid fall in blood ammonia levels in the initial phase may have been influenced by the absorption of glucose from the dialysis fluid (blood glucose rising to 340 mg./100 ml. at this stage). Brown *et al.* (1967) have shown a correlation between ammonia and glucose metabolism in patients with liver disease, particularly those with a porto-systemic shunt. They noted a fall in blood ammonia associated with a rise in blood glucose concentration, and suggested that this was due to utilization of ammonia by increased activity of the tricarboxylic acid cycle. This could act by producing more aspartate to combine with citrulline, or by increased conversion of α-ketoglutaric acid to glutamic acid and glutamine. The deficiency of ornithine transcarbamylase in our patient would limit the amount of citrulline available and make this first mechanism unlikely to be contributory.

However, the initial fall in blood ammonia levels could represent the combination of peri-
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toneal clearance and increased conversion of ammonia via \( \alpha \)-ketoglutaric acid.

Summary and Conclusion

An 18-month-old infant with hyperammonaemia due to ornithine transcarbamylase deficiency is described in whom peritoneal dialysis rapidly reduced blood ammonia from extremely high levels to lower levels which had previously not been associated with coma. At these levels, similar to those seen in hepatic coma (i.e. approximately 300 \( \mu \)g./100 ml.), a stable state was maintained using peritoneal dialysis even in the face of continuing high ammonia production.

Though removal of ammonia at lower levels is less efficient, peritoneal dialysis may be of value in the control of a hyperammonaemic crisis, as in this case, or where the condition leading to the high blood ammonia level is reversible.

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