Therapy of Bicarbonate-losing Renal Tubular Acidosis

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Donckerwolcke, R. A., van Stekelenburg, G. J., and Tiddens, H. A. (1970). Archives of Disease in Childhood, 45, 774. The therapy of bicarbonate-losing renal tubular acidosis. A 2-year-old girl with severe bicarbonate-losing renal tubular acidosis was treated successively with bicarbonate, THAM, and two diuretics, hydrochlorothiazide and frusenide. Only with hydrochlorothiazide was adequate correction of the acid-base balance achieved. The relative importance of changes induced by this treatment in the extracellular fluid volume and in chloride depletion was assessed.

In treating patients with bicarbonate-losing renal tubular acidosis, Rodriguez Soriano and co-workers (1967a, b) studied bicarbonate medication and pointed out the necessity of raising the plasma [HCO₃⁻] to a normal level. Since in this condition the renal threshold for HCO₃⁻ is lowered, this course leads to bicarbonaturia. The bicarbonate depletion can be limited, and the total dose required thus reduced, by frequently administering small amounts; in this way Rodriguez Soriano et al. (1967a, b) was able to achieve adequate correction in patients with only a slightly lowered HCO₃⁻ threshold, but no patients with a substantially lowered threshold were studied. Rampini et al. (1968) used diuretics in treating the impaired bicarbonate reabsorption in a patient with the Toni-Debré-Fanconi syndrome. Three diuretics were tried, hydrochlorothiazide, frusenide, and triamterene, but only with hydrochlorothiazide was adequate correction of acid-base balance achieved. They were unable to explain the difference in effect.

This paper presents the results of treating a patient with renal acidosis using bicarbonate, tris-hydroxymethyl-amino-methane (THAM), and two different diuretics.

Patient and Methods

The study concerns a 2-year-old girl described in an accompanying paper (Donckerwolcke, van Stekelenburg, and Tiddens, 1970). She was weighed daily and measured three times per week, using methods recommended by Tanner, Whitehouse, and Takaishi (1966). The pH, Pco₂, HCO₃⁻, and total CO₂ values were determined in finger-prick blood by the Astrup method. Blood volume was determined with radioactive iodine (¹³¹I) bound to human serum albumin. The bromide volume was determined in addition. The two parameters were measured simultaneously (de Planque et al., 1965), using the Volémétron. The extracellular (ECF) volume could be calculated from bromide volume and blood volume.

Therapeutic methods. A low salt diet was given (0.75 g. NaCl/24 hours) throughout the period of treatment. The patient was treated successively with sodium bicarbonate, a 15% THAM solution buffered with citric acid to pH 8.2, and two different diuretics, hydrochlorothiazide and frusenide.

Results

Treatment with bicarbonate. Initially, treatment with citrate had to be discontinued because of gastro-intestinal upset; sodium bicarbonate medication was then started at a daily dosage of 3 g., increased to 15 g., administered in eight fractional doses distributed evenly over 24 hours. When the total daily dose reached 15 g. (22 mEq/kg.), this medication also had to be abandoned because of gastro-intestinal symptoms. There was a negligible effect on serum pH, HCO₃⁻, and electrolytes (Fig. 1).

Treatment with THAM. This compound was given orally in increasing doses up to 1-25 g./kg. per day divided into six doses. After three weeks it had to be discontinued because of severe diarrhoea. Only a slight effect on serum pH, Pco₂, standard HCO₃⁻ and total CO₂ was observed (Table I).

Treatment with two different diuretics. The optimal dosage was established empirically for both compounds. To prevent hyponatraemia, salt was added to the low sodium diet, first in the form

Received 6 May 1970.
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TABLE I

Effect of Oral Treatment with THAM

<table>
<thead>
<tr>
<th>Date (September 1969)</th>
<th>1st</th>
<th>2nd</th>
<th>5th</th>
<th>10th</th>
<th>17th</th>
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<tr>
<td>Blood chemistry</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>7-12</td>
<td>7-14</td>
<td>7-17</td>
<td>7-19</td>
<td>7-16</td>
</tr>
<tr>
<td>Pco2 (mm. Hg)</td>
<td>25-0</td>
<td>26-4</td>
<td>20-4</td>
<td>26-5</td>
<td>30</td>
</tr>
<tr>
<td>St. HCO3 (mEq/l.)</td>
<td>9-7</td>
<td>10-6</td>
<td>10-6</td>
<td>11-2</td>
<td>11-8</td>
</tr>
<tr>
<td>Total CO2 (mEq/l.)</td>
<td>8-5</td>
<td>9-4</td>
<td>8-0</td>
<td>9-1</td>
<td>11-2</td>
</tr>
<tr>
<td>Treatment</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>THAM (g./kg. per day)</td>
<td>0</td>
<td>0-5</td>
<td>0-75</td>
<td>1</td>
<td>1-25</td>
</tr>
</tbody>
</table>

of NaCl (1.5 g. daily) and then as NaHCO3 (3 g. daily). The effects of the two diuretics are shown in Fig. 2 and 3. Hydrochlorothiazide led to a good correction of the serum pH and HCO3 but was accompanied by severe hypopotassaeemia, such that substantial amounts of KHCO3 (up to 10 g. daily) had to be added to the therapy.

The initial effective dose of hydrochlorothiazide was 20 mg. daily; once correction was achieved, this could be reduced to 10 mg. daily (distributed over the day in four doses). No complications occurred during five months of this medication, though plasma potassium remained very low in spite of ingesting large amounts of KHCO3. No associated ECG changes were observed. By contrast, with frusemide no clear effect was obtained though the daily dosage was increased to 30 mg. (Fig. 3).

Discussion

Treatment with bicarbonate. On theoretical grounds it would be predicted that, taking into account the GFR (27 mg./min. per 0.3 m.2) and the HCO3 threshold (10.5 mEq/L), a daily dose of 375 mEq or 31.5 g. NaHCO3 would have had to be administered to attain a constant serum level of 24 mEq/L. So large an amount of sodium bicarbonate as is thus required by patients with a very low bicarbonate threshold and low HCO3 reabsorption cannot be administered orally; hence this mode of

![Fig. 1.—Serum acid-base and electrolyte values during treatment with sodium bicarbonate.](http://adc.bmj.com/)

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treatment is practicable only in patients with a minor lowering of the threshold.

**Treatment with THAM.** Vert et al. (1968) administered THAM to patients with severe acidosis resulting from various renal affections, with a favourable effect in a patient with distal tubular acidosis (type Lightwood).

The failure of this therapy in our patient, with pronounced proximal tubular acidosis, can be explained as follows. Administration of THAM reduces [H+] and thus leads to a change in the relative concentration of all proton donor and proton acceptor pairs operating as buffer systems in the biological pH range (Strauss, 1968). For the HCO3⁻/H2CO3 buffer system, this implies a decrease in [H₄CO₃] and an increase in [HCO₃]. In our patient an increase in serum [HCO₃] led to increased excretion, which greatly reduced the effect. Thus the therapy also seems to be of little value in patients with a severely lowered bicarbonate threshold.

**Treatment with diuretics.** Only with hydrochlorothiazide did we achieve correction of the acid-base balance; frusemide at the dosage used (3 mg./kg.) was ineffective. These findings corroborate Rampini's observations.

Diuretics can cause an increase in bicarbonate reabsorption by three different mechanisms of action: (1) by reduction of the extracellular volume; (2) by causing Cl⁻ depletion; and (3) by increased K⁺ excretion, resulting in intracellular K⁺ depletion. Several investigators have studied these various factors. The effect of changes in extracellular volume on renal HCO₃⁻ reabsorption was studied by Kunau et al. (1966), who demonstrated that expansion of the extracellular volume reduced HCO₃⁻ reabsorption in the proximal tubule. Oetliker and Rossi (1969) demonstrated in two patients with Lowe's syndrome that the renal HCO₃⁻ threshold was directly dependent on the plasma volume. Reduction of the extracellular volume gives rise to increased sodium avidity, whereupon sodium reabsorption increases due to raised Na⁺/H⁺ exchange, resulting in increased HCO₃⁻ reabsorption. The specific effect of chloride on the acid-base balance was studied by Schwartz et al. (1961), Gulyassy, van Ypersele de Strihou, and Schwartz (1962), and Needle, Kaloyanides, and Schwartz (1964), who showed that selective depletion of the chloride reserves led to metabolic alkalosis, that this
alkalosis was maintained by chloride restriction in the diet, and that chloride administration was essential in restoring the acid-base balance to normal. However, these conditions of chloride depletion are associated with a reduction of the extracellular volume (Seldin and Wilson, 1966), and the resulting raised Na⁺/H⁺ exchange can explain the alkalosis. Cohen (1968) suggested that, in alkalosis induced by diuretics, the restoration of the acid-base balance was achieved by restoration of the extracellular volume and chloride repletion, and that these two factors were interdependent. Though Rector, Bloomer, and Seldin (1964) have indicated that in hypopotassemia the proximal reabsorption of HCO₃⁻ increases, the hypopotassemia in his experimental arrangement was explicable as a result of increased Na⁺/cation exchange (Schwartz, van Ypersele de Strihou, and Kassirer, 1968).

Our study has shown that, during treatment with diuretics at the dosage used (by us and by Rampini et al., 1968), the extracellular volume and blood volumes decrease only during hydrochlorothiazide medication (Table II).

In addition we determined the relative importance

TABLE II

Extracellular Fluid Volume during Treatment with Diuretics

<table>
<thead>
<tr>
<th>Extracellular Fluid Volume Measurements</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>None</td>
</tr>
<tr>
<td>Br-volume (L)</td>
<td>2.93</td>
</tr>
<tr>
<td>Plasma volume (L)</td>
<td>0.50</td>
</tr>
<tr>
<td>Calculated blood volume (L)</td>
<td>0.71</td>
</tr>
<tr>
<td>Calculated ECF volume (L)</td>
<td>2.49</td>
</tr>
<tr>
<td>Haematocrit (%)</td>
<td>33</td>
</tr>
</tbody>
</table>
of chloride deficiency, during a situation of increased HCO₃⁻ reabsorption and reduced extracellular volume. To a regimen consisting of a low-sodium (88 mg./day), low-chloride (232 mg./day) diet, with a daily administration of 8 mg. hydrochlorothiazide and 2 g. NaHCO₃, potassium (100 mEq) in the form either of KHCO₃ or of KCl was added successively. The effect of chloride on the acid-base balance and on the extracellular volume (measured by changes in body weight) was thereby studied.

Addition of chloride led to recurrence of acidosis and to expansion of the extracellular volume (Fig. 4). Rampini et al., 1968 has shown that substituting potassium phosphate for potassium citrate in the course of treatment with hydrochlorothiazide did not alter the acid-base balance, so that the recurrence of acidosis in our patient was unlikely to have been the result of the withdrawal of bicarbonate.

These findings corroborate Cohen's (1968) contention that chloride depletion and reduction of the extracellular volume are interdependent.

Conclusions
Severe forms of bicarbonate-losing renal tubular acidosis cannot be corrected by means of oral sodium bicarbonate or buffer solutions owing to the low threshold for and consequent losses of HCO₃⁻.
Diuretic medication is a good alternative in these conditions, with a diuretic that exerts its influence mainly on the distal nephron. The resulting rise in proximal Na+/H+ exchange can lead to restoration of the acid-base balance.

With a diuretic-induced augmentation of Na+/H+ cation exchange, chloride depletion and extracellular volume are closely correlated. Consequently, attempts to correct hypokalaemia and alkalosis by giving potassium chloride (Schwartz et al., 1968) may prove ineffective. Correction of changes in the acid-base balance in our experiments also led to a diminution of the effect of the diuretic on extracellular volume.

REFERENCES


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Arch Dis Child 1970 45: 774-779
doi: 10.1136/adc.45.244.774

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