CONJUGATION REACTIONS IN THE NEWBORN INFANT: THE METABOLISM OF PARA-AMINOBENZOIC ACID

BY

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It has been shown that the liver of the newborn infant has a limited capacity to perform certain transformation or conjugation reactions when compared to older subjects (Driscoll and Hsia, 1958; Kretchmer, Levine, McNamara, and Barnett, 1956). This limitation is evident in the formation of glucuronides (Brown and Zuelzer, 1958; Vest, 1958) and plays an important role in the genesis of neonatal jaundice. There are indications that other conjugation or detoxification mechanisms are also carried out in a different way from that of later life. The observation that after administration of sodium benzoate the excretion as hippuric acid is incomplete and delayed during the first three to four months of life (Vest, 1959) prompted us to investigate hippurate formation and acetylation in this age period.

Material and Methods

Para-aminobenzoic acid (PAB), in the form of the sodium salt (NaPAB), was injected intravenously in a dose of 100 mg. per kg. body weight into four children 8 to 11 years old, three newborn premature infants of 2 to 6 days, two newborn full-term infants, and two 5- and 8-week-old infants. All subjects were male. Fluid intake was not restricted during the test.

Blood, 1-2 ml., was collected 5, 15, and 30 minutes, 1, 2, 3, 4, and, if necessary, 6 and 8 hours after the administration of the test dose. Urine was collected over 24 hours, divided in three 4-hour and a final 12-hour period. The urine was kept at 4°C. when assayed within a few hours, otherwise it was frozen at minus 20°C.

The sodium-p-aminobenzoate solution for intravenous use was prepared as follows: 10% NaPAB was dissolved in distilled water (w/v). The water was boiled before use and nitrogen bubbled through to remove carbon dioxide; 10 ml amounts were put into brown ampoules which were filled with nitrogen just before sealing. The ampoules were sterilized by autoclaving at 120°C for a quarter of an hour.

The method of Deiss and Cohen (1950) was used for the estimation of PAB and para-aminohippuric acid (PAH). PAH was estimated after extraction of PAB with ether. After extracting mixtures of known composition, on the average 1% PAB and 98% PAH remained. For the estimation of acetylated PAB and PAH (acetylated-benzoic acid plus acetylated-hippuric acid) in the serum aliquots of the deproteinized supernatant were hydrolysed by boiling with 0.05 volume of 4 N HCl for 20 minutes. In the case of urine better results were obtained by hydrolysing for 45 minutes. In some experiments the hydrolysis of acetylated compounds was carried out by heating at 96°C. for three and a half hours as suggested by Smith, Finkelstein, Aliminosa, Crawford, and Graber (1945). After addition of the diazo reagents, the samples were read against a reagent blank at 540 μm in a Coleman junior spectrophotometer. The amounts were determined from a calibration curve made with various concentrations of PAH. All results are expressed in micromols (μM) of PAH.

The colour intensity obtained without hydrolysis is equal to the free PAB and PAH. By extracting PAB, PAH alone can be measured. After hydrolysis the colour development is equal to the sum of PAH, PAB, and acetylated compounds. From this value the amount of total acetyl-metabolites (acetylated PAB and PAH) can be calculated by subtracting free PAB and PAH. Acetylated PAH is found by extracting the hydrolysate with ether (which removes the free and the formerly acetylated PAB) and subtracting the value of free PAH.

In two infants and one child the urinary metabolites were separated by counter-current distribution techniques. With each plate 5 ml. ethyl acetate was used as organic, and 5 ml. 1·6 M acetate buffer pH 3·4, as aqueous phase (Way, Smith, Howie, Weiss, and Swanson, 1948; Tabor, Freeman, Baily, and Smith, 1951). 50 to 100 plate distributions were run and both layers analysed for PAB, PAH, acetylated PAB and PAH, and glucuronides. For analysis the organic layer was first taken to dryness. A drop of ethanol facilitated dissolving the residue in water. The acetate phase was analysed directly. Glucuronide was estimated by the carbazole method of Dische (1947).

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The identity of the urinary metabolites separated by counter-current distribution was studied by running aliquots on paper chromatograms. An ascending system of butanol-glacial acetic acid and water in the proportions 4:1:2 (v/v) was used with Whatman No. 1 paper. Standards of PAH, PAB, acetylated PAH, and acetylated PAB were run simultaneously. Ehrlich reagent (p-dimethylaninobenzaldehyde, 10% in concentrated HCl solution (w/v) 1 vol., acetone 4 vol.) gave an immediate yellow colour with PAB, PAH, and PAB-glucuronide. With acetylated PAB and PAH the colour development was delayed. Compounds with a free amino group can also be distinguished from the acetylated metabolites by the Ekmann reagent specific for diazotizable amines. Glucuronyl derivatives were identified by the naphthothesorcinol and aniline-diphenylamine reagent (Smith, 1960).

**Results**

**PAB Metabolites in the Serum.** Fig. 1 shows typical examples of the course of the various PAB acid metabolites in the serum in the different age-groups. In the 2-day-old premature infant (Fig. 1A) free PAB disappears slowly from the circulation. PAH is the first conjugate to appear. It reaches a peak of 0.12 μM per ml. serum 3 hours after injection of NaPAB. Acetylated PAB is the chief conjugate formed. It reaches a peak concentration of up to 0.6 μM per ml. at 6 hours. Besides PAH and acetyl-PAB small amounts of acetylated p-aminohippuric acid are formed. This conjugate is the last to appear in significant amounts.

Compared to the newborn infants, free PAB decreases quicker in the 5-week-old and more so in the 2-month-old infant (1B and C). At an age of 5 weeks acetyl-PAB is still the major conjugate formed (Fig. 1B). In the 2-month-old infant PAH formation is more prominent, though it is still less than that found in older children (Fig. 1C). Acetylated PAB is present to a greater degree than in older children, but much less than at 5 weeks. Some acetylated PAH is also present. In the 8½-year-old child (Fig. 1D) there is a rapid decrease of free PAB. The major conjugate now is PAH which reaches a peak of 0.25 μM per ml. at 1 hour. Acetyl-PAB appears later and its concentration remains lower than that of PAH. Some acetyl-PAH is also present. Both acetylated metabolic products are still present 4 hours after injection, when PAB and PAH have already disappeared. No attempt has been made to investigate glucuronide conjugation in the serum.

Fig. 2 shows the average plasma disappearance curves of total PAB metabolites at different ages. In the newborn period it is rather flat; at 8 weeks it reaches the slope found in older children. The 5-week-old infant takes an intermediate position.
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The time elapsing until the plasma concentration of total PAB has fallen to 50% of the initial level, is about 6 hours for the newborns, 3 hours 40 minutes in the 5-week-old infant, and 1 hour and 40 minutes in the 8-week-old infant and in the children.

PAB Metabolites in the Urine. The excretion of PAB derivates in the urine reflects the situation in the serum. In Fig. 3 the cumulative urinary excretion of the 4 metabolites is given as a percentage of the injected dose of sodium PAB for the same children whose serum concentrations are shown in Fig. 1. In the newborn infant (Fig. 3A), the metabolites appear slowly. After 4 hours only about 10% of the given PAB is excreted. In the serum (Fig. 1A) the PAH formed has been largely excreted after 6 to 8 hours. The same is evident in the cumulative urinary excretion, i.e. after 8 to 12 hours the amount excreted as PAH hardly increases any further. The chief metabolites, acetylated PAB and PAH on the other hand, appear more slowly and are excreted predominantly between 8 to 24 hours. Free PAB accounts for less than 7% of the total excreted.

Fig. 3B and C show the transition from the pattern prevailing in the newborn period to the one later in childhood. Not only does the total 24-hour recovery increase but the excretion accelerates. In the 5-week-old infant the chief metabolites are still acetyl-PAB and -PAH, but at 8 weeks PAH is the major conjugate. Even at this age however, much more acetyl-PAB is excreted than later in life.

In the child (Fig. 3D), the excretion is much more rapid, so that 70 to 85% is recovered within 4 hours after injection. The chief conjugate is p-aminohippuric acid, which accounts for about 50% of the total PAB recovered. Free p-aminobenzoic acid generally takes the second place. Acetylated p-aminohippuric acid accounts for less than 10% of the recovered PAB.
**Table 1**

PERCENTAGE DISTRIBUTION OF THE DIFFERENT PAB DERIVATIVES IN THE URINE IN RELATION TO AGE-GROUPS

<table>
<thead>
<tr>
<th>Name</th>
<th>Age</th>
<th>Total 24-hour Recovery (%)</th>
<th>PAH (%)</th>
<th>PAB (%)</th>
<th>Acetyl PAH (%)</th>
<th>Acetyl PAB (%)</th>
<th>Glucuronides (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature new-borns</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M.P.</td>
<td>2 days</td>
<td>65-2</td>
<td>12-8</td>
<td>2-5</td>
<td>19-2</td>
<td>30-7</td>
<td>7-9</td>
</tr>
<tr>
<td>Sch. A.</td>
<td>4 days</td>
<td>47-7</td>
<td>14-6</td>
<td>1-4</td>
<td>15-5</td>
<td>16-2</td>
<td>7-7</td>
</tr>
<tr>
<td>L.R.</td>
<td>6 days</td>
<td>70-6</td>
<td>10-2</td>
<td>5-9</td>
<td>21-1</td>
<td>33-4</td>
<td>6-2</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>61-1</td>
<td>12-5</td>
<td>3-2</td>
<td>18-6</td>
<td>26-7</td>
<td>7-2</td>
</tr>
<tr>
<td>Full-term new-borns</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>M.R.</td>
<td>3 days</td>
<td>72-8</td>
<td>29-0</td>
<td>2-2</td>
<td>20-5</td>
<td>21-1</td>
<td>23-5</td>
</tr>
<tr>
<td>A.D.</td>
<td>5 days</td>
<td>62-3</td>
<td>24-5</td>
<td>2-3</td>
<td>12-2</td>
<td>23-3</td>
<td>13-7</td>
</tr>
<tr>
<td>Infants</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F.R.</td>
<td>5 wk.</td>
<td>78-4</td>
<td>16-9</td>
<td>6-0</td>
<td>17-0</td>
<td>38-5</td>
<td>16-9</td>
</tr>
<tr>
<td>S.J.</td>
<td>8 wk.</td>
<td>85-6</td>
<td>39-4</td>
<td>1-9</td>
<td>11-7</td>
<td>32-6</td>
<td>21-9</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td>87-2</td>
<td>47-2</td>
<td>15-7</td>
<td>8-2</td>
<td>16-0</td>
<td>16-8</td>
</tr>
</tbody>
</table>

Glucuronic Acid Conjugates. Because glucuronic acid conjugates are present in the urine even without PAB administration, it is difficult to estimate the percentage of PAB glucuronides accurately. If the quantity of glucuronic acid found in a 24-hour urine sample before the test is subtracted from the amount found after PAB injection, it is possible to get an estimate of the percentage of PAB excreted as a glucuronyl conjugate. As will be shown later, two glucuronyl conjugates, p-aminobenzoyl- and p-acetamido-benzoyl glucuronide, are present in various proportions depending on age. Because their solubility is not very different from that of the glycine conjugates, they stay behind when PAB is extracted. In the colorimetric assay these conjugates are therefore estimated along with PAH and acetyl-PAH respectively. They can, however, be measured separately by the Dische (1947) method. Analysis of the consecutive urine samples shows that in the newborn the glucuronides follow the same course as

![Figure 4](http://adc.bmj.com/)

Fig. 4.—Result of a 100 plate counter-current distribution of an aliquot of the first 4-hour urine sample in a 10-year-old child after intravenous PAB injection. The concentration of the various metabolites is given as μM PAH per plate.
the other PAB conjugates, i.e. they appear much slower than in the group of children.

In Table 1 the percentage of the various PAB derivatives in the urine is given for the different age groups. The value for the glucuronides is given separately but is already included in the column headed Total 24-hour Recovery, because, as mentioned above, they form a part of PAH and acetyl-PAH.

Separation of PAB Metabolites by Counter-current Distribution. Fig. 4 shows the results of the counter-current distribution in a 10-year-old child. A distribution of an aliquot of the first four-hour urine sample was run. A complete separation of PAH from acetyl-PAH and of PAB from acetyl-PAB could not be achieved, but the glucuronides were clearly separated from PAH and acetyl-PAH respectively and the latter from PAB and acetyl-PAB. The results agree with those obtained by colorimetric estimation. The chief metabolite present is PAH, with free PAB, acetylated PAB, and acetylated PAH following in decreasing order. There are two glucuronic acid conjugates: PAB- and acetyl-PAB-glucuronide. These do not quite account for all the glucuronic acid in the first plates. This is to be expected as other glucuronides are present in the urine. Planimetry of the curve results in the following percentage distribution:

- PAH 36·7, acetyl-PAH 2·7, PAB 28·8, acetyl-PAB 19·4, p-aminobenzoyl glucuronide 7·7, and acetamidobenzoyl glucuronide 4·7.

The result in a newborn infant is shown in Fig. 5. It was obtained from an aliquot of the urine collected from 8 to 24 hours and confirms the predominance of acetylated PAB and acetylated PAH over PAH, and the presence of PAB-glucuronides. In contrast to older children, p-acetamidobenzoyl glucuronide, not p-aminobenzoyl glucuronide is the main glucuronic acid conjugate in infants. The percentage distribution is as follows: PAH 3·7, acetyl-PAH 26·7, PAB 0·7, acetyl-PAB 54·8, p-aminobenzoyl glucuronide 2·2, acetamidobenzoyl glucuronide 11·9.

In Table 2 the ratio of distribution of the four standard compounds between ethyl acetate and buffer is compared to that calculated from the counter-current distribution. They agree only to a limited extent.

Chromatography of PAB Metabolites. Fig. 6 shows a chromatogram of the main peaks obtained by counter-current distribution in child E. J. Plate No. 24 has the same value (Rf 0·62) as PAH, 81 as acetyl-PAB (Rf 0·89), and 97 as PAB (Rf 0·83). In No. 2, which gives the highest reading for glucuronic acid, there are two spots the Rf values of which are

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**Table 2**

<table>
<thead>
<tr>
<th>PAB-derivative</th>
<th>Constant</th>
<th>Child E. Plate No.*</th>
<th>Constant</th>
<th>Plate No.†</th>
<th>Constant</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAH</td>
<td>0·29</td>
<td>24</td>
<td>0·31</td>
<td>22</td>
<td>0·30</td>
</tr>
<tr>
<td>PAB</td>
<td>11·5</td>
<td>92</td>
<td>11·5</td>
<td>PAB too low for calculation</td>
<td></td>
</tr>
<tr>
<td>Acetyl-PAH</td>
<td>0·31</td>
<td>24</td>
<td>0·31</td>
<td>19</td>
<td>0·25</td>
</tr>
<tr>
<td>Acetyl-PAB</td>
<td>7·85</td>
<td>86</td>
<td>6·15</td>
<td>85</td>
<td>8·5</td>
</tr>
</tbody>
</table>

* 100 plates. † 95 plates.
below that of the standards used. The lower one is diazotizable and gives an immediate Ehrlich reaction, i.e. it has a free amino group. Both react with naphthoresorcinol and aniline-diphenylamine. It is likely therefore that the lower spot (Rf 0.28) is PAB-glucuronide, whereas the second from the baseline (Rf 0.45), which gives a delayed reaction with Ehrlich reagent, is acetyl PAB-glucuronide. 

In Fig. 7 the same is shown for the newborn, M.P. Here, in plate No. 2, the upper spot, acetamido-benzoyl glucuronide (Rf 0.46) is much more prominent than PAB-glucuronide (Rf 0.30). Plate No. 17 corresponds to acetyl-PAH (Rf 0.79) and 84 to acetyl-PAB (Rf 0.89). As the counter-current distribution was done on the 8- to 24-hour urine sample no detectable PAH is present.

Direct chromatography of the urine shows the same metabolites (see Fig. 8). The 0-4 hour urine sample of child, E.J., and the 4 to 8 hour sample of the infants, M.P. and L.R., are shown together with both infants there is a spot for acetyl-PAH (Rf 0.76) beside the one for PAH (Rf 0.63). From the PAH glucuronides only the upper spot corresponding to acetyl-PAB glucuronide (Rf 0.46) is present.

**Discussion**

In children, intravenously administered NaPAB is rapidly cleared from the plasma mainly by conjugation with glycine, and to some extent by acetylation and glucuronidation. Nearly 50% of the administered dose is excreted as p-aminohippuric acid.
Other important metabolites are free and acetylated PAB (about 15% each) and the glucuronides of PAB and acetyl-PAB (about 16%). Little is excreted as acetylated PAH. This agrees with the results in human adults (Deiss and Cohen, 1950; Tabor et al., 1951), whereas in animals different proportions of the various metabolites have been found (Riggs and Christensen, 1951; Williams, 1959).

In newborn infants PAB disappears, more slowly from the circulation. There is diminished and delayed conjugation with glycine. This confirms the observation of Brandt (1960) who found that the activity of the hippurate synthesizing system of the liver in foetal and newborn rats is diminished. Instead of hippurate formation, acetylation at the amino group is much more prominent than in the children. This is to a certain extent at variance with the report of a limited acetylation of sulphonamides in the newborn period (Fichter and Curtis, 1955). With PAB there is, however, not only a predominance of acetyl-PAB over free PAB, but also of acetyl-PAH over PAH and acetamidobenzoyl glucuronide over p-aminobenzoyl glucuronide. In accordance with previous results (Brown and Zuelzer, 1958; Vest, 1958), the extent of glucuronide formation is also lower in the newborn premature infants than later in life.

It is difficult to establish to what extent a diminish-
105 ml. In the child the inulin clearance was 111, PAH 535, PAB 80, acetyl PAH 260, and acetyl PAB 430 ml./min./1·73 sq. m. The low PAB clearance accounts for the small amount of free PAB excreted by the newborn infants. The predominance of the acetylated metabolites cannot, however, be explained by the clearance studies.

Every alteration of these reactions or their rate could influence the extent of hippurate formation. So it is conceivable that the glycine-condensing enzyme matures more slowly than the acetylase.

Lack of available glycine could contribute to the decreased PAH formation. Administration of 1 g. glycine to newborn infants one hour before the

The reason why acetylation is enhanced in the newborn at the expense of hippurate formation is not clear. Hippuric acid and PAH synthesis requires an activated form of benzoic acid, benzoyl co-enzyme A. The later is then able to acylate glycine (Schachter and Taggart, 1953). This reaction is catalysed by the enzyme glycine-N-acylase (Schachter and Taggart, 1954). Similarly acetylation requires the previous formation of 'active acetate', i.e. acetyl co-enzyme A. In the case of aromatic amines the acetylating enzyme is called arylamine acetylase.

PAB injection actually leads to increased PAH concentration in the plasma and to a rise in the PAH excretion in the urine as it does in adults, but PAH formation is still much inferior to that found in children (Vest and Rossier, 1963).

Summary

The metabolism of p-aminobenzoic acid (PAB) was studied by injecting a dose of 100 mg. per kg. of the sodium salt into 3 newborn premature infants, 2 full-term newborns, 2 infants 5 and 8 weeks old,
and 4 children 8½ to 11 years old. In the children PAB is rapidly cleared from the serum, chiefly by formation of p-aminohippuric acid (PAH). The 24-hour recovery in the urine lies between 76 and 93%, of which 68 to 85% are excreted within the first 4 hours. The chief conjugate in the urine is PAH, which accounts for nearly half of the total. Other metabolites are acetyl-PAB, free PAB, acetyl-PAH, PAB- and acetyl-PAB-glucuronides. In the newborn infants PAB is eliminated tardily from the circulation. The main conjugate formed is acetyl-PAB. PAH, acetyl-PAH, and free PAB are other metabolic products. The excretion in the urine is also delayed so that only about 10% of the given dose appears within the first 4 hours. The 24-hour recovery amounts to between 48 and 70%. Again the major metabolite is acetylated PAB with acetyl-PAH, PAH, and free PAB following in decreasing order. Glucuronides account for only about 7%, acetylatedbenzoyl glucuronide prevailing over p-aminobenzoyl glucuronide. As the two older infants show, the transition from the pattern in the newborn period to that in children takes place around the 8th week of life.

References


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