Urinary amino-acetone excretion in lead workers

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SYNOPSIS  Urinary amino-acetone concentrations were determined in a group of 51 persons with industrial exposure to lead and the findings compared with those obtained from a control group of 27 persons having no exposure to lead.

A good correlation was found between delta-aminolaevulinic acid and amino-acetone excretion in the control group but this correlation was not so well defined in the lead-exposed group. In this group excretion of amino-acetone seemed to be better correlated with excretion of delta-aminolaevulinic acid for values less than 0·6 mg. % than with values greater than 0·6 mg. % . All the men concerned in the investigation were employed in dusty occupations. Hypotheses are put forward to attempt to explain the differences in excretion of amino-acetone noted amongst lead workers with high excretion of delta aminolaevulinic acid.

Little correlation was found between amino-acetone excretion and that of either porphobilinogen or coproporphyrin.

Amino-acetone is formed by the condensation of glycine with acetyl coenzyme-A (Gibson, Laver, and Neuberger, 1958; Kikuchi, Kumar, and Shemin, 1959; Urata and Granick, 1961) or by the oxidation of the hydroxyl group of L-threonine (Elliott, 1959; Neuberger and Tait, 1960; Urata and Granick, 1961). Its role in metabolism may be as part of a cycle by which glycine and threonine can be oxidized, and such a cycle was postulated by Elliott (1959) as an alternative (Fig. 1) to the succinate-glycine cycle of Shemin, Russell, and Abramsky (1955).

Amino-acetone was first reported to be present in the urine of normal individuals by Mauzerall and Granick (1956) and subsequently was found to be absent from the urine of animals with experimental porphyria (de Matteis, 1962) and of patients with acute intermittent porphyrja (de Matteis and Rimington, 1962). An hypothesis was put forward by these authors to explain the metabolic defect in acute intermittent porphrya on the basis of preferential metabolism of glycine through the succinate-delta-aminolaevulinic acid cycle rather than through the acetate-amino-acetone cycle (Fig. 2). It is well known that the symptoms of lead poisoning and those of acute intermittent porphyrja are very similar, so much so that it has been suggested that the mechanism of the production of the symptoms

FIG. 1. The amino-acetone-acetate cycle.

FIG. 2. Glycine-delta-aminolaevulinic acid and amino-acetate cycles. $C = \alpha$ carbon atoms of glycine to purines, formate, etc.
of the two diseases might have a common metabolic pathway (Goldberg, Smith, Lockhead, and Dagg, 1962). Although recent reports in which aminoacetone was found in the urine of animals with experimental porphyria (Granick and Urata, 1963) and of patients with acute intermittent porphyria (Tschudy, Welland, Collins, and Hunter, 1963) have cast some doubts on the hypothesis of de Matteis and Rimington, it was thought worthwhile to conduct a survey amongst workers exposed to lead to see whether any differences in their excretion of aminoacetone were noted as compared with that of a normal control group.

**METHOD**

Urinary concentrations of aminoacetone were measured in a group of 51 workers with exposure to lead and, as a control, in a group of 27 persons with no history of exposure to lead. In addition delta-aminolaevulinic acid and porphobilinogen were determined on each urine specimen, which, as was aminoacetone, were estimated by a modification of the method of Urata and Granick (1963) as follows.

**REAGENTS**

**RESINS** All the resins were stored wet before use.

- **Dowex 2 × 8, 200-400 mesh** Small particles were removed from the resin and the resin was converted to the acetate form by washing first with 3 M sodium acetate until the eluate was chloride-free and then with water until the eluate was neutral to litmus.

- **Dowex 50 × 8, 200-400 mesh** After removal of small particles the resin was converted to the sodium form by allowing it to stand overnight with 2 N sodium hydroxide. It was then washed until neutral and reconverted to the acid form by washing with 1 vol. 4 N HCl, 6 vol. 2 N HCl, 6 vol. 1 N HCl, and 6 vol. water.

- **Amberlite IRC 50, 100-200 mesh** The resin was washed in succession with 10 vol. 1 N NaOH, water, 10 vol. 1 N HCl, water, 10 vol. acetate buffer (pH 4-6), and water.

**ACETIC ACID** 1 M and 0-2 M.

**SODIUM ACETATE** 0-5 M and 3 M.

**ACETATE BUFFER (pH 4-6)** Glacial acetic acid, 57 ml., and 136 g. of sodium acetate trihydrate were made up to 1 litre with water.

**HYDROCHLORIC ACID, 2 N**

**ACETYLACETONE**

**MODIFIED EHRlich REAGENT** p-Dimethylaminobenzaldehyde, 1 g., was dissolved in 20 ml. of glacial acetic acid and 8 ml. of 72% perchloric acid added. The solution was diluted to 50 ml. with acetic acid. The reagent was prepared freshly each time.

**PROCEDURE**

Urine, 1 ml., was introduced onto a column of the prepared Dowex 2 resin 2 × 1 cm. The urine flowed through into a test tube, A, and the column was washed with 4 ml. of water, also collected in tube A. Porphobilinogen was eluted from the resin with 2 ml. of 1 M acetic acid and 2 ml. of 0-2 M acetic acid. The combined eluates were diluted to 10 ml. with water and equal volumes of this solution and the Ehrlich reagent were allowed to react together for exactly 15 minutes. The optical density of the solution was then measured in the spectrophotometer at 553 mμ against a reagent blank in cells of 1 cm. optical path. The concentration of porphobilinogen was read off from a calibration graph prepared from porphobilinogen isolated from the urine of a patient with acute intermittent porphyria by the method of Cookson and Rimington (1954).

The contents of tube A were passed through a column of Amberlite resin (1 × 7 cm.) to absorb the aminoacetone. The eluate was collected in a separate tube, B. The Amberlite column was washed with 10 ml. of water which was also collected into tube B. Aminoacetone was eluted from the column with 9 ml. of 2 N HCl and neutralized to pH 4-6 with 5 ml. 3 M sodium acetate. The volume of the solution was made up to 15 ml. with water and 2 ml. was incubated with 0-1 ml. acetylacetone for 10 minutes at 100°C. to convert the aminoacetone to 2,4-dimethyl-3-acetylpyrrole which gives a red coloured compound with the Ehrlich reagent. After cooling, equal volumes of the solution and the Ehrlich reagent were mixed and the optical density of the solution determined after exactly 15 minutes at 556 mμ. Concentrations of aminoacetone were read off from a graph prepared from aminoacetone toluene-p-sulphonate.

The eluate in tube B was passed through a column of Dowex 50 (1 × 2 cm.) and the column was washed with 16 ml. of water to free the column of urea. The column was washed with 3 ml. 0-5 M sodium acetate and deltaaminolaevulinic acid eluted with 7 ml. 0-5 M sodium acetate. Acetylacetone, 0-25 ml., was added to the eluate and the volume made up to 10 ml. with acetate buffer (pH 4-6). The solution was incubated for 10 minutes at 100°C. to convert the delta-aminolaevulinic acid to 2-methyl-3-acetyl-4-propionic acid pyrrole. After cooling, 2 ml., of the solution was mixed with an equal volume of Ehrlich reagent and the optical density of the solution determined at 553 mμ after exactly 15 minutes. Concentrations of delta-aminolaevulinic acid were read off from a calibration chart prepared from commercially obtained delta-aminolaevulinic acid · HCl (L. Light & Co.).

The pyrroles formed by the reaction with acetylacetone were identified by the chromatographic method of Urata and Granick (1963).

Urinary coproporphyrin concentrations were determined using a slight modification of the chromatographic method of Schlenker, Davis, and Kitchell (1963).

This material was obtained from Professor C. Rimington to whom I am most grateful.
RESULTS

The mean concentration of urinary amino-acetone in the control group was found to be 0.22 mg.\% (range 0.15-0.31 mg.\%). The mean delta-aminolaevulinic acid concentration in this group was 0.28 mg.\% (range, 0.06-0.51 mg.\%). There was a good correlation between concentrations of amino-acetone and delta-laevulinic acid as shown in Figure 3. The first results from the test group indicated that there was some correlation between their excretions with values for delta-aminolaevulinic acid up to about 0.6 mg.\% but above this value it seemed that the values of amino-acetone became much more widely scattered. To test this impression further, urine was obtained from a number of workers who were known to be excreting large amounts of delta-aminolaevulinic acid. Results obtained from these workers showed that although there was a tendency for amino-acetone values to rise simultaneously with delta-aminolaevulinic acid, occasionally values for amino-acetone were obtained which were lower than might have been expected (Fig. 4.)

The results of the determinations of amino-acetone for the lead-exposed group are given in Table I. Figures are shown for the whole group and also for workers with delta-aminolaevulinic acid concentrations greater than 0.6 mg.\% and less than 0.6 mg.\%. It will be seen from Table I that although the mean amino-acetone values for the workers with a level of delta-aminolaevulinic acid >0.6 mg.\% was higher than that of the control group, the value obtained from the workers with a level <0.6 mg.\% was almost identical with that of the control group. Coproporphyrin values in the lead-exposed group were higher than in the control group, as one would expect. There was some tendency for amino-acetone values to rise with increasing coproporphyrin levels, but at the higher values of coproporphyrin the scatter of those values was very wide (Fig. 5). The mean coproporphyrin concentration found in the control group was 31 \(\mu\)g./l. (range 5-80 \(\mu\)g./l.).

Little difference was observed between the mean porphobilinogen values found in the two groups. The mean result for the control group was 0.6 mg.\% (range, 0.0-12 mg.\%); for the test group the mean was 0.08 mg.\% (range, 0.0-36 mg.\%). In three workers the level was greater than 0.2 mg.\%. In one man with a porphobilinogen level of 0.22 mg.\% the amino-acetone level was 0.54 mg.\% and delta-aminolaevulinic acid 5.96 mg.\%, and in the two other cases, the porphobilinogen values were 0.28 and 0.36 mg.\% associated with amino-acetone values of 0.97 and 0.82 mg.\% and delta-aminolaevulinic acid values of 7.71 and 5.19 mg.\% respectively. Generally no correlation between porphobilinogen and amino-acetone was found.

### Table I

<table>
<thead>
<tr>
<th></th>
<th>Whole Group (51)</th>
<th>Workers with Delta-aminolaevulinic Acid &lt;0.6 mg.% (22)</th>
<th>Workers with Delta-aminolaevulinic Acid &gt;0.6 mg.% (20)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean</td>
<td>0.30</td>
<td>0.21</td>
<td>0.37</td>
</tr>
<tr>
<td>Range</td>
<td>0-0.97</td>
<td>0.15-0.30</td>
<td>Nil-0.97</td>
</tr>
</tbody>
</table>

FIG. 3. Correlation between delta-aminolaevulinic acid and amino-acetone excretion in control group.

FIG. 4. Correlation between delta-aminolaevulinic acid and amino-acetone excretion in lead-exposed group. Line parallel to abscissa = normal upper limit of amino-acetone excretion. Line parallel to ordinate = normal upper limit of delta-aminolaevulinic acid excretion.
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The correlation between amino-acetone and delta-aminolaevulinic acid excretion was not so well defined in the lead workers as in the control group. In the lead workers, the correlation was rather better for delta-aminolaevulinic acid values less than 0·6 mg.% than for the values above this figure (the upper limit of normal quoted by Haeger-Arionsen, 1960). Above the level of 0·6 mg.% the excretion of amino-acetone becomes very erratic.

Of the workers with an excretion of delta-aminolaevulinic acid >0·6 mg.%, 16 had a concentration of amino-acetone above 0·31 mg.%, the upper limit of normal found, whereas the remaining 13 workers had values of 0·31 or less. In one worker with an excretion of delta-aminolaevulinic acid of 3·63 mg.% no amino-acetone was found at all. The mean value of amino-acetone for those workers with a level of >0·6 mg.% of delta-aminolaevulinic acid, however, was noticeably higher than that of the control group and the very highest values were all accompanied by high excretion of amino-acetone.

None of the workers examined was suffering from frank clinical plumbism, but in one case of lead poisoning examined by Schlenker (1964) the level of amino-acetone was found to be slightly decreased and those of delta-aminolaevulinic acid and porphobilinogen were elevated. The finding of raised amino-acetone levels amongst some of the lead workers was unexpected in the light of enzymatic studies performed in animals by Tschudy (1964) who was of the opinion, as a result of this work, that lead would be unlikely to increase excretion of amino-acetone.

The explanation to account for the differences in excretion of amino-acetone among the workers with evidence of lead absorption is at present obscure although two theories might be put forward. It may be that in some cases glycine is preferentially metabolized through the succinate-delta-aminolaevulinic acid cycle, as postulated by de Matteis and Rimington (1962), and this would account for the low amino-acetone values found. On the other hand, lead may impose a partial blockage in the synthesis of other metabolites in the glycine-amino-acid-delta-aminolaevulinic acid cycles which could result in the high levels of both these amino-acids found. It is known that the carbon atoms bearing the amino-group in both amino-ketones are precursors of C\textsuperscript{2} and C\textsuperscript{8} of guanine and uric acid and also of formate (Nemeth, Russell, and Shemin, 1957; Kikuchi, Kumar, and Shemin, 1959), so this theory might be confirmed if concentrations of these substances were found to be diminished in persons with high amino-ketone excretion. A mechanism of this kind has been observed by Talman, Case, Nevé, Labbe, and Aldrich (1955), who found a decrease in purine synthesis which ran parallel with an increase in porphyrin in chick embryos given experimental porphyria with Sedormid. A great deal more work would need to be undertaken, however, before either of these hypotheses could be verified.

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