Conventional voltage electrophoresis for formiminoglutamic-acid determination in folic acid deficiency

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SYNOPSIS  A new method for the determination of urinary formiminoglutamic acid (FIGLU) using conventional electrophoresis at 200 to 500 v. on cellulose acetate strips is reported. Experience in 166 determinations on 137 patients shows the method to be a simple, practical, and apparently sensitive one for the determination of FIGLU in the urine.

Results of the application of the measurement of urinary FIGLU with histidine loading as a test for folic acid deficiency are also reported.

In 1951, Bakerman, Silverman, and Daft detected a precursor of glutamic acid in the urine of folic acid-deficient rats. This material was found to be formiminoglutamic acid (FIGLU) and was shown to be an intermediate product of histidine metabolism (Borek and Waelsch, 1953; Tabor, Silverman, Meehler, Daft, and Bauer, 1953; Tabor and Meehler, 1954; Seegmiller, Silverman, Tabor, and Meehler, 1954). It was also found that for the further metabolism of FIGLU to glutamic acid, folic acid in the form of tetrahydrofolic acid is required (Miller and Waelsch, 1956 and 1957; Tabor and Rabinowits, 1956). The metabolism of histidine and the relationship to folic acid is as follows:

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<table>
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<tr>
<th>Folic Acid</th>
<th>Histidine</th>
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<tr>
<td>Dihydrofolic Acid</td>
<td>Urocanic Acid</td>
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<tr>
<td>Tetrahydrofolic Acid</td>
<td>4-Imadazolone-</td>
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<td></td>
<td>5-Proprionic Acid</td>
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<tr>
<td></td>
<td>Formiminoglutamic Acid</td>
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<td></td>
<td>5-Formimino-tetrahydrofolic Acid</td>
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<td></td>
<td>Glutamic Acid</td>
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Therefore in the presence of folic acid deficiency FIGLU is not further metabolized to glutamic acid and appears in increased amounts in the urine. Increased urinary FIGLU has been demonstrated in patients with amethopterin-treated leukaemia (Broquist, 1956; Hiatt, Goldstein, and Tabor, 1958; Broquist and Luhby, 1959) and in megaloblastic anaemias due to folic acid deficiency (Broquist and Luhby, 1959; Luhby, Cooperman, and Teller, 1959a and 1959b; Spray and Witts, 1959). The use of histidine loading increases the sensitivity of the test (Luhby, et al., 1959b).

Although the measurement of urinary FIGLU was considered to be a sensitive test for folic acid deficiency, clinical application has been limited by the complexities of the methods required for the determination of FIGLU. These methods include a microbiological assay using Lactobacillus arabinosus (Silverman, Gardiner, and Bakerman, 1952; Broquist, 1956), an enzymatic method (Tabor and Wyngarden, 1958), and a combined enzymatic-microbiological technique (Silverman, Gardiner, and Condit, 1958). Paper chromatography (Silverman et al., 1952) has also been used, but was not regarded to be sufficiently sensitive. Knowles, Frankerd, and Westall (1960) introduced the use of high-voltage electrophoresis for FIGLU determination but this method requires expensive equipment and the use of 2,000-6,000 volts.

Recently we introduced the use of conventional voltage electrophoresis on cellulose acetate strips as a simple and practical method for the determination of FIGLU (Kohn, Mollin, and Rosenbach, 1961). We now wish to report the method in detail and its application as a test for folic acid deficiency.

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**METHOD**

**PRINCIPLE** Formiminoglutamic acid (FIGLU) in the urine is separated by means of small-scale cellulose acetate electrophoresis (Kohn, 1959, 1960) in a pyridine-acetic acid buffer. It is then demonstrated by staining with ninhydrin after exposure to ammonia, which converts FIGLU to glutamic acid. Two parallel aliquots of the patient’s urine (after administration of histidine) are run at the same time and under the same conditions, together with aliquots of a ‘marker’ urine containing a known quantity of FIGLU. By comparing the corresponding spots on the ammoniated strip with those on the non-ammoniated one the presence of FIGLU in the urine can be detected and its approximate concentration estimated. With a current of 200 v. the mobilities of FIGLU and glutamic acid are almost identical and their spots overlap to a great extent. The interpretation of the results is, however, quite easy due to the fact that FIGLU can only be demonstrated on the ammoniated strip.

Using higher voltages, 400-500 v., a satisfactory separation of FIGLU from glutamic acid can be achieved without overlapping. The choice of the method depends mainly on the degree of experience and whether a suitable power pack is available. For routine purposes, however, 200 v. is recommended as being simpler and more reliable.

**GLASS CAPILLARY AUTOMATIC PIPETTES** These are most useful, and can be easily prepared in the laboratory (Dacie, 1956). They should deliver 6.5 to 7 µl. It is recommended to make the capillary part from thick-walled glass so as to obtain a fair length. The pipettes are washed out with the buffer between applications.

**APPARATUS**

A horizontal tank suitable for cellulose acetate electrophoresis is required. All our separations were performed on a Shandon universal (Kohn) tank. The bridge gap, i.e., the distance between the shoulder pieces, is set at 10 cm. With a 12 x 5 cm. strip an overlap of 1 cm. is thus obtained.

**CURRENT**

Current is supplied by a constant voltage power pack capable of delivering 200 v. With the buffer recommended a current of 2-2.5 m.amp. per strip would be obtained. Due to heat generation the current tends to rise during the run.

**HISTIDINE ADMINISTRATION AND COLLECTION OF URINE SAMPLE**

L-Histidine monohydrochloride, 15 g., is given by mouth to the patient in a fasting state and food is withheld until one hour after the administration of the dose. As the histidine is only slightly soluble it is administered by mixing it with water in a glass and then the residue in the glass is washed down with water until all the histidine has been taken.

Three hours after taking the dose the patient voids urine and this is discarded. All urine passed over the next five hours (three to eight hours after histidine) is collected in a bottle to which has been added 1 ml. of concentrated hydrochloric acid and a few thymol crystals. The urine volume is measured and an aliquot taken for analysis. The collection period of between three and eight hours is used as the determination of hourly FIGLU excretion in patients has shown the peak excretion of FIGLU to occur during this time.

**TECHNIQUE OF ASSAY**

The technical electrophoretic details are essentially the same as recommended for cellulose acetate electrophoresis in general (Kohn, 1959, 1960).

1 Mark the cellulose acetate strips with a soft pencil or preferably with non-diffusible ink with the name of patient, date, and with dots indicating the application sites in the following pattern:

   - Figlu marker
   - Patient’s urine
   - Patient’s urine
   - Figlu marker

This pattern is arbitrary, of course, and any other arrangement meeting some particular requirements can be adopted.
2 Impregnate the strip with the buffer solution. It is essential to float the strip first and to submerge only after it has soaked up from underneath.

3 After impregnation, blot the strip lightly on both sides between two sheets of filter paper. No excess moisture should be seen.

4 Place the strip in position in the apparatus.

5 Apply 6-5 to 7 μl samples of patient's urine to the two previously marked centre spots and the FIGLU marker urine on the lateral spots. This is best performed by means of a glass capillary automatic pipette.

6 Close the lid and apply a constant voltage of 200 v. for 30 minutes. The timing may vary somewhat, depending upon conditions. The optimal time will soon be found by experience.

7 After the run is completed, remove the strips and dry them in an oven at 80 to 100°C. For 10 to 15 minutes until they are completely dry. An oven with a fan would be preferable.

8 Cut the strip into two identical halves lengthwise. Each cut part will now be a mirror image of the other.

9 Place one half strip into a container with ammonia (strip hung from the lid and a beaker with ammonia in the bottom of the container) for about 30 minutes.

10 After removing the strip from the ammonia atmosphere, dry it again at 90 to 100°C. For a few minutes, in order to drive off the ammonia.

11 STAINING Place the ninhydrin solution in a small evaporating basin or Petri dish and pass the strip slowly through the solution. It will start drying almost immediately and, at this stage, tends to curl slightly. Place it, therefore, flat between two sheets of filter paper lying between two pieces of cardboard; clip the cardboard pieces together with two Bulldog clips and put it back into the oven at 90 to 100°C, for about five minutes. The strips can then be inspected, but the colour does not develop fully until about 30 minutes. Inspect the strip by transmitted light using a bright lamp.

12 EVALUATION It has to be remembered that the whole procedure is essentially based on the comparison between the ammoniated and the non-ammoniated strips. The patterns obtained are shown in Figs. 1 and 2. Three spots can be seen on each strip; the spot nearest the cathode corresponds mainly to histidine, the one near the application line in the centre contains glycine, and the spot towards the anode is glutamic acid, which in most urines is usually very faint. If FIGLU is present in the urine it can be detected only on the ammonia-exposed strip as a spot corresponding to the mobility of the FIGLU marker and merging with the glutamic acid spot. A semi-quantitative estimation of the amount of FIGLU present can be obtained by comparing the intensity of the colour of the marker spot with the patient's FIGLU spot. If the FIGLU spot is very intense the patient's urine should be diluted in such a manner that the resulting colour intensity will be close to that of the marker urine. The dilutions should be made with acidified urine (preferably the patient's own urine before histidine was given or with any normal urine). Diluting with water is not recommended. An approximate estimation can also be performed by interpolation between two known standards.

With careful technique and some experience the presence of as little as 20 μg of FIGLU per ml of urine can be detected. Pure FIGLU can be detected in even smaller concentrations.

The amount of FIGLU in the urine is then expressed in milligrams per hour over the five-hour collection period (between the third and the eighth hour after the administration of 15 g. histidine).

SEPARATION AT 400 V.

This is an alternative procedure which permits a clearer separation of the FIGLU and the glutamic acid spots (Fig. 3). The separation can be achieved by the application of a higher voltage, i.e., 400 up to 500 v. The time of running is 20 minutes. Under these conditions the basic amino-acids (histidine group) run off the strip into the cathode buffer. The sample should be applied as an
elliptical drop, moving the tip of the pipette slightly at right angles to the direction of the run. The current should not exceed 4-5 m.amp. if overheating is to be avoided. A rapid fall in the amperage indicates overheating and in this case semi-transparent areas can be observed. In all other respects the technique is exactly the same as with 200 v.

It should be pointed out, however, that the conditions of this method are somewhat more critical. The technique requires more experience and a power pack capable of delivering up to 500 v. The danger of overheating the strip is greater, particularly with a higher ambient temperature. Sometimes a syphoning effect, probably due to the excessive evaporation in the centre of the strip, may cause small areas or bands of discoloration after ninhydrin staining. With some experience, however, there is no difficulty in the interpretation of the results.

INVESTIGATIONS AND DIAGNOSTIC CRITERIA

All of the FIGLU determinations were carried out by one of us to whom the urine samples were sent without any clinical information on the patients. Therefore, any possible element of bias in the determination was eliminated.

Each of the patients tested had a complete blood count, and a serum vitamin B₁₂ level was measured by microbiological assay with Euglena gracilis (Hutner, Bach, and Ross, 1956) and a serum folic acid level with the modified L. casei method (Waters and Mollin, 1961). A bone marrow biopsy was performed on all of the anaemic patients, and when indicated vitamin B₁₂ and folic acid absorption studies were done. In most of the patients with a megaloblastic anaemia, the haematological response to 0-4 mg. folic acid or 2 μg. vitamin B₁₂ a day was followed.

Patients diagnosed as idiopathic steatorrhoea had evidence of malabsorption with a typically abnormal jejunal biopsy. Patients were considered to have nutritional folic acid deficiency if there was a megaloblastic anaemia with a normal serum B₁₂ level, a poor dietary history, a normal jejunal biopsy, no evidence of malabsorption of B₁₂ or folic acid, and if they had not been taking anticonvulsant drugs. Pernicious anaemia was diagnosed by the usual criteria and confirmed with radioactive B₁₂ absorption tests. Patients designated as having a post-gastrectomy megaloblastic anaemia due to B₁₂ deficiency had a partial or total gastrectomy, a megaloblastic bone marrow, a low serum B₁₂ level, and radioactive B₁₂ absorption studies similar to those of pernicious anaemia.

RESULTS

One hundred and sixty-six tests have now been carried out on 137 patients with various disease states. In Table I are summarized the results on the normal controls and patients with idiopathic steatorrhoea, nutritional folic acid deficiency, vitamin B₁₂ deficiency, iron deficiency, and a few miscellaneous conditions.

Most normals showed no detectable FIGLU in the urine by this test. In an occasional normal subject a trace of FIGLU may be present but the calculated excretion has never been greater than 2 mg./hour.

In all patients with clinical evidence of folic acid deficiency the excretion has been greater than 3-5 mg./hour.

The highest levels of FIGLU with a range of 30 to 80 mg./hour were most consistently obtained in the untreated cases of idiopathic steatorrhoea, all of which showed haematological evidence of folic acid deficiency. Three patients who were well controlled on gluten-free diets for periods of one to two years showed a lower but still elevated urinary FIGLU excretion with a range of 4 to 8 mg./hour.

Several patients with idiopathic steatorrhoea were treated with intramuscular injections of 0-4 mg. of folic acid a day, and in spite of an excellent haematological response serial FIGLU determinations showed only a gradual return towards normal. In one patient, for example, the FIGLU excretion after 30 days of treatment with 0-4 mg. of folic acid a day had decreased from 44 mg./hour to 15 mg./hour. A large dose of folic acid, 10 mg. per day, was then instituted, and the FIGLU excretion became normal within another seven days.
The six cases of nutritional folic acid deficiency also had elevated urinary FIGLU levels but of lower magnitude, ranging from 4 to 20 μg./hour. In all of these patients with idiopathic steatorrhoea and nutritional folic acid deficiency the serum folic acid level was low.

Of the 14 cases of vitamin B₁₂ deficiency (seven with pernicious anaemia, six with post-gastrectomy megaloblastic anaemia, one with a terminal ileum resection), 10 showed no increase in urinary FIGLU. However, four patients, three with pernicious anaemia and serum B₁₂ levels between 40 and 65 μg./ml. and the other with a post-gastrectomy megaloblastic anaemia and a serum B₁₂ level of 45 μg./ml., had increased FIGLU in the urine. In all 14 of these cases with vitamin B₁₂ deficiency the serum folic acid level was normal. One of the patients with untreated pernicious anaemia was given 0·4 mg. of folic acid daily for 14 days without a significant haematological response.

The urinary FIGLU was not increased in eight patients with uncomplicated iron-deficiency anaemia.

In addition, increased urinary FIGLU was detected in three patients with acute leukaemia on amethopterin therapy, one case of megaloblastic anaemia related to anticonvulsant therapy, and in one patient with a megaloblastic anaemia in association with clinical scurvy. Increased FIGLU in the urine has also been present in a number of cases of widely disseminated carcinoma, haemolytic anaemia, and active liver disease. Six of seven patients with sarcoidosis also showed raised urinary FIGLU levels.

These results will be reported and discussed in greater detail elsewhere.

**DISCUSSION**

This method of FIGLU determination using conventional voltage electrophoresis on cellulose acetate strips is simple and practical. Our experience suggests that it is sufficiently sensitive for clinical and diagnostic purposes.

In regard to the application of the urinary excretion of FIGLU to the diagnosis of folic acid deficiency, experimental work has shown that animals made folic acid-deficient excrete increased amounts of FIGLU in the urine (Bakerman et al., 1951; Silverman et al., 1952; Tabor et al., 1953). The majority of workers have reported a good correlation of urinary FIGLU excretion with clinical folic acid deficiency (Broquist and Luhby, 1959; Luhby et al., 1959a and 1959b; Spray and Witts, 1959; Knowles et al., 1960). However, others have claimed that the test is not uniformly reliable. These reports, not as yet published in detail, indicate that some patients with pernicious anaemia, responding completely to vitamin B₁₂ alone, have significant amounts of FIGLU in their urine and some patients with folic acid deficiency do not have FIGLU in their urine (Rucknagel, La Du, Laster, Seegmiller, Daft, Silverman, and Pitney; Marshall, Jandl, Castle, and Hiatt; Herbert, Baker, Frank, Pasher, Sobotka, and Wasserman; as cited in Herbert, Baker, Frank, Pasher, Sobotka and Wasserman, 1960).

In our experience so far, all patients with clinical and microbiological evidence of folic acid deficiency, have had an increased urinary excretion of FIGLU (Table 1). However, four of 14 patients with vitamin B₉ deficiency have also shown an increased urinary FIGLU excretion. The explanation for this finding would be that an elevated FIGLU...
is uncertain but it may possibly represent concomitant or secondary folic acid deficiency (Chanarin, Mollin, and Anderson, 1958). Nevertheless, the serum folic acid level was normal in all four of these cases as well as in the other 10 cases of vitamin B₁₂ deficiency (Table I). In all four cases the excessive excretion of FIGLU disappeared when the patients were treated with vitamin B₁₂. It is therefore possible that the excessive excretion of FIGLU by these patients was due to a defect in folic acid metabolism caused by the deficiency of vitamin B₁₂.

A number of patients with disseminated carcinoma, active liver disease, and sarcoidosis, many of whom showed no megaloblastic change in the marrow, had an elevated urinary excretion of FIGLU. Whether this represents tissue depletion of folic acid or whether it is due to deficiency of FIGLU transferase or folic acid-reducing enzymes, as suggested in alcoholic cirrhosis of the liver (Carter, Schaffner, and Heller, 1960), is not known. In this connexion the persistence of the increased FIGLU excretion in patients with folic acid deficiency while on treatment with 0-4 mg. of folic acid daily, in spite of an excellent haematological response, suggests that the test may be very sensitive to depletion of folic acid in the tissues. However, until more is known about the specificity and sensitivity of FIGLU excretion, the meaning of these findings is uncertain.

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REFERENCES