Enzymatic release of folate activity from the red cells in megaloblastic anaemia of pregnancy

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SYNOPSIS The content of folate activity precursors in washed red cells and the enzymatic plasma factor activity, necessary for the release of folate from the precursors, were studied in normal subjects and in patients with megaloblastic anaemia of pregnancy. Subjects with megaloblastic anaemia of pregnancy had a significantly reduced folate activity precursor content, and 14 subjects (58%) had significantly low plasma factor activity, which in at least four subjects may have been due to the presence of inhibitors. This study indicates that impaired activity of the plasma factor may play a part in the aetiology of megaloblastic anaemia of pregnancy.

In recent years the method using *Lactobacillus casei* for determining serum folate activity has been widely used for assessing folic acid status in various conditions in man (Baker, Herbert, Frank, Pasher, Hutner, Wasserman, and Sobotka, 1959; Herbert, Baker, Frank, Pasher, Sobotka, and Wasserman, 1960; Waters and Mollin, 1961; Cooper and Lowenstein, 1961; Ball and Giles, 1964). Izak, Rachmilewitz, Sadovsky, Bercovici, Aronovitch, and Grossowicz (1961) and Grossowicz, Mandelbaum-Shavit, Davidoff, and Aronovitch (1962), measuring folate in both whole blood and serum, reported that serum contains only about 10% of the whole blood activity and that the bulk of the folate is located in the red cells.

Previously, Toennies, Usdin, and Phillips (1956) and Toennies and Phillips (1959) showed that the red cells contain large amounts of folate activity precursors, and that plasma and serum contain equal amounts of enzymatic factors (plasma factors) capable of releasing folate activity from folate activity precursors.

Since megaloblastic anaemia of pregnancy is characterized by low levels of folic acid in serum (Herbert *et al.*, 1960; Waters and Mollin, 1961; Ball and Giles, 1964) and in whole blood (Izak *et al.*, 1961; Grossowicz *et al.*, 1962), it was felt advisable to investigate a possible deficiency of either plasma factor or folate activity precursors in these subjects compared with normal controls. Accordingly, plasma factor was measured in serum and folate activity precursor in erythrocytes.

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MATERIALS AND METHODS

We studied 22 healthy control subjects, men and women who were not pregnant at the time; 25 pregnant women with low haemoglobin concentrations (less than 10 g./100 ml.) and megaloblasts visible in marrow preparations; eight pregnant women with low haemoglobin concentration (less than 10 g./100 ml.) and a normoblastic marrow picture; and 24 normal pregnant subjects with a haemoglobin level above 12 g./100 ml.

Venous blood (20 ml.) was collected from the subjects, preferably in a fasting state or at least two hours after a light meal. After clotting, serum was separated and stored at −20°C. without the addition of ascorbic acid. It was usually tested within two weeks.

The remaining blood clot was gently shaken with 6 ml. acid citrate dextrose, and the suspended erythrocytes were decanted and centrifuged at 1,400 g for 20 min. in graduated tubes. The supernatant was removed and the packed cells were resuspended in 3 volumes of acid citrate dextrose. The suspension was either used immediately or stored at 4°C. for not more than one week.

STANDARDS A standard preparation of folate activity precursors was made according to the method of Toennies *et al.* (1956). The partially purified powder was stored in a desiccator at room temperature. Fresh batches were prepared as required and compared with the existing batch before use.

For the standard plasma factor, serum obtained at intervals from the same normal subject was used. The enzymatic activity of this serum was frequently checked with a standard preparation of folate activity precursor and remained apparently constant.

ASSAY FOR FOLATE ACTIVITY Folate activity was determined with *L. casei* using a modification of the methods...
of Waters and Mollin (1961) and Herbert (1961). Dehydrated Difco *L. casei* medium was used. The buffers employed were 0·1 M phosphate buffer of pH 6·1, containing 200 mg.％ ascorbic acid (0·1 M phosphate ascorbate buffer), and 0·05 M phosphate buffer of pH 6·1, containing 100 mg.％ ascorbic acid (0·05 M phosphate ascorbate buffer). Serum was diluted in 20 with 0·1 M phosphate ascorbate buffer before assay.

**Folate Activity Precursor Content of the Red Cells**

The content of folate activity precursors was expressed as the amount of folate activity released from the preparation of washed red cells by an amount of standard plasma factor known to release all the folate activity from the amount of folate activity precursor likely to be present under the standard conditions.

The erythrocyte suspension was recentrifuged for 20 min. at 1,400 g and a fresh suspension (1 in 4) in acid citrate dextrose was made. It was haemolysed by diluting it 1 in 10 with 0·05 M phosphate ascorbate buffer, and then recentrifuged. To 0·8 ml. of the clear red haemolysate (corresponding to 0·02 ml. of the red cells), 19·0 ml. of 0·05 M phosphate ascorbate buffer was added, followed by 0·2 ml. of the standard plasma factor preparation. A plasma factor blank was prepared by adding 2·2 ml. of standard plasma to 19·8 ml. of 0·05 M phosphate ascorbate buffer, and a folate activity precursor blank by adding 0·8 ml. of red cell haemolysate to 19·0 ml. of 0·05 M phosphate ascorbate buffer and 0·2 ml. of water. The three samples were incubated at 37°C for 90 min. before assay.

The folate activity precursor content of the red cells was calculated by subtracting the folate activity of the two blanks from the test and multiplying by 50, and was expressed in ng./ml. of erythrocytes.

**Plasma Factor Activity of Serum**

Plasma factor activity was expressed as the amount of folate activity released by 0·2 ml. of serum under test from an amount of standard preparation of folate activity precursor known to saturate at least 0·5 ml. of the standard serum.

The serum under test (0·2 ml.) was added to 3 mg. standard folate activity precursor in 19·8 ml. of 0·05 M phosphate ascorbate buffer. Serum and folate activity precursor blanks were also prepared. The three samples were incubated at 37°C for 90 min. before assay.

The plasma factor activity was calculated by subtracting folate activity levels of the two blanks from the test and multiplying by 5, and was expressed in ng./ml. serum.

In experiments discussed later, 6 mg. of the standard folate activity precursor was used with 0·2 to 1·0 ml. of the test serum, made up to a final volume of 20 ml. with 0·05 M phosphate ascorbate buffer. Further incubation and assay was as before.

**Effect of Serum Dialysates on Standard Folate Activity Release from Folate Activity Precursors**

Sera (5 ml.) from different subjects were dialysed overnight at room temperature, using Visking tubing, against 50 ml. of distilled water. Samples, each of 9·5 ml. of the dialysates, were included in the incubation mixture for standard folate activity release from folate activity precursor. The composition of the final mixture was otherwise unchanged. Incubation and final assay were as usual. The activity corrected for blanks was expressed as the percentage difference from the activity in the absence of dialysate.

**Results**

**Folate Activity Precursor Content of the Red Cells**

Table I shows the serum folate activity and the folate activity precursor content of the red cells in control subjects, in women with megaloblastic anaemia of pregnancy, and in subjects with normoblastic anaemia of pregnancy.

The content of folate activity precursor in erythrocytes is significantly lower in megaloblastic anaemia of pregnancy compared with the controls (P < 0·001) or pregnant women with normoblastic anaemia (P = 0·001). The latter two groups differ significantly (P = 0·02). Only three subjects with megaloblastic anaemia of pregnancy had values greater than 325 ng./ml., which was the lowest value in control subjects.

**Plasma Factor Activity of Serum**

Plasma factor was determined in the three conditions previously investigated. A second control group of normal pregnant women has now been included. Individual results appear in Fig. 1 and grouped results in Table II.

Subjects with megaloblastic anaemia of pregnancy have significantly lower plasma factor activity than control or normal pregnant subjects (P < 0·001) or pregnant women with normoblastic anaemia (P = 0·001). Normal pregnant women have significantly higher plasma factor activity than control subjects (P < 0·001) or women with normoblastic anaemia (P = 0·001). In the control group there is no

**Table I**

<table>
<thead>
<tr>
<th>Condition</th>
<th>No. of Cases</th>
<th>Serum Folate (ng./ml.)</th>
<th>Folate Activity Precursor Content (ng./ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Range</td>
<td>Mean</td>
<td>S.D.</td>
</tr>
<tr>
<td>Control</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Megaloblastic anaemia of pregnancy</td>
<td>22</td>
<td>6·7-14·6</td>
<td>9·20</td>
</tr>
<tr>
<td>Normoblastic anaemia of pregnancy</td>
<td>18</td>
<td>0·7-3·8</td>
<td>2·03</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>4·8-10·0</td>
<td>6·35</td>
</tr>
</tbody>
</table>
difference in plasma factor between male (8) and female (14) subjects (P < 0.1). Women with normoblastic anaemia do not differ from control subjects (P > 0.1).

It can be seen in Fig. 1 that subjects with megaloblastic anaemia do not form a homogenous group. They can be further divided into 10 subjects with plasma factor activity (range 17-0-21-5 ng./ml.; mean 19-4; S.D. 1-35) not significantly different from controls or patients with normoblastic anaemia of pregnancy but significantly lower (P < 0.001) than normal pregnant women, and 14 subjects with plasma factor activity (range 6-0-15-0 ng./ml.; mean 11-1; S.D. 2-29) significantly lower (P < 0.001) than all other groups.

CORRELATION BETWEEN SERUM FOLATE ACTIVITY, PLASMA FACTOR, AND FOLATE ACTIVITY PRECURSOR

In 22 control subjects not pregnant, significant correlation was found between serum folate activity and folate activity precursor (r = 0.50; 0.02 > P > 0.01) and plasma factor and folate activity precursor (r = 0.46; 0.05 > P > 0.02) but no correlation between serum folate activity and plasma factor (r = 0.09; P > 0.1).

In 24 normal pregnant subjects no significant correlation was found between serum folate activity and plasma factor (r = 0.36; P > 0.05). No figures were available for folate activity precursor to calculate the correlation between folate activity and its precursor.

In eight pregnant subjects with normoblastic anaemia no significant correlation was established between folate activity and its precursor (r = 0.07; P > 0.1), or between folate activity and plasma factor (r = 0.15; P > 0.1). There are, however, fewer subjects in this group.

In 17 subjects with megaloblastic anaemia of pregnancy there was a very significant correlation between serum folate activity and plasma factor (r = 0.85; P < 0.001), and a less significant one between folate activity and its precursor (r = 0.51; 0.05 > P > 0.02). There was no correlation between plasma factor and folate activity precursor (r = 0.13; P > 0.01).

In non-pregnant subjects serum folate activity depends on the amount of its precursor. In megaloblastic anaemia of pregnancy serum folate activity seems to depend both on its precursor and on plasma factor. Thus low values for plasma factor found in 14 subjects (58%) with megaloblastic anaemia of pregnancy may be partially responsible for low serum folate activity and cumulatively for megaloblastic erythropoiesis.

EFFECT OF INCREASING SERUM PLASMA FACTOR ON FOLATE ACTIVITY RELEASE FROM FOLATE ACTIVITY PRECURSORS

Figure 2 shows the effect of increasing the concentration of plasma factor in the incubation mixture on folate activity release from a constant amount of folate activity precursors.

Curve A represents the mean response for five normal subjects. Curves B, C, D, and E are examples of the effect obtained with sera from subjects with megaloblastic anaemia of pregnancy. In all four curves the amount of folate activity release is less than the normal.

Although curves B and E suggest deficiency of enzymes, curves C and D are more difficult to

<table>
<thead>
<tr>
<th>Table II</th>
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<table>
<thead>
<tr>
<th align="right">Condition</th>
<th align="right">No. of Cases</th>
<th align="right">Serum Folate (ng./ml.)</th>
<th align="right">Plasma Factor Activity (ng./ml.)</th>
</tr>
</thead>
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<td align="right">---</td>
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</tr>
<tr>
<td align="right">Control</td>
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<td align="right">6-7-14-6</td>
<td align="right">16-0-23-5</td>
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<tr>
<td align="right">Megaloblastic anaemia of pregnancy</td>
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<td align="right">0-6-3-8</td>
<td align="right">6-0-21-5</td>
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<tr>
<td align="right">Normal pregnant</td>
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<td align="right">5-2-11-1</td>
<td align="right">19-0-28-0</td>
</tr>
<tr>
<td align="right">Normoblastic anaemia of pregnancy</td>
<td align="right">8</td>
<td align="right">4-8-10-0</td>
<td align="right">16-0-25-0</td>
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</tbody>
</table>
Interpret. The shape may result from the presence of enzyme inhibitors in the serum, but as the system is so complex, and the number of factors in plasma factor uncertain, further speculation is unprofitable.

Altogether sera from nine subjects with megaloblastic anaemia of pregnancy were tested. Five showed responses similar to curves B and E, and four were like curves C and D.

**EFFECT OF SERUM DIALYSATES ON STANDARD FOLATE ACTIVITY RELEASE FROM PRECURSORS**

Dialysates from the sera of three control subjects had a negligible effect on the standard reaction. The percentage differences were -1·1, +1·2, and +7·0.

Serum dialysates from seven subjects with megaloblastic anaemia of pregnancy gave a wider range of percentage differences, viz., -22·1, -16·2, -13·3, -11·1, -2·4, +2·2, and +8·2.

Thus in four subjects with megaloblastic anaemia of pregnancy, serum dialysates depressed the standard reaction by more than 10%, suggesting the presence of inhibitors.

**DISCUSSION**

Toennies et al. (1956), Toennies and Phillips (1959), and Noronha and Aboobaker (1963) showed that the bulk of folate activity resides in the red cells as folate activity precursors. The former workers have further indicated that human plasma contains a factor capable of releasing folate activity from its precursors. Thus serum folate activity, shown by Herbert, Larrabee, and Buchanan (1962) to be mainly 5-methyl-tetrahydrofolic acid (5-CHO-THFA), may be derived from the action of plasma factor on folate activity precursors released by the red cells when they undergo destruction at the end of their life span. In addition serum may contain folate factors directly derived from the diet.

That both dietary folic acid and folate derived from the red cells play a part in folate metabolism in man is suggested by the results of Herbert (1962). In this study, a healthy physician developed folate deficiency megaloblastic anaemia after four and a half months on a diet deficient in folic acid; at that time his erythrocyte 'folate' stores were exhausted.

A possible scheme for folic acid turnover in blood, shown in Fig. 3, indicates that folate deficiency in anaemic subjects may be due to dietary lack of folic acid, abnormal release of folate activity from its precursors in the erythrocyte, or disturbed conversion and utilization of folate compounds.

The red cell appears to be the main storage site for folate activity precursors in blood. In anaemic subjects the red cell mass, and thus folate activity precursor, is reduced. The anaemia may be caused in the first instance by any erythropoietic disturbance or by haemorrhage as well as by folic acid depletion alone. If the anaemia is not corrected or supplemented with folic acid, the diminishing circulating red cell mass will release less folate, further depressing red cell formation. The cumulative effect may lead to depletion of folate stores and eventually to a megaloblastic marrow.

In subjects with megaloblastic anaemia of pregnancy, low folate levels are well recognized but the cause of this reduction is problematic. Since there is significant correlation between serum folate activity and its precursor in the erythrocyte, both in control subjects and in women with megaloblastic anaemia of pregnancy, low serum folate activity values may indicate a decreased total folate activity precursor content due to reduced red cell volume or to diminished deposits of precursor within the red cells. The reduced concentration of precursor found in the erythrocytes of subjects with megaloblastic anaemia of pregnancy (Table I) suggests that folate deprivation has existed for some time. Folate enters only young erythrocytes (Herbert and Zalusky, 1962) and it takes about 120 days for a normal red cell to be completely replaced by a folate-deficient cell (Herbert, 1964).

The initial cause of folate depletion in pregnant subjects is uncertain. Dietary deficiency, excessive demands of the foetus, and impaired absorption of folate have been implicated. The foetus obtains all
its folic acid from the mother and its demands may precipitate megaloblastic anaemia if the mother already suffers from folic acid deficiency, either from malnutrition or malabsorption. In the cases of Izak et al. (1961), foetal demand seemed an important factor. In apparently normal pregnant subjects, lower serum folic acid (Ball and Giles, 1964), and more rapid clearance of folic acid from the serum (Chanarin, Anderson, and Mollin, 1958a; Chanarin, Mollin, and Anderson, 1958b; Chanarin, Mac-Gibbon, O'Sullivan, and Mollin, 1959) suggest that in pregnancy turnover of folic acid is increased. Low vitamin B12 levels found in some subjects with megaloblastic anaemia of pregnancy (Baker et al., 1959; Cox, Matthews, Meynell, Cooke, and Gaddie, 1960; Izak, et al., 1961; Ball and Giles, 1964) may also contribute to folic acid depletion, since vitamin B12 is involved in folic acid conversion and utilization (Herbert and Zalusky, 1962; Larrabee, Rosenthal, Cathou, and Buchanan, 1963; Waters and Mollin, 1963; Cooper and Lowenstein, 1964).

Our experiments with plasma factor activity indicate that yet another factor may be responsible for folic acid deficiency in megaloblastic anaemia of pregnancy. Although no correlation could be established between serum folic acid and plasma factor in non-pregnant or normal pregnant controls, in subjects with megaloblastic anaemia of pregnancy the correlation is highly significant.

Of 24 subjects with megaloblastic anaemia of pregnancy, 14 had significantly reduced plasma factor activity compared with non-pregnant subjects, and all had diminished activity compared with normal pregnant controls. Subjects with depressed plasma folic acid activity may be less able to utilize folic acid precursor released from the red cells, thus reducing the amount of the 'available' folic acid in circulation.

It is difficult to establish if a low level of plasma factor in subjects with megaloblastic anaemia of pregnancy is due to the diminished production of enzymes, deficiency of enzyme activators, or the presence of inhibitors. The possibility exists that plasma factor may be produced according to demand, and a low level may merely reflect a diminished content of folic acid precursor. Thus in normal pregnancy plasma factor may rise to cope with an increased total folic acid precursor, due to the increment in total red cell mass.

However, curves C and D in Fig. 1, and the depressive effect of serum dialysates from four out of seven subjects studied, suggest that in some subjects with megaloblastic anaemia of pregnancy low plasma factor activity may be due to the presence of inhibitors.

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