Microbiological assay on microtitre plates of folate in serum and red cells

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Abstract

**Aims:** To develop a simple microbiological assay for serum and red cell folates on 96-well microtitre plates, suitable for use in routine clinical diagnosis.

**Methods:** Use of a chloramphenicol resistant organism (NCIB 10463) saved time by avoiding aseptic precautions. Use of plate sealers facilitated mixing. Evaluation of assay performance included estimations of folate recovery, assay reproducibility, and response to reduced folate. Results obtained on sera (193) and red cell folates (150) were compared with those obtained using a traditional microbiological assay.

**Results:** Good recovery of folic acid added to serum and also good interassay and intra-assay precision were obtained with both serum (CV% of <5) and red cell folate pools (CV% of <5).

Equimolar assay responses were obtained with folic acid, 5-formyltetrahydrofolate (L-form), and 5-methyltetrahydrofolate (L-form). The microassay correlated well with a traditional assay for estimation of folate in both serum (n = 193, r = 0.975) and red cells (n = 150, r = 0.96).

**Conclusions:** This assay is more compact and less time consuming than the traditional assay. It is extremely economical and is easy to perform in a routine clinical laboratory.

Estimations of blood folate, whether in serum or red cells, remain crucial in the differential diagnosis of nutritional anaemia. Although most clinical laboratories assay folates using commercially available techniques incorporating competitive protein binding, a few prefer to use the microbiological assay with Lactobacillus casei, which is the reference method. In 1987 improvements in microtitre plate technology facilitated the development of a microbiological assay for food folates on these plates.1 Subsequently Horne and Patterson1 reported a similar plate assay which was simpler, with cryopreservation of the organism (ATCC 7469) and a shortened incubation time. This assay could be used to estimate serum folates. We now report a microbiological assay for folates on microtitre plates using a chloramphenicol resistant strain of L casei (NCIB 10463). The assay can be used to determine concentrations of both serum and red cell folate of patient samples. It is technically simpler than previously published techniques.

**Methods**

*Lactobacillus casei* (NCIB 10463) was obtained from Torrey Research Station (Aberdeen, Scotland). Vitamin folic acid assay broth was obtained from Merck (Darmstadt, Germany). Flat-bottomed microtitre plates (0·396 ml capacity) were from A/S Nunc (Roskilde, Denmark) and mylar plate sealers were from Dynatech Laboratories (Chantilly, Virginia 22021, USA). Chloramphenicol was from Parke Davis & Co. (Pontypool, Gwent, Wales) and β-lactamase (EC 3·5·2·6) ex Bacillus cereus was supplied by Koch Light Ltd (Suffolk, England). Microtubes (2 ml capacity) for storing cryopreserved organisms were obtained from Sarstedt Ltd (Wexford, Ireland). Ascorbic acid, sodium ascorbate, folic acid [PteGlu], 5-5-methyltetrahydrofolate (sodium salt), and folic acid (calcium salt) were from Sigma (London). Folate standard concentrations were verified spectrophotometrically7 using a Beckman model 35 UV/Vis spectrophotometer. Culture tubes (13 × 100 mm) were from Corning (UK). Stericol disinfectant was obtained from Sterling Medicare (Guildford, Surrey, England). Additions of dilute sera, whole blood haemolysates, and folates were made with a Stepper semiautomated pipette (Socorex, Renens, Switzerland). Inoculated assay medium was dispensed using an eight-channel Titertek dispenser and plates were read on a Titertek Multiscan Plus Mark 2 plate reader, both from Flow Laboratories (Ayrshire, Scotland). All other chemicals were of reagent grade and distilled water was used throughout.

**Preparation of Samples**

Lysates of red cell folates were prepared by dilution of whole blood anticoagulated with EDTA 1 in 10 with freshly prepared 1% ascorbic acid,4 mixing thoroughly, and incubating at 37°C for 30 minutes. These lysates could then be stored at −20°C until assay. The standard precautions against infection by high risk samples were taken with all samples, both in sample preparation and throughout the assay.

**Preparation of Assay Organism**

*L. casei* (NCIB 10463) was cryopreserved by
the methods of Grossowitz as described for *L casei* (ATCC 7469), and modified by Wilson and Horne. This cryopreserved culture was stored at −70°C in 1 ml aliquots. Reconstituted assay medium was seeded by thawing a phial of cryopreserved culture rapidly (in a water bath at 37°C), adding to the bulk assay medium at the concentration of 200 µl/100 ml, and mixing very thoroughly using a magnetic stirrer.

**MIGHTTITRE PLATE ASSAY PROCEDURE**
A fresh solution of sodium ascorbate (0–5%) was made and used to dilute serum samples 1 in 20 (100 µl up to 2 ml) and red cell lysates (as described above) were diluted 1 in 40 (50 µl up to 2 ml). Dilute sera were mixed and added to four plate wells (100 µl to each of two wells and 50 µl to each of the other two). A compensating volume of 50 µl of fresh 0–5% sodium ascobrate was added to make the volume in all wells up to 100 µl. Dilute red cell lysates were mixed and added to six wells (100 µl to each of three wells and 50 µl to each of the remaining three). The additional well at each dilution was to act as a subtractable blank value to compensate for the colour contribution of lyse haemoglobin. Compensating volumes of 0–5% sodium ascobrate were added to make all wells up to 100 µl volume.

**ASSAY STANDARDS**
A working standard solution of folic acid (500 pg/ml) was made by dilution of a stock standard in 0–5% sodium ascobrate. Additions of this standard were made to the wells to give a range of concentrations from 0–50 pg/well (0–100 µl of standard). All well volumes were made up to 100 µl with 0–5% sodium ascobrate.

**ASSAY MEDIUM**
Medium was reconstituted in distilled water by adding 5–7 g vitamin folic acid assay broth, 3 mg choromphenicol, and 30 µl Tween 80 per 100 ml medium. After mixing, heating to boiling, and cooling, ascobic acid (75 mg/100 ml) was added. Finally, the assay organism was added as described above. Medium was stirred continuously while being dispensed into the microtitre plate wells at the rate of 200 µl/well. An addition of 10 µl of dilute Stericil disinfectant (1 in 25 dilution) was made to several zero tubes of the standard curve to act as assay blanks. Similar additions of disinfectant were made to the blank wells, one at each dilution. This addition of disinfectant had the effect of preventing organism growth without contributing to the optical density of the blanks. It thus avoided the need for dispensing unseeded medium into these blanks with the hazard of cross-contamination with organism from adjacent wells. All plates were sealed firmly with plate sealers, mixed by inversion, and incubated in a warm air cupboard (not humidified) at 37°C in the dark for 42 hours. They were then mixed thoroughly by inversion, the plate sealers removed and read at 570 nm. It is important to ensure that there are no bubbles over the wells while reading, as these can alter the reading and, if they burst, can soil the optics of the reader.

**CALCULATION OF RESULTS**
The results lend themselves readily to data reduction. The appropriate dilution factors were used to calculate serum folate (1 in 200 and 1 in 400) and whole blood folate (1 in 400 and 1 in 800) concentrations. Red cell folates were calculated from the values for whole blood folate, serum folate, and from the packed cell volume (PCV). The results were expressed as folate concentration per ml of packed red cells.

**Results**

**EVALUATION OF ASSAY RESPONSE TO REDUCED FOLATE MONOGLUTAMATES**
The response of the assay organism to the reduced folate monoglutamates folic acid (5-formyl tetrahydrofolate) and 5-methyltetrahydrofolate were compared with that of folic acid (PteGlu). Reduced folate standards were diluted in sodium ascobrate (500 mg/100 ml) to equimolar concentrations of biologically active l-isomer. Comparison of the growth responses of these folates with that of folic acid under the conditions of the assay gave similar results (fig 1).

**ASSAY PERFORMANCE**
The reproducibility of the plate assay was monitored using control sera and red cell lysates (1 in 10, whole blood:ascobrate). These were stored at −20°C. Reproducibility of these pools within and between assays were estimated. The results show good reproducibility and are expressed in tables 1 and 2. When folic acid was added to serum at five different concentrations (*n* = 10 in each case), its recovery ranged from 94% to 102% of the expected value, with a mean of 98.4%.

**COMPARISON OF RESULTS USING A TRADITIONAL ASSAY**
The results of assays for serum and red cell folate with the methods of Grossowitz and modified by Wilson and Horne were compared with those from the microtitre plate method. Overall these results demonstrated that the methods were directly comparable with the exception that the microtitre plate assay results were consistently higher than those from the traditional assay. The reasons for this are not clear.
Table 1 Between assay reproducibility of control values for folate in serum and whole blood (n = 10 in each case)

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<thead>
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<th>Serum folates (µg/l)</th>
<th>Whole blood folates (µg/l)</th>
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<tbody>
<tr>
<td>Mean</td>
<td>CV%</td>
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<tr>
<td>1.6</td>
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<td>4.5</td>
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Table 2 Intra-assay reproducibility of control values for folate in serum and red cells (n = 12 in each case)

<table>
<thead>
<tr>
<th>Serum folates (µg/l)</th>
<th>Whole blood folates (µg/l)</th>
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<tbody>
<tr>
<td>Mean</td>
<td>CV%</td>
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<tr>
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Folates by a microtitre plate assay were compared with those obtained using a traditional tube assay. The tube assay was completed in soda glass culture tubes (13 × 100 mm) using the same cryopreserved organism. The results showed good correlation for both serum folates (n = 193, r = 0.975) and red cell folates (n = 150, r = 0.96) (fig 2). The similarity of results obtained between these assays meant that no adjustment of the normal ranges of blood folate values was necessary, and in this laboratory we consider serum folate values of <2.4 µg/l and red cell folate values of <100 µg/l of packed cells to be deficient. Each laboratory, however, should calculate its own ranges of values.

EFFECT OF ANTIBIOTICS ON THE ASSAY

Antibiotics can interfere with microbiological assays. Some 77 (2%) of 3860 sequential diagnostic serum folate assays showed evidence of such interference and seven of these were due to methotrexate. These cases were identified by either the complete inhibition of organism growth or a partial inhibition which created a discrepancy in the results of two dilutions. No interference was noted with red cell folate assays over this period, probably due to the pre-assay dilution factor for red cell folates being some 20-fold higher than that for serum. We have since noted occasional inhibition due to methotrexate. Serum preincubation with a β-lactamase preparation had been shown to eliminate interference caused by penicillins, augmentin, and third generation cephalosporins in a vitamin B12 microbiological assay using L leichmanii. The efficiency of β-lactamase in eliminating antibiotic interference in this assay was evaluated in a pilot study by determining the recovery of a spike of folic acid after lactamase treatment. Sera suspected of containing interfering antibodies were diluted in three separate tubes for repeat assay. Tube A contained serum (50 µl), tube B contained serum (50 µl) with β-lactamase stock solution (20 µl), and tube C was similar to tube B with an added spike of folic acid. Tubes were mixed, incubated (15 minutes at 37°C), diluted to 1 ml with sodium ascorbate (500 mg/100 ml) and assayed. The result of tube A confirmed the antibiotic interference and a good recovery of folate from tube C confirmed the validity of the serum folate result of tube B. A study of 30 such samples which arrived sequentially in the laboratory showed good recovery of added folic acid (mean 99-5%, CV% 15). This confirmed the potential of β-lactamase in treating samples containing antibiotics and such treatment of selected samples (or all samples) can save delays caused by having to repeat assays. Further studies using patient sera and in vitro additions of antibiotics to normal sera will be necessary to optimise this procedure.

Discussion

The recent emergence of microbiological assays for vitamins on microtitre plates owes much to advances in microtitre plate technology and to improved optical qualities of the plates. The obvious advantages of such miniaturisation of the assays include not only speed of reading and reduced reagent costs but also these simplified assays have become less exclusive and are incorporated more readily into the modern clinical or research laboratory.

In introducing such an assay for food folate analysis Newman and Tsai suggested that use of organism cryopreservation and also assay with an antibiotic resistant organism would further simplify the assay. The folate assay subsequently introduced by Horne and Patterson took advantage of organism cryopreservation but retained the traditional organism (ATCC 7469) which necessitated sterilisation of reagents by filtration and stringent aseptic precautions. The assay presented here is simpler than previously published assays and is suitable for routine diagnostic estimation of folate in serum and red cells. Use of a chloramphenicol-resistant strain of L casei (NCIB 10463) allows it to be completed openly on the laboratory bench. The plates are covered with
Assay of serum and red cell folate

plate sealers which avoids the use of a humidified incubator and permits easy mixing by inversion both before incubation and reading. The results of this assay correlate well with those of a traditional assay in the estimation of both serum (r = 0.975, n = 193) and red cell folates (r = 0.97, n = 150). An equimolar assay response was obtained for both folic acid (pteroylglutamic acid) and the reduced folate monoglutamates; 5-formyltetrahydrofolate (folinic acid), and 5-methyltetrahydrofolate, signifying that the assay will neither under- nor overestimate these reduced derivatives. The assay also shows acceptable reproducibility of values for the folate pool of both serum and red cells (tables 1 and 2) and good recovery of folic acid (98.4%) from serum. A pilot study indicates that it may be possible to assay certain occasional samples which contain high concentrations of antibiotics. This will be investigated further.

Microbiological assay using L casei has long been accepted as the reference method for routine diagnostic investigations of folate concentration. However, such microbiological assays have been difficult to carry out routinely in the general clinical laboratory setting. The assay described is within the competence of any routine clinical laboratory, has extremely low operating costs, and is particularly well suited to handling large volumes of samples.