Aseptic addition method for *Lactobacillus casei* assay of folate activity in human serum

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**SYNOPSIS** An 'aseptic addition' method is described for microbiological assay with *Lactobacillus casei* of folate activity in human serum. It has the following advantages over the previously reported 'standard' method. 1 The manipulations involved in the assay are halved, by deleting autoclaving of serum in buffers. 2 The use of 1 g. % ascorbate better preserves serum folates than the lower amounts of ascorbate which are the maximum quantities usable in the standard methods. 3 Only 0·3 ml. of serum is required (0·1 ml. for one sample; 0·2 ml. for its duplicate).

Herbert, Wasserman, Frank, Pasher, and Baker in 1959 reported that folate deficiency could be measured in man using microbiological assay of serum folate activity with *Lactobacillus casei*. Many other workers have confirmed this work (see review by Herbert, 1965). Various minor modifications of the methodology described by Herbert, Frank, Pasher, Hutner, Wasserman, and Sobotka (1959) and Herbert (1961) have been reported, including several which appeared in this Journal (Waters and Mollin, 1961; Chanarin and Berry, 1964; Spray, 1964).

It was indicated in the recent past (Herbert, 1964) that the 'aseptic addition' method for microbiological assay with *L. casei* of serum folate has a number of advantages over the 'standard' method. These advantages include halving the manipulations involved in the assay by deleting the step involving autoclaving of serum in buffer; allowing the use of 1 ml. of 1 g. % ascorbate, which better preserves serum folates than do the lower amounts of ascorbate used in the standard method, requiring only 0·3 ml. of serum (0·1 ml. for one sample; 0·2 ml. for its duplicate), an especially valuable point when infants are under study.

**METHOD**

**PREPARATION OF SERUM SAMPLES** Blood is obtained from fasting subjects using either acid-washed sterile syringes, disposable plastic syringes, or Vacutainers® no. 3200 to ensure freedom from contamination with traces of folate. The blood is allowed to stand for approximately three hours at room temperature (in the original Vacutainer

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1Becton, Dickinson and Co., Rutherford, N.J.

or after transfer of blood from syringes to acid-washed screw-top tubes). The clots are 'rimmed' with glass rods or wooden applicator sticks, the tubes centrifuged for five minutes at 3000 r.p.m. and the supernatant serum aspirated with acid-washed or disposable pipettes and frozen at –20°C. until assay. On the day of assay, the sera are thawed. A 0·1 ml. and a 0·2 ml. aliquot of serum is added aseptically to autoclaved solutions containing 5 ml. of double-strength medium (see Table I), 4 ml. of de-ionized water4, and 1 ml. of phosphate buffer (containing 1 g. ascorbic acid per 100 ml. buffer) (see Table II).

**MAINTENANCE OF ASSAY ORGANISM AND PREPARATION OF INOCULA L. casei, ATTC (American Type Culture Collection) no. 7469 (obtainable for $5 in either lyophilized or agar slant culture from American Type Culture Collection, 2112 M Street, Washington, D.C.) is maintained in the medium described in Table III and stored as a liquid culture at 4°C. We have carried our current culture in liquid medium for six years, transferring it every two weeks to fresh maintenance medium, and no mutation has yet occurred. Liquid culture is technically simpler than the agar-slant culture used by a number of investigators.

De-ionized water5 is used for preparation of all media and buffers. It is free of folate and more easily prepared than distilled water. Furthermore, the heavy metal ions in distilled water have the undesirable ability to accelerate oxidation of the ascorbic acid in the buffer.

Five hundred ml. of maintenance medium (Table III) is generally prepared at one time and dispensed in 10 ml. aliquots into 50 screw-capped tubes. The tubes, with screw caps loosely affixed, are autoclaved for 30 minutes at 118°C. They are then allowed to cool, the screw caps

4Tap water passed through a Barnstead Bantam standard demineralizer cartridge no. 0802 (Barnstead Still and Sterilizer Co., Boston, Mass.)
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TABLE I

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Amount for 4 l. of Double-strength Medium (mg.%)</th>
<th>Concentration in Final Volume (g./100 ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiamine HCl</td>
<td>0-8 g.</td>
<td>1-0</td>
</tr>
<tr>
<td>Tween 80</td>
<td>4 ml.</td>
<td>1-0</td>
</tr>
<tr>
<td>L-asparagine H₂O</td>
<td>1-0 g.</td>
<td></td>
</tr>
<tr>
<td>Nicotinic acid</td>
<td>0-5 ml.</td>
<td></td>
</tr>
<tr>
<td>Ca Pantothenate</td>
<td>0-1 ml.</td>
<td></td>
</tr>
<tr>
<td>Pyridoxine HCl</td>
<td>0-5 ml.</td>
<td></td>
</tr>
<tr>
<td>Para-aminobenzoic acid</td>
<td>0-1 ml.</td>
<td></td>
</tr>
<tr>
<td>L-cysteine HCl</td>
<td>1-0 g.</td>
<td></td>
</tr>
<tr>
<td>KH₂PO₄</td>
<td>2-00 ml.</td>
<td></td>
</tr>
<tr>
<td>NaCl</td>
<td>2-00 ml.</td>
<td></td>
</tr>
<tr>
<td>FeSO₄-7H₂O</td>
<td>0-5 ml.</td>
<td></td>
</tr>
<tr>
<td>MgSO₄-7H₂O</td>
<td>4-00 ml.</td>
<td></td>
</tr>
<tr>
<td>Yeast extract (Difco)</td>
<td>0-04 ml.</td>
<td></td>
</tr>
<tr>
<td>Protease peptone (Difco)</td>
<td>0-04 ml.</td>
<td></td>
</tr>
<tr>
<td>Glucose</td>
<td>0-10 ml.</td>
<td></td>
</tr>
<tr>
<td>KH₂PO₄</td>
<td>0-05 ml.</td>
<td></td>
</tr>
<tr>
<td>Tomato juice filtrate</td>
<td>0-05 ml.</td>
<td></td>
</tr>
<tr>
<td>Tween 80</td>
<td>0-05 ml.</td>
<td></td>
</tr>
<tr>
<td>L-cysteine HCl</td>
<td>0-05 ml.</td>
<td></td>
</tr>
</tbody>
</table>

1Adjust to pH 6-8 to 7 with 1% KOH. This medium, without the L-cysteine, has been previously described.

2Canned or vacuum-bottled tomato juice passed through coarse filter paper; straw-coloured filtrate adjusted to pH 7 with 10% KOH.

3See footnote 8, Table I.

are tightened and the tubes incubated overnight at 37°C.

The following morning they are examined for clarity (indicating absence of bacterial contamination) and, if sterile, are then stored at 4°C until used.

At a maximal interval of once every two weeks, 1 drop of stored liquid culture is added to 10 ml. of fresh maintenance medium, incubated for 18 hours at 37°C, and then stored at 4°C. During the afternoon of the day before an assay, 1 drop of the latest stored culture is added to 10 ml. of maintenance medium and incubated for 18 hours at 37°C. The next morning, 0-5 ml. of this fresh 18-hour culture is added to 10 ml. of maintenance medium and incubated for six hours at 37°C. The inoculum for the assay is prepared by adding 0-5 ml. of this fresh six-hour culture to 10 ml. of single-strength basal medium. One drop of inoculum is added to each assay flask.

Volatile preservative, consisting of 1 part ethylene dichloride, 1 part monochlorobenzene, and 2 parts l-chlorobutane, was then added to all stored solutions of assay medium and of buffer by spraying a small aliquot from a wash bottle into each solution after each use before returning to storage.

STANDARDS Three standard solutions of folic acid (pteroylmonoglutamic acid, P.G.A.) were prepared in 50 ml. aliquots, and stored between usages at -20°C (no appreciable deterioration in two months): 10⁻⁴, 10⁻¹⁰ g. per ml. Crystalline P.G.A., 10 mg., is dissolved in 100 ml. 20% ethanol containing 1 ml. of

TABLE II

<table>
<thead>
<tr>
<th>Flask</th>
<th>Double-strength Basal Medium (ml.)</th>
<th>Ion-free Water (ml.)</th>
<th>Buffer (ml.)</th>
<th>Folic Acid Standard (ml.)</th>
<th>Serum (ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number</td>
<td>Nature</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1, 8</td>
<td>Control</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>2, 9</td>
<td>Standard</td>
<td>5</td>
<td>3</td>
<td>1 (10⁻¹⁵ g./ml.)</td>
<td>0</td>
</tr>
<tr>
<td>3, 10</td>
<td>Standard</td>
<td>5</td>
<td>1</td>
<td>1 (10⁻¹⁵ g./ml.)</td>
<td>0</td>
</tr>
<tr>
<td>4, 11</td>
<td>Standard</td>
<td>5</td>
<td>3</td>
<td>1 (10⁻¹⁵ g./ml.)</td>
<td>0</td>
</tr>
<tr>
<td>5, 12</td>
<td>Standard</td>
<td>5</td>
<td>1</td>
<td>1 (10⁻¹⁵ g./ml.)</td>
<td>0</td>
</tr>
<tr>
<td>6, 13</td>
<td>Standard</td>
<td>5</td>
<td>3</td>
<td>1 (10⁻¹⁵ g./ml.)</td>
<td>0</td>
</tr>
<tr>
<td>7, 14</td>
<td>Standard</td>
<td>5</td>
<td>1</td>
<td>1 (10⁻¹⁵ g./ml.)</td>
<td>0</td>
</tr>
<tr>
<td>15, 16</td>
<td>Unknown serum 1</td>
<td>5</td>
<td>4</td>
<td>1 (no. 15)</td>
<td>0</td>
</tr>
<tr>
<td>17, 18</td>
<td>Unknown serum 2</td>
<td>5</td>
<td>4</td>
<td>0 (no. 17)</td>
<td>0</td>
</tr>
</tbody>
</table>

1Sodium phosphate buffer, 0-05 M, pH 6-1, containing 1% ascorbic acid.

TABLE III

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Amount for 500 ml. Medium (mg.)</th>
<th>Concentration in Final Medium (g./100 ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yeast extract (Difco)</td>
<td>3-75 g.</td>
<td>0-75</td>
</tr>
<tr>
<td>Protease peptone (Difco)</td>
<td>3-75 g.</td>
<td>0-75</td>
</tr>
<tr>
<td>Glucose</td>
<td>0-5 ml.</td>
<td>1-0</td>
</tr>
<tr>
<td>KH₂PO₄</td>
<td>5 ml.</td>
<td>0-2</td>
</tr>
<tr>
<td>Tomato juice filtrate</td>
<td>10 ml.</td>
<td></td>
</tr>
<tr>
<td>Tween 80</td>
<td>0-5 ml.</td>
<td>0-1</td>
</tr>
<tr>
<td>L-cysteine HCl</td>
<td>0-5 g.</td>
<td>0-1</td>
</tr>
</tbody>
</table>
0.1 N NaOH, to yield a 10⁻⁴ g. per ml. solution of P.G.A. This concentrated stock solution is stored at −20°C, and is diluted with water to provide new aliquots of the standard solutions as they are used up; it does not deteriorate appreciably in a year.

With an automatic pipette, 5 ml. of double-strength basal medium was added to each 10 ml. micro-Fernbach flask (10 ml. Erlenmeyer flasks are less expensive, equally good, but more susceptible to tipping over; alternatively, test tubes may be used, but growth in them is slower). From 1 to 4 ml. of de-ionized water was next added to each flask, the volume being such as to bring the total volume in each flask to 10 ml. when all ingredients had been added.

Table I illustrates the assay protocol of the 'aseptic addition' method for the determination of folate activity in serum. After all ingredients except serum are added, the flasks are autoclaved for 30 minutes at 105 to 110°C, then allowed to cool. Of each serum to be assayed, 0.1 ml. is added to one 'unknown' flask and 0.2 ml. to the other. The flasks are then all inoculated with L. casei.

One drop of inoculum is added to each flask, and the flasks incubated for 16 to 18 hours at 37°C. Each flask is then vigorously shaken and its content transferred to a Klett (or other) colorimeter tube. Growth density is then measured with a Klett-Summerson or other photoelectric colorimeter using a red filter (λ = 640 to 700 mμ) to reduce error due to the colour of the medium. The optical densities representing growth of the standards are plotted on the geometric ordinate scale of a semilogarithmic paper and the folate content of the standard is plotted on the logarithmic abscissa scale. The unknown sera then can have their folate content determined from this curve. The standards are run in duplicate, the results averaged, and the average values plotted. The curve should rise in slightly concave fashion (concave side up) until the 1 × 10⁻⁸ standard, and then should level off; for this reason the 3 × 10⁻⁸ standard need not be plotted on the graph.

ASSAY MEDIUM The preparation of the assay medium is described in Table I; it is, however, now commercially available. These 'dry mixes' must be stored refrigerated to avoid deterioration; to prepare the assay medium from them, just add water and stir.

PHOSPHATE BUFFER The sodium phosphate buffer is prepared as follows: dissolve 27.6 g. of Na₉H₄PO₄·H₂O in 1 l. distilled water (solution A); dissolve 71.6 g. Na₂HPO₄·12H₂O in 1 l. distilled water (solution B). To 212.5 ml. of solution A add 37.5 ml. of solution B; dilute to 1 l. with de-ionized water. The pH should be 6.1. Store solutions at room temperature with volatile preservative added.

SUMMARY OF TECHNIQUE

All glassware used in the assay must be folate-free, i.e., all flasks, beakers, etc., must have been boiled for 30 minutes in Hemosol solution (or 7 × solution or other strong blood-dissolving detergent) and then must have been rinsed 12 times with tap water and three times with distilled water.

The maximum convenient number of flasks for one assay is 114, as that is the number of 10 ml. micro-Fernbach flasks that fit into three 2- quart no. 232 Pyrex baking trays; 10 ml. Erlenmeyer flasks are acceptable, as are test tubes. Micro-Fernbach flasks are preferred because they tip over less easily.

1 The afternoon before the planned assay, inoculate tube with L. casei for 16-18 hour growth as follows: from the last 16-18 hour growth (stored in refrigerator at 4 to 6°C) (safe storage period three weeks) inoculate 1 drop into 10 ml. of fresh maintenance medium. Place in the incubator at approximately 5 p.m. and let it remain overnight at 37°C.

2 First thing in the morning, prepare six to eight hour growth as follows:—Take 0.5 ml. of the 16-18 hour growth and inoculate that amount into a fresh tube of 10 ml. of maintenance medium. Shake. Store the 16-18-hour growth in a rack in a refrigerator. Incubate the newly inoculated tube for six to eight hours, i.e., till approximately 4 p.m. It is from the six-to-eight-hour growth that the assay flasks will be inoculated.

3 Thaw the three P.G.A. standards, which should be always kept in amber or brown plastic bottles, by removing from freezer and placing in the incubator at 37°C. Remove from incubator before completely thawed, i.e., with a sliver of ice still present, and complete thawing at room temperature. Standards, i.e., solutions with known amounts of pterolymonoglutamic acid, contain 1 × 10⁻⁸, 1 × 10⁻⁹, 1 × 10⁻¹⁰ g./ml.

4 Number flasks (10 ml. micro-Fernbachs) as follows: mark one 'LC' (this will be the L. casei dilution flask), number flasks 1-14 (for standard curve in duplicate), number two flasks for each unknown, i.e., 15, 16 for first unknown, 17, 18 for second unknown, etc.

5 Make up buffer-ascorbate by adding 1 g. ascorbic acid just before use, because ascorbate in solution deteriorates rapidly, to each 100 ml. of phosphate buffer (which is stored on a shelf at room temperature).

6 With automatic syringe set at 1 ml. put 1 ml., of ascorbate buffer into each flask except the dilution (LC) flask. Rinse syringe with distilled water.

7 Add 5 ml. double-strength assay medium (stored in brown plastic bottle in refrigerator, marked '2 × L. casei') to each micro-Fernbach flask. Add volatile preservative if necessary to brown bottle before returning to refrigerator after use. (Sniff mouth of flask; if strong odour of volatile preservative is not present, add it.)

Each micro-Fernbach flask gets 5 ml. medium. For this use an automatic syringe. Use large (folate-free) beaker, and fill with medium. Try not to overestimate the quantity since more will have to be made up sooner if you do. Add 5 ml. medium to each flask.

8 Add de-ionized water with automatic syringe as follows: 5 ml. to dilution (LC) flask; 4 ml. to flasks nos. 1 and 8 (standard curve); 4 ml. to unknown; 3 ml. to standard curve flasks nos. 2, 9, 4, 11, 6, 13; 1 ml. to standard curve flasks nos. 3, 10, 5, 12, 7, 14.

9 Add standard solutions, i.e., known amounts of

1Baltimore Biological Laboratory, 2201 Aisquith Street, Baltimore 18, Maryland, and Difco Laboratory, Detroit, Michigan, U.S.A.

2Meinecke and Co., New York, N.Y.
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pteroylmonoglutamic acid. Be sure that they are thawed but not warm; a tiny remaining sliver of ice should have just dissolved when solutions are used and then returned to the freezer.

1 Flasks no. 2, 9 get 1 ml. of $1 \times 10^{-18}$ standard
Flasks no. 3, 10 get 3 ml. of $1 \times 10^{-18}$ standard
Flasks no. 4, 11 get 1 ml. of $1 \times 10^{-18}$ standard
Flasks no. 5, 12 get 3 ml. of $1 \times 10^{-18}$ standard
Flasks no. 6, 13 get 1 ml. of $1 \times 10^{-18}$ standard
Flasks no. 7, 14 get 3 ml. of $1 \times 10^{-18}$ standard

1 For pipetting standard solutions and adding standards to flasks, use 10 ml. blow-out pipettes. Fill to the 8 ml. mark with the standard you are using. Put 1 ml. in each of the two flasks that get 1 ml.; put 3 ml. in the two flasks getting 3 ml. each.

10 Get out three Pyrex dishes (2-quad trays). Place the flasks in numerical order in the trays and cover with metal or glass caps.

11 Autoclave the flasks in the Pyrex trays for 30 minutes at 225°F. Set time at 30 min., temp. dial on O of 230°. Check water level of autoclave and if necessary fill with distilled (de-ionized) water. Turn on autoclave. After 30 min., at 225 to 230°F., turn off autoclave.

12 Let trays cool for approximately half an hour.

13 Get sera (unknowns) which you are running from freezer. Sera take approximately 45 minutes to thaw.

14 When the sera have thawed and trays have cooled, add 0.1 ml. of sera (unknown) to first unknown flask and 0.2 ml. to second unknown flask. For this, use 0.5 ml. pipettes (blow-outs); fill to 0.3 ml. and place 0.1 ml. in one of the two flasks and 0.2 ml. in the other.

15 Incubate between 3 p.m. and 5 p.m. (remember six-to-eight-hour growth requires six to eight hours).

Put 0.5 ml. of the six-to-eight-hour growth into flask marked ‘LC’. This must be done with aseptic technique. Mix thoroughly by rotating flask, or stirring with pipette (sterile). Take sterile 1 ml. pipette, flame tip (briefly). Let tip cool. Fill pipette with solution in freshly-stirred ‘LC’ flask. Place one drop in each flask including standard curve flask.

16 Since L. casei falls into the facultative anaerobe category, the following steps should be taken before incubation overnight: the Pyrex trays with the capped flasks must be covered by an inverted Pyrex tray. Tape the sides of the trays together with masking tape. Place the trays in the incubator overnight.

17 The following morning glance at the bottoms of standard curve flasks. If the flasks have been incubated long enough a definite increase in the amount of growth will be evident in the flasks with each stepwise higher concentration of pteroylmonoglutamic acid; if this increase is not evident to the naked eye, incubate for two or more hours.

18 Read the assay. The colorimeter must be turned on approximately 20 minutes before using (warm-up necessary). Colorimeter must be properly adjusted with blank. Take out a colorimeter tube. Pour contents of the flasks one by one immediately after vigorously shaking flask into colorimeter tube, read. Pour contents back into flasks. As you go, record as indicated in the data book.

19 Graph the standard curve.

20 Calculate the unknowns in terms of the standard curve.

DISCUSSION

The aseptic addition method here described has been in use since 1960 in our studies in Boston and New York, and is also now being used by people to whom we have supplied the details in other American cities, Puerto Rico, South America, and England. The validity of the assay is unaffected by various minor changes in assay method or medium, provided contamination of glassware or ingredients by folate is excluded. (The commercial grade ingredients, especially casein and glucosamine, from many drug firms are contaminated with sufficient folate to invalidate the assay.) Both the assay medium and the maintenance medium we use (Baker, Herbert, Frank, Pasher, Hutner, and Wassermann, 1959; Herbert, 1961) are almost identical to prior media such as described by Jukes in 1955. The validity of the assay resides not in minor changes in method or medium, but rather in the use of L. casei as the assay organism and in the use of a reducing agent to protect folate against oxidative destruction. This is because the major folate in human serum appears to be N5 methyltetrahydro folic acid (Herbert, Larrabee, and Buchanan, 1962), on which L. casei grows but Streptococcus faecalis and Pedicoccus cerevisiae (Leuconostoc citrovorum) do not grow.

The use of a reducing agent such as ascorbate to protect serum L. casei folate activity from destruction during the course of the assay is crucial because serum L. casei folate activity is highly labile. In our experience (Herbert, 1962), as well as that of Davis and Kelly (1962), the addition of ascorbate to serum before storage by freezing at $-20°C$. is not necessary to preserve folate. Waters and Mollin (1961) found some decline in serum L. casei folate activity when serum was stored at $-20°C$. without added ascorbate; Chanarin and Berry (1964) also reported some decline without added ascorbate. Spray (1964) confirmed our report (Herbert, 1961) that ascorbate stimulates the growth response of L. casei, and also presented data supporting the contention that there may be loss of folate activity of serum stored for 12 weeks at $-15°C$. Unlike these workers, we do not add 5 mg. ascorbate per millilitre of serum during storage periods preceding assay, partly because we have not found any deterioration after storage up to two years and partly because added ascorbate renders specimens less suitable for vitamin $B_{12}$ assay because of the destructive effect of ascorbate on the natural forms of vitamin $B_{12}$ present in human serum (Smith, 1965).
A very important methodological consideration is cleanliness to avoid contamination with folate. Contamination with micro-organisms is rarely a problem because of the rapidity with which L. casei grows and makes lactic acid. Also important is refrigeration of media to avoid deterioration, and the aforementioned preservation of serum against oxidative destruction.

Since 1960, we have assayed 4,500 sera by the aseptic addition method. Interpretation of results is as indicated in Table IV. While the aseptic addition method generally gives results essentially identical to the standard method, it occasionally yields significantly higher results. We interpret this as due to the greater preservation of highly labile serum folates, present to variable degree in different sera, by the larger ascorbate concentration, and also to the lack of exposure of the serum to autoclaving such as occurs in the standard method.

Tetracycline antibiotics suppress L. casei growth, yielding false low results. This fact must be kept in mind when interpreting results with sera from patients receiving antibiotic therapy.

**ADDENDUM**

Harper (1965) has recently reported a modified aseptic addition assay procedure for the measurement of serum folate activity. His paper confirmed much of our work. The author modifies our assay procedure in two ways. He reduces the final volume from 10 ml. to 4 ml.; this has value when only 0.04 ml. of serum can be obtained instead of 0.1 ml. for assay. He deletes phosphate buffer, stating that the buffering capacity of the assay medium is sufficiently good and the extra buffer is superfluous. He also states that 'the assay medium of Waters and Mollin with additional L-tryptophan is used instead of the assay medium of Herbert'. In fact this 'assay medium of Waters and Mollin' with additional L-tryptophan is our assay medium. They used our medium but deleted tryptophan. By again adding tryptophan, Harper has made the assay medium as we originally described it.

I am deeply indebted to the following technician co-workers: Becky Fisher Dunn, Becky Jo Koontz, Nancy Cunneen Boardman, Brenda Conti Dicken, Mary Ellen Spector, Mary Small, Virginia Chapin, Peter Mason, and Leona Bandel.

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