Rapid protein A assay for intrinsic factor and its binding antibody

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SUMMARY A simple and rapid method for the measurement of cobalamin bound intrinsic factor (Cbl-IF) complex and intrinsic factor binding antibody is described. The method is based on the principle of affinity chromatography and adapted to a batch separation technique. A specific ligand staphylococcal protein A was coupled to Sepharose to form a convenient solid phase matrix. The intrinsic factor binding antibody in patients with pernicious anaemia was used to form an immune complex with Cbl-IF. This complex was adsorbed on to staphylococcal protein A. Gastric juice from control subjects and patients with pernicious anaemia was assayed for intrinsic factor activity and the results correlated very closely with two other established methods. Sera from 30 control subjects were assayed for binding intrinsic factor antibody and all were found to be negative; of 15 patients with pernicious anaemia, six were positive. These patients were selected with blocking antibody. The method does not require technologically advanced protein separation techniques and could therefore be applied in any clinical laboratory using radioisotopes. It could also be adapted to assay cobalamin in body fluids or in food.

The measurement of cobalamin bound intrinsic factor (Cbl-IF) complex is important in the study of the physiology and pathology of Cbl absorption. Although there are simple and rapid methods for measuring free intrinsic factor, the available methods for measuring Cbl-IF are either tedious or not suited to assaying large numbers of samples. A simple and specific method for measuring Cbl-IF was therefore developed, adapting the principle of affinity chromatography to a batch assay separation technique. The specific ligand used was staphylococcal protein A, which binds specifically to the Fc region of IgG molecules. There are two types of intrinsic factor antibodies which are directed towards different antigenic sites on the intrinsic factor molecule: a blocking antibody which blocks the attachment of Cbl to intrinsic factor; and a binding antibody which binds to intrinsic factor or IF-Cbl complex. The binding antibody to intrinsic factor found in some patients with pernicious anaemia was used to form an immune complex with IF-Cbl. This complex is adsorbed on to staphylococcal protein A agarose gel, which forms a convenient solid phase matrix. This method can be used to measure intrinsic factor, antibody to intrinsic factor, and Cbl.

This paper describes the staphylococcal protein A method for the assay of IF-Cbl and for the screening of patients with pernicious anaemia for the presence of binding antibody to intrinsic factor. The intrinsic factor assay results were compared with those from two established methods, one using the blocking antibody to intrinsic factor and another using cobinamide (Cbi) which binds almost exclusively to R-protein and not to intrinsic factor. Control samples of gastric juice were also assayed for intrinsic factor by gel filtration chromatography.

Preliminary results of this work were presented at the International Society of Haematology congress in Sydney in 1986.

Material and methods

PROTEIN A-SEP哈RASE CL-4B
Staphylococcal protein A covalently coupled to Sepharose by the cyanogen bromide method (Pharmacia Fine Chemicals AB, Sweden) was swollen in 10 mM phosphate buffered saline (PBS), pH 8-0, containing 0-1% sodium azide and stored at 4°C. Before use the gel was washed in a sintered funnel with 0-1 M sodium citrate/citric acid (pH 3-0), containing azide 0-1%, to free any bound material and was then equilibrated at pH 8-0. The gel can be used repeatedly without appreciable deterioration. The protein A content of the swollen gel was 2 mg/ml and the binding capacity for human IgG was about 25 mg IgG/ml gel.

Accepted for publication 1 December 1987
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**ALBUMIN COATED CHARCOAL**

Charcoal suspension was prepared as described. Activated acid washed charcoal was obtained from BDH Chemicals Ltd and 20% solution of bovine albumin from the Blood Group Reference Laboratory, Oxford.

**GASTRIC JUICE**

Gastric juice specimens from 10 healthy subjects and two patients with pernicious anaemia were used. The latter two specimens were obtained without pentagastric stimulation during routine investigations. Eight of the 10 specimens from healthy subjects were kindly donated by Dr Russell Bayley (Amersham International); the remaining two, which were used as controls, were collected from two healthy volunteers who were members of staff at St Bartholomew’s Hospital. Gastric juice from healthy subjects was aspirated at 15, 20, and 45 minutes after an intramuscular injection of pentagastrin (6 μg/kg) was given. The juice was filtered and the pH adjusted to 10-0 with 1N NaOH to inactivate the pepsin and 30 minutes later brought down to pH 7-4 with 1N hydrochloric acid. The juice was stored in small aliquots at −20°C until further use.

**INTRINSIC FACTOR ANTIBODY**

Sera from five patients with pernicious anaemia, all containing blocking antibody to intrinsic factor, were screened for binding antibody to intrinsic factor and two were found to be positive. One of these was used throughout this study.

Sera from 30 healthy subjects and 15 patients with pernicious anaemia were used.

**SEPHACRYL-S200 COLUMN CHROMATOGRAPHY**

Column chromatography was used to measure intrinsic factor activity in control gastric juice. The separation of intrinsic factor from R-protein is improved by incubating the gastric juice with serum containing binding antibody to intrinsic factor before loading the sample on to the column. The Cbl-intrinsic factor-antibody complex then elutes in the excluded volume before the R-protein. Gastric juice (100 μl) was incubated with CN-(Co)Cbl one and a half times its predetermined binding capacity for 20 minutes at room temperature; 200 μl of antibody serum was then added and incubated for a further 20 minutes; the mixture was then cooled and applied to 2.5 × 100 cm column of Sephacryl-S200 (Pharmacia Fine Chemicals).

Columns were equilibrated and eluted at 4°C with 0.05 M sodium phosphate buffer (pH 7-4) containing 0.5 M sodium chloride and 0.02% sodium azide. Fractions (4 ml) were collected and the radioactivity of each fraction was determined in a well type gamma scintillation spectrometer. The amount of radioactivity in the protein peaks was converted to picograms of band CN-(Co)Cbl using the appropriate specific activity.

**PRELIMINARY INVESTIGATIONS**

**Combination of binding antibody with Cbl-IF complex**

Adding increasing amounts of Cbl-IF complex to a constant amount of antibody. These experiments were carried out using small amounts of Cbl-IF complex which are not sufficient to saturate the amount of antibody used. Four tubes were set up for each amount of Cbl-IF tested. Buffer at pH 8-0 (400 μl), 5 μl of 20% bovine albumin, and 80 μl of CN-(Co)Cbl were added to each tube to saturate all binding sites of gastric juice. Increasing amounts of gastric juice 5, 10, 15, 20, 25, 30 μl were pipetted and the tubes incubated at room temperature for 20 minutes. To two of the four tubes 35 μl of serum containing antibody was added, and normal serum or buffer was added to the remaining two tubes. These served as control tubes to determine the non-specific binding. After the addition of 500 μl of staphylococcal protein A the tubes were incubated at 4°C for 20 minutes washed twice in buffer (pH 8-0) and the radioactivity of the pellet gel counted.

Adding increasing amounts of antibody to a constant amount of Cbl-IF complex. The effect of adding excessive amounts of antibody to a limited amount of Cbl-IF was tested. The tubes were set up in triplicate and 400 μl of buffer, 5 μl of 20% albumin, and 25 μl of CN-(Co)Cbl were added. After the addition of 10 μl of gastric juice the tubes were incubated at room temperature for 20 minutes. To a duplicate set of tubes 5, 10, 15, 20, 30, 35, 40, 45, 50 μl of serum containing antibody was added and control serum was added to the remaining set of tubes. The subsequent steps were as described above.

**ASSAY OF INTRINSIC FACTOR USING STAPHYLOCOCCAL PROTEIN A**

The scheme for the assay of Cbl-IF using staphylococcal protein A-Sepharose is summarised in table 1. The tubes were incubated at room temperature for 20 minutes after each addition of CN-(Co)Cbl and serum containing antibody. The added CN-(Co)Cbl and antibody were both in excess of the anticipated...
binding sites of the gastric juice and intrinsic factor, respectively. To facilitate pipetting, staphylococcal protein A was diluted 1/2 in buffer before being added to the assay tubes. All the tubes were subsequently incubated at 4°C for 20 minutes, centrifuged, and the gel washed twice in the buffer. The gel-pellet was counted in a well-type scintillation counter. The average counts of tubes 5 and 6 containing control serum were subtracted from the average counts of tubes 3 and 4 containing the antibody serum. The amount (ng) of bound CN(57)CoCbl was calculated as follows:

\[
\frac{\text{Sample counts (antibody) } - \text{control counts}}{\text{Total counts } - \text{background}} \times 3 \text{ng}
\]

This represents the amount of CN(57)CoCbl bound to intrinsic factor in the sample.

Intrinsic factor in gastric juice samples was measured by two methods—that of Gottlieb et al.² and that of Begley and Trachenberg.³ These methods require the blocking antibody to intrinsic factor and Cbi, respectively. Serum from one patient with pernicious anaemia, with a high titre of blocking antibody, and Cbi Dicyanide (Sigma Chemicals) were used. The two control gastric juice samples were also assayed for intrinsic factor using column chromatography.⁵

**ASSAY OF BINDING ANTIBODY TO INTRINSIC FACTOR IN SERUM**

To 400 μl of buffer (pH 8-0) was added 5 μl of 20% bovine albumin, 20 μl gastric juice, and 50 μl CN-(57)CoCbl. The tubes were incubated for 20 minutes at room temperature and 50 μl of serum was added (positive reference with high titre antibody, negative control, or the unknown sample). Staphylococcal protein A slurry (500 μl) was added and the tubes incubated for 20 minutes at 4°C, centrifuged at 1000 g, the gel washed once in buffer, and the pelleted gel counted. When the average counts obtained with the patient sample was at least twice as high as the counts obtained with the negative control, the assay was positive for the presence of intrinsic factor antibody in the patient serum. If a quantitative measurement is needed then two washes should be performed and the non-specific binding taken into account. The average counts of the negative control were subtracted from the average counts containing the antibody serum. The amount (ng) of CN(57)CoCbl bound to intrinsic factor was calculated as follows:

\[
\frac{\text{Sample counts (antibody) } - \text{control counts}}{\text{Total counts } - \text{background}} \times 5 \text{ng}
\]

Fig 1 shows that over the range tested, the relation between antibody and Cbl-IF complex was linear, so that Cbl-IF measurements could be made without performing a series of standards. Furthermore, excess antibody did not appreciably affect the results (fig 2). The non-specific binding in the tubes containing control serum or buffer was similar. This was usually less than 1% and never exceeded 2% of the total count when the gel was washed twice. After one wash the non-specific binding averaged 3% and did not exceed 3.5%.

**Results**

**COMPARISON OF STAPHYLOCOCCAL PROTEIN A, BLOCKING ANTIBODY, AND CBL ASSAY**

Table 2 shows the results obtained by these three
different intrinsic factor assays. There was good agreement between the results. The correlation between staphylococcal protein A and blocking intrinsic factor antibody methods (r = 0.99, p < 0.001) was excellent, and the calculated regression line (y = 0.88 x + 7.9) was not significantly different from the line of identity (45°). The correlation of the results between staphylococcal protein A-Sepharose and Cbi methods was similar (r = 0.97, p < 0.001), and the calculated regression line (y = 0.85 x + 8.5) was not significantly different from the line of identity (45°).

The two control gastric juice samples were assayed repeatedly. Using four different assays based on completely different principles, the results obtained were similar (table 2).

SCREENING SERA FOR BINDING ANTIBODY
Thirty sera from control subjects were screened and found to be negative. Of the 15 patients with intrinsic factor blocking antibody in their serum, six had intrinsic factor binding antibody.

Discussion
In this study staphylococcal protein A coupled to Sepharose was used to measure intrinsic factor and its binding antibody. The antibody interaction with antigen was linear at low concentrations, which agrees with the results of previous work done in this laboratory and is similar to the interaction of the blocking antibody to intrinsic factor. Furthermore, excess antibody, up to 10-fold the amount required to saturate intrinsic factor hardly influenced the results of intrinsic factor activity in gastric juice from normal subjects and patients with pernicious anaemia measured by this assay closely correlated with the results of two established assays. This is particularly relevant as the two assays are complementary: one measures the intrinsic factor activity in gastric juice by determining the drop of Cbl binding in gastric juice using blocking antibody which blocks the attachment of Cbl to intrinsic factor; the other uses cobinamide which blocks the attachment of Cbl to R-protein present in gastric juice and measures the binding of intrinsic factor directly.

In a group of 15 patients with pernicious anaemia with positive blocking antibody six (40%) were also found to have the binding antibody, which was slightly
higher than the average incidence (33%) expected in patients with pernicious anaemia tested at random. The binding antibody could not be detected in any of the control subjects.

The advantage of this method of assay is that it can measure radioactive and potentially non-radioactive Cbl-bound intrinsic factor which facilitates binding studies of Cbl to intrinsic factor, carried out under different conditions, such as pH and proteolytic enzymes. This may be of particular value in investigating further the aetiology of Cbl malabsorption in pancreatic insufficiency and the mechanism by which bicarbonate and pancreatic proteases are necessary to degrade partially R-protein and enable Cbl to be bound exclusively to intrinsic factor. Furthermore, screening patients for binding antibody to intrinsic factor eliminates false positive results caused by high serum Cbl concentrations, which can be a problem with the blocking intrinsic factor antibody test. It is predicted that in patients in whom serum IgG is greatly increased, falsely low results may be obtained. Work is in progress to develop an assay for serum Cbl using this technique.

The method is simple, sensitive, and accurate and suitable for use in any clinical laboratory using radioisotopes.

I thank Drs John Burman and John Amess of St Bartholomew’s Hospital for providing the clinical data and blood samples. I also acknowledge Professor Alan Waters (St Bartholomew’s Hospital) for many useful discussions and Mr James A Begley of the Veterans’ Administration, Albany, New York for his helpful comments and suggestions.

References


