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Research Article

Human intrinsic factor (*IF*) saturated with ⁶⁰Co-labeled cyanocobalamin (⁶⁰CoB₁₂) was purified and then iodinated with ¹²⁵I to yield ¹²⁵I-labeled IF-⁶⁰CoB₁₂ preparations of high specific activity. Sephadex G200 and DEAE-cellulose chromatography of the iodinated IF-⁶⁰CoB₁₂ complex showed coincidence of the major ¹²⁵I and the ⁶⁰Co radioactivity peaks. During starch-gel electrophoresis ⁶⁰Co radioactivity from noniodinated and iodinated complexes migrated to the same extent while ¹²⁵I radioactivity from the iodinated complex migrated slightly further anodally than did the ⁶⁰Co radioactivity. After the iodinated complex was mixed with antibody to the IF-B₁₂ complex (antibody II) the ¹²⁵I and ⁶⁰Co radioactivity were: (*a*) precipitated in similar amounts by antiglobulin serum. (*b*) eluted coincidentally in the 19S region on Sephadex G200, and (*c*) excluded to the same extent from starch gel during electrophoresis. After equilibrium exchange of IF "blocking" antibody (antibody I) for ⁶⁰Co-vitamin B₁₂ on ¹²⁵I-labeled IF. ¹²⁵I radioactivity from the IF-antibody I complex: (*a*) was precipitated by antiglobulin serum, (*b*) was eluated in the 19S region on Sephadex G200 gel filtration, and (*c*) migrated slowly towards the anode on starch-gel electrophoresis. Urinary excretion of ⁶⁰Co radioactivity in pernicious anemia patients after oral administration of ⁶⁰Co-vitamin B₁₂ bound to freshly prepared ¹²⁵I-labeled IF was similar to that obtained with noniodinated intrinsic factor.

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Radioiodination of Human Intrinsic Factor

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ABSTRACT Human intrinsic factor (IF) saturated with 60Co-labeled cyanocobalamin (60CoB12) was purified and then iodinated with ^{125}I to yield $^{125}\text{I}\text{--labeled IF-}^{60}\text{CoB}_{12}$ preparations of high specific activity. Sephadex G200 and DEAE-cellulose chromatography of the iodinated IF-60CoB₁₂ complex showed coincidence of the major ¹²⁵I and the ⁶⁰Co radioactivity peaks. During starch-gel electrophoresis ⁶⁰Co radioactivity from noniodinated and iodinated complexes migrated to the same extent while ¹²⁵I radioactivity from the iodinated complex migrated slightly further anodally than did the ⁶⁰Co radioactivity. After the iodinated complex was mixed with antibody to the IF-B₁₂ complex (antibody II) the ¹²⁵I and ⁶⁰Co radioactivity were: (a) precipitated in similar amounts by antiglobulin serum, (b) eluted coincidentally in the 19S region on Sephadex G200, and (c) excluded to the same extent from starch gel during electrophoresis. After equilibrium exchange of IF "blocking" antibody (antibody I) for 60Co-vitamin B₁₂ on ¹²⁵I-labeled IF, ¹²⁵I radioactivity from the IF-antibody I complex: (a) was precipitated by antiglobulin serum, (b) was eluated in the 19S region on Sephadex G200 gel filtration, and (c) migrated slowly towards the anode on starch-gel electrophoresis. Urinary excretion of ⁶⁰Co radioactivity in pernicious anemia patients after oral administration of ⁶⁰Co-vitamin B₁₂ bound to freshly prepared ¹²⁵I-labeled IF was similar to that obtained with noniodinated intrinsic factor.

These results show that iodination of IF-60CoB₁₂ complex does not markedly alter the chromatographic, electrophoretic, antigenic, or absorption-promoting properties of IF.

INTRODUCTION

Ignorance of the precise role of intrinsic factor (IF) in the physiological absorption of vitamin B₁₂ persists

despite nearly 40 yr of investigation. Of particular importance in preventing progress in this field is the fact that presently available methods of tracing IF action depend on binding of IF to vitamin B₁₂ labeled with a radioactive marker. Dependence on binding to radioactive vitamin B₁₂ prevents examination of the metabolism of IF itself, and only an independently labeled IF can overcome this difficulty.

In an attempt to obtain a suitably labeled IF preparation, human IF was first purified by the method of Chosy and Schilling (3) and then iodinated with ¹²⁵I by the technique of Greenwood, Hunter, and Glover (4). To determine whether iodination had altered the IF-B₁₂ complex, the chromatographic, electrophoretic, antigenic, and absorption-promoting properties of purified preparations of IF were compared before and after iodination. The results indicate that ¹²⁵I-labeled IF-B₁₂ is sufficiently similar to unlabeled IF to serve as a tracer of IF.

METHODS

Purification of human intrinsic factor. Human IF possessing a high degree of purity was prepared from human gastric juice by a modification of the method of Chosy and Schilling (3). Gastric juice was obtained from patients with peptic ulcer during routine gastric secretory studies. Specimens were collected in iced containers after intramuscular injection of 1.5 mg of Histalog/kg body weight. The crude juice was filtered through glass wool, raised to pH 10 with NaOH to inactivate pepsin (18), titrated to pH 7.0 with HCl and stored in 2-liter pools at — 20°C. To 500 ml of pooled gastric juice was added sufficient [®]Co-labeled cyanocobalamin ([®]CoB₁₂) to saturate all the vitamin B₁₂ binding sites of the gastric juice. The specific activity of the [®]CoB₁₂ was 0.1 mc/mg. After incubation at room temperature, the gastric juice—B₁₂ was ultrafiltered through Visking casing (Union Carbide Corp., Visking Div., Chicago, Ill.) to a volume of about 10 ml.

All procedures involved in purification, except radioactivity counting, were performed at 4°C. 4.0-ml fractions were collected from each column directly into radioactivity counting tubes.

As the first purification step, the ultrafiltered sample was applied to a 1.8×20 cm column of Amberlite CG-50 cation exchange resin of mesh size 200–400. The resin was prepared

Preliminary reports of this work have appeared in abstract form (1, 2).

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by the method of Hirs, Moore, and Stein (5), equilibrated and packed in 0.05~M sodium acetate buffer, pH 5.4, and eluted with 0.58~M sodium acetate buffer, pH 5.4. Two protein-bound radioactivity peaks were consistently found, the first comprising about 10% of the total $^{\infty}\text{CoB}_{12}$ -binding material. Since Chosy and Schilling (3) had previously shown that only the second of these two peaks has in vivo IF activity, 4.0-ml fractions from this peak were pooled and ultra-filtered.

The second step in the purification process consisted of downward flow Sephadex G200 gel filtration using a 50×3.8 cm column and 0.05 M sodium phosphate buffer, pH 7.5. A single protein-bound radioactivity peak was eluted in the albumin region from the ultrafiltered pooled fractions from the Amebrlite column. 4.0-ml fractions from this peak were pooled and concentrated by ultrafiltration.

The final purification step was ion-exchange chromatography on a 1.8×18 cm column of DEAE-cellulose prepared by cycling through HCl and NaOH and equilibrated in 0.02 M sodium phosphate buffer, pH 7.5, at 4°C. The concentrated sample obtained from gel filtration was allowed to enter the DEAE-cellulose column which was then washed with 40 ml of the equilibrating buffer. A gradient elution was applied to the column with rising molarity and falling pH from 0.02 M sodium phosphate buffer, pH 7.5, to 0.2 M sodium dihydrogen phosphate, pH 4.3. A single protein-bound radioactivity peak was eluted, and after removal of an aliquot for nitrogen determination (6), the final pooled sample was stored at -20°C until used.

Radioiodination of human IF. The chloramine T method of Greenwood et al. (4) was used to label the purified IF-\(^{\text{0}}\)COB₁₂ preparation (IF-\(^{\text{0}}\)COB₁₂) with radioiodine \(^{125}\)I. The appropriate quantity of chloramine T was selected by estimating the destructive effect of chloramine T on the \(^{\text{0}}\)COB₁₂ binding capacity of the purified IF. Concurrently, as a basis for comparison between the destructive effect of chloramine T on IF-\(^{\text{0}}\)COB₁₂ and on crude dialyzed human gastric juice-\(^{\text{0}}\)COB₁₂ (GJ-\(^{\text{0}}\)COB₁₂), equal concentrations of chloramine T solution were also added to GJ-\(^{\text{0}}\)COB₁₂. Equivalence of \(^{\text{0}}\)COB₁₂ binding by the two preparations was obtained by dilution in saline of the more concentrated IF-\(^{\text{0}}\)COB₁₂. As a result, both preparations bound 17 ng \(^{\text{0}}\)COB₁₂/ml, but the nitrogen content of the GJ-\(^{\text{0}}\)COB₁₂ was more than 300 times greater than that of IF-\(^{\text{0}}\)COB₁₂.

Serial dilutions containing 80.0-0.009 mg/ml of chloramine T were then added in a final volume of 0.1 ml solution to 1.0 ml either of the GJ-60CoB12 or of the IF-60CoB12 preparation. This mixture was incubated for 60 min at room temperature. To remove any free 60CoB12 released by exposure to chloramine T, 1.0 ml of bovine serum albumincoated charcoal (7) was added to the mixture. After adsorption of free 60CoB12, the charcoal was separated by centrifugation, and the supernate containing the remaining protein-bound 60 CoB12 was decanted. Radioactivity in the supernate was compared with that of IF-60CoB12 and GJ-60CoB12 standards which had not been exposed to chloramine T. Table I shows per cent destruction of vitamin B₁₂ binding for each amount of chloramine T added. Release of ⁶⁰Co radioactivity from the GJ-60CoB12 complex occurred only with amounts of chloramine T greater than 500 µg. In contrast, 125 µg chloramine T was sufficient to destroy 14% of the ⁶⁰CoB₁₂ binding of purified IF. This difference in chloramine T sensitivity of the two preparations may have been related to the difference in their protein concentration, but this possibility was not tested. because of the large quantities of purified material required for such an investigation.

TABLE I

Per cent Destruction of B₁₂ Binding*

Chloramine	Purified IF-60CoB12	Crude dialyzed GJ-60CoB12
T added	(0.135 μg N)	(44 μg N)
μg	%	%
8000	98 ± 0.0	92 ± 5.0
4000	97 ± 1.4	85 ± 1.4
1000	96 ± 1.4	25 ± 11.3
500	73 ± 13.5	8 ± 2.0
250	28 ± 13.5	2 ± 5.4
125	14 ± 10.6	3 ± 2.3
62.5	3 ± 3.0	3 ± 3.0
31.2	7 ± 3.6	1 ± 0.5
3.9	5 ± 5.0	1 ± 1.4
0.9	2 ± 2.2	2 ± 1.4

^{*} Mean ±sp of two experiments.

Since some denaturation of the IF might occur even with amounts of chloramine T insufficient to cause significant damage to ⁶⁰CoB₁₂ binding, 31.2 µg of chloramine T, one-fourth the amount that caused demonstrable destruction, was used for radioiodine labeling.

IF-⁶⁰CoB₁₂ containing 5-8 μg nitrogen and 0.5 μg ⁶⁰CoB₁₂ was thawed, transferred to a shortened disposable plastic counting tube, and diluted to 2.0 ml with 0.05 M sodium phosphate buffer, pH 7.5. The tube was then sealed with a rubber cap and placed in a small lead pot in a hood. Through the rubber cap, 1.0 ml of NaI solution containing from 5 μc to 1.0 mc ¹²⁶I and 31.2 μg of freshly prepared chloramine T in 0.1 ml distilled water was then injected with a tuberculin syringe. 20 min later the reaction was stopped by the injection of 1.0 ml of potassium metabisulphite (96 mg/ml) and 1.0 ml potassium iodide (800 mg/ml). The lead pot containing the tube and reagents was shaken gently after each addition.

To separate the iodinated IF-®COB₁₂ (128I-IF-®COB₁₂) from the iodinating reagents, the contents of the tube were removed through the rubber cap with a long needle attached to a disposable syringe. The syringe was then emptied into a 2 × 40 cm column of Sephadex G50 in bead form, packed in 0.05 M sodium phosphate buffer, pH 7.5. The sample was eluted with the same buffer at a flow rate of 40 ml/hr. The distribution of ¹²⁸I radioactivity allowed easy identification of radioactivity appearing immediately after the void volume. This protein-bound radioactivity was consistently well separated from free ¹²⁸I. The ¹²⁸I-IF-®COB₁₂ was then pooled, ultrafiltered if necessary, and stored at — 20°C until use. To ensure complete removal of small amounts of free iodine, ¹²⁸I-IF-®COB₁₂ was dialyzed against running tap water for 12 hr immediately before use.

Evaluation of the effect of iodination on human intrinsic factor. Since the only difference between ¹²⁵I-IF-⁶⁰CoB₁₂ and IF-⁶⁰CoB₁₂ was the exposure of the former to the iodination reaction, the chromatographic, electrophoretic, antigenic, and absorption-promoting properties of the two preparations were compared to evaluate the effect of iodination of IF-⁶⁰CoB₁₂.

DEAE-cellulose chromatography was performed by the method already described. Gel filtration was carried out using a reverse flow 2.0×103 cm column of Sephadex G200.

Vertical electrophoresis was performed in alkaline starch borate gel by the method of Smithies (8). Electrophoresis was carried out for 18 hr at 130 v either at room temperature or at 4°C. The gel was then cut into 0.5 cm slices and assayed for radioactivity.

IF antibody serum was obtained from patients with adult type pernicious anemia (P.A.) established by the presence of histamine-fast achlorhydria and abnormal vitamin B₁₂ absorption which was corrected by human IF (9). Each serum was tested for antibody I or "blocking" activity by the charcoal test of Gottlieb, Lau, Wasserman, and Herbert (7) and for antibody II or "binding" activity by the antiglobulin coprecipitation technique of Taylor, Roitt, Doniach, Couchman, and Shapland (10) as well as by the electrophoretic retention test of Jeffries, Hoskins, and Sleisenger (11). P.A. sera containing IF antibody II (12) (AbII) precipitated IF-⁶⁰COB₁₂ with antiglobulin serum and also caused retention of the IF-⁶⁰COB₁₂ at the origin of the starch gel. Although certain P.A. sera contained only AbI, sera with AbII activity always contained, in addition, AbI.

Reactions between ¹²⁶I-IF-⁶⁰CoB₁₂ and AbII were demonstrated by three techniques: (a) a modification of the antiglobulin coprecipitation technique of Taylor et al. described in detail by Schade, Abels, and Schilling (12), (b) the Sephadex G200 gel filtration method of Imrie and Schilling (13), and (c) the electrophoretic retention test of Jeffries et al. (11). Reactions between ¹²⁶I-IF-⁶⁰CoB₁₂ and AbI were demonstrated by incubating the ¹²⁶I-IF-⁶⁰CoB₁₂ for 24 hr at 37°C with AbI to allow the AbI to exchange (14) with the ⁶⁰CoB₁₂ bound to ¹²⁶I-labeled IF. The same three techniques described above were then used to test for the ¹²⁶I-IF-AbI complex obtained by this prolonged incubation.

Rabbit anti-human globulin serum obtained from rabbits injected with ethanol-fractionated, DEAE-cellulose-separated, human gammaglobulin was kindly supplied by H. L. Deutsch and R. L. Johnson.

The urine radioactivity test described by Schilling (9) was used to compare absorption of ¹²⁶I-IF-⁶⁰CoB₁₂ or IF-⁶⁰CoB₁₂ containing 0.5 μg of ⁶⁰CoB₁₂ in totally gastrectomized or pernicious anemia patients. When ¹²⁶I-IF-⁶⁰CoB₁₂ was used, subjects received Lugol's solution of iodine three drops t.i.d. for 3 days before and 7 days after the test to block thyroidal uptake of radioiodine. The dose of ¹²⁶I administered in these studies never exceeded 1.0 μc, and at least 50% of the ingested ¹²⁶I radioactivity was excreted in the urine within 24 hr. The tests were performed in the following sequence: free ⁶⁰CoB₁₂, IF-⁶⁰CoB₁₂, ¹²⁶I-IF-⁶⁰CoB₁₂. After the test with IF-bound ⁶⁰CoB₁₂, two additional flushing doses of 1000 μg nonradioactive cyanocobalamin were injected intramuscularly and urinary radioactivity was allowed to return to background before the second test with ¹²⁵I-IF-⁶⁰CoB₁₂ was performed.

Urine radioactivity was determined by counting 500-ml aliquots of urine in a well counter. All other radioactivity was assayed with a Autogamma detector (Packard Instrument Co., Inc., Downers Grove, Ill.) with one channel adjusted to count ⁶⁰Co radioactivity and another to count ¹²⁵I radioactivity. With this arrangement ⁶⁰Co counted approximately 1000 counts/nc above background and ¹²⁵I contributed to the ⁶⁰Co channel less than 0.05% of the counts detected in the ¹²⁵I channel. ¹²⁵I counted approximately 500 counts/nc above background and to this channel ⁶⁰Co contributed 5.3% of the counts detected in the ⁶⁰Co channel. To estimate ¹²⁵I radioactivity, counts attributable to ⁶⁰Co were subtracted from the total counts detected in the ¹²⁵I channel.

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RESULTS

IF isolated from human gastric juice by ion-exchange chromatography and Dextran gel filtration bound 1.0 µg $^{60}\text{CoB}_{12}/10\text{--}16~\mu\text{g}$ nitrogen. When compared to crude dialyzed gastric juice, this respresents nearly a 300-fold increase in purity. These relatively pure IF preparations were consistently obtained in yields of 10-20%. Iodination of the purified IF-60CoB12 preparation by the modification of the method of Greenwood et al. (4) resulted in ¹²⁵I-IF-⁶⁰CoB₁₂ preparations with specific activities from $0.4 \mu c$ to $50 \mu c^{125} I/\mu g$ nitrogen. When fresh ¹²⁵ I-labeled NaI was used, 50% of the 125 I radioactivity added to the reaction mixture consistently became protein bound. On the other hand, when 125I-NaI that had decayed through two or more half-lives was used, as little as 2% of the added 125 I radioactivity became protein bound. The specific activity of the iodinated complex was directly related to the radioactivity of the iodinating reagents. Since virtually all of the ⁶⁰Co radioactivity present was eluted from the Sephadex G50 column in the protein peak, ⁶⁰CoB₁₂ was not released from IF during iodination.

Column chromatography and starch-gel electrophoresis. Chromatography of ¹²⁶I-IF-⁶⁰CoB₁₂ on DEAE-cellulose resulted in the elution of a major ⁶⁰Co radioactivity peak that was coincident with the major ¹²⁶I radioactivity peak (Fig. 1). Immediately following and incompletely separated from the major peak, a second small peak of coincident ¹²⁶I and ⁶⁰Co radioactivity was consistently found. With gel filtration on Sephadex G200

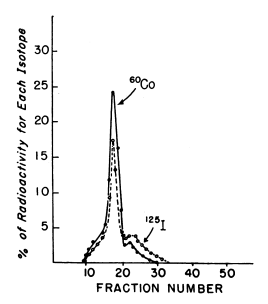


FIGURE 1 Fractionation of ¹²⁶I–IF-⁶⁰CoB₁₂ on DEAE-cellulose. The major ⁶⁰Co radioactivity coincides with the major ¹²⁶I radioactivity peak. A small peak of coincident ⁶⁰Co and ¹²⁵I radioactivity follows, and is incompletely separated from the major peak.

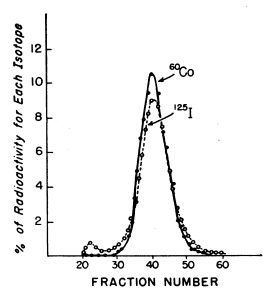


FIGURE 2 Fractionation of ¹²⁸I-IF-⁶⁰CoB₁₂ complex on Sephadex G200. The ⁶⁰Co radioactivity peak coincides with the major ¹²⁶I peak. A small ¹²⁶I radioactivity peak precedes, and is incompletely separated from, the major peak.

(Fig. 2), coincidence of a single ⁶⁰Co radioactivity peak with the major ¹²⁵I radioactivity peak was seen. With this separation a second smaller ¹²⁵I peak consistently preceded the major ¹²⁵I peak.

Simultaneous electrophoresis in starch gel of ¹²⁶I-IF-⁶⁰CoB₁₂ and IF-⁶⁰CoB₁₂ showed coincidence at the same anodal distance of the ⁶⁰Co radioactivity from both complexes (Fig. 3). Although considerable overlap occurred, the ¹²⁶I radioactivity peak from ¹²⁶I-IF-⁶⁰CoB₁₂ consistently migrated 0.5 cm further anodally than did the ⁶⁰Co radioactivity peak.

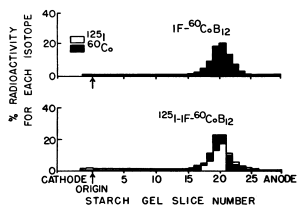


FIGURE 3 Vertical electrophoresis in alkaline starch borate gel for 18 hr at 130 v. The ⁶⁰Co radioactivity peaks both from noniodinated IF-⁶⁰CoB₁₂ (above) and iodinated IF-⁶⁰CoB₁₃ (below) are coincident at the same anodal distance. The ¹²⁵I radioactivity is seen one gel slice further towards the anode than the coincident ⁶⁰Co radioactivity.

Reactions with intrinsic factor antibody II. Incubation of either 125I-IF-60CoB12 or IF-60CoB12 with normal human serum resulted in precipitation of only a very small proportion of the ⁶⁰Co radioactivity from each complex when rabbit anti-human globulin serum (antiglobulin serum) was added (Table II). The percentage of 125 I radioactivity precipitated, however, was consistently somewhat greater. At present it is not at all clear why in the presence of normal serum, somewhat more 125 I radioactivity than ⁶⁰Co radioactivity should be precipitated by antiglobulin serum. When AbII was incubated with either 125 I-IF-60 CoB12 or IF-60 CoB12, antiglobulin serum precipitated 20 times more ⁶⁰Co radioactivity from each preparation than after incubation with normal serum. Although the percentage of 125 I radioactivity precipitated by AbII and antiglobulin serum was only 4-fold greater than with normal serum and antiglobulin serum, it was still consistently greater than 50% of the total ¹²⁵I radioactivity present.

Gel filtration on Sephadex G200 of ¹²⁸I–IF-⁶⁰CoB₁₂ incubated with AbII consistently showed a strikingly different elution pattern from that obtained after incubation with normal human serum (Fig. 4). Whereas coincident ⁶⁰Co and major ¹²⁸I radioactivity peaks were eluted in the albumin region with normal serum (peak B, Fig. 4, above), after incubation with AbII, the major ⁶⁰Co and ¹²⁸I radioactivity peaks were instead eluted in the 19S region (peak A, Fig. 4, below). Moreover, in the albumin region a small peak of ¹²⁸I radioactivity was eluted alone (peak B, Fig. 4, below) and in the third peak only ⁶⁰Co radioactivity was found (peak C, Fig. 4, below). This third peak, which consisted of dialysable free ⁶⁰CoB₁₂, was not seen after incubation of ¹²⁶I–IF-⁶⁰CoB₁₂ with normal serum.

Table II

Antigenic Activity of Iodinated Intrinsic Factor-Vitamin B₁₂

Complex towards IF Antibody II Serum

	Per cent of total radioactivity of each isotope precipitated by antiglobulin serum* 126I 60Co		
	%	%	
Noniodinated IF-60CoB ₁₂ +normal serum	_	3.0 ± 1.5	
Noniodinated IF-60CoB ₁₂ + IF antibody II serum		58.4 ± 6.8	
Iodinated IF-60CoB ₁₂ + normal serum	13.3 ± 1.9	2.0 ± 2.0	
Iodinated IF-60CoB ₁₂ + IF antibody II serum	52.8 ± 1.4	54.9 ±1.9	

^{*} Mean ±sp of six experiments.

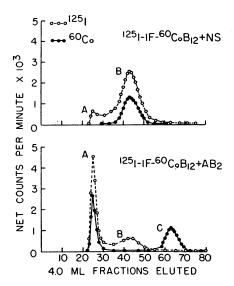


FIGURE 4 Elution pattern from Sephadex G200 of ¹²⁶I-IF- ⁵⁰CoB₁₂ incubated with normal serum (NS, above) and IF antibody II (below). Coincidence of the major ¹²⁶I peak and the ⁵⁰Co peak in the region B which contains serum albumin and a small ¹²⁶I peak A in the 19S region is seen after incubation with normal serum (above). After incubation with AbII the major coincident ¹²⁶I and ⁵⁰Co peak, A, is eluted in the 19S region (below). A small ¹²⁶I peak, B, is eluted in the albumin region without coincident ⁵⁰Co radioactivity and a small ⁵⁰Co peak, C, without coincident ¹²⁶I is seen in the small molecular region.

Incubation of ¹²⁵I–IF-⁶⁰CoB₁₂ with AbII serum abolished the anodal electrophoretic migration both of ¹²⁵I and ⁶⁰Co radioactivity (Fig. 5, below) that was seen after incubation with normal serum (Fig. 5, above). Cathodal migration of ⁶⁰Co radioactivity similar to that observed (12, 14) during electrophoresis of free ⁶⁰CoB₁₂ was seen only after incubation of ¹²⁵I–IF-⁶⁰CoB₁₂ with AbII.

Reactions with intrinsic factor antibody I. Only small amounts of ¹²⁵I radioactivity were precipitated by antiglobulin serum immediately after mixing ¹²⁵I–IF- ⁶⁰CoB₁₂ either with normal serum or with AbI serum (Table III). Incubation of ¹²⁵I–IF- ⁶⁰CoB₁₂ with normal serum for 24 hr at 37°C produced no change in the ¹²⁵I radioactivity precipitated by antiglobulin serum. However, incubation of ¹²⁵I–IF- ⁶⁰CoB₁₂ with AbI for 24 hr at 37°C consistently resulted in a marked increase in the ¹²⁵I radioactivity precipitated. This increase was similar to the precipitation observed when antiglobulin serum was added after AbII was mixed with ¹²⁵I-IF- ⁶⁰CoB₁₂.

Gel filtration on Sepadex G200 immediately after mixing ¹²⁶I–IF-⁶⁰CoB₁₂ with AbI resulted in the same elution pattern as was seen after mixing with normal serum (Figs. 4, above, and 6, above). After incubation with AbII for 24 hr, ¹²⁶I and ⁶⁰Co radioactivity peaks were

eluted together in the 19S region (peak A, Fig. 4, below). After incubation with AbI, however, the 19S region (peak A, Fig. 6, below) contained only ¹²⁶I radioactivity. The effects of AbII and AbI differed in another important respect. When ¹²⁶I–IF-⁶⁰CoB₁₂ was incubated with AbII and subjected to gel filtration the albumin region (peak B, Fig. 4, below) contained ¹²⁶I radioactivity only. When AbI was used, however, the albumin region (peak B, Fig. 6 below) contained coincident ⁶⁰Co and ¹²⁶I radioactivity. In both instances, however, the third peak (peak C, Figs. 4, below, and 6, below) contained only ⁶⁰Co radioactivity.

Starch-gel electrophoresis of ¹²⁵I-IF-⁶⁰CoB₁₂ immediately after the addition of AbI (Fig. 7, above) showed the same migration of the ⁶⁰Co and ¹²⁵I radioactivity towards the anode as was observed when ¹²⁵I-IF-⁶⁰CoB₁₂ was mixed with normal serum (Fig. 5, above). In addition a small peak of ¹²⁵I radioactivity was usually retained at the origin, and a small peak of ⁶⁰Co radioactivity consistently migrated towards the cathode. On

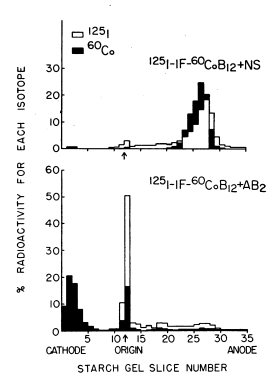


FIGURE 5 Vertical electrophoresis in alkaline starch borategel of ¹²⁶I-IF-²⁰CoB₁₂ incubated with normal serum (above) and with IF antibody II (below). The black bars show the per cent of total ²⁰Co radioactivity and the open bars the per cent of total ¹²⁶I radioactivity in each gel slice. The anodal migration of ¹²⁶I and ²⁰Co radioactivity peaks seen after incubation with normal serum (above) is abolished after incubation with IF antibody II (below). In addition a peak of ²⁰Co radioactivity migrating towards the cathode is seen after IF antibody II incubation.

TABLE III

Antigenic Activity of Iodinated Intrinsic Factor-Vitamin B₁₂

Complex towards IF Antibody I Serum

	Per cent of total 125I radioactivity precipitated by antiglobulin serum* added:		
	Immediately	After 24 hr incubation	
Iodinated IF-B ₁₂ + normal serum	10.5 ±3.2	16.3 ±1.7	
Iodinated IF-B ₁₂ + IF antibody I	16.1 ± 4.1	53.7 ± 8.7	
Iodinated IF-B ₁₂ + IF antibody II	51.1 ± 6.2	51.9 ± 5.5	

^{*} Mean ±sp of four experiments.

the other hand, incubation of ¹²⁵I-IF-⁵⁰CoB₁₂ for 24 hr at 37°C with AbI resulted in an electrophoretic pattern (Fig. 7, below) different from that obtained after similar incubation with either normal serum or AbII (Fig. 5, above and below). Whereas incubation with AbII abolished anodal electrophoretic migration of ⁵⁰Co and

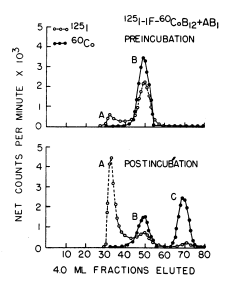


FIGURE 6 Elution pattern from Sephadex G200 of ¹²⁸I-IF- ⁶⁰CoB₁₂ immediately after exposure to IF antibody I (above, preincubation) and after 24 hr incubation at 37° with IF antibody I (below, postincubation). As shown above, the elution pattern after immediate incubation (preincubation) with IF antibody I is similar to that seen after incubation with normal serum. 24 hr incubation with IF antibody I results in a different pattern from that observed both after incubation with normal serum and IF antibody II. As shown below, in the 19S region a single ¹²⁵I peak (A) without coincident ⁶⁰Co is eluted. In the albumin region a small coincident peak (B) of ¹²⁶I and ⁶⁰Co is seen. In peak C, the small molecular region, only ⁶⁰Co is found.

¹²⁵I radioactivity, incubation with AbI resulted in the anodal migration of two ¹²⁵I peaks (Fig. 7, below). The first of these contained only ¹²⁶I radioactivity and moved only a short distance into the gel; the second contained both ⁶⁰Co and ¹²⁵I radioactivity and migrated to the same anodal distance as was observed during electrophoresis immediately after AbI had been mixed with ¹²⁵I–IF-⁶⁰CoB₁₂. After the 24 hr incubation with AbI, the cathodally migrating ⁶⁰Co radioactivity peak consisting of free ⁶⁰CoB₁₂ increased greatly.

In vivo activity. Urinary excretion of [∞]Co radioactivity after oral administration of purified but noniodinated IF-[∞]CoB₁₂ to patients with pernicious anemia or total gastrectomy was consistently greater than after free [∞]CoB₁₃ (Table IV). Enhancement of [∞]Co urinary radioactivity excretion was also seen after the oral administration of ¹²⁶I-IF-[∞]CoB₁₂. With freshly prepared ¹²⁶I-IF-[∞]CoB₁₃, the urinary excretion of [∞]Co was not significantly different from that observed with noniodinated IF-[∞]CoB₁₃. After storage for 1-2 months, however, the urinary [∞]Co excretion after ¹²⁶I-IF-[∞]CoB₁₂ fell to less than half of that obtained with noniodinated

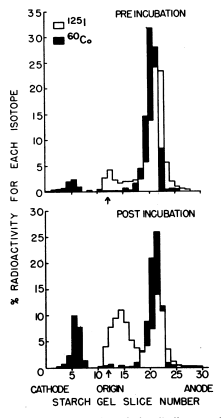


FIGURE 7 Vertical electrophoresis in alkaline starch borategel at 4°C of ¹²⁵I–IF-⁸⁰CoB₁₂ immediately after exposure to AbI (above, preincubation) and after 24 hr incubation at 37°C with AbI (below, postincubation).

Table IV
In vivo Activity of Iodinated Intrinsic
Factor-Vitamin B_{12} Complex

Chin	11 15	4 D	F. C		E 17
Subject	н. Е.	A. R.	Е. С.	в. к.	E. K.
0.5 μg Co ⁶⁰ B ₁₂	1.1	0.7	0.6	1.8	0.5
0.5 μg Co ⁶⁰ B ₁₂ -	13.0	21.1	9.5	16.8*	12.6‡
noniodinated IF					
0.5 μg Co ⁶⁰ B ₁₂ -	15.0	14.5	7.4	6.0*	6.2‡
iodinated IF					

All values expressed in terms of per cent of oral dose excreted in the urine in $24\,\mathrm{hr}$.

- * IF preparations stored for 1 month at -20°C.
- ‡ IF preparations stored for 2 months at -20 °C.

IF-®CoB₁₂ in the two trials performed. Nevertheless, excretion was still greater than that observed after free ⁶⁰CoB₁₂. Nonetheless, simultaneous electrophoresis of these stored preparations of ¹²⁵I-IF-⁶⁰CoB₁₂ and IF-⁶⁰CoB₁₂ incubated both with normal serum and with AbII demonstrated no change in their electrophoretic or antigenic properties.

DISCUSSION

Radioiodination of IF-60CoB12 consistently resulted in preparations of 125 I-IF-60 CoB12 which contained readily measureable quantities of 125 I radioactivity. Vitamin B12 was not released from the IF-60CoB12 complex during iodination. Although the vitamin B12-binding activity of the IF was thus apparently not damaged by the procedure, it was important to determine that other characteristics of the IF molecule had not been altered since many macromolecular substances are able to bind vitamin B12, but unlike IF, do not enhance vitamin B12 absorption by the distal small intestine (16) and presumably do not exhibit antigenicity to IF antibody. Since the affinity of vitamin B₁₂ for IF (17) differs markedly from the affinity of IF-B₁₂ complex for the intestinal cell surface (18), it is likely that at least two distinct active binding sites are present on the IF molecule: (a) a vitamin B12 binding site present in IF as well as other macromolecules and (b) an ileal mucosal receptor binding site and antibody binding sites which are unique to IF. Because of their apparently distinct nature, integrity of one does not necessarily mean integrity of the others. It was therefore essential to determine the effect of the iodination process on the unique antigenic and biologic properties of IF, despite the fact that the vitamin B₁₂binding activity of IF was not impaired.

The coincidence of the ⁶⁰Co radioactivity peak with the major ¹²⁶I radioactivity peak both on DEAE-cellulose and Sephadex G200 indicates that iodination of IF-⁶⁰CoB₁₂ occurs without significant damage to its chromatographic or gel filtration characteristics. Furthermore, the coincidence at the same anodal distance of the ⁶⁰Co radioactivity peaks from iodinated and noniodinated IF-⁶⁰CoB₁₂ after electrophoresis on starch gel demonstrates the absence of damage to the electrophoretic characteristics of IF-⁶⁰CoB₁₂. During electrophoresis, however, there was a consistent though very slight difference in the migration of ⁶⁰Co and the ¹²⁶I radioactivity from iodinated IF-⁶⁰CoB₁₂. Further investigations are necessary to explain this observation.

The chromatographic and electrophoretic behavior of iodinated IF-[®]CoB₁₂ provides a means of testing the effect of iodination on the antigenicity of IF-[®]CoB₁₂. For this purpose one can use the antibody that binds the IF-[®]CoB₁₂ complex (AbII) as well as the antibody that blocks B₁₂ binding by IF (AbI). The serum used as the source of AbII activity in these studies always contained AbI activity in addition to AbII activity. The AbI serum used, however, contained only AbI activity.

Several findings indicate that antigenicity of IF-60CoB₁₂ is not altered by the iodination procedure: (a) the similar precipitation of ⁶⁰Co radioactivity from iodinated and noniodinated IF-60CoB12 by AbII and antiglobulin serum (Table II), (b) the transfer of the major ⁶⁰Co radioactivity peak on Sephadex G200 from the albumin region with normal serum to the 19S region after AbII was mixed with iodinated IF-60CoB12 (Fig. 4, above and below), and (c) the abolition by AbII of the anodal electrophoretic migration of ⁶⁰Co radioactivity both from iodinated (Fig. 5, above and below) and noniodinated IF-60CoB₁₂. In addition, several observations suggest that 125 I radioactivity is attached to the IF-vitamin B12 complex rather than to any contaminating substances: (a) the 125 I radioactivity precipitated by antiglobulin serum from iodinated IF-60CoB12 incubated with AbII is four times greater than that precipitated after incubation with normal serum (Table II, (b) the 125I and 60Co radioactivity eluted from Sephadex G200 coincided in the albumin region when iodinated IF-60CoB12 was incubated with normal serum and in the 19S region when incubated with AbII (Fig. 4, above and below), and (c) anodal electrophoretic migration both of ⁶⁰Co and ¹²⁵I radioactivity from iodinated IF-60CoB12 was abolished by incubating with AbII (Fig. 5, below).

The consistent appearance of cathodally migrating ⁶⁰Co radioactivity seen on starch-gel electrophoresis (Fig. 5) when iodinated IF-⁶⁰CoB₁₂ was mixed with AbII serum suggests that this serum, which has both AbI and AbII activity, has an effect in freeing ⁶⁰CoB₁₂ from its binding to IF. This effect of AbII serum was also seen during gel filtration on Sephadex G200. Free ⁶⁰CoB₁₂ was eluted in the small molecular region (Fig. 4). On the other hand, release of ⁶⁰CoB₁₂ from IF by serum that contains AbI activity alone occurs only after prolonged incubation (Table III, Fig. 6, above and below,

Fig. 7, above and below). Since release by AbI requires prolonged incubation (14) and since ${}^{\infty}\text{CoB}_{12}$ was released shortly after IF- ${}^{\infty}\text{CoB}_{12}$ was mixed with AbII serum, it is possible that this release was an effect of the AbII activity rather than any AbI activity present in the AbII serum. On the other hand, since AbII sera tend to contain high titers of AbI, the observed rapid release might have resulted from the presence of these high titers. To distinguish unequivocally release due to AbI from that due to AbII, serum containing AbII alone would be required, but this type of IF antibody is found very rarely in pernicious anemia patients.

One might expect that when an antiglobulin coprecipitation test was performed with purified IF-60CoB12 and AbII serum, all or nearly all of the IF-bound ⁶⁰Co radioactivity would have been precipitated. Instead, the amount of **CoB12 bound to purified IF, precipitated (Table II), was similar to that found by other workers using radioactive vitamin B12 bound to crude gastric juice (10). Crude gastric juice contains, in addition to IF, non-IF vitamin B₁₂-binding macromolecules that presumably are not antigenic to AbII and thus are not precipitated by the addition of AbII and antiglobulin serum. Purified IF-60CoB12 does not contain these non-IF vitamin B₁₂-binding macromolecules (15) so that the presence of nonantigenic vitamin B₁₂ binders in the supernate after antiglobulin precipitation cannot account for the unprecipitated vitamin B12 radioactivity. The fact that some **CoB12 is released from the IF-**CoB12 complex in the presence of AbII serum may explain why in the test performed with relatively pure IF-60CoB12 all of the ⁶⁰CoB₁₂ was not precipitated by AbII and antiglobulin serum. Unfortunately, the amounts of radioactivity used in these experiments were insufficient to determine accurately whether this unprecipitated ⁶⁰CoB₁₂ was in fact free vitamin which had been released from

The results of gel filtration and electrophoresis studies indicated that prolonged incubation of AbI with 125 I-IF-60 CoB12 resulted in an exchange of AbI for 60 CoB12 on ¹²⁵I-IF. As a result of this exchange free ⁶⁰CoB₁₂ was released and 125I-IF-AbI complex was formed. Several observations support the occurrence of this exchange and indicate that the 126 I radioactivity was attached to IF rather than to cyanocobalamin: (a) after exchange a 4-fold greater precipitation of 125I radioactivity occurred when antiglobulin serum was added to the reactants than occurred before exchange (Table III); (b) 128 I radioactivity was eluted from Sephadex G200 as a peak in the albumin region coincident with "Co radioactivity (Figs. 4 above, and 6 above), and after exchange, as a single peak in the 19S region without coincident **Co radioactivity (Fig. 6, below); and (c) only after exchange were two anodally migrating peaks of 125I radioactivity, one of which was associated with

⁶⁰Co radioactivity, consistently seen on starch-gel electrophoresis (Fig. 7, below).

The results of gel filtration and electrophoresis after prolonged incubation of ¹²⁸I–IF-⁶⁰CoB₁₂ with AbI are best explained on the basis of an equilibrium (14) between ¹²⁸I–IF-AbI and ¹²⁸I–IF-⁶⁰CoB₁₂. The ¹²⁸I–IF-AbI is eluted in the 19S region on Sephadex G200 (peak B, Fig. 4, below) and migrates a short distance into the gel on starch-gel electrophoresis (Fig. 7, below). The ¹²⁸I–IF-⁶⁰CoB₁₂ is eluted in the albumin region on Sephadex G200 (peak B, Fig. 6, below) and migrates into the starch gel to the same anodal distance as noniodinated IF-⁶⁰CoB₁₂ (Fig. 7, below). The free ⁶⁰CoB₁₂ displaced by AbI appears in the third radioactivity peak from Sephadex G200 (peak C, Fig. 4, below) and migrates cathodally in the starch gel (Fig. 7, below). The equilibrium reaction may be written as follows:

AbI
+

$$^{125}I-IF-^{60}CoB_{12}$$
 \rightleftharpoons $^{125}I-IF$ + $^{60}CoB_{12}$
↓↑
 $^{125}I-IF-AbI$

These studies provide additional evidence that AbI and AbII are distinct entities as proposed by Schade, Feick, Imrie, and Schilling (14). In contrast to AbII which precipitated similar percentages of ⁶⁰Co and ¹²⁵I radioactivity from iodinated IF-60CoB12 in the presence of antiglobulin serum, AbI precipitated no ⁶⁰Co radioactivity and 125I radioactivity only after exchange had taken place (Table III). Furthermore, AbII shifted the coincident 125 I and 60 Co radioactivity peak obtained in the albumin region on gel filtration of iodinated IF-⁶⁰CoB₁₂ (Figs. 2, 4, and 6 above) to another coincident peak in the 19S region (Fig. 4, below), while AbI moved only the 125I radioactivity peak to the 19S region (Fig. 6, above). Finally, AbII blocked anodal electrophoretic migration of 125 I as well as 60 Co radioactivity (Fig. 5, below), and AbI after exchange resulted in two anodally migrating 125 I radioactivity peaks, only one of which was associated with 60Co radioactivity (Fig. 7, below).

The results of this study indicate that the iodinated complex is not entirely homogeneous. Iodination of nitrogen-containing impurities and/or aggregation and denaturation of some IF molecules that are still bound to ${}^{\infty}\text{CoB}_{12}$ during the iodination process may explain the small but persistent ${}^{126}\text{I}$ radioactivity peaks eluted without coincident ${}^{60}\text{Co}$ radioactivity from Sephadex G200 (Fig. 2, above, and 6, above) and with coincident (Figs. 2, 4, above, and 6, above) and with coincident ${}^{60}\text{Co}$ radioactivity from DEAE-cellulose (Fig. 1). The incomplete transfer of ${}^{126}\text{I}$ radioactivity from the albumin region to the 19S region on Sephadex G200 by AbII

(Fig. 4) also suggests that either iodinated protein is present as a contaminant or that some IF molecules have been altered and have lost antigenic activity. Nevertheless, the weight of evidence from studies with three complementary techniques using two distinct IF antibodies indicates that the predominant ¹²⁸I-labeled molecule in this heterogenous mixture is in fact IF.

The in vivo trials of iodinated IF-[®]CoB₁₂ tested the absorption-promoting activity of IF. The three studies with freshly prepared material suggest that the iodination procedure did not alter the IF-[®]CoB₁₂ complex from its readily absorbable form. Absorption was decreased, however, after prolonged storage although no alteration in antigenicity to AbII could be demonstrated by starchgel electrophoresis. Nonetheless, absorption was still greater than after the administration of free [®]CoB₁₂.

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