THE ABSORPTION OF RADIOIRON LABELED FOODS AND 
IRON SALTS IN NORMAL AND IRON-DEFICIENT 
SUBJECTS AND IN IDIOPATHIC 
HEMOCHROMATOSIS

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It is generally accepted that, in the absence of 
bleeding or pregnancy, approximately 1 mg. of 
iron is lost per day from the body (1–11). As a 
corollary, the quantity of iron in the body is largely 
determined by the amount of iron absorbed from 
the gastrointestinal tract. Fundamental knowl-
edge of iron absorption has been obtained by bal-
ance studies (2, 4, 12–18), by determination of 
the increase in the serum iron level (19) or circu-
lating hemoglobin (20–21) after oral iron admin-
istration, and by evaluation of the percentage of 
orally administered radioiron incorporated into 
hemoglobin (22–26) or accounted for in both he-
moglobin and feces (7, 27).

The present investigation was undertaken to 
compare the absorption of egg and vegetable iron 
with that of iron salts fed to normal subjects, to 
patients with iron-deficiency, and to patients with 
idiopathic hemochromatosis. The method of study 
used was similar to that introduced by Dubach, 
Callender, and Moore (27). These investigators 
showed that the quantity of iron used in hemo-
globin formation may not always be a true index 
of iron absorption (27, 28). Since a negligible 
quantity of iron is excreted into the intestinal tract 
(1–3, 7–8, 11) and since stool iron is almost en-
tirely unabsorbed dietary iron (2), additional in-
formation concerning iron absorption may be ob-
tained by determining both the percentage of orally 
administered radioiron incorporated into hemo-

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globin and the percentage recovered in the feces. 
The percentage of the oral dose not recovered in 
hemoglobin and feces may be considered an ap-
proximate measure of the quantity of iron de-
posited in tissue stores. In certain patients this 
figure may be subject to a significant experimental 
error due particularly to incomplete stool collection.

EXPERIMENTAL SUBJECTS AND METHODS

Three groups of patients were studied:
1. Normal subjects. This group consisted of 32 male 
patients with uncomplicated dermatitis, psychosomatic or 
psychiatric illnesses, or neurologic diseases. All had 
normal hematologic and serum iron values. Iron stores 
were not specifically measured by tissue biopsy (bone 
marrow or liver). Blood loss was excluded by history 
and stool examination. Systemic diseases that might in-
fluence iron absorption, utilization, or the level of iron 
stores were excluded by appropriate laboratory and clini-
cal evaluation.

2. Subjects with deficient iron stores. This group com-
prised 15 patients with evidence of chronic blood loss. 
The red blood cells were hypochromic and microcytic, 
serum iron levels were less than 50 micrograms per 100 
ml. and a definite history of blood loss was obtained in 
each case.

3. Subjects with excess iron stores. This group of 9 
patients had idiopathic hemochromatosis. The diagnosis 
was established by clinical features, elevated serum iron 
level, and by liver or skin biopsy.

The number of absorption studies and the form in 
which the iron was administered to these subjects are 
summarized in Table I.

Red blood cell indices were determined for each sub-
ject using equipment certified by the Bureau of Standards. 
Serum iron was measured by the method of Kitzes, 
Elvehjem, and Schuette (29). The whole blood, red cell, 
and plasma volumes were estimated using a radiophos-
phorus tagged red cell method (30).

Following these studies, tracer doses of radioiron were 
given by mouth to the fasting subject either as ferrous or
ferric chloride or as a radioiron labeled food. Ferrous chloride was prepared by the reduction of ferric chloride with powdered ascorbic acid or, in 2 studies, with cysteine. The oral dose of Fe\(^{56}\) used in the iron salt absorption studies ranged from 12 to 50 microcuries, and from 7 to 23 microcuries in the food iron absorption studies.

In all but 7 studies evaluating the absorption of ferrous chloride, ascorbic acid tablets (total of 0.5 to 1.0 gm.) were administered concomitantly with the radioactive iron salt. A variable quantity of non-radioactive carrier iron ranging from 4 or 5 mg. to 80 mg. was usually added to the tracer dose. The smaller carrier doses were used if the absorption of food iron had been previously studied in the subject, since this dose range approximated the quantity of iron administered in the labeled foods. Larger carrier doses were used in the remaining iron salt absorption studies.

The Fe\(^{56}\) labeled foods were chicken eggs and vegetables. The techniques developed for preparation of these labeled foods will be reported separately (31). The quantity of iron in a given oral dose of labeled food was determined by the method of Kitzes, Elvehejm, and Schuette (29) on an aliquot of the food substance after Kjeldahl digestion with concentrated nitric, sulfuric, and perchloric acid. Representative iron content of the various foods used is tabulated in Table II.

Radioiron labeled foods were administered after a night's fast. Two pieces of bread (containing approximately 1 mg. of iron) without butter, and black coffee (no iron) with sugar were eaten with the labeled food. Eggs were served scrambled and vegetables were boiled and served with the cooking liquid. Eight ounces of orange juice were given with scrambled eggs in a single study.

After iron administration, stools were collected until less than 1 per cent of the oral tracer dose was recovered in a 24-hour collection. Samples of blood were obtained at 2 to 5-day intervals. In most instances, blood samples were obtained until a plateau of constant activity was reached.

The preparation of the collected materials for counting was relatively simple. Water was added to feces in the large collecting bottles, total weight determined, and the mixture homogenized by an Osterizer or an Equipoise shaker. After mixing, three aliquots by weight were transferred to screw cap bottles holding 25 ml. of material. Since the quantity of radioiron present in plasma after the first 24 hours was not significant, blood samples were processed for assay by pipetting 25 ml. of whole blood into similar bottles. Appropriate standards representing the administered oral dose of radioactive iron salt or food iron were similarly prepared. The standard for iron salts consisted of a portion of the solution taken per mouth by the subjects while that for labeled foods was made up from weighed aliquots of scrambled eggs or cooked vegetables. These were digested in concentrated nitric acid and diluted to a 25-ml volume in vials like those used for counting the stool and blood samples.

Gamma radiation of the prepared samples was quantitatively measured by Geiger-Mueller counting tubes. Initially, a Sylvania GG306, all metal, bismuth cathode tube was used. This heavily shielded tube was mounted in a horizontal position in a plastic frame and the sample bottles were placed as close as possible beneath the tube, with the long axis of the sample bottle parallel to the tube. Counts were recorded on a Tracerlab Autoscaler. The counting efficiency of this arrangement for Fe\(^{56}\) was 0.4 per cent and 180 cps represented 1 microcurie. Statistical analysis of the whole sampling, positioning, and counting technique revealed a potential error that did not exceed ±5 per cent.

Later another type of Geiger-Mueller tube was used, the Texas Well Counter. This tube was approximately 8 to 10 times as sensitive as the GG306. All food absorption studies utilized the Texas Well Counter, counts being recorded by a Berkeley Decimal Scaler.

\* Welch-Allyn Company, Skaneateles, New York.

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**TABLE I**

**Summary of iron absorption studies**

<table>
<thead>
<tr>
<th>Group</th>
<th>No.</th>
<th>As FeCl(_3)</th>
<th>As FeCl(_2)</th>
<th>Eggs Vegetables</th>
<th>Total no. of studies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>32</td>
<td>25</td>
<td>2</td>
<td>14</td>
<td>6</td>
</tr>
<tr>
<td>Iron deficient</td>
<td>15</td>
<td>14</td>
<td>2</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Hemochromatosis</td>
<td>9</td>
<td>13</td>
<td>0</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Total</td>
<td>56</td>
<td>52</td>
<td>4</td>
<td>18</td>
<td>10</td>
</tr>
</tbody>
</table>

**TABLE II**

**Iron content of labeled food products**

<table>
<thead>
<tr>
<th>Food product</th>
<th>No. of assays</th>
<th>Iron content (mg./100 gm.)</th>
<th>Iron administered (mg.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Range Average</td>
<td>Range Average</td>
</tr>
<tr>
<td>Chicken eggs</td>
<td>16</td>
<td>2.8-4.7 3.8</td>
<td>3.9-10.0 6.0</td>
</tr>
<tr>
<td>Swiss chard</td>
<td>5</td>
<td>0.6-0.9 0.8</td>
<td>2.2-4.4 2.9</td>
</tr>
<tr>
<td>Beet greens</td>
<td>5</td>
<td>1.1-2.5 1.8</td>
<td>2.0-4.8 3.0</td>
</tr>
</tbody>
</table>

\* The radioactive iron used in these studies was prepared in the atomic pile at Oak Ridge. The specific activity ranged from 455 to 4,237 mc. per gm. Fe.
The accuracy of the gamma counting methods used for the assay of Fe in fecal and blood samples was determined by in vitro recovery experiments. A known quantity of Fe in the form of FeCl was added directly to 4 different fecal collections. This material was homogenized, processed as above, and the amount of Fe present determined by gamma counting techniques. Data obtained were compared with activity observed in a water solution of Fe counted under identical circumstances. These data are summarized in Table III and reveal that 96 to 101 per cent of the added Fe were accounted for in this study. The presence of solid material in feces did not significantly alter the counts recorded.

### RESULTS

**Absorption of iron by normal subjects (Figures 1, 2, 3, and 4)**

Figures 1 and 2 reveal a significant difference between the absorption of ferrous chloride and food iron. Recovery results in Figure 1 show approximately 4 to 12 per cent of radioiron were absorbed and incorporated into hemoglobin when Fe was administered as ferrous chloride along with 27 to 41 mg. of carrier ferrous iron.

### Hematologic values for subjects

<table>
<thead>
<tr>
<th></th>
<th>Range</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hematocrit (%)</td>
<td>43.0–51.5</td>
<td>46.5</td>
</tr>
<tr>
<td>Hemoglobin (gm. %)</td>
<td>13.3–16.0</td>
<td>14.4</td>
</tr>
<tr>
<td>MCHC (%)</td>
<td>28.0–33.0</td>
<td>31.9</td>
</tr>
</tbody>
</table>

(MCHC—Mean corpuscular hemoglobin concentration)
seven to 96 per cent were recovered in feces and 1 to 22 per cent were not recovered in blood or feces. The percentage of Fe\(^{99}\) incorporated into hemoglobin is indicative of the quantity of carrier iron definitely absorbed and used to form hemoglobin (0.9 to 4.6 mg.). As the legend to Figure 1 states, the unrecovered percentage of radioiron in this group and in all subsequent groups represents a combination of three factors: absorbed iron not synthesized into hemoglobin but deposited in tissue stores, the error of the determinations, and failure by the subject to make absolutely quantitative collection of stools.

The percentage of absorption and utilization of food iron (Figure 2) was significantly less than that of ferrous chloride even though the amounts of iron given were considerably smaller. With a single exception only 0.5 to 2.3 per cent of egg and vegetable radioiron were incorporated into hemoglobin. In one study 5 per cent of labeled egg iron was used to form hemoglobin. Sixty-five to 99.5 per cent were recovered in the feces and 4 to 28 per cent were not recovered in either the feces or blood. The addition of orange juice in one study did not increase the absorption of egg iron. The percentage of Fe\(^{99}\) absorbed and incorporated into hemoglobin indicated the absorption of from 0.03 to 0.3 mg. of the egg or vegetable iron administered. Calculations of the percentage of iron not recovered suggest that as much as 0.3 to 1.3 mg. of iron could have been added to tissue stores.

The greater absorption of the iron salt, ferrous chloride, is more apparent when compared with the absorption of similar quantities of ferrous chloride and egg iron (Figure 3). Seven normal subjects, who received Fe\(^{99}\) labeled eggs containing from 4 to 8 mg. of iron, absorbed and incorporated 0.5 to 5.0 per cent into hemoglobin, representing...
the absorption of from 0.08 to 0.3 mg. of iron. When a similar quantity of ferrous chloride was given, 7.8 to 74.5 per cent of the dose appeared in hemoglobin, representing the absorption of 0.32 to 3.0 mg. of iron. The increased absorption and incorporation of this iron salt into hemoglobin ranged from 2 to 100 times that of egg iron.

The effect of egg itself upon the absorption of iron was studied by giving inorganic Fe\(^{56}\), as ferrous chloride, to five subjects along with non-labeled scrambled eggs (Figure 4). Under this experimental condition a considerably smaller percentage of iron was absorbed and utilized for hemoglobin formation than when a similar amount of ferrous chloride was administered alone. However, when compared to the absorption and utilization of egg iron, there was usually an appreciably greater absorption and use of the ferrous iron salt even in the presence of eggs. This difference ranged from 1.5 to 10 times the amount of food iron absorbed and incorporated into hemoglobin.

These data as summarized in Table IV indicate that iron in eggs and certain vegetables is not nearly as well absorbed as the iron salt ferrous chloride, and that the concomitant presence of food such as egg may decrease significantly the absorption of the iron salt.

### Absorption of iron by subjects with deficient iron stores (Figures 5 and 6)

Representative results summarized in Figure 5 show that with one exception more than 20 per cent of the iron was absorbed and incorporated into hemoglobin when Fe\(^{56}\) was administered as ferrous or ferric chloride. Nearly all of the remaining radioiron was recovered in the stools except in a single patient with a complicating chronic infection (Figure 5, Studies 7 and 8).

The absorption of Fe\(^{56}\) labeled food by subjects with deficient iron stores is summarized in Figure 6. Food iron was not as well absorbed as iron
administered as ferrous chloride. Iron-deficient subjects receiving Fe\(^{65}\) labeled chicken eggs absorbed and utilized 10.3, 20.5, and 0.6 per cent of Fe\(^{65}\), while those receiving Fe\(^{65}\) labeled vegetables absorbed and utilized 3.6 and 30 per cent. Most of the remaining Fe\(^{65}\) not appearing in hemoglobin was recovered in the stools.

### TABLE IV

Per cent of radioiron incorporated into hemoglobin in normal subjects

<table>
<thead>
<tr>
<th>Form of iron</th>
<th>No. of studies</th>
<th>Oral dose (mg.)</th>
<th>Per cent in hemoglobin</th>
<th>Range</th>
<th>Mean</th>
<th>Median</th>
</tr>
</thead>
<tbody>
<tr>
<td>FeCl(_2)</td>
<td>2</td>
<td>27.0-31.0</td>
<td>0.8-0.9</td>
<td>0.85</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td>FeCl(_3)</td>
<td>12</td>
<td>20.0-41.0</td>
<td>3.9-11.6</td>
<td>8.73</td>
<td>9.80</td>
<td></td>
</tr>
<tr>
<td>FeCl(_4)</td>
<td>8</td>
<td>4.0-6.4</td>
<td>7.8-74.5</td>
<td>31.40</td>
<td>23.25</td>
<td></td>
</tr>
<tr>
<td>Egg + FeCl(_2)</td>
<td>5</td>
<td>7.9-8.4</td>
<td>1.3-43.2</td>
<td>12.60</td>
<td>3.90</td>
<td></td>
</tr>
<tr>
<td>Egg</td>
<td>14</td>
<td>4.0-8.6</td>
<td>0.5-5.0</td>
<td>1.40</td>
<td>0.60</td>
<td></td>
</tr>
<tr>
<td>Swiss chard</td>
<td>4</td>
<td>2.3-4.4</td>
<td>0.8-1.6</td>
<td>1.20</td>
<td>1.25</td>
<td></td>
</tr>
<tr>
<td>Beet greens</td>
<td>2</td>
<td>2.6-3.0</td>
<td>0.8-0.9</td>
<td>0.85</td>
<td>0.85</td>
<td></td>
</tr>
</tbody>
</table>

The quantity of carrier iron absorbed and incorporated into hemoglobin by patients with deficient iron stores ranged from 1.0 mg. when 2 to 10 mg. of food iron were given, to from 4.8 to 18.5 mg. when 31 to 80 mg. of iron were administered as ferrous or ferric chloride. These figures cannot be compared since similar quantities of iron were not administered. However, iron-deficient patients absorbed a much larger percentage of food iron than did normal subjects who received a similar quantity of food iron.

Multiple studies were carried out on two iron-deficient subjects to compare the absorption of ferrous chloride and labeled food. One patient (Figure 6, Studies 2 and 3) absorbed 20 per cent of Fe\(^{65}\) incorporated in eggs and 100 per cent of a comparable dose (5.0 mg.) of ferrous chloride. The second patient (Figure 6, Studies 4 through
7) was a female who had had a subtotal gastric resection three years previously for a bleeding ulcer. Subsequent to this operative procedure she developed a profound iron deficiency anemia in the absence of further bleeding other than that associated with her menses. This anemia was reported to be refractory to therapy with ferrous sulfate. When she received labeled eggs and chard in two separate studies she was able to absorb and incorporate into hemoglobin only 0.6 and 3 per cent, respectively. Virtually all the remaining Fe\textsuperscript{59} was recovered in the stools so that less than 0.1 mg. of iron was actually absorbed. In striking contrast, when a comparable quantity of Fe\textsuperscript{59} was administered as ferrous chloride (4.0 mg. of iron), 88 per cent was absorbed and incorporated into hemoglobin, representing the absorption of 3.5 mg. of iron. When the carrier dose of ferrous chloride was increased to approximate the 80 mg. of iron received daily on an average therapeutic regimen of ferrous sulfate, 32 per cent of the dose was absorbed and incorporated into hemoglobin, representing the absorption of 26 mg. of iron.

In summary, iron as ferrous chloride is absorbed in larger quantity by iron-deficient subjects than is the iron in eggs and vegetables. However, food iron is absorbed and used to a much greater extent by patients with iron deficiency than by normal subjects.

**Absorption of iron by subjects with excess iron stores (Figures 7 and 8).**

Sixteen absorption studies have been performed on 9 patients with the excess iron stores of idiopathic hemochromatosis. Seven of these patients were males and two\textsuperscript{7} were females past the menopause. The comparative absorption of ferrous chloride and Fe\textsuperscript{59} labeled eggs was evaluated in one patient with hemochromatosis. The absorption of iron before and after an intensive venesection program was measured in one patient with

\textsuperscript{7}We wish to thank Drs. Alexander Marble and Frank Gardner, Boston, Massachusetts, for the opportunity to study these patients.
hemochromatosis, and only after venesection in another.

It is evident from Figure 7 that patients with well-established, untreated idiopathic hemochromatosis of long duration, with elevated serum iron levels and saturated iron stores did not absorb either ferrous chloride or food iron to any greater extent than did normal subjects studied under similar conditions. Patients with hemochromatosis absorbed and incorporated into hemoglobin 0.5 to 11.1 per cent of Fe⁵⁹ administered as ferrous chloride. This represented the absorption of 0.1 to 3.3 mg. of the 24 to 41 mg. of carrier iron administered. Almost all of the remaining iron was recovered in the feces. Additional iron beyond that incorporated into hemoglobin may have been absorbed by the two female patients studied (Figure 7, Studies 7 and 8). The total quantity of iron possibly absorbed by these two female subjects was 28 and 29.7 per cent, or approximately 11 mg. of iron by each patient.

Absorption of food iron by untreated patients with hemochromatosis was evaluated in two patients who received Fe⁵⁹ labeled beet greens containing 2.8 and 4.8 mg. of iron (Figure 7). Only 1.0 and 1.1 per cent of iron was absorbed and incorporated into hemoglobin. This represents absorption of a quantity of food iron which is essentially the same as that obtained in normal subjects. If all the unrecovered iron was absorbed and deposited in tissue, the total absorption could be increased to 22 and 18.1 per cent, or 0.6 and 0.9 mg. of iron.

Absorption of Fe⁵⁹ labeled eggs was not evaluated in any untreated patient with hemochromatosis. However, such a study was completed in one subject with hemochromatosis after the removal of 25 liters of blood over an 18-month period (Figure 8). Multiple iron absorption studies were completed before and after the venesection program. Initially the serum iron ranged between 235 and 311 micrograms per 100 ml. When studied at this stage he absorbed and incorporated into hemoglobin 4.7 to 6.2 per cent of a 40-mg. dose of ferrous chloride. Ninety-four per cent of the dose was recovered in stools. After extensive venesections, during which period his hemoglobin remained at essentially normal levels and approximately 13 gm. of iron were removed from his stores, his serum iron was reduced to 60 micrograms per 100 ml. At this time he absorbed and used 20.2 per cent of 9.4 mg. of food iron, equivalent to the absorption of 1.9 mg. of iron. This result approximates closely that obtained in subjects with iron deficiency who received labeled eggs. In contrast, when Fe⁵⁹ was administered as ferrous chloride along with non-radioactive eggs, he absorbed and incorporated

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**Fig. 7. Recovery of Radioiron in Patients with Hemochromatosis Following Administration of Ferrous Chloride or Labeled Beets**

These patients had not been venesected. The symbol ♀ identifies a female patient.
into hemoglobin 80.4 per cent, or 6.7 mg. of an 8.3 mg. dose of iron. Subsequently, with a serum iron of 54 micrograms per 100 ml he absorbed and used 42 per cent or 15.2 mg. of a 40-mg. dose of ferrous chloride. The additional 30 per cent not recovered, if absorbed and added to tissues, would increase the total iron absorbed under these conditions to 29 mg. Without any additional venesections the serum iron gradually increased to 120 micrograms per 100 ml. At this time he absorbed and used 54.5 per cent, or 22 mg. of a 40-mg. dose. Most of the remaining iron was recovered in the feces except for 9 per cent, some of which may have been absorbed and deposited in tissue stores.

A second patient with hemochromatosis was studied only after extensive venesections. He received FeSO₄ without added carrier iron, and absorbed and utilized 85.5 per cent of the administered iron for the formation of hemoglobin. The remainder was recovered in stools.

In summary, neither beet iron nor the iron salt, ferrous chloride, is absorbed to any demonstrably greater extent by patients with well-established untreated hemochromatosis than by normal subjects. An intensive venesection program will significantly lower the serum iron and remove iron from the body stores. Under these circumstances, absorption of both ferrous chloride and egg iron is greatly increased, approximating that usually encountered in iron-deficient subjects.

DISCUSSION

Our studies indicate that normal subjects, iron-deficient patients, and patients with idiopathic hemochromatosis absorb ferrous chloride more readily than food iron. In many of these studies a solution of ferrous chloride was administered to the subject in a quantity that considerably exceeded the amount of iron usually eaten by an individual in a single average meal. Chemical or physical substances that might decrease iron absorption were excluded since the patients were fasting. In most cases a supplementary reducing agent such as ascorbic acid was given with the ferrous chloride. There is evidence that the administration of ascorbic acid may increase the absorption of ferrous iron salts (19). However, in the present study there was no apparent difference between the absorption of iron by 7 subjects who did not receive supplementary ascorbic acid and comparable subjects who received this reducing agent with ferrous chloride. No systematic comparison was made of the absorption of ferrous iron salts administered to the same patient with and without a reducing agent.

Iron in considerable excess of the daily loss of 1 mg. may be absorbed when ferrous chloride is administered to normal subjects under the described experimental conditions. These results agree with those reported by Dubach, Callender, and Moore (27). Within the limitations imposed by inaccuracies in stool collection our data may be interpreted as lending support to their suggestion that normal subjects may absorb and deposit in tissue stores additional iron beyond that which is incorporated into hemoglobin.

Normal subjects absorbed much less iron after a single feeding of radioiron labeled eggs or vegetables. These observations were obtained under physiological conditions more closely approximating the ingestion of an average meal containing 5 mg. of iron. Only 0.3 mg. of iron at best, or approximately one-third of the daily iron requirement was absorbed and incorporated into hemoglobin. This observation agrees with previously reported studies (26). Although fecal recovery data suggest that additional iron in food may possibly be absorbed and deposited in body stores, the limited absorption of food iron observed supports the suggestion (26) that the daily adult requirement of 12 to 15 mg. of dietary iron recommended by the National Research Council (32) may be barely sufficient to maintain the body iron stores.

The present studies indicate that patients with iron deficiency usually absorb significantly more food iron than normal subjects. However, the amount of iron absorbed from food was not much greater than the quantity lost daily from the body in the absence of bleeding or pregnancy. Moore and Dubach, on the other hand, found only a few iron-deficient subjects who absorbed more food

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8 This subject was studied through the cooperation of Dr. Wayne Rundles, Duke University School of Medicine.

9 These 7 subjects include 5 normal subjects, 1 iron-deficient patient, and 1 patient with idiopathic hemochromatosis.
iron than did normal individuals, although the co-administration of a reducing substance such as ascorbic acid significantly increased the quantity of iron absorbed (26). In our studies markedly limited absorption of food iron was observed in an iron-deficient patient with a subtotal gastrectomy. Iron as ferrous chloride was as readily absorbed by this patient as by the other iron-deficient patients. Partial gastrectomy may have influenced the absorption of food iron since there is some evidence that dietary iron is ionized, reduced to the ferrous form, and better absorbed at an acid pH (33–37).

The limited absorption of food iron in these studies suggests that it would be difficult for an iron-deficient patient to replenish his stores by diet alone. Supplementary medicinal iron seems indicated for individuals who undergo chronic blood loss, have increased physiological requirements, or have an absorptive defect due to an operative procedure, e.g., subtotal gastrectomy, or steatorrhea (7, 26, 38, 39). Therapy should be continued after the hemoglobin level has been restored to normal since it has been shown that depleted iron stores are not easily reconstituted (40).

A different situation is observed in hemochromatosis. Although our studies indicate that neither ferrous chloride nor food iron is absorbed to any greater extent in well-established hemochromatosis than in normal subjects, it appears necessary to assume that excess iron must be absorbed during the developmental phase of this disease. By no other means can one satisfactorily explain the huge iron stores found in a patient with idiopathic hemochromatosis. This assumption is perhaps supported by our observation that absorption of both ferrous chloride and food iron was markedly increased in patients with hemochromatosis after they had undergone an intensive venesection program. After 12 to 15 grams of iron had been removed by weekly or bi-weekly 500-ml. venesections, the absorption of iron was approximately that seen in iron deficiency. It is significant that this increased absorption was observed even after the serum iron concentration had returned to normal (120 micrograms per 100 ml.).

Studies showing increased radioiron absorption in younger patients with hemochromatosis have been reported recently which lend support to the concept that iron absorption must be increased during the developmental stage of idiopathic hemochromatosis (41–43). External measurements indicated most of the absorbed iron was stored in the liver (43, 44). Increased absorption of a lesser degree has also been observed in three older female patients (43). Of 9 patients with hemochromatosis in our studies, the two women were the only patients who appeared to absorb an increased quantity of iron. Menstrual loss of blood may have had an effect similar to that of repeated, small venesections, so that full development of the disorder was delayed. Our studies after venesection as well as those of Peterson and Ettinger (41) do show that extensive blood loss will modify the absorption of iron even after hemochromatosis has fully developed. Increased absorption may then occur even with a normal serum iron concentration. These results suggest iron stores may reaccumulate in idiopathic hemochromatosis unless venesections are continued at intervals as recommended by Finch and Finch (45).

Although the present study has demonstrated greater absorption of iron salts than of food iron, it does not provide a clear explanation why this occurs. Multiple studies in normal subjects evaluating the comparative absorption of labeled egg iron and of ferrous chloride alone and with non-labeled eggs suggest that the presence of egg will decrease the absorption of the iron salt. This decreased absorption may be related to the solid content of the test meal (46), or formation of an insoluble compound of iron with a chemical constituent of egg or bread. Hegsted, Finch, and Kinney (47) have shown that a high concentration of added phosphate will decrease absorption of iron by rats on a corn grit diet, presumably by formation of an insoluble iron phosphate. Similarly, soluble phytates may also interfere with iron absorption (46, 48). Conversely, rats on a corn grit diet with added iron will absorb large amounts of iron and produce progressive hemosiderosis of the tissues (49). Studies by Hegsted, Finch, and Kinney indicate that the low level of dietary phosphate attained with a corn grit diet was primarily responsible for increased absorption of iron (47). Such increased absorption of iron observed on a phosphate-deficient diet with excess iron might account for development of dietary hemosiderosis.
observed in malnourished pellagrins in South Africa (50, 51).

The influence of phosphates on iron absorption may explain the present observations that egg iron was poorly absorbed and that addition of egg decreased absorption of ferrous chloride. Halkett, Peters, and Ross found that egg yolk iron is in the ferric state and is strongly complexed to the phosphate of yolk phosphoproteins (52). Formation of such an iron phosphate complex occurs both in the biological production of eggs and when iron is added to eggs in vitro. They further observed that egg yolk iron is not removed by peptic digestion and acidity unless a reducing agent is present. It appears egg iron is not readily available for absorption and it is not surprising that so little was absorbed in the present study as well as in previous animal studies (53). In contrast it has been shown that a 10 to 20 fold increase in the absorption of food (egg) iron generally occurs in iron-deficient subjects when large amounts of ascorbic acid are administered with the iron (7, 26). This effect is presumably dependent upon the reduction of iron to the ferrous form and may be accomplished by other reducing substances in food.

It is of interest to consider how results of the present studies may relate to the theory that the intestinal mucosa is an important regulator of iron absorption (24, 54–57). According to Granick (56), iron is transferred from the intestinal lumen to blood by a protein, apoferitin, present in cells of the intestinal mucosa. Iron is taken up by the mucosal cells until all apoferitin is converted to ferritin. No more iron may then be absorbed until ferritin has given up iron to plasma. This is the concept of the "mucosal block" originally suggested by Hahn, Bale, Ross, Balfour, and Whipple (24).

Dubach, Callender, and Moore (27) have presented evidence that this block is at best a partial one and that in certain conditions, such as refractory anemia, pernicious anemia in relapse, or hemolytic anemia, the mucosal block does not prevent iron from being absorbed in spite of adequate body iron stores. We have observed a similar increase in iron absorption in thalassemia minor and in renal anemia (44). Moreover, it would appear that the block must fail significantly during the developmental stage of idiopathic hemochromatosis and again in this disease after an extensive course of phlebotomies. Other examples of the alteration of the mucosal block in animals and men have been discussed previously in relation to factors that influence absorption of iron.

Finally, our data in normal subjects, as well as those of others (27) suggest that the "mucosal block" does not prevent absorption of an increased quantity of iron salt when it is administered in a single feeding to a fasting subject under optimal conditions, i.e., as a solution of ferrous chloride with ascorbic acid, or when given in large quantities (4). Such excess iron absorption has also been observed after average doses of iron salts have been orally administered for a period of years (58, 59).

**SUMMARY AND CONCLUSIONS**

1. The iron salt, ferrous chloride, is absorbed far more readily and in greater quantity by normal subjects and by patients with deficient and excess iron stores than is iron present in certain foods (eggs, vegetables).

2. Egg and vegetable iron are not absorbed sufficiently to supply iron in the face of increased loss or increased physiological requirements.

3. Absorption of certain food iron and ferrous chloride in patients with well-established hemochromatosis is approximately equal to that observed in normal subjects. However, it can be assumed that excess quantities of iron must be absorbed during the developmental phase of this disease.

4. After removal of blood by multiple venesections absorption of ferrous chloride and egg iron by patients with idiopathic hemochromatosis is markedly increased.

5. Absorption of iron salts is significantly influenced by dietary factors which may modify the form and solubility of iron in the lumen of the gastrointestinal tract.

6. Further evidence has been presented to support previous data in the literature which indicated that the "mucosal block" to iron absorption is only relatively complete and may not uniformly prevent the excess accumulation of iron in the body.

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