Bioavailability of caseinophosphopeptide-bound iron
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Iron deficiency, one of the main worldwide nutritional deficiencies, results from the low bioavailability of most dietary iron, including cow milk. Hydrolysis of the cow milk protein casein produces low molecular weight caseinophosphopeptides (CPPs). Binding of iron to CPPs keeps it soluble in the digestive tract and prevents the formation of high molecular weight ferric hydroxides, which are poorly absorbed. Previous experimental studies have shown that iron bound to the phosphopeptide containing the first 25 amino acids of β-casein, or β-CN (1-25), is well absorbed and corrects efficiently iron deficiency. We sought to assess in vivo iron absorption and uptake by tissues involved in iron metabolism and storage (liver, spleen, bone marrow), using radiolabeled iron. β-CN (1-25)-Fe displayed better absorption and tissue uptake by the vascularized rat loop model compared with a control substance, ferric ascorbate. The metabolism of β-CN (1-25)-Fe labeled with iron 59, added to cow milk, was also studied in young women. Although the absorption of β-CN (1-25)-Fe was not significantly higher than that of ferrous sulfate, it displayed significantly higher tissue uptake. This increase was transient and had disappeared by the 14th day of the study, suggesting that iron was used for metabolic purposes.

Abbrevations: CN = casein; CPP: caseinophosphopeptide; β-CN (1-25)-Fe = first 25 amino acids of β-casein–bound iron; β-CN (1-25) = first 25 amino acids of β-casein
mination of radioactivity incorporated into circulating red blood cells, as well as tissue uptake, in young women.

METHODS

Preparation of \( \beta\)-CN (1-25)-bound iron. \( \beta\)-CN (1-25) (purity > 85\%) was purified from tryptic hydrolysate of \( \beta\)-CN as previously described.\(^8\)

We bound iron by mixing \( \beta\)-CN (1-25) with a \( \text{FeCl}_2 \) solution for 1 hour at 37\( ^\circ \)C and a pH of 6.5. The resulting solutions were then ultrafiltered and diafiltered on a 3-kD membrane to remove free minerals. The amount of iron complexed to phosphopeptides and the presence of traces of calcium and sodium were determined with the use of atomic-absorption spectrometry (Model AA 1275; Varian; Les Ulis, France) on freeze-dried samples. The final product had an iron/phosphopeptides molar ratio of 4 and contained less than 1 mg sodium and 0.1 mg calcium/g.

Labeling of iron. \( \gamma\)-Emitting \( ^{59}\)Fe was purchased from Amersham (Amersham Pharmacia Biotech, Orsay; specific activity 3.7 MBq/mL). Perfusion solutes were extrinsically labeled in the laboratory with \( \text{FeCl}_2 \), and mixed with cold ferric ascorbate at a ratio of 0.001\%, whereas the complex of \( \beta\)-CN (1-25)–bound iron was intrinsically labeled during its preparation.

For the human study, milk was labeled either with extrinsically tagged \( \text{FeSO}_4 \) with \( ^{59}\)Fe citrate or with the intrinsically labeled \( \beta\)-CN (1-25)–Fe.

Each 250-mL meal contained the same amount of radioactivity (74 kBq).

Experimental study. In an experimental study of perfused rat duodenal loop, four groups (each group \( n = 6 \)) of adult Sprague-Dawley rats were studied after being fasted overnight, as previously described.\(^9\) They were perfused with ferric ascorbate or \( \beta\)-CN (1-25)-Fe. We tagged iron with \( ^{59}\)Fe so that we might assess iron tissue uptake in addition to its absorption.

The composition of the perfusion solute was adapted from Ringer-Lavosier solute; its pH was adjusted to that of the proximal duodenum (5.5) and contained 100 \( \mu \)mol/L iron in either form.

Duodenum of anesthetized rats was perfused at a delivery rate of 0.16 mL/min; every element of the perfusion device had been washed with a solution of Triton X-100 (1 g/L) to prevent contamination. We kept the perfusion solute at 37\( ^\circ \)C with a thermostatic control and added a nonabsorbable marker (polyethylene glycol 4000) to assess actual net water flux. After 2 hours of perfusion, the animal was killed with an overdose of Dolethal; then the perfused loop was withdrawn and washed with saline solution.

Tissues were digested in nitric acid. The radioactivity of gut mucosa, blood, liver, and spleen were measured on a scintillation counter.

Human study. Our human study was approved by the local committee of ethics. Ten subjects gave their written informed consent to participate: All were 20- to 30-year-old female students of the university, and all stated that they were in good health, had no recent history of digestive or inflammatory disease, or infection; were taking no medication or mineral supplementation; were using a chemical method of contraception; and had negative results on pregnancy testing.

Before the administration of the first test meal, blood was drawn for measurement of red blood cell count, serum ferritin, C-reactive protein (a marker of recent inflammation or infection), and background radioactivity.

The test subjects drank radioisotopically labeled milk after an overnight fast; nothing but water was allowed for the next 4 hours. All subjects drank 250 mL of sterilized milk containing 3 mg iron (12 mg/L), either as \( \text{FeSO}_4 \) or \( \beta\)-CN (1-25)-Fe, labeled with \( ^{59}\)Fe. The two meals were given at random. The first was given on day 1; on day 14, blood was drawn to measure the increase in red blood cell radioactivity. External counting of liver, spleen, and blood marrow (sacrum) areas was performed on days 7 and 14 with the use of a solid scintillation counter equipped with a thick crystal for external isotope tissue measurement (CGR Nucléaire Médecine GM2C; Paris, France); counting lasted 10 minutes. The heart area was used as control, and background radioactivity was subtracted.\(^{11}\)

The second meal was given after a washout period of 2 weeks (day 28), after residual red blood cell and body radioactivity had been determined. External counting was performed on days 35 and 42; erythrocyte radioactivity was measured on day 42.

Percentage absorption of iron was calculated on the basis of blood volume estimated from weight and height\(^{12}\) and an assumed hemoglobin incorporation of absorbed iron of 80\%.\(^{13}\) In addition, data were corrected for background radioactivity and for the radioisotope decay of the residual radioactivity of tissues and red blood cells.

Statistical methods. We used Student’s \( t \) test to analyze data from the experimental study. Human data were subjected to analysis of variance and and Fisher’s exact test. \( P \) values of less than .05 were considered statistically significant.

RESULTS

Experimental study. The absorption of ferric ascorbate and \( \beta\)-CN (1-25)-Fe by isolated perfused rat duodenal loop system is shown in Table I. \( \beta\)-CN (1-25)-Fe displayed greater gut uptake and net absorption, greater spleen uptake, and increased blood radioactivity compared with ferric ascorbate.

Human study. Hematologic data and results of iron-absorption testing are given in Table II. Three subjects were iron-deficient (ferritin concentration < 12 g/L).

Mean iron absorption was similar in the two groups, yet paired comparisons showed a nonsignificant trend toward better absorption of \( \beta\)-CN (1-25)-Fe.

Results of external counting are given in Fig 1. A significant difference was displayed at day 7 between \( \text{FeSO}_4 \) and iron-labeled CPP for liver (\( P = 0.05 \)), spleen (\( P = 0.05 \)), and sacrum (\( P = 0.03 \)); the total radioactivity of these organs was different between groups (\( P = 0.01 \)) at day 7. No difference was observed at day 14.
DISCUSSION

Iron deficiency remains a worldwide health problem, mainly involving growing infants and women of childbearing age.\textsuperscript{14} In addition, the side effects of iron supplementation limit adherence to its prescription,\textsuperscript{15} necessitating a search for highly bioavailable sources of iron, free of digestive interactions.

CPPs are produced during digestion of CN; they bind divalent cations and keep them soluble at luminal pH.\textsuperscript{7,16,17} The strength of iron binding to CPP is about 100 times greater than that of calcium and others cations.\textsuperscript{3,18}

When assessed in rats on the basis of metabolic balance and direct measurement of tissue storage, the bioavailability of iron bound to $\beta$-CN (1-25) is higher than that of reference iron salts gluconate or ferrous sulfate.\textsuperscript{9,10}

In this study, the absorption rate and tissue uptake of Fe\textsuperscript{59} were similar to those noted in recent reports.\textsuperscript{19} Our findings showed that absorption, blood content, and spleen uptake of $\beta$-CN (1-25)–bound iron by duodenal rat loop during the experiment were better than that of inorganic iron. The better absorption and tissue uptake of $\beta$-CN (1-25) bound iron supports the results of tissue analysis after a 4-week repletion period.\textsuperscript{9,10}

Results varied between the rat and human groups. Differences in experimental methods could explain the isotopic data in the human group, in which iron-absorption rates were close to those obtained from studies that employed cow milk as source of iron.\textsuperscript{6,20-23} As expected,\textsuperscript{24} these results differed quantitatively from those of the rat group but displayed the same trends; some differences between the two experiments could explain the lower significance in the human trial. First, iron was given in milk and not in a pure solution as in the rat group. This decreases iron absorption and blunts differences between dietary sources of iron.\textsuperscript{14} Second, physiologic changes, such as the onset of menses, could have occurred during the time elapsed between the two tests in the human group. Use of $^{55}$Fe and $^{59}$Fe on consecutive days could have alleviated these potential variations, but the use of $\beta$-emitting $^{55}$Fe precludes external counting.

Although stable isotopes are valid tools in the assessment of iron absorption from food,\textsuperscript{25,26} they do not supply information on tissue kinetics, as radioisotopes

| Subject | Ferritin (g/L) | Hemoglobin (g/dL) | Iron absorption (%) | Ratio of $\beta$-CN (1-25)-Fe to FeSO$_4$ |
|---------|---------------|------------------|--------------------|----------------------------------------|
|         |               |                  | $\beta$-CN (1-25)-Fe | FeSO$_4$                              |                                       |
| 1       | 36            | 14.4             | 24.3               | 19.6                                  | 1.24                                 |
| 2       | 24            | 13.2             | 4.4                | 6.9                                   | 0.65                                 |
| 3       | 45            | 12.4             | 19.8               | 19.3                                  | 1.03                                 |
| 4       | 5             | 14.8             | 14.6               | 18.4                                  | 0.80                                 |
| 5       | 48            | 13.3             | 4.2                | 2.1                                   | 2.05                                 |
| 6       | 31            | 14.1             | 10.2               | 9.5                                   | 1.08                                 |
| 7       | 11            | 11.6             | 9.7                | 26.5                                  | 0.37                                 |
| 8       | 40            | 12               | 7.7                | 1.8                                   | 4.37                                 |
| 9       | 4             | 11.4             | 22.6               | 16.6                                  | 1.36                                 |
| 10      | 33            | 14.3             | 5.3                | 2.8                                   | 1.92                                 |
| Mean±SD | 27.7 ± 16.1   | 13.1 ± 0.4       | 9.8 ± 6.1          | 9.9 ± 7.1                             | 1.49 ± 0.36*                         |

*Wilcoxon’s test not significant.
do. Our results suggest that tissue uptake of β-CN (1-25)–bound iron was significantly enhanced; this assumption is supported by the findings of our experimental study of the rat group, in which tissues were assessed immediately after absorption. The transient increase of iron concerned every organ involved in iron metabolism (bone marrow, liver, and spleen). Yet the function of iron uptake remains unclear; it may merely reflect the difference of radioactivity in the blood because blood was not removed before counting; however, counting of the blood-filled myocardial tissue did not display significant differences. Stored iron was quickly used: no difference was found 1 week later.

However, these data suggest that measuring only iron radioactivity incorporated into circulating erythrocytes could miss some differences in iron bioavailability, as noted by Fomon in newborn infants27; it is not known whether the observed differences concern iron absorption or kinetics.

Our findings show that in experimental models, iron bound to β-CN (1-25) displayed better absorption and better tissue uptake than inorganic salts; similarly, the findings of external counting of organs was temporarily enhanced in human subjects after intake of β-CN (1-25)–bound iron.

These beneficial results may partly explain the high iron bioavailability of breast milk, which is rich in β-CN.

Further human studies are needed to precisely assess the bioavailability of β-CN (1-25)–bound iron, particularly the lack of interaction with other minerals previously shown in experimental studies.

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