The bioavailability of iron picolinate is comparable to iron sulfate when fortified into a complementary fruit yogurt: a stable iron isotope study in young women

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Abstract

Purpose A technological gap exists for the iron (Fe) fortification of difficult-to-fortify products, such as wet and acid food products containing polyphenols, with stable and bioavailable Fe. Fe picolinate, a novel food ingredient, was found to be stable over time in this type of matrix. The objective of this study was to measure the Fe bioavailability of Fe picolinate in a complementary fruit yogurt.

Methods The bioavailability of Fe picolinate was determined using stable iron isotopes in a double blind, randomized cross-over design in non-anemic Swiss women (n = 19; 25.1 ± 4.6 years). Fractional Fe absorption was measured from Fe picolinate (2.5 mg ⁵⁷Fe per serving in two servings given morning and afternoon) and from Fe sulfate (2.5 mg ⁵⁴Fe per serving in two servings given morning and afternoon) in a fortified dairy complementary food (i.e. yogurt containing fruits). Fe absorption was determined based on erythrocyte incorporation of isotopic labels 14 days after consumption of the last test meal.

Results Geometric mean (95% CI) fractional iron absorption from Fe picolinate and Fe sulfate were not significantly different: 5.2% (3.8–7.2%) and 5.3% (3.8–7.3%) (N.S.), respectively. Relative bioavailability of Fe picolinate versus Fe sulfate was 0.99 (0.85–1.15).

Conclusion Therefore, Fe picolinate is a promising compound for the fortification of difficult-to-fortify foods, to help meet Fe requirements of infants, young children and women of childbearing age.

Keywords Iron bioavailability · Stable isotopes · Iron picolinate · Complementary fruit yogurt · Women · Iron sulfate

Introduction

Iron deficiency is a major nutritional problem worldwide today. It affects hundreds of millions of people and is especially prevalent in infants, young children and women of child bearing age [1]. Although the highest prevalence of iron deficiency in young and school-aged children is found in developing countries [2], it is also reported in industrialized areas [3]. In children, iron deficiency can have adverse effects on cognition, decrease motor activity, social attention and school performance, and increase susceptibility to infection [4].

To overcome micronutrient malnutrition, the World Health Organization [5] proposes three approaches i.e. food diversification and education, supplementation and fortification. Food fortification with iron is generally regarded as the most cost effective and sustainable long-term approach for reducing the prevalence of iron deficiency. Iron compounds used for fortification have to be carefully selected considering bioavailability and reactivity upon formulation. Available iron compounds for food fortification are classified according to their solubility. In general, the water-soluble ones are bioavailable but often cause unacceptable sensory changes in the product. Conversely, the water insoluble compounds create less sensory problems but their bioavailability is generally lower [6]. Many food products can be fortified...
with available forms of iron, but there is still a technologi-
cal gap for the iron fortification of products having high
moisture, low pH, containing polyphenols and having a long
shelf life, such as shelf-stable yogurt containing fruits [7].
Iron picolinate, a novel food ingredient, is stable over time
in this type of product (internal unpublished data). Fe pico-
linate is made from the complexation of iron with picolinic
acid. Picolinic acid is an endogenous metabolite of trypto-
phan and an isomer of niacin (vitamin B3) and, therefore, a
normal constituent of the human body and diet. The use of
the parent compounds (i.e. chromium and zinc picolinate)
has already been evaluated by the European Food Safety
Authority (EFSA) as a source of minerals in food supple-
ments for adults and children [8]. EFSA concluded that there
was no safety concern for picolinate intakes ≤ 1.6 mg/kg
body weight per day (maximal estimated combined exposure
from chromium and zinc picolinate) as long as the upper
limits of chromium and zinc are not exceeded. Therefore,
no safety concerns are expected for its application in both
young children (i.e. from 6 up to 36 months) and in the
general population.

To our knowledge, the bioavailability of Fe picolinate has
not been evaluated in humans. While the target population of
the potential application is young children, Hurrell et al. and
Harrington et al. have previously demonstrated that results
obtained in adults on iron absorption can be extrapolated to
them [9, 10]. Thus, the objective of this work was to deter-
mine the iron bioavailability of iron picolinate in comparison
to iron sulfate from a complementary food (i.e. shelf-stable
yogurt containing fruits) using stable iron isotope tech-
niques in healthy young women. Our hypothesis was that
there would be no significant difference in iron absorption
from iron picolinate versus iron sulfate when fortified into
this food matrix.

Methodology and trials

Subjects

Twenty women were selected from an initial screening of
59 women among the staff of the Nestlé Research Center
and the Ecole Polytechnique of Lausanne. Subject inclu-
sion criteria were healthy women, aged between 18 and
40 years old, with a weight below 65 kg and a plasma ferritin
(PF) < 50 µg/L. Exclusion criteria were pregnancy or
lactation, major chronic diseases or food allergies, infection
in the 4 weeks before the study, smokers (> 5 cigarettes/
day), alcohol consumption above 2 units a day, significant
blood loss in the 6 months before the study and/or who had a
significant weight loss within the 3 months before the study
(10% and more). Intake of vitamin/mineral supplements in
the 3 weeks before the study was not allowed. The study was
conducted at the Metabolic Unit of the Clinical Develop-
ment Unit, Nestlé Research Center, Lausanne, Switzerland
according to the guidelines laid down in the Declaration
of Helsinki. The study protocol was approved by the Com-
mission for Ethical Research on Human Beings of Canton
de Vaud, Switzerland (approval no.: 276/15 dated 1 Sept
2015) and written informed consent was obtained from all
subjects. The study was registered at clinicalTrials.gov under
the reference NCT02585661 (https://clinicaltrials.gov/ct2/
show/NCT02585661).

Study design

A controlled randomized, double blind cross-over design
was used. One group (n = 10) started the study with the con-
sumption of 54Fe sulfate fortified shelf stable yogurt, while
the other group (n = 10) consumed 57Fe picolinate fortified
shelf stable yogurt. On day 1, the first Fe labelled shelf sta-
bile yogurt was administered to the fasting subjects twice at
8:00 am and 3 pm, and they received a standardized lunch at
noon. The following day, the second Fe labelled shelf stable
yogurt was administered according to the same procedure.
Apart from the standardized lunch, the diet was unrestricted
during the study. During the baseline screening, 3 weeks
before the labeled test meals were given, a venipuncture
blood sample was collected to determine clinical chemis-
try parameters, hemoglobin (Hb), plasma ferritin and CRP
(as an inflammation marker). Body weight and height were
measured and subjects were asked to complete a short ques-
tionnaire about dietary habits. On the day of the first stable
isotope administration (day 1) a venous blood sample was
drawn under fasting conditions to measure Hb to be used for
iron absorption calculation. A final venous blood sample was
drawn 14 days after intake of the last test meal. The blood
samples were sent to the ETH Zurich, Switzerland for the
measurement of stable isotope ratios and calculation of Fe
absorption.

Isotopic labels

After an overnight fast, subjects consumed either the intrin-
sically labeled 57ferrous picolinate or 54ferrous sulfate mixed
into the shelf-stable yogurt. Isotopically labeled 57Fe picolo-
nate and 54Fe sulfate were prepared by Dr. Paul Lohmann
GmbH (Emmerthal, Germany) and Innophos (NJ, USA),
respectively, from isotopically enriched elemental iron (Iso-
flex, CA, USA). Compounds were prepared using a down-
scaled procedure that follows closely the process employed
for the production of commercially available products. Isotopic enrichments of 57Fe picolinate and of 54Fe sulfate
were 94.5% and 99.6%, respectively. Iron concentration in
the labeled 57Fe picolinate and in the unlabeled compound
was determined after mineralization in a CEM Microwave
digested with an HNO\textsubscript{3}/H\textsubscript{2}O\textsubscript{2} digestion system according to official method EN 13805:2014 [11].

\textsuperscript{54}Fe sulfate and unlabeled commercial Fe sulfate dried were dissolved in 0.5 M HNO\textsubscript{3}. Fe in digests and dissolved samples was analyzed by ICP-AES according to the Association of Official Analytical Chemists (AOAC) official method 2011.14. Solubility of the \textsuperscript{57}Fe picolinate, \textsuperscript{54}Fe sulfate as well as of unlabeled reference compound was measured at pH 1.0 as proposed by Lynch et al. [12]. Aliquots of Fe compounds containing 20 mg Fe were weighed into 500-mL conical flasks, and 250 mL of 0.1 M HCl warmed to 37 °C, were added. Flasks were placed in a shaking water bath at 37 °C and gently shaken at a rate of 1 Hz for 30 min. Aliquots of 2 mL were taken and centrifuged for 2 min at 1000g. Iron was analyzed in supernatants by inductively coupled plasma atomic emission spectrometry (ICP-AES) (Varian Vista MPX, Varian AG, Mulgrave, Australia) as described above.

**Test meal and stable isotope administration**

Test meals consisted of a shelf stable yogurt (Jogolino® strawberry, Nestlé) specifically developed for children aged from 6 months up to 3 years old. The serving size was 100 g of product containing 3 g of protein, 3.4 g of fat, 15.7 g of carbohydrate, 3.5 g strawberry puree, 150 mg of calcium, fortified with 30 mg of magnesium and 1.5 mg of zinc as indicated on the label. Isotopically labeled Fe (2.50 mg per serving) were added in powder form to the yogurt 30–60 min before consumption by the subjects. Compounds were pre-weighed in individual portions into pre-weighed glass vials with Teflon caps at an accuracy of ±0.05 mg. Administered doses varied within ±5% (1SD) between individuals. The shelf-stable yogurt fortified with one labeled Fe compound was consumed with 200 mL low mineralized water containing less than 130 mg/L Total Dissolved Solid (TDS). Part of the water was used to rinse the yogurt container, i.e. three times with 15 mL of water, and subjects consumed the rinse water; they were not allowed to drink or eat for 3 h following the test meals. A standardized meal was provided at noon consisting of pasta (200 g) served with tomato sauce (30 g), a green salad (50 g) served with a balsamic sauce (30 g) and a sorbet (100 g), served with mineral water. Afterwards, subjects were asked to fast for 3 h. Then, they received a second serving of the product fortified with the same label as in the morning meal using identical procedures.

**Blood collection and analysis**

Blood samples were drawn by experienced nurses using EDTA-coated vacutainers (monovettes®, Sarstedt GmbH, Nümbrecht, Germany). Hb was measured using an automated Counter (ACT 5 diff counter, Beckman Coulter International S.A., Nyon, Switzerland). A control material of 3 levels (Coulter AC-T5 diff Control Plus) was analyzed with each set of measurements of Hb. Plasma was separated for ferritin and CRP measurements by centrifugation (Sorvall RC6 + centrifuge, ThermoFisher Scientific, Osterode, Germany). Ferritin and CRP were measured with the Siemens Dimension® Clinical Chemistry system using Flex® reagent cartridges. Ferritin was measured using the enzyme immunoassay method and CRP using the C-Reactive Protein Extended Range (RCRP) method based on a particle enhanced turbidimetric immunoassay (PETIA) technique. Two level serum control material (Liquid Assayed Multi-Qual Premium) were analyzed with each ferritin and CRP measurements.

**Isotopic analysis of the blood samples**

Each isotopically enriched blood sample was analyzed in duplicate for its isotopic composition. Whole blood was mineralized by microwave digestion, and Fe was separated by anion exchange chromatography and a subsequent precipitation step with ammonium hydroxide [13]. Fe isotope ratios were determined by an MC-ICP-MS instrument (Neptune; Thermo Finnigan).

**Calculation of iron absorption**

Fractional iron absorption was calculated based on the incorporation of enriched \textsuperscript{57}Fe and \textsuperscript{54}Fe from absorbed isotopically labeled Fe compounds into red blood cells. Amounts of \textsuperscript{57}Fe and \textsuperscript{54}Fe isotopic label present in the blood 14 days after test meal administration were calculated based on measured shifts in the iron isotope ratios in the blood samples compared to natural iron abundance (i.e. baseline) and the amount of iron circulating in the body. The natural iron abundance was borrowed from a historical control of comparable women at ETH Zurich. Calculations were based on isotope dilution principles as previously described [14]. Fractional absorption expressed as percentage of dose was calculated by dividing the amount of absorbed iron label by the administered dose multiplied by 100. Circulating iron was calculated based on blood volume and hemoglobin concentration [15]. Blood volume calculations were based on height and weight [16]. For the calculation of fractional iron absorption, 80% incorporation of the absorbed iron into red blood cells was assumed [17].

**Food analysis**

Fe and calcium content of the shelf-stable yogurt and the water were determined by ICP-AES after mineralization as described above.
Statistics

The sample size calculation was based on data from a previous trial [18]. The within subject standard deviation of fractional absorption and the relative bioavailability was approximately 2.5% and 25%, respectively (estimated by the mixed model). Therefore, with eighteen subjects, the fractional iron absorption and the relative bioavailability can be measured with approx. \( \frac{25\%}{18^{0.5}} = 6\% \) standard error which was considered as sufficiently precise. Anticipating a potential dropout of two subjects, the sample size was increased to 20. Fractional iron absorption was approximately log-normally distributed. The natural logarithm was used for the transformation. Descriptive statistics of fractional iron absorption from both compounds was displayed by geometric means and by the back transformed means ± standard deviations. Log-transformed fractional absorption was analyzed by a mixed model. Fixed-effect was compound (Fe-picolinate or Fe-sulfate), random-effect was subject. The model-based geometric mean is the exponent of the model based estimate of the predicted mean. The relative bioavailability is the exponent of the model-based treatment difference. To draw conclusions on bio-equivalence, criteria provided by FDA were used [19].

Results

Iron compounds and in vitro solubility

Iron contents of \( ^{54}\text{Fe} \) sulfate and \( ^{57}\text{Fe} \) picolinate were 28.4 and 19.3%, respectively. The experiments conducted at pH = 1 after 30 min showed similar solubility between labelled and unlabeled compounds i.e. 94.9 ± 1.4% for \( ^{54}\text{Fe} \) sulfate, 96.9 ± 6.6% for commercial Fe sulfate, 97.8 ± 4.6% for \( ^{57}\text{Fe} \) picolinate and 90.7 ± 0.6% for unlabeled Fe picolinate.

Subject characteristics

Age, anthropometric and baseline characteristics for Hb and plasma ferritin for the twenty enrolled subjects are shown in Table 1. None of the subjects were anemic. Only three subjects had a plasma ferritin concentration below or at borderline of the cut-off value of 15 µg/L that defines iron deficiencies [5]. The mean (± SD) serum ferritin of the subjects was 29.9 ± 9.8 µg/L. This represents adequate but still low iron stores. At screening, all subjects had a CRP value < 4 mg/L. Plasma ferritin and Hb were not different at screening and on the first day of stable isotope administration. The analysis of the questionnaire about dietary habits revealed that none of the subjects was vegan or vegetarian. Nineteen subjects completed the study. One subject had an accident during the study (i.e. twisted ankle) not related to the study product or study procedures and did not provide the last blood sample.

| Characteristics | Summary value (n = 20) |
|-----------------|-----------------------|
|                 | Mean  | SD   | Min–Max   |
| Age (years)     | 25.1  | 4.6  | 20–30     |
| Weight (kg)     | 55.8  | 4.4  | 49–62     |
| Height (cm)     | 164.8 | 4.4  | 157–174   |
| BMI (kg/m\(^2\))| 20.5  | 1.4  | 18.7–23.7 |
| Hb (g/L)        | 132.3 | 7.3  | 120.0–147.0|
| PF (µg/L)       | 29.9  | 9.8  | 10.8–50.9 |

Hemoglobin and plasma ferritin were measured on the morning of the day of the first stable isotope administration.

PF plasma ferritin

Food analysis

The results of mineral analysis showed that the quantity of iron and calcium brought by the shelf-stable yogurt was < 0.05 and 159 ± 1.9 mg/serving, respectively. The quantity of iron and calcium present in the mineral water consumed on the test day was below the limit of detection and 1.2 ± 0.1 mg/serving, respectively.

Fe absorption

The geometric mean fractional iron absorption from the \( ^{54}\text{Fe} \) sulfate and \( ^{57}\text{Fe} \) picolinate are presented in Fig. 1. The geometric mean fractional iron absorption from the shelf stable strawberry yogurt fortified with \( ^{57}\text{Fe} \) picolinate and \( ^{54}\text{Fe} \) sulfate was 5.2% (95% CI 3.8–7.2%) and 5.3% (95% CI 3.8–7.3%). The relative iron absorption (RBV) from Fe picolinate in this study was 99% (95% CI 85.2–115.0) of \( ^{54}\text{Fe} \) sulfate. This value is within the boundaries for bio-equivalence according to FDA: (90% CI 0.80 and 0.125) [19].

In the mixed models, a time effect (day 1 versus day 2) and the effect of plasma ferritin on iron absorption was investigated. The time effect was significant \((p = 0.04)\) and the point estimate was 0.84 indicating the fractional absorption on day 2 was 0.84 times less than on day 1. Absorption values and RBV corrected for this time effect were 5.2% (3.8–7.1%), 5.4% (3.8–7.3%) and 96% (84–110.5%) for \( ^{57}\text{Fe} \) picolinate, \( ^{54}\text{Fe} \) sulfate and the RBV, respectively, and were not significantly different than the time-effect unadjusted values. Log-plasma ferritin was also significant \((p = 0.03)\) (Fig. 2) and the point estimate was 0.418 suggesting that as plasma ferritin increases from 10 to 27 µg/L, fractional
iron absorption falls from 7 to 2.9%. Individual iron absorption values were corrected to a serum ferritin concentration of 30 μg/L according to Cook [20], to allow comparison to published data. This led to a geometric mean (95% CI) fractional absorption from Fe sulfate and Fe picolinate of 5.0% (3.67–6.37) and 4.8% (3.67–6.37) (N.S.), respectively.

Discussion

This study represents the first published evaluation of iron absorption from iron picolinate in humans. When added to a shelf-stable yogurt containing fruits, iron absorption from iron picolinate was not significantly different than iron absorption from iron sulfate. This observation is in agreement with findings from the in vitro Caco-2 cell model coupled with simulated digestion (internal, unpublished data). Other picolinate salts (zinc picolinate or chromium picolinate) have been shown to be well absorbed in humans or in rats. The comparative oral absorption of zinc picolinate and zinc gluconate, an organic salt having similar bioavailability than zinc sulfate, was studied in healthy human volunteers. At the end of 4 weeks supplementation periods hair, urine and erythrocyte zinc levels was found significantly increased compared to the placebo treatment, without significant difference between zinc picolinate and gluconate [21]. The oral bioavailability of chromium from picolinate and chromium chloride was evaluated in rats using radiolabelling. The absorption of chromium picolinate was twice as high as from chromium chloride. However, 1–3 days after administration, the relative distribution of 51Cr from both compounds was similar in all tissues, indicating that both compounds con-
reported fractional iron absorption in the range of 3.7–7.8% from a breakfast fortified with 2.8–4.2 mg iron and consisting of coffee or tea, wheat rolls made from 40 g unfortified white wheat flour, 10 g margarine and either orange marmalade or 15 g cheese [27]. Nevertheless, the fractional iron absorption from ferrous sulfate in our study is significantly lower than from fresh cheese consumed alone (23.4%) (iron absorption adjusted to a serum ferritin of 30 µg/L) [28]. A part of the explanation could be the higher concentration of calcium in the shelf stable yogurt (i.e. 156 mg per serving) when compared to the fresh cheese (i.e. 70 mg per serving). Calcium inhibits iron absorption in single meal studies; however, calcium is considered a low-level inhibitor when compared to phytic acid and phenolic compounds [29]. Despite the relatively high calcium content in iron-fortified milk products, they have been shown to improve iron status in efficacy studies [30–34]. The phytate and polyphenol content in the shelf stable yogurt containing 3.5% strawberries used in our study was not measured. Although the phytate content of strawberries is negligible, the concentration of polyphenols in strawberries puree is variable and may decrease during processing (http://phenol-explorer.eu/food-processing/foods). Thus, polyphenols content of the yogurt would be relatively low compared to doses reported to impair iron absorption [35–40]. It should be noted that not only the polyphenol concentration but also the type of polyphenols present in a food matrix can affect iron absorption, and this has not been widely investigated in humans. In addition, although in single meal studies polyphenols decrease iron absorption, longer term intake studies show less of an inhibiting effect of polyphenols on Fe absorption [41]. Alternatively, the low rate of iron absorption from the yogurt might be due to its protein content. Few studies have shown the impact of casein and whey on iron absorption [42, 43], but caution might need to be used when considering these results as they were generated with semisynthetic meals and low number of subjects. Finally, it is important to remind that the present study has been performed without the addition of ascorbic acid in the yogurt. Ascorbic acid enhances iron absorption in a dose-dependent manner and is thought to exert its enhancing effect by reducing ferric to ferrous iron and by binding iron in a soluble form available for absorption [44]. Depending on the molar ratio of ascorbic acid to iron it can raise iron absorption from iron fortified milk-based food by two- to fourfold [29, 44, 45].

Young children and toddlers are particularly vulnerable to iron deficiency due to their high iron requirements to meet the needs for rapid growth and development [46]. This is particularly true after 6 months of age, due to the decreased iron liver stocks constituted during the gestational phase. Thus, when time of food diversification arrives, appropriate iron-rich food need to be introduced in their diet [47]. Many young children do not consume large quantities of foods rich in iron, and the iron-rich food need to be introduced in their diet [47].
in bioavailable iron, such as red meat. In addition, theoretical dietary models show that it is difficult to reach the recommended intakes of iron with a diet following available food guides for infants and young children [48]. Iron fortification of complementary food can be an effective mean to reduce the risk of iron deficiency and iron deficiency anemia [29]. Nevertheless, iron added in this type of product must be provided at adequate quantity and must be bioavailable. Our study was performed in young women with low iron stores, and suggests that iron picolinate is as bioavailable as iron sulfate from a complementary yogurt. A higher fractional absorption of iron picolinate would be likely if co-ingested with ascorbic acid by iron deficient children. In conclusion, iron picolinate could be a promising compound for the fortification of difficult-to-fortify foods, to help meet Fe requirements of infants, young children and women of childbearing age.

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Author contributions The authors’ responsibilities were as follows: MS, DG, MB, PK, DM, CZ and MBZ: designed the study; KG, LFG and MB: conducted the study; LFG: analyzed biochemical indicators; JR, SD, PK and MS: performed the solubility test, contributed to the preparation of the labeled salts and, to the meal and stable isotope analysis; CZ, DM and MBZ: performed the stable isotopes ratio analysis and calculation of absorption and relative bioavailability; EH proposed and developed iron picolinate as iron fortificant for dairy and suggested to Innophos its production; DG: performed statistical analyses; MS, DG, MB and MBZ wrote the first draft of the manuscript; all authors: read and edited the manuscript.

Compliance with ethical standards

Conflicts of interest DM, CZ and MBZ declared no conflicts of interest. MS, DG, MB, LFG, KG, PK, SD, JR, and EH are employed at the Nestlé Research Center.

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