Iron deficiency intravenous substitution in a Swiss academic primary care division: analysis of practices

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Background: Iron deficiency is a common problem in primary care and is usually treated with oral iron substitution. With the recent simplification of intravenous (IV) iron administration (ferric carboxymaltose) and its approval in many countries for iron deficiency, physicians may be inclined to overutilize it as a first-line substitution.

Objective: The aim of this study was to evaluate iron deficiency management and substitution practices in an academic primary care division 5 years after ferric carboxymaltose was approved for treatment of iron deficiency in Switzerland.

Methods: All patients treated for iron deficiency during March and April 2012 at the Geneva University Division of Primary Care were identified. Their medical files were analyzed for information, including initial ferritin value, reasons for the investigation of iron levels, suspected etiology, type of treatment initiated, and clinical and biological follow-up. Findings were assessed using an algorithm for iron deficiency management based on a literature review.

Results: Out of 1,671 patients, 93 were treated for iron deficiency. Median patients’ age was 40 years and 92.5% (n=86) were female. The average ferritin value was 17.2 μg/L (standard deviation 13.3 μg/L). The reasons for the investigation of iron levels were documented in 82% and the suspected etiology for iron deficiency was reported in 67%. Seventy percent of the patients received oral treatment, 14% IV treatment, and 16% both. The reasons for IV treatment as first- and second-line treatment were reported in 57% and 95%, respectively. Clinical and biological follow-up was planned in less than two-thirds of the cases.

Conclusion: There was no clear overutilization of IV iron substitution. However, several steps of the iron deficiency management were not optimally documented, suggesting shortcuts in clinical reasoning.

Keywords: iron deficiency, iron substitution, general practice, quality of care, cross-sectional design

Introduction

Anemia is a global public health problem affecting both developing and developed countries. Overall, 24.8% of females and preschool children around the world suffer from anemia, and iron deficiency is considered to be among the most important contributing factors.¹ Iron deficiency without anemia affects 10%–30% of menstruating females in Europe and the US.²,³ Although prevalence of iron deficiency depends on age, sex, and socioeconomic status,⁴ it is a common problem in primary care and is an underrecognized cause of fatigue and other symptoms in females of childbearing age.⁵

Oral iron administration is the first-line therapy to manage the symptoms of iron deficiency and increase the ferritin value. However, oral administration presents
disadvantages, such as low absorption of iron and the high incidence of gastrointestinal side effects. In addition, it should be used with caution in patients with advanced renal failure or chronic heart failure because of decreased gastrointestinal absorption and increased iron loss, and is not indicated during flare-up of inflammatory bowel diseases.\(^6\)\(^-\)\(^9\) As a result, intravenous (IV) iron formulations have been approved in over 50 countries, including Switzerland, to correct iron deficiency among primary care patients. IV formulations may be indicated, for example, if oral therapy does not bring the expected results despite good compliance or is not tolerated by the patient.\(^10\)\(^-\)\(^14\)

Increased use of IV iron has been reported in Switzerland\(^15\) despite the lack of large-scale, good quality studies examining the effectiveness of oral and/or IV treatment for iron deficiency without anemia.\(^16\) Most studies assessing the effectiveness of IV versus oral iron treatment were conducted in specific populations, such as patients with chronic renal insufficiency, inflammatory diseases, neoplastic diseases, during or after pregnancy, but not among primary care patients.\(^17\)\(^-\)\(^21\) In addition, hypersensitivity reactions after iron infusion are not negligible and can be life threatening. Transient decreases in serum phosphate below the normal range are frequent. Although these are generally asymptomatic, the long-term consequences of repeated iron infusions on bone metabolism are not known.\(^22\)

Several countries have developed initiatives to promote the appropriate and efficient use of the relevant tests and procedures.\(^23\)\(^-\)\(^26\) Since it is the mission of training institutions to ensure both the most rational use of medical resources and a high quality of patient care, we decided to conduct a quality improvement project on the use of oral and IV iron substitution in a division of primary care. The aim of the project was to evaluate whether iron deficiency has been managed appropriately in clinical practice 5 years after the commercialization of ferric carboxymaltose in Switzerland.

**Methods**

**Study design and setting**

A retrospective observational study was conducted in the Division of Primary Care at the Geneva University Hospitals, Switzerland. The mission of this division includes delivering primary medical care to patients, providing pre-and postgraduate training in primary care, and conducting research in community-based clinics in Geneva, which is a canton of 450,000 inhabitants. The division is spread out over seven geographically distinct medical sites over the city but most clinical activities are conducted in the general outpatient clinic at the University Hospital. Each year, the clinic provides 13,000 scheduled medical consultations. It also trains 22 medical residents in ambulatory care. The patient catchment includes the general population of Geneva, particularly the socioeconomically deprived sector, including many vulnerable legal and illegal immigrants.

**Selection of study subjects**

All patients treated for iron deficiency in this clinic in March and April 2012 were included, excepting those with a hemoglobin level inferior to 80 g/L or a renal clearance inferior to 30 mL/min. According to our laboratory values, iron deficiency anemia was defined as a hemoglobin level of <120 g/L for females and <130 g/L for males, with a ferritin level of <100 μg/L. Ferritin reference values for iron deficiency vary according to sex, age, and the presence of several chronic conditions (12 and 200 μg/L) and a recent review suggested to consider serum ferritin <100 μg/L as an indicator of iron deficiency for most conditions.\(^3\) However, it is usually agreed that serum ferritin <10–15 μg/L represents exhausted reserves and that ferritin <30 μg/L indicates iron deficiency for patients without coexistent inflammatory disease.\(^27\)

**Measurement and outcomes**

Subjects were identified by reviewing all computerized medical files of patients who had visited the general outpatient clinic during March and April 2012 and had a documented iron supplementation prescribed. Nurses’ records were also reviewed for identification of patients who had received IV iron. A team of two residents and three chief residents developed a grid to analyze the relevant medical files and collect the following data: sociodemographic characteristics; iron, ferritin, and C-reactive protein levels; documented rationale for measuring iron levels (or presumed rationales according to symptoms or risk factors documented); suspected underlying etiology of iron deficiency (or presumed etiology according to the complementary investigations that were requested); treatment schedule; rationale for IV treatment; and, finally, the planned clinical and biological follow-up. This chart was first tested on ten medical files to evaluate the clarity and consistency of the topics and refine the rules of coding accordingly.

We reviewed the literature on iron deficiency detection and management and developed an algorithm based on results of scientific studies and national recommendations.\(^2\)\(^,\)\(^3\)\(^,\)\(^13\)\(^,\)\(^20\)\(^-\)\(^32\) At the end of our study, we distributed this algorithm into our Division of Primary Care medicine in order to optimize our medical practices (Figure 1).
Iron deficiency and substitution practices

Indication for blood transfusion?
- Hb <70 g/L
- Hb <80 g/L if >65 years, CVD, pulmonary disease
- Hb <90 g/L if active CVD

Indication for intravenous treatment?<sup>10,30,32</sup>
- CRD (CrCl <30 mL/min)
- Flare-up of inflammatory bowel disease
- Malabsorption syndrome (including bypass)

Oral iron treatment:
- Optimal administration (3 months, mornings with vitamin C)
- Clinical follow-up
- Biological follow-up 15 days after end of oral treatment

Blood transfusion (1 BU = 250 mg iron, increases Hb 10 g/L)
+ iron treatment

IV iron treatment:
- Presence of iron deficiency without anemia: iron reserve replacement (=500 mg if weight >35 kg) or 1,000 mg if ferritin <10 µg/L
- Presence of anemia: Body weight (kg) × (Hb goal – Hb measured) (g/dL) × 2.4 + iron reserve replacement (=500 mg if weight >35 kg)<sup>31</sup>
- Biological follow-up 8 weeks after end of IV treatment

Failure?
- Compliance (duration, vitamin C)
- Treatment of underlying pathology
- Etiological diagnostic differential

Reevaluate:
- Clinical and biological follow-up if etiology persists.

Repeat oral treatment if:
- Prescription related problem
- Persistence of mild iron deficiency or anemia
- Mild side effects (change of product)

Second-line IV treatment if:<sup>10–12</sup>
- Persistence or worsening of anemia despite appropriate treatment
- Severe side effects with oral treatment

Figure 1 Algorithm for iron deficiency management.

Note: Developed in collaboration with Dr Favrat (Policlinique médicale universitaire, CHUV) and Dr Sami (Service of Hematology, HUG).

Abbreviations: Hb, hemoglobin; Ht, hematocrit; MCV, mean corpuscular volume; MCHC, mean corpuscular hemoglobin concentration; CRP, C-reactive protein; creat, creatinine; ALAT, aspartate transaminase; CVD, cardiovascular disease; BU, blood unit; CRD, chronic renal disease; IV, intravenous.
The approval of the Research Ethics Committee of the Geneva University Hospitals and patient consent was not sought because the study was deemed an improvement activity and not a human subject research.

Statistical analysis
Participants’ sociodemographic data, ferritin levels and documented rationale for measuring iron levels, suspected etiologies, treatment schedule, reasons for choosing IV treatment, and planned clinical and biological follow-up were analyzed descriptively using mean and/or percentages. SPSS software Version 18 (SPSS Inc., Chicago, IL, USA) was used for the analysis.

Results
Out of 1,671 patients seen in March and April 2012, 97 patients were initially identified. Four patients met the exclusion criteria and so 93 patients were included in the study: 86 females and seven males. Median age was 40.2 years (standard deviation 11.8 years). In all, 81.7% (n=81) of the prescribers were residents and 12.9% (n=12) senior residents or attending physicians. Ferritin was measured in 97% (n=90) of patients and its mean value was 17.2 μg/L (standard deviation 13.3 μg/L).

The rationale for measuring iron levels was documented in 81.7% (n=76) medical files. The rationale for measuring C-reactive protein levels was documented in 30.1% (n=28) of medical files. The main reasons included presence of symptoms associated with iron deficiency, such as fatigue, or presence of risk factors for iron deficiency (mainly gynecological blood loss) (Table 1). The suspected etiology for iron deficiency was documented in 66.7% (n=62) of all cases. Gynecological and digestive blood losses were the main suspected causes. Malabsorption or low iron intakes were mentioned to a lesser extent (Table 1).

The majority of patients received an oral treatment (Table 2). Out of 28 patients treated with IV iron, only seven received it as first-line and 21 as second-line treatment. The rationale for using IV iron as first-line treatment was documented in half of the cases: malabsorption (one), noncompliance (one), very low ferritin (7 μg/L) (one), and low hemoglobin (86 g/L) (one). When IV iron was used as a second-line treatment, the rationale was mentioned in 95% (n=20) of the cases: persistence of iron deficiency despite oral treatment correctly taken (n=14; 70%) and/or adverse effects of oral treatment (n=7; 35%).

Clinical and biological follow-up at 3 months was planned in more than two-thirds of the patients but not all took place (Table 3).

Discussion
The study shows that there was no clear overutilization of IV iron substitution in an academic division of primary care medicine 5 years after the approval of ferric carboxymaltose for treatment of iron deficiency in Switzerland. Residents generally prescribed oral iron in first-line treatment of iron deficiency. However, several steps of the iron deficiency management were not optimally documented.
Residents did not systematically note the rationale for checking iron levels. Iron deficiency screening has not been proven to be useful as a routine procedure. Iron status should generally be requested in the presence of suggestive symptoms, such as fatigue, faintness, weakness, hair loss, loss of concentration, reduced physical endurance, and restless leg syndrome in females of childbearing age. It should also be determined in the presence of risk factors of iron deficiency, such as low dietary iron (vegetarian diet, socioeconomic difficulty), particularly during growth and pregnancy; diseases that impair iron absorption; gastrointestinal loss (chronic inflammatory disease, celiac disease, and cancer); gynecological or urological loss; renal or cardiac insufficiency; regular blood donations; and recent surgery.

Documentation of the suspected etiology was made in approximately two-thirds of medical files. The main risk factor for iron deficiency is low intake when iron requirements are especially high. However, more chronic and severe illnesses, such as gastrointestinal blood loss from an ulcer, carcinoma, or inflammatory bowel disease, or a uterine fibroma, for example, can be misdiagnosed and thus lead to poor therapeutic management.

According to Swiss recommendations, there was no clear overutilization of IV iron substitution as a first-line treatment. The majority of IV iron substitution was prescribed as a second-line treatment after a documented oral treatment failure or intolerance. However, the indication of first-line IV therapy was documented in only half of cases, suggesting that the rationale of IV treatment may not always be based on evidence. It is noted that a few studies show that IV formulations are more effective than oral forms in presence of iron deficiency from both biological and clinical perspectives (better physical or cognitive performances, decreased symptoms, decreased need for transfusions or erythropoietin). However, oral iron substitution is still considered to be an economic and effective way to treat iron deficiency if it is taken appropriately over several months and early treatment is implemented before total exhaustion of reserves.

Finally, 70% of second-line IV therapy was prescribed for persistent iron deficiency attributed to the ineffectiveness or intolerance of oral treatment. Given the length of oral treatment and frequency of side effects, the persistence of an iron deficiency could be due to poor compliance not disclosed by the patient or a suboptimal oral intake (eg, not taken while fasting or combined with ascorbic acid). In case of intolerance, a change of formulation of oral iron substitution or time of dosing may be an appropriate strategy before considering IV treatment.

Physicians may also be put under pressure by patients’ demands for a quick IV treatment. As with the use of IV or intramuscular analgesia in case of acute pain, it may be easier to meet the demands of patients to have an IV treatment, rather than to persuade them that it is not needed or does not bring any additional benefit. Technological progress brings more and more opportunities for interventions which quickly become “state of the art” without any evidence of true superiority.

Limitations
There are several limitations to this study. First, it is descriptive and retrospective, not an interventional study. Therefore, we do not know whether the guidelines created and distributed after this work improved residents’ documentation and management practices of iron deficiency. Second, we based our analysis on a review of medical files and therefore relied on documented data to assess whether clinical reasoning and practice patterns were correct. There were indeed several missing data on oral treatment noncompliance or rationale for second-line treatment. It is possible that some residents may have had correct clinical reasoning regarding measuring iron levels and subsequent management but omitted to document them sufficiently. Finally, because the study took place only in one clinical setting, our results may not be applicable to other outpatient settings. They may not be relevant for countries where IV iron supplementation for iron deficiency without anemia is neither common nor recommended, for example.

Conclusion
This study shows that in a country where IV iron substitution is recommended as a second-line treatment for iron deficiency without anemia, there was no clear overuse of it in an academic primary care division. However, several steps of iron deficiency management were not appropriately documented, suggesting lack of knowledge or clinical reasoning shortcuts that may lead to misdiagnoses of chronic diseases. More training is required in this field.

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Disclosure

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