A prospective study on the efficacy and safety of ferric carboxymaltose in correcting anaemia in patients with heavy menstrual bleeding

Abstract

**Background:** More than 50% of women in the developing countries are affected with iron deficiency anaemia. Ongoing blood loss in women with heavy menstrual bleeding can pose a difficulty to combat anaemia in these patients. Ferric carboxymaltose (FCM) is a novel molecule which can be safely administered at large doses in a very short time.

**Objective:** This prospective study was designed to study the efficacy and safety of ferric carboxymaltose in treating anaemia in women with heavy menstrual bleeding.

**Material and method:** This hospital based prospective study included 65 anaemic (Hb<10 gm%) women with heavy menstrual bleeding. Diagnosis of iron deficiency anaemia was established by peripheral blood smear and serum ferritin level. About 1000mg FCM was given by infusion in 15 minutes and improvement in haemoglobin and serum ferritin was assessed 14 days after the infusion. Any side effects were recorded during the study. Results were statistically analysed by paired t test.

**Result:** The mean haemoglobin before and after administration of FCM infusion were 7.71±0.66 gm% and 10.33±1.21 gm% respectively. The mean difference in haemoglobin was 2.62 gm%. (p < 0.0001). The mean ferritin level before and after FCM infusion were 26.77±22.0ng/ml and 254.75±70.00 ng/ml respectively. Mean difference in ferritin was 227.97ng/ml (p <0.0001). No major adverse reactions were seen.

**Conclusion:** This study demonstrated the safety and efficacy of FCM in treating anaemia in patients with heavy menstrual bleeding and should be considered in such women to avoid the possible need of blood transfusion.

**Keywords:** ferric carboxy maltose, iron deficiency anaemia, efficacy, heavy menstrual bleeding

Introduction

According to WHO reports, anaemia is one of the most common and wide spread disorders in the world affecting about 2 billion people and resulting in an estimated global prevalence of about 25%. In developing countries, anaemia affects more than half of school children and women and this incidence is at least 30-40% in developed world. 

Among the several causes of anaemia, iron deficiency is undoubtedly the most common cause and oral iron is the first mode of therapy with iron deficiency anaemia (IDA) but its ability to replenish iron stores is limited by patient compliance problems due to gastrointestinal side effects. Treatment with parenteral iron could present some advantage over oral iron by leading to a rapid and higher increase of hemoglobin levels and body iron stores.

Ferric carboxymaltose (FCM) is a new intravenous iron formulation. It is a polynuclear iron (III)-hydroxide carbohydrate complex designed to mimic physiologic ferritin. It is a water-soluble compound with molecular weight of 150000 Dalton containing approximately 1000 iron atoms which correspond to an iron content of 24-32%. FCM complex has neutral pH (5-7) and has physiologic osmolarity. FCM does not contain dextran or modified dextran and does not react with dextran antibodies thus having a very low immunogenic potential and therefore does not predispose to anaphylactic reaction. FCM has a ferric hydroxide core stabilized by a carbohydrate shell. There is controlled delivery of iron to the cells of reticuloendothelial system and subsequent delivery to iron binding protein with minimal risk of large amount of ionic iron being released into the circulation. These properties permit the administration of iron in large doses (15 mg/kg, maximum of 1000mg/infusion) during a single and rapid session (15 minute infusion) without the requirement of test dose.4,5

The management of iron deficiency anaemia includes the treatment of underlying cause and replacement of body iron stores. However, in patients with heavy menstrual bleeding, treatment of anaemia can be difficult due to ongoing blood loss. These patients often require preoperative or postoperative blood transfusion. Intravenous iron preparation can be given to replace iron and replenish iron stores rapidly. To date, few clinical studies showed the efficacy of FCM in correcting anaemia in patients with heavy menstrual bleeding.4,9 There is no study which investigates the efficacy and safety of FCM in North eastern population of India. This study was designed to see the efficacy and safety of FCM in correcting iron deficiency anaemia of Indian women with heavy menstrual bleeding.
Materials and methods

The study was a prospective study done over a period of two years at the obstetrics and gynaecology department of a tertiary health centre in the northeastern India. Our primary objective was to evaluate the efficacy of FCM in correcting iron deficiency anaemia in women with heavy menstrual bleeding. Our secondary objective was to specify the safety profile of FCM treatment. Approval was taken from the institutional ethical committee. All patients with gynaecological disorder and haemoglobin less than 10 g/dl, which provided informed consent and agreed upon complete follow up, were included in this study. The exclusion criteria were anaemias unrelated with iron deficiency, receipt of blood transfusion, and known allergy to parenteral iron therapy. Complete haemogram, serum ferritin and peripheral blood smear for cell morphology were done before administering ferric carboxymaltose by infusion and the same parameters except peripheral blood smear were repeated after two weeks. A total of 65 patients were given 1000mg ferric carboxymaltose (Orofer FCM, Emcure Pharma) in 250ml of normal saline over 15 minutes. Adverse effects like nausea, vomiting, hypotension, headache, fever, pain, tingling sensation or itching at the injection site during the infusion of ferric carboxymaltose were noted.

Statistical analysis

Collected data were expressed as mean ± standard deviation using descriptive statistics. Paired samples t test was performed to compare the mean values determined before and after transfusion. All p values less than 0.05 within 95% confidence interval and at 5% level of significance were considered to be statistically significant.

Results

This study included a total of 65 patients with various gynaecological disorders and haemoglobin less than 10 g/dl. The baseline demographic profiles of the patients are shown in Table 1. The mean age and parity of the study population were 38.96±9.77 years and 2.35±1.75 respectively. The underlying gynaecological disorders associated with anaemia are enlisted in Table 2. The mean haemoglobin before and after administration of FCM were 7.71±0.66 g/dl and 10.33±1.21 g/dl respectively (p < 0.0001). The mean difference in haemoglobin was 2.62 g/dl. The mean ferritin level before and after FCM infusion were 26.77±22.08 ng/ml and 254.75±70.00 ng/ml respectively (Table 3). The mean increase in serum ferritin level after FCM infusion was statistically significant (227.97 ng/ml, p < 0.0001). No major adverse reactions were observed except pain at the site of injection in three cases, nausea in two cases and headache in one patient (Table 4).

Table 1 Demographic & baseline data of women

| Parameters       | Mean      |
|------------------|-----------|
| Age (years)      | 38.96±9.77|
| Parity           | 2.35±1.75 |
| Hemoglobin (g/dl)| 7.71±0.66 |
| Ferritin (ng/ml) | 26.77±22.08|

Discussion

Present study confirmed the efficacy and safety of FCM infusion in treating IDA in women with heavy menstrual bleeding. Fourteen days after the infusion of FCM, the mean increase in haemoglobin and ferritin values were 2.62 g/dl and 227.98 ng/ml respectively and this difference was statistically significant (p<0.0001 for both). Our findings were consistent with two other previously published studies which were performed in women with heavy menstrual bleeding related IDA.8,9

A study by Van Wyck et al showed a mean increase of 2 g/dl or more in haemoglobin within 42 days and anaemia was corrected in about 73% of the women (Haemoglobin >12g/dl). The same study showed a mean increase up to 3 g/dl in haemoglobin after 14 days of FCM administration.

The efficacy and safety of FCM treatment in the correction of IDA was demonstrated in patients with various medical disorders like...
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A prospective study on the efficacy and safety of ferric carboxymaltose in correcting anaemia in patients with heavy menstrual bleeding. 14, 15 In these studies FCM was

Two retrospective studies mentioned the efficacy and safety profile of FCM in pregnant women with IDA. 14, 15 In one study with 457 patients, after 6 weeks of FCM infusion.

In one study in pregnancy, ferric carboxy maltose compared with oral iron Ferrous sulfite (FS). Hb levels improved at comparable rates across both treatment. However significantly more woman achieved anemia correction with FCM vs FS (Hb≥11g/dl, 84% vs 70% P=0.031) and within shorter time frame (median 3.4 vs 4.3 weeks). Treatment related adverse events were experienced in FCM was 11% as compare to FS (15%) 17

In one more study by patricia FCM was compared with IS in antenatal women. Mean rise of Hb value was 15.4g/l for FCM and 11.7g/dl for IS. The incidence of drug induced adverse events was low and mild in both the groups. 18

There are several studies which documented the efficacy and safety of FCM treatment in postpartum anaemia. 19–22 A study by Von Wyck et al. demonstrated an earlier rise of haemoglobin (≥2g/dl) with FCM when compared to oral iron (7 days compared with 14 days) and an increase ≥3g/dl in hemoglobin was noted after 7 days in 86.3% of postpartum women. While comparing the efficacy and safety of FCM with iron sucrose, Sharma et al reported an increase of 3.14/dl in mean haemoglobin and an increase of 125.91ng/ml in mean ferritin values after 14 days after they treated postpartum anaemia with FCM infusion. The FCM treatment is also found to be efficacious in improving haemoglobin values of anaemic patients who are to undergo surgery. 21

In one study with 457 patients, after 6 weeks of FCM infusion, increase in Hb level by ≥:or=2g/dl was attained by 41% of all the patient,40% in the disease of digestive system group,55% in the disease of genitourinary system groups,26% in the neoplasm groups and 29% in the disease of circulatory system. Hb increase of ≥:or=3g/dl after 6 weeks after FCM dose was attained by 20% of all patients,24% of the patients in the IDA group,22%in the diseases of digestive system,26%in the disease of genitourinarysystem,115 in the neoplasm group and 16%in the diseases of circulatory system Groups. 24

In one systematic review, FCM when compared to other iron formulation, performed better in the achievement of a rapid and consistent Hb response. A higher increase in serum ferritin was also observed in patients receiving FCM comparing to other therapy. 25

Another study compared FCM with iron dextran (ID) and Iron sucrose (IS) in iron deficiency anaemia. There was increase in Hb concentration significantly greater in both IS and FCM group compared to ID (p=0.04and, 0.01 respectively).However there was no statistically significant difference between the groups treated in IS and FCM. There was significant increase in both mean serum ferritin concentration and mean MCV after treatment in all groups. 25

In the present study, adverse effects associated with FCM infusion were mild and temporary. All the patients had uneventful recovery after receiving FCM infusion. Various adverse effects like nausea, injection site pain, headache, hypertension, constipation, fatigue, dizziness and vomiting were minimal and momentary. Adverse effects relted with FCM treatment resemble those observed in other studies. 9, 19–22

Surgery or blood transfusion or both may be required several times in patients with menstruation. Red cell transfusion is not only costly and short in supply but also has its well described risks and side effects. In moderate to severe anaemia related with heavy menstrual bleeding, the administration of FCM infusion can avoid the need of blood transfusion as it results in a rapid rise in hemoglobin levels. This study displays the safety and efficacy of FCM infusion in treating the anaemia of patients with heavy menstrual bleeding and, thu the need for blood transfusion could be avoided.

The power of the present study was limited by relatively small number of women and shorter follow up period of patients.

Conclusion

This study demonstrated the safety and efficacy of FCM infusion in treating anaemia in patients with heavy menstrual bleeding. The FCM treatment should be considered in such women to avoid the possible need of blood transfusion.

Acknowledgments

None.

Conflict of interests

Authors declare that there is no conflict of interest.

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