Policy for the use of intravenous Iron Dextran
(CosmoFer ®)

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<td>Lead author and designation</td>
<td>Dr K Rege, Martin Drury and Kaye Bowen</td>
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Policy for the use of intravenous Iron Dextran (CosmoFer ®)

Background

Therapy for iron deficiency anaemia includes treatment of its underlying cause and restoration of normal haemoglobin concentrations and iron stores. This can be accomplished by the oral or parenteral route. Although the oral route is usually preferred, clinical situations exist where the parenteral route is indicated (see point 1.2 below).

There are two preparations of IV iron available. IV iron sucrose (Venofer ®) given as divided dosages, and low molecular weight iron dextran (CosmoFer ®). This can be given as divided dose or as a total dose infusion (CosmoFer ® may also be given intramuscularly).

Choice of preparation is dependent on patient and physician choice and how immediate the requirement to complete the dose i.e. iron dextran (CosmoFer ®) can be given as a total dose infusion, iron sucrose (Venofer ®) is given as divided doses.

THIS POLICY REFERS TO LOW MOLECULAR WEIGHT IRON DEXTRAN (COSMOFER ®)

Purpose and scope

This policy exists to give guidance on the prescription and use of intravenous Iron Dextran (CosmoFer®). It is to be used by all members of staff involved in the prescription and administration of the product.

1. General information

1.1 Investigations required

Patients with anaemia should be fully investigated as appropriate. The following blood investigations are required prior to starting treatment with IV iron:
- Full blood count + film, reticulocyte count
- Iron profile
- CRP

In addition B12 and folate levels may be indicated.

1.2 Clinical indications

Intravenous iron is indicated for the treatment of iron deficiency in the following situations:
- Demonstrated intolerance to oral iron preparations
- A clinical need to deliver iron rapidly to replenish iron stores
- Active inflammatory bowel disease where oral iron preparations are not tolerated or contraindicated
- Patient non-compliance with oral iron therapy
- Patients over 14 years of age
Oral iron must not be administered concomitantly with a course of IV iron or until 5 days after the last dose.

1.3 Contraindications

- Anaemia not attributable to iron deficiency
- Iron overload
- A history of hypersensitivity to parenteral iron preparations
- History of cirrhosis of the liver
- Acute or chronic infection
- Active rheumatoid arthritis
- First trimester of pregnancy
- Acute renal failure
- Patients with a history of severe asthma, eczema or other atopic allergy
- Drug hypersensitivity including mono- or di- saccharide complexes and dextran

1.4 Response

Due to iron metabolic pathways, a rise in reticulocyte count will occur during the second week and thereafter, provided bleeding is not excessive, one can expect a rise in haemoglobin of approximately 1.5g/dL/week.
1.5 Flowchart for the use of iron in confirmed iron deficiency anaemia

Is patient taking oral iron?
Oral iron replacement therapy should usually be continued for 4 months to replenish stores fully

Yes

Is anaemia resolving?

Yes
Continue & monitor

No

Is diagnosis secure?

Yes

Re-investigate

No

Intolerant?

Change preparation
↑vitamin C intake to aid absorption

Tolerated?

Yes

Continue & monitor

No

? Compliance
? Ongoing bleeding
Poor absorption

IV iron

Tolerant

Yes

Continue & monitor

No
2. Protocol for administration of intravenous iron dextran - CosmoFer®

2.1 Dosage - Total dose infusion:

The dose calculation for CosmoFer® is based on patient’s body weight according to the table below and is diluted in 500 mls of 0.9% sodium chloride.

If calculating the dose for an obstetric patient, use the pre-pregnancy weight. For obese patients (BMI > 30) use ideal body weight.

2.2 How to select the correct dose of CosmoFer®

In the left hand column, find the body weight closest to the patient’s body weight, read across this row to the column headed by the patient’s current haemoglobin value, values for body weight and haemoglobin must be rounded up or down to the nearest stated value. The number at this point is the dose required (in milligrams of iron). The target Hb using the table shown is 13g/dL.

**Note:** If the dose is shaded in grey, it exceeds the total upper limit for total dose infusion (20mg/kg body weight) and must be administered as a divided dose. (See section 2.3)

**A test dose must be given prior to the infusion.**

**Facilities for cardiopulmonary resuscitation/treatment of anaphylaxis, should be available.** (See section 2.4)

CosmoFer® must be added to 500mLs of Sodium Chloride 0.9% IV infusion and infused over 4 to 6 hours. There is no upper concentration limit for the infusion to maintain product stability. Once made up, the infusion should be used immediately but is stable for 24 hours. The patient should be observed for 1 hour after completion of the infusion.

CosmoFer® must be administered via an IV infusion pump

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<th>8</th>
<th>9</th>
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Example  74kg, current Hb 8.2g/dL – use body weight 75kg and Hb 8g/dL dose is 1400mg iron (28mLs Cosmofer® injection) in 500mLs Sodium Chloride 0.9%

2.3 Dosage – Total dose infusion exceeding 20mg/Kg
If the required dosage exceeds 20mg/kg, then it should be given on two separate days. This can be done by:

- Giving half the dose on each day
- Giving up to 20mg/kg in the first infusion, then the remainder in the second infusion.

An interval of one week for every 600mg of iron given in the first infusion should be allowed between the first and second doses. For example if 1200mg of iron was given in the first dose, then the second infusion containing the remaining iron should be given 2 weeks later.

2.4 Test dose
The first infusion of CosmoFer® must include a test dose; facilities for cardiopulmonary resuscitation/treatment of anaphylaxis, should be available. (Refer also to section 3 – Adverse events) 25mg of CosmoFer® should be infused over a period of 15 minutes. The patient should then be observed for 45 minutes. If no adverse reactions are seen, give the remaining dose. For subsequent doses the 25mg test dose must still be given, however there is no requirement for the 45 minute observation period.

2.5 Administering the test dose
The test dose should be given by adding CosmoFer® 25mg to Sodium Chloride 0.9% 100mls and infusing over 15 minutes. If no reaction is noted after a further 45 minutes, proceed with the remainder of the infusion (i.e. the total calculated dose minus the 25mg test dose).

2.6 Continuing the infusion
The remainder of the infusion should be given at the following rate:

- 50mLs/hr for the first hour
- 100mL/hr for the next hour
- 150mL/hr until infusion complete

2.7 Patient monitoring
Blood pressure and pulse should be monitored prior to the infusion and every 15 minutes during the test dose and observation period. For the remainder of the infusion, BP and pulse should be monitored every 30-60 minutes or as clinically indicated.

3. Adverse events
Adverse reactions are rare. However, facilities for dealing with anaphylaxis and cardiopulmonary resuscitation should be available.

It is recommended that the anaphylaxis box be kept in the close vicinity of the patient receiving IV iron. Administration should be carried out by nurses or midwives who are IV certified and who have attended the Trusts anaphylaxis study day or received training in the management of anaphylaxis.

- Acute, severe anaphylactoid reactions are uncommon. They usually happen within the first few minutes of administration and are generally characterised by the sudden onset of respiratory difficulty and/or cardiovascular collapse.
CAUTION: You must refer to the intranet for the most recent version of this policy

- Less severe manifestations of immediate hypersensitivity are also uncommon and include urticaria, rashes, itching, nausea and shivering.
- Delayed reactions are characterised by arthralgia, myalgia and sometimes fever. Symptoms may last 2-4 days and settle spontaneously or following the use of simple analgesics such as paracetamol.
- Exacerbation of joint pain in rheumatoid arthritis can occur
- Local reactions such as phlebitis around the administration site may occur.

3.1 Management of adverse events
In the event of a serious anaphylactic or allergic reaction stop the infusion, IM adrenaline should be administered and appropriate resuscitation measures initiated. Mild allergic reactions should be managed by stopping the infusion and administering antihistamines. Hypotensive episodes may occur if administration is too fast, so decrease infusion rate as clinically indicated.

4. Treatment of obstetric patients with CosmoFer ®
CosmoFer® should not be used during the first trimester but can be used during the second and third trimester and during lactation if oral iron therapy is ineffective or impracticable. Dosing for antenatal patients should be based on pre-pregnancy weight (or ideal body weight if obese prior to pregnancy).

5. Follow-up
Full blood count, reticulocyte and iron profile should be checked 3-4 weeks after the CosmoFer® infusion.

References

CosmoFer® model protocol and data sheet