IRON PRODUCT CHOICE AND DOSE CALCULATION FOR ADULTS

Guidance for Australian Health Providers

MARCH 2016
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The content obtained from this document or derivative of this work must be attributed as the Iron Product Choice and Dose Calculation.

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IRON PRODUCT CHOICE AND DOSE CALCULATION - ADULTS

This guide has been developed to assist clinicians determine the appropriate formulation and dosage of iron replacement therapy for adults who have been diagnosed with iron deficiency (ID) and/or iron deficiency anaemia (IDA).

Iron requirements for infants, children and adolescents are included in Chapter 4 of the Patient Blood Management Guidelines: Module 6 Neonatal and Paediatric (in development).

For information on aetiology, prevention, diagnosis and investigation of iron deficiency refer to Appendix 1.

IRON FORMULATIONS

All patients with ID should have iron supplementation to correct anaemia and/or replete body stores.¹-³

Patients should be provided with information brochures about the iron therapy prescribed.

Diet

Increase in dietary iron may be valuable for secondary prevention of iron deficiency¹ but should not be relied upon as treatment. Consider referral to an Accredited Practising Dietician.

Oral therapy

Oral iron therapy is suitable and effective as first line therapy in most patients with iron deficiency or iron deficiency anaemia.¹ When given at equivalent elemental iron doses, different oral iron salts have similar efficacy and tolerability.¹

Intramuscular therapy

Intramuscular (IM) iron is effective but painful and may be associated with permanent skin staining. It is no safer than IV infusion.¹ Its use is discouraged.¹,²

Intravenous therapy

Three preparations of intravenous iron (IV) are approved for IV use in Australia:

- Ferric carboxymaltose: Ferinject®
- Iron polymaltose: Ferrosig®
- Iron sucrose: Venofer®

See Appendix 2 for indications and comparison information.
Making the choice

Oral iron therapy is suitable and effective as first line therapy in most patients, including most obstetric patients, with iron deficiency or iron deficiency anaemia. Indications for intravenous iron include:

- contraindications to oral iron, or compliance or tolerance (side effect) issues
- pregnancy (beyond the first trimester) and postpartum if oral iron not suitable or effective, or to prevent physiological decompensation
- comorbidities which may impact on absorption (eg. intestinal mucosal disorders), or bone marrow response
- chronic renal impairment receiving erythropoiesis-stimulating agent therapy
- ongoing iron losses that exceed absorptive capacity
- requirement for rapid iron repletion (eg. prevention of physiological decompensation or preoperatively for non-deferrable surgery)

Although the initial rise in haemoglobin (Hb) is more rapid with parenteral iron, the rise in Hb at 12 weeks is similar to that observed during oral iron therapy.

Availability of IV preparations, dosing schedules and facilities for administration are also important considerations.

IRON DOSE

Oral therapy

The usual recommended dose in adults is 100–200 mg of elemental iron daily, in 2 to 3 divided doses. Lower doses may be as effective and better tolerated. In maternity patients with iron deficiency without anaemia, a low dose of elemental iron (eg. 20-80 mg daily) may be considered, and may be better tolerated than higher doses.

More than 100 preparations containing iron are available over the counter in Australia; however few contain sufficient elemental iron to treat IDA effectively. Appropriate formulations are outlined in the Oral preparations for treatment of iron deficiency anaemia (IDA) chart (Appendix 3). Multivitamin-mineral supplements should not be used to treat IDA as iron content is low and absorption may be reduced.

After therapeutic doses of oral iron, reticulocytosis should occur within 72 hours, and Hb levels should rise by about 20 g/L every 3 weeks. Oral iron should be continued for 3 months after anaemia has been corrected to replenish stores. Inadequate response to oral iron therapy can be due to a number of factors, such as inadequate intake or absorption, ongoing losses, coexisting conditions or incorrect diagnosis, with more than one often being involved.
Calculating total body iron deficit

The patient’s total body iron deficit (cumulative amount of iron required to replete body iron stores) is NOT the same as the allowable iron dose per infusion which is DIFFERENT for each product. Calculate the deficit to determine how many doses of the desired preparation is required. Refer to the specific product information and local administration guidelines for information on the maximum iron dose per infusion for each product (Appendix 2).

The cumulative dose for repletion of iron is based on the patient’s Hb and body weight and should not be exceeded. There are two methods for determining the cumulative dose – the Ganzoni formula and the Simplified Method.

Ganzoni formula:

\[
\text{Total body iron deficit/cumulative iron dose (mg)} = \text{body weight} \times (\text{target Hb} - \text{actual Hb in g/L}) \times 0.24 + \text{iron depot (mg)}
\]

*Use ideal body weight in overweight patients. If underweight, use actual body weight

**The factor 0.24 = 0.0034 \times 0.07 \times 1,000:

For this calculation the iron content of haemoglobin = 0.34%,

blood volume = 7% of the bodyweight, and

1,000 is the conversion from g to mg

***Iron depot:

<35 kg body weight: iron depot = 15 mg/kg body weight

\geq 35 kg body weight: iron depot = 500 mg

For example a 70 kg female with Hb 80 g/L has an iron deficit of:

\[
70 \times (150 - 80) \times 0.24 + 500 = 1676 \text{ mg i.e. approx. 1700 mg}
\]

Note that the target Hb may vary according to patient population. The UK guidelines on the management of iron deficiency in pregnancy recommend a target Hb of 110 g/L, based on prepregnancy weight, in women from the second trimester onwards and postpartum period, with iron deficiency anaemia who fail to respond to, or are intolerant of, oral iron. Refer to local policies and procedures.
Simplified Method:

To date only the product information (PI) of Ferinject® incorporates the Simplified Method. However, expert practice and published localised drug guidelines now reflect this change. The following table can be used for estimating the cumulative amount of iron required to replete body iron stores (for adult patients of body weight ≥ 35 kg).

Estimated cumulative iron dose

<table>
<thead>
<tr>
<th>Hb g/L</th>
<th>Body weight 35 kg to &lt;70 kg*</th>
<th>Body weight ≥70 kg*</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;100 g/L</td>
<td>1,500 mg</td>
<td>2,000 mg</td>
</tr>
<tr>
<td>≥100 g/L</td>
<td>1,000 mg</td>
<td>1,500 mg</td>
</tr>
</tbody>
</table>

*Use ideal body weight in overweight patients. If underweight, use actual body weight

Caution is recommended with the Simplified Method as it is based on experience in a single clinical trial in adults with inflammatory bowel disease with a median Hb 104 g/L (range 61-146 g/L) and body weight ≥ 35 kg. Seek expert advice from a haematologist if in doubt.

**ADMINISTRATION**

Infusion rates, maximum dose per infusion and dilution are NOT interchangeable between IV iron products. Refer to the specific product information and administration guidelines.

Oral iron is not required after IV iron is given if the total iron deficit has been (or will be) repleted with IV iron therapy.

A “total-dose” infusion (where iron stores can be repleted in a single treatment episode) can be administered with iron polymaltose and in mild cases of IDA with ferric carboxymaltose, however iron sucrose requires multiple small intermittent doses over days to weeks.

Anaphylaxis may occur with IV iron and resuscitation facilities should be available. It would appear that iron polymaltose may have a higher incidence of severe systemic reactions than iron sucrose and ferric carboxymaltose.

Hypophosphataemia has been reported with all three intravenous iron preparations and may be more common and severe with ferric carboxymaltose. Caution should be taken with patients at risk.

All three iron preparations have similar mild adverse event profiles.
REFERENCES


APPENDIX 1: IRON THERAPY RESOURCES

Iron deficiency anaemia guidelines/references:


Iron deficiency anaemia education/information/tools:

BloodSafe eLearning Australia:
- Iron deficiency anaemia algorithm app (iPhone, iPad, Android)
- Iron deficiency anaemia course
  Available at: https://www.bloodsafelearning.org.au
- Australian Red Cross Blood Service information about Iron deficiency anaemia:
  - Treatment Options for Iron Deficiency Anaemia
  - Major Reasons for Inadequate Response to Oral Iron Therapy
  - Oral Iron Therapy Interactions and Management
  - Oral Iron Therapy Side Effects and Management
  - Spectrum of iron deficiency
  Available at: http://www.transfusion.com.au

Intravenous iron references

Product information for intravenous iron preparations available in Australia:
- Ferric carboxymaltose: Ferinject®
- Iron polymaltose: Ferrosig® and Ferrum H®
- Iron sucrose: Venofer®
  Available at: http://www.ebs.tga.gov.au
• Australian Injectable Drugs Handbook (AIDH)
  • Ferric carboxymaltose
  • Iron polymaltose complex
  (note: AIDH requires subscription, however the above pages are accessible)

• Guiding principles for the development of intravenous (IV) iron infusion practice developed
  for Victorian health services
  • Additional resources from Victorian health services
    o Ballarat Health 2010 Iron Polymaltose infusion policy
    o Iron carboxymaltose Administration guidelines 2012 : Peter MacCallum Cancer
      centre
  principles_iron_infusion.htm

• Fremantle Hospital and Health Service, Department of Pharmacy. Specialised Drug
  Guidelines regarding intravenous iron use:
  • Iron carboxymaltose
  • Iron polymaltose
  • Iron sucrose
  Available at: http://www.health.wa.gov.au/bloodmanagement

  Perinatal Practice Guidelines – iron infusion.
  Available at: http://www.sahealth.sa.gov.au

• National Prescribing Service (NPS). Ferric carboxymaltose (Ferinject) for iron deficiency
  anaemia.
  Available at: http://www.nps.org.au

• Gozzard D. When is high-dose intravenous iron repletion needed? Assessing new treatment

• Auerbach M, Ballard H. Clinical use of intravenous iron: administration, efficacy, and safety.
  pubmed.gov
Guiding Principles for the use of off-label medicines:

Healthcare professional resources
- Dieticians Association of Australia
  Available at: http://daa.asn.au/
- Patient Blood Management Guidelines
  - Module 2 Perioperative
  - Module 3 Medical
  - Module 5 Obstetrics and Maternity
  Available at: http://www.blood.gov.au
- BloodSafe Oral iron therapy dosing chart
  Available at: http://www.bloodsafe.sa.gov.au

Resources for patients
- BloodSafe Iron therapy brochures for patients (oral)
  Available at: http://www.bloodsafe.sa.gov.au
- BloodSafe: Intravenous (IV) iron infusions
  Available at: http://www.bloodsafe.sa.gov.au
- Intravenous (IV) iron infusions: Fremantle Hospital and Health Service
  Available at: http://docs.health.vic.gov.au/docs/doc/Intravenous-iron-infusions-Fremantle-Hospital-and-Health-Service
- Australian Red Cross Blood Service patient website: iron deficiency anaemia for patients
  Available at: http://mytransfusion.com.au
### APPENDIX 2: INTRAVENOUS IRON PREPARATIONS COMPARISON INFORMATION (AS PER PRODUCT INFORMATION)

<table>
<thead>
<tr>
<th></th>
<th>Ferric carboxymaltose</th>
<th>Iron sucrose</th>
<th>Iron polymaltose</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Indication</strong></td>
<td>Treatment of (laboratory diagnosed) iron deficiency when oral iron preparations are ineffective or cannot be used.</td>
<td>The treatment of (laboratory diagnosed) iron deficiency anaemia in patients undergoing chronic haemodialysis and who are receiving supplemental erythropoietin therapy.</td>
<td>Treatment of iron deficiency anaemia when oral therapy is contraindicated, enteric absorption of iron is defective or when patient non-compliance or persistent gastrointestinal intolerance makes oral therapy impractical.</td>
</tr>
<tr>
<td><strong>Iron concentration (mg/mL)</strong></td>
<td>50 mg/mL</td>
<td>20 mg/mL</td>
<td>50 mg/mL</td>
</tr>
<tr>
<td><strong>Presentation</strong></td>
<td>2 mL (100 mg) or 10 mL (500 mg) vial</td>
<td>5 mL ampoules</td>
<td>2 mL ampoules</td>
</tr>
<tr>
<td><strong>Maximum dose in a single administration for patients ≥35 kg</strong></td>
<td>1000 mg (not more than 1000 mg per week) (max 20 mg iron/kg body weight*)</td>
<td>100 mg not more than 3 times per week Most patients will require a minimum cumulative dose of 1000mg</td>
<td>2500 mg</td>
</tr>
<tr>
<td><strong>Frequency of administration</strong></td>
<td>IV infusion: do not give more than 1000mg iron per week. Do not exceed calculated cumulative iron dose or 20 mg iron/kg body weight. IV bolus injection: max dose of 200mg no more than 3 times a week.</td>
<td>100mg no more than three times per week.</td>
<td></td>
</tr>
<tr>
<td><strong>Rate of administration</strong></td>
<td>IV infusion:</td>
<td>Intravenous infusion 100 mg over 15 minutes. For haemodialysis patients may be given slow IV injection into the venous limb of the dialysis line at 1 mL (20 mg iron) per minute (ie. 5 minutes per ampoule)</td>
<td>The first 50 mL should be infused slowly at 20-40 mL/hour If tolerated may be increased to 120 mL/hour Note: see links below for rapid infusion information</td>
</tr>
<tr>
<td><strong>Total dose single infusion</strong></td>
<td>No (Unless total body iron deficit is &lt;1000 mg)</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td><strong>Link to Product information</strong></td>
<td><strong>Ferinject®</strong></td>
<td><strong>Venofer®</strong></td>
<td><strong>Ferrosig® and Ferrum H®†</strong></td>
</tr>
</tbody>
</table>

*Use ideal body weight in overweight patients. If underweight, use actual body weight
†Whilst considered brand equivalent to Ferrosig®, Ferrum H® is not licensed for intravenous use in Australia – refer to Guiding Principles for the use of off-label medicines.1

See over for important considerations
Important considerations

Always refer to local health service guidelines/protocols and product information for the specific iron preparation. Maximum doses per infusion, infusion rates and dilution are not interchangeable between the different IV iron preparations.

Emerging data and regional practice indicate that alternative dosing and infusion schedules may be practical and safe.\(^1\)\(^-\)\(^3\)

There is also evolving evidence regarding the appropriateness of the three IV iron preparations in varying clinical circumstances. If in doubt, refer to local guidelines/protocols or consult with an expert haematologist.

Note: TEST DOSE NOT REQUIRED - The European Medicines Agency’s (EMA) Committee for Medicinal Products for Human Use (CHMP)\(^4\) has considered that the current practice of first giving the patient a small test dose is not a reliable way to predict how the patient will respond when the full dose is given. A test dose is therefore no longer recommended but instead caution is warranted with every dose of intravenous iron that is given, even if previous administrations have been well tolerated.\(^4\)

Current IV iron Product Information (PI) can be found on the Therapeutic Goods Administration website: [www.ebs.tga.gov.au](http://www.ebs.tga.gov.au) - click on “Public TGA Information” in menu bar on left then click on Product Information and enter product name in search box.

References:


# APPENDIX 3: ORAL PREPARATIONS FOR TREATMENT OF IRON DEFICIENCY ANAEMIA (IDA) IN AUSTRALIA*

<table>
<thead>
<tr>
<th>NAME (Manufacturer)</th>
<th>TABLET</th>
<th>FORMULATION</th>
<th>ELEMENTAL IRON CONTENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>FERRO-LIQUID (AFT pharmaceuticals) PBS listed</td>
<td></td>
<td>Ferrous Sulphate Oral solution</td>
<td>30 mg/5 mL</td>
</tr>
<tr>
<td>FEFOL® Iron and folate supplement (Pharm-a-care)</td>
<td></td>
<td>Ferrous Sulphate 270 mg Folic acid 300 mcg Delayed release capsule</td>
<td>87.4 mg</td>
</tr>
<tr>
<td>Ferro-f-tab (AFT pharmaceuticals)</td>
<td></td>
<td>Ferrous Fumarate 310 mg Folic acid 350 mcg</td>
<td>100 mg</td>
</tr>
<tr>
<td>Ferro-tab (AFT pharmaceuticals)</td>
<td></td>
<td>Ferrous Fumarate 200mg</td>
<td>65.7 mg</td>
</tr>
<tr>
<td>FERRO-GRADUMET (Abbott)</td>
<td></td>
<td>Ferrous Sulphate 325 mg Modified release tablet</td>
<td>105 mg</td>
</tr>
<tr>
<td>FERRO-GRAD C (Abbott)</td>
<td></td>
<td>Ferrous Sulphate 325 mg Ascorbic acid 500 mg Modified release tablet</td>
<td>105 mg</td>
</tr>
</tbody>
</table>
**NAME (Manufacturer)**

<table>
<thead>
<tr>
<th>TABLET FORMULATION</th>
<th>ELEMENTAL IRON CONTENT</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>FGF (Abbott)</strong></td>
<td>250 mg Ferrous Sulphate Modified release tablet</td>
</tr>
<tr>
<td><strong>#Maltofer (Aspen Pharmacare)</strong></td>
<td>Iron polymaltose 370 mg</td>
</tr>
<tr>
<td><strong>#Maltofer Syrup (Aspen Pharmacare)</strong></td>
<td>Iron polymaltose 185 mg Oral solution</td>
</tr>
</tbody>
</table>

*Response to oral iron polymaltose (Maltofer) may be slower than with ferrous iron. Maltofer is licenced in Australia for treatment of iron deficiency in adults and adolescents where the use of ferrous iron supplements is not tolerated, or otherwise inappropriate.*


See below for dosing and considerations.

**Dosing and considerations:**

- Usual ADULT dose for IDA is around 100–200 mg elemental iron daily in divided doses
- Ideally give 1 hr before or 2 hrs after food
- GI upset may be reduced by taking tablet with food or at night & increasing dose gradually
- Consider giving supplement with Vitamin C (eg. orange juice) to improve absorption
- When a rapid increase in Hb is not required, intermittent dosing (1 tablet 2–3 times a week) or lower doses of iron (e.g. 30–60 mg of elemental iron, increasing to twice daily or three times a day if tolerated: try Ferro-tabs or titrate liquid) may reduce GI upset
- Multivitamin-mineral supplements should not be used to treat IDA as iron content is low and absorption may be reduced
- Iron overdose may be fatal – keep medication out of reach of children
- Based on limited available data, controlled-release iron formulations appear to have fewer GI side effects, but similar discontinuation rates and comparable efficacy; release of iron distal to the site of maximal intestinal absorption may theoretically limit response in some patients.