**Hypersensitivity reactions to intravenous iron: guidance for risk minimisation and management**.

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**Abstract.**

Intravenous (iv) iron is widely used for the treatment of iron deficiency anemia when oral iron is inappropriate, ineffective or poorly tolerated. Acute hypersensitivity reactions during iron infusions are very rare but can be life-threatening: this paper reviews their frequency, pathogenesis and risk factors and provides recommendations about their management and prevention. Complement activation-related pseudo-allergy triggered by iron nanoparticles is probably a more frequent pathogenetic mechanism in acute reactions to current formulations of iv iron than is an immunologic IgE-mediated response. Major risk factors for hypersensitivity reactions include a previous reaction to an iron infusion, a fast iron infusion rate, multiple drug allergies, severe atopy and possibly systemic inflammatory diseases. Early pregnancy is a contra-indication to iron infusions, while old age and serious co-morbidity may worsen the impact of acute reactions if they occur. Management of iron infusions requires meticulous observation, and, in the event of an adverse reaction, prompt recognition and severity-related interventions by well-trained medical and nursing staff.

**Introduction.**

Intravenous (iv) iron is increasingly used for the treatment of iron deficiency anemia (IDA) when oral iron is ineffective or poorly tolerated and when it or blood transfusion is inappropriate 1,2. It is also indicated in combination with erythropoiesis-stimulating agents in chronic kidney disease and chemotherapy-induced anemia. While acute reactions during iron infusions are very infrequent, they can be life-threatening.

In 2013, the European Medicines Agency (EMA) published a report of their 2-year investigation of the adverse drug reactions to all iv iron drugs available in Europe 3 (Table 1). The formulations considered were sodium ferric gluconate (Ferrlecit), iron sucrose (Venofer), iron (III)-hydroxide dextran complex (Cosmofer), ferric carboxymaltose (Ferinject) and iron (III) isomaltoside 1000 (Monofer).

The aims of the present article are:

* to outline the frequency and outcomes of reactions to iv iron;
* to summarise current views about the pathogenesis of such reactions;
* to indicate the risk factors for reactions to iv iron; and
* to provide detailed guidance on risk minimisation and management of iron infusions and acute reactions to them

We are unaware of any existing guidance on how to prevent and manage hypersensitivity reactions (HSRs) to this increasingly used treatment, and intend this paper primarily for healthcare professionals, whether doctors or nurses, who prescribe and administer iv iron. Our aim is to offer advice which has been developed from a comprehensive literature search and iterative expert review about best practice before, during and after administration of iv iron to patients with IDA.

**Terminology.**

Current nomenclature relating to acute adverse reactions to iv drugs is confusing, inconsistent and sometimes contradictory. In this report, we refer to all acute reactions to iv iron as hypersensitivity reactions (HSRs), sub-dividing them into mild, moderate or severe/life-threatening, depending on their clinical presentation. We have adopted the World Allergy Organisation proposal that the term "anaphylaxis" is reserved for severe HSRs 4, irrespective of pathogenesis, and avoid the term "anaphylactoid", which historically has been used loosely to denote either non-immunologic, or even mild HSRs.

**Methods.**

We undertook literature searches in PUBMED and EMBASE using the search terms "intravenous", "anaphylaxis", "anaphylactic", "anaphylactoid", " iron" as major subject headings or occurring in the title/abstract. These searches were supplemented by refined drug-class specific searches with the term "infusion reactions". Secondary searches were done manually by screening articles retrieved by the online searches. Articles giving experimental data on adverse reactions to iv drugs were selected along with major anaphylaxis guidelines.

We concluded from these searches that (a) no existing anaphylaxis treatment guideline is strictly evidence-based, since the very rare occurrence of severe acute infusion reactions precludes randomised clinical trials as to how they should be managed; and (b) there is so little information specifically relating to iv iron reactions that indirect evidence relating to acute reactions to other intravenous drugs had to be considered in preparing the present guidance.

We therefore assembled a panel of experienced clinicians from fields of medicine in which IDA is common (gastroenterology, hematology, immunology, internal medicine, nephrology, obstetrics and gynecology), as well as experts in the pharmacology of drug reactions and iv iron. Our recommendations result from development of a consensus in which the working group went through an iterative process of literature review and discussion of current clinical practice about each measure proposed.

**Preparation of paper and declaration of interests**.

One of us (DSR) initiated this review but considered that it could not be undertaken without administrative and financial support to assemble the necessary international clinical expertise. Pharmacosmos (which manufactures Monofer and Cosmofer) funded two one-day meetings in Copenhagen for initial preparation of the paper. All available iv iron drugs were considered and no distinctions have been made between them in terms of safety or efficacy (see below). The pharmaceutical company had no editorial influence and were not offered approval of the manuscript. No medical writer or unacknowledged authors were involved.

**Frequency and outcomes of hypersensitivity reactions to intravenous iron infusions.**

Unlike previous authors, who had used a range of methodologies and assessed products which included poorly tolerated high molecular weight dextran preparations which are no longer available 5-8, EMA were unable to differentiate between current iv iron products in relation to the risk of severe HSRs. In this context, Wysowski et al 9 also concluded, from US data, that "because of under-reporting, possible differential reporting, absence of iron dextran brand names, and incomplete use (denominator) data, incidence rate and relative risk estimates cannot be calculated". Death and other severe long-term sequelae arising from use of iv iron are very rare. In the largest such study to date, death certificate data from the US National Center for Health Statistics between 1979-2005 showed that there were about 3 deaths/year ascribed to iron infusions in the US, approximating to 1 for every 5 million doses of iv iron sold 9,10.

**Pathogenesis of hypersensitivity reactions to intravenous iron.**

Mechanisms by which iron infusions induce adverse reactions may vary with the iron preparation used and with the pre-existing morbidity of the recipient. They cannot be distinguished by their clinical presentation.

The two main possibilities are immunologic IgE-mediated responses, for example to the dextran component of iv iron preparations containing this molecule, and complement activation-related pseudo-allergy (CARPA) 11. There is however no data to support the concept that IgE-mediated hypersensitivity accounts commonly for reactions to current formulations of iv iron12,13. CARPA may be the most common mechanism of acute HSRs provoked by any infusion containing nanoparticles, of which all existing iv iron preparations consist14. The final common pathway of these processes is likely to include activation of mast cells and basophils, either directly, or via anaphylatoxins (C3a and C5a) that rise in blood as a consequence of complement activation. The secretion products of these cells, which include histamine, thromboxanes, leukotrienes and platelet activating factor 11, trigger smooth muscle contraction, increased capillary permeability and loss of fluid from the intravascular space. Subsequent bronchospasm, laryngeal oedema, tachycardia, hypo- or hypertension, hypoxia and reduced tissue perfusion can culminate, in severe HSRs, in loss of consciousness, circulatory collapse (shock) and cardiac and respiratory arrest15.

A fast iron infusion rate is a well-recognised risk factor, one possible explanation being a rapid rise of labile free iron in this situation16. However, prevention of HSRs by reducing the speed of infusion is an effective practice not only with iv iron but also with other reactogenic drugs, so the phenomenon is unlikely to be solely due to higher levels of free iron. For example, it is also possible that, after rapid injection of iv iron, the clearance rate from the blood of anaphylatoxins by carboxypeptidase N and by uptake by blood and other cells is exceeded by their rate of production, leading to exacerbation of the CARPA pathogenic sequences described above11,15.

**Risk factors for hypersensitivity reactions to intravenous iron.**

Several factors have been suggested, on an evidence base of varying robustness, as predisposing to, on the one hand, an increased risk of an HSR occurring in patients given iv iron, and on the other, to a reaction, which if it occurs, has a worse outcome in the iron recipient. These factors have recently been restated by EMA3*,* and represent a relative contra-indication to the administration of iv iron to patients displaying them (Table 2). If iv iron is to be given to individuals with any of these risk factors, an extremely slow infusion rate and meticulous observation is recommended (see below).

Some of the factors, such as a previous adverse reaction to iv iron or other drugs, a fast iron infusion rate (see above), a history of severe atopy and systemic mastocytosis, appear to increase both the incidence and severity of HSRs 3,11. .

In contrast, pre-existing severe respiratory or cardiac disease, old age and the use of beta-blockers or ACE inhibitors, may worsen the outcome of an HSR if it occurs17. In pregnancy, iv iron is contra-indicated in the first trimester 3,18 since there are no trials confirming its safety during this time: existing data suggest that its use should be confined to the second or third trimester of pregnancy if the benefit is judged to outweigh the risks for both mother and foetus 18,19. While earlier anecdotal reports suggested that iron dextran may worsen disease activity in patients with rheumatoid arthritis and lupus20, more recent data indicate that iv iron could even have a beneficial effect on the underlying disease21.

Lastly, it has been suggested that anxiety on the part of healthcare professionals giving iv drugs increases the risk of HSRs 22.

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**Management of intravenous iron infusions and HSRs.**

**1. How can we reduce the risk of HSRs to intravenous iron?**

HSRs may occur in anyone given iv iron, and it is essential that every effort is made to prevent these being poorly managed if they occur, whether due to inadequate facilities or staff being undertrained. The following factors require attention before and during any iv iron infusion.

**a. Location.** Iron infusions should be given only in appropriately staffed sites equipped with resuscitation facilities 3. If iv iron is to be given outside hospital, there should be arrangements in place for immediate treat-and-transfer to an intensive care facility in the event of a severe reaction. EMA3 states that iron should not be infused in the home.

**b. Personnel.** In most countries, iv iron is given by nursing staff with immediate access to on-site medical help in the event of an adverse reaction. All staff should have regularly updated training in management of iv infusions and adverse reactions to them. The confidence and competence provided by regular training should help reduce any anxiety on the part of HCP and, conceivably, the risk of HSRs22. The healthcare professional (HCP) administering the iron infusion should be in the infusion area and easily accessible by the patient throughout its course, as HSRs can develop rapidly.

**c. The patient.** Information should be provided about the risk of an HSR before the iron infusion, if possible using a visual aid to indicate its rarity23. The relevant symptoms should be described, with advice to tell the HCP administering the infusion immediately if any occur.

**d. Administration of intravenous iron.** This should be preceded by a (re-)check for risk factors for an HSR (Table 2), the general condition of the patient and baseline observations including pulse rate, blood pressure and respiratory rate. The infusion should be prepared as stated in the manufacturer's instructions. No test dose of iv iron is necessary, because it can give false reassurance about the safety of the subsequent therapeutic infusion 3. However, on the basis of clinical experience, we recommend that the iron infusion should be initiated at less than 50% of the rate recommended by the manufacturer and not increased to the recommended rate until it is clear that it is being well-tolerating (usually 10-15 minutes). We suggest checking observations every 15 minutes and for 30 minutes after the infusion finishes.

**e. Risk minimisation.** While anyone having iv iron should be regarded as susceptible to HSR, some people are at higher risk than others (Table 2). In such individuals, the prescribing clinician should take a carefully considered decision as to whether the potential risks associated with an iron infusion are outweighed by its benefits3. If so, in addition to the requirements itemised above, it is recommended that, in case of urgent need by the HCP giving the infusion, an experienced doctor is in close proximity throughout the infusion. The infusion should be given at 10% of the recommended rate for the first 15 minutes. Monitoring should continue for 30-60 minutes after the infusion.

A previous HSR to iv iron increases the risk of an adverse response to a subsequent iron infusion. In line with EMA's conclusions 3, we recommend that, unless their previous reaction was due to exceptional circumstances (eg very rapid iron infusion), noone having had a severe HSR to iv iron should have a subsequent infusion of any iron products. After a mild-moderate previous HSR, the same iv iron preparation should not be used again, and a different one used only after careful consideration by the responsible clinician and the patient, as to whether the potential benefits exceed the risks3.

**2. How are HSRs diagnosed and graded?**

For ease of recognition and prompt and appropriate management, acute HSRs to iron infusions, as to other iv drugs, are best classified as mild, moderate or severe/life-threatening (or anaphylactic) (Fig 1)24,25.

Reactions can be identified on the basis of subjective symptoms, objective signs and bedside monitoring. Diagnosis of an HSR does not require the presence of every feature shown in Fig 1. Mild reactions can progress rapidly through moderate to severe ones; the latter can also occur from the outset without progression through the milder syndromes. Symptoms such as metallic taste and mild headache are part of the normal pharmacological response to iv iron and are not of clinical significance.

A further mild acute adverse reaction has been described by Fishbane 1,26. This occurs in about 1/100 those given intravenous drugs and is characterised by transient flushing and truncal myalgia (pains in the back and chest) with joint pains. Its pathogenesis has not been investigated.The symptoms abate spontaneously over a few minutes and do not usually recur on rechallenge.

**3. How should hypersensitivity reactions to intravenous iron be managed?**

Management of an HSR to iv iron depends on the severity of the event and is outlined in the algorithm shown in Fig 1. As stated at the outset, there is scanty evidence relating specifically to management of iron infusions: the recommendations made below are drawn from other contexts in which iv drugs are given17,27-31.

The selection of individual drugs for treatment of HSRs, and their doses and routes of administration, varies and can be modified according to local practice.

**a. Mild HSR**

The features of a mild HSR tend to be more subjective than objective (Fig 1). The patient often feels increasingly anxious: calm explanation of what is being felt will provide reassurance and realisation that the HCP is experienced and knows how to proceed (Fig 1).

Our stepwise advice to the HCP dealing with a mild HSR is:

* stop the infusion for at least 15 minutes and assess the response.
* if not medically qualified, immediately alert the attending physician, recheck the vital signs and watch for progression or resolution of the HSR.
* if there is an improvement over a few minutes, cautiously resume the iron infusion after 15 minutes at no more than 50% of the initial infusion rate.
* if all goes well, complete the infusion and continue observations for at least 1 hour to ensure there are no recurrent symptoms.
* if there is no resolution in 5-10 minutes, or if at any time the symptoms and signs worsen, manage as for a moderate HSR (Fig 1)

**b. Moderate HSR.**

Moderate HSRs may develop from mild reactions, or start without any prodrome.

Transient cough is a common initial feature. The symptoms include those of a mild HSR, with more marked chest tightness and shortness of breath (Fig 1). The pulse rate may rise and blood pressure fall.

Management is as for a mild HSR with immediate additional measures(Fig 1):

* stop the iron infusion immediately if it is still running
* if there has been a rise in heart rate and fall in blood pressure, give an isotonic fluid load (eg 500ml 0.9% saline, Ringer’s or Hartmann’s solution) and lie the patient horizontally. Although not clearly evidence-based, an iv corticosteroid (eg hydrocortisone 100-500mg) can be considered.
* if there is improvement, continue monitoring for at least an hour
* in the event of deterioration, implement the measures applying to management of a severe HSR immediately.

While previously recommended by some, both as premedication against and treatment of HSRs, the role of iv H1 blockers has recently been questioned30. Their therapeutic benefit is unclear, and indeed, by sometimes potentiating tachycardia, hypotension and somnolence, H1 antagonists can make an HSR appear more hemodynamically significant.

**c. Severe/life-threatening HSR (anaphylaxis).**

An anaphylactic reaction may be of sudden onset, or occur as a rapid worsening of the features of a moderate HSR. There will be increasing wheeze, due to bronchospasm, sometimes with stridor associated with laryngeal oedema. Increasing tiredness and distress will occur, and periorbital edema may develop. Increasing hypoxia leads to confusion. If the HSR worsens, pallor, clamminess, cyanosis and loss of consciousness progress quickly to cardiac and respiratory arrest. During this time, the pulse accelerates and the blood pressure and oxygen saturation fall.

A severe HSR is a major medical emergency:

* if the HSR occurs in a hospital, call the emergency response team immediately.
* if hypotension is severe, give iv adrenaline (epinephrine)(0.1mg (1ml) as a 1/10000 solution over 5 minutes); continuous ECG and blood pressure monitoring is essential in case of arrhythmia or a hypertensive response to the adrenaline.
* if the iron infusion is being given outside a hospital, adrenaline by intramuscular injection into the anterolateral thigh (0.3-0.5mg (0.3-0.5ml) 1/1000 solution) may be safer26.
* give oxygen at a high rate (>10 l/min) by face mask initially, with a nebulised β2-adrenergic agonist and/or ipratropium to combat wheezing.
* rapidly volume load with 1-2 l 0.9% saline or similar isotonic fluid (see b above).
* give an iv corticosteroid (see b above) if not already administered as part of management of a moderate HSR
* the site’s advanced cardiac and life support (ACLS) team should implement standard protocols in the event of cardiac or respiratory arrest.

If the patient responds well to initial measures, they should be observed carefully for at least 4 hours after resolution; if they are elderly, frail or in high risk categories (Table 1) observation for up to 24 hours may be necessary. While there is a theoretical risk of a biphasic event in patients having HSRs, this has not been clearly described after iron infusions.

Where response is not immediate and complete, prompt transfer for further management to an appropriate high dependency or intensive care facility is necessary.

**d. Documentation of hypersensitivity reaction after resolution.**

In every HSR, the responsible clinician should be notified promptly and the event carefully documented using a proforma designed along the lines suggested in Table 3. The report should be filed in the patient’s case records: it will help the clinician to decide how to treat the anemia in the future. A report of every HSR should also be submitted to the appropriate national regulatory body.

**Conclusions.**

There is a paucity of evidence about how to manage HSRs to iron infusions. The rarity of HSRs means that there will never be a formal clinical trial to assess optimal therapeutic measures. Areas in which further research is needed and could be productive, however, include clarification of the pathogenesis of HSRs, risk definition in individual patients and in different diseases, and the role of premedication and risk reduction protocols in high risk patients.

Hypersensitivity reactions to iv iron are rare but potentially life-threatening. They are at least partly preventable by implementation of risk minimisation measures. Their management requires prompt recognition and grading of severity, together with meticulous monitoring and immediate treatment. All staff involved in giving iron infusions need regular training to ensure that when these rare events develop they are dealt with calmly and expeditiously.

***Authorship****:*

DSR conceived the report and drafted the manuscript; JOF undertook the systematic literature review; all the authors participated in the iterative literature review and guidance development, and contributed to the planning, revision and final approval of the manuscript

***Disclosures.***

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* All iv iron preparations carry a small risk of adverse reactions which can be life-threatening if not treated promptly.
* Nevertheless, the benefits of iv iron outweigh its risks in the treatment of iron deficiency when the oral route is insufficient or poorly tolerated.
* Iv iron products should be administered only when staff trained to evaluate and manage anaphylactic reactions, as well as resuscitation facilities, are immediately available.
* A test dose is not appropriate as it may give false reassurance.
* Patients should be closely monitored for signs of hypersensitivity during and for at least 30 minutes after each administration.
* All iv iron products are contraindicated in patients with known serious hypersensitivity to any parenteral iron product.
* Iv iron should not be given to pregnant women in the first trimester. Careful risk/benefit evaluation is required before use in the second or third trimester.
* Special precautions are needed if iv iron is to be given to patients with known allergies (including drug allergies), severe atopy or systemic inflammatory diseases (eg systemic lupus erythematosus, rheumatoid arthritis).

**Table 1.** Summary of conclusions of EMA report (2013)3 on iv iron products

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* Previous reaction to intravenous iron
* Fast iron infusion rate
* History of other drug allergy or allergies
* Severe asthma or eczema
* Mastocytosis
* Severe respiratory or cardiac disease
* Old age
* Treatment with beta-blockers, ACE inhibitors
* Pregnancy (first trimester)\*
* Systemic inflammatory disease (eg rheumatoid arthritis, lupus)\*\*
* Anxiety (patient or staff)

**Table 2.** Factors increasing risk and/or severity of hypersensitivity reactions (HSRs) in patients given iron infusions. \*iv iron is contra-indicated in early pregnancy. \*\*evidence equivocal with current iv iron preparations. ACE: angiotensin converting enzyme

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* Severity of reaction (mild, moderate or severe/life-threatening)
* Previous iv iron preparations given, dates, doses, number of previous infusions, infusion rates
* Risk factors for HSR
* Initial symptoms and course of progression
* Interventions, timing, patient response
* Timing of symptom onset and resolution
* Discharge instructions or transfer to intensive care
* Responsible clinician and regulatory bodies to whom this information has been sent

**Table 3.** Information to be recorded in patients' case records immediately after any iv iron-related hypersensitivity reaction (HSR) (adapted from Vogel 2010 29). This information helps to determine future treatment strategy.

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**Legend to Fig 1. .**Algorithm outlining grading and management of acute hypersensitivity reactions to intravenous iron infusions. Details are given in the text

**Fig 1**

***Mild HSR***

itching, flushing, urticaria, sensation of heat, slight chest tightness, hypertension,

back/ joint pains

***Severe/life-threatening HSR***

Sudden onset and rapid aggravation of symptoms + wheezing/stridor, periorbital edema, cyanosis, loss of consciousness, cardiac/respiratory arrest

***Moderate HSR***

As in ***Mild reaction***+ transient cough, flushing, chest tightness, nausea, shortness of breath, urticaria, tachycardia, hypotension

***Management***

Stop iron infusion for ≥15 mins

Inform doctor

Monitor pulse, BP, resp rate, O2 saturation

Wait and watch

***Treat as for mild reaction AND***

Stop iron infusion

Call doctor

Consider volume load (eg iv 0.9% saline 500ml), iv corticosteroid (eg hydrocortisone 200mg)

***Treat as in moderate reaction AND***

Call fast response team

Stop iron infusion

Adrenaline im (0.5mg 1/1000) or iv (0.1mg 1/10000)

Nebulised Β2 agonist

Further isotonic volume load

iv corticosteroid

O2 face mask

ACLS (if necessary)

***Patient no better in***

***5-10 mins, or deteriorating***

***Patient well***

Observe for ≥1-4 hr

Document event

Consider future treatment strategy

***Patient no better***

Transfer quickly to intensive care unit

***Patient deteriorating***

***Symptoms recur***

Stop iron infusion

Manage as above

Document event

***Patient better***

Restart iron infusion at reduced rate (eg 50%)