FERAHEME is indicated for the treatment of iron deficiency anemia (IDA) in adult patients:

- who have intolerance to oral iron or have had unsatisfactory response to oral iron or
- who have chronic kidney disease (CKD)

The recommended dose of FERAHEME is an initial 510 mg dose followed by a second 510 mg dose as early as 3 days and up to 8 days later, each dose infused over at least 15 minutes while the patient is in a reclined or semi-reclined position.

### Feraheme® (ferumoxytol injection) Important Safety Information

**WARNING: RISK FOR SERIOUS HYPERSENSITIVITY/ANAPHYLAXIS REACTIONS**

Fatal and serious hypersensitivity reactions including anaphylaxis have occurred in patients receiving Feraheme. Initial symptoms may include hypotension, syncope, unresponsiveness, cardiac/cadiorespiratory arrest.

- Only administer Feraheme as an intravenous infusion over at least 15 minutes and only when personnel and therapies are immediately available for the treatment of anaphylaxis and other hypersensitivity reactions.
- Observe for signs or symptoms of hypersensitivity reactions during and for at least 30 minutes following Feraheme infusion including monitoring of blood pressure and pulse during and after Feraheme administration.
- Hypersensitivity reactions have occurred in patients in whom a previous Feraheme dose was tolerated.

Please see additional **Important Safety Information** in this document and accompanying full **Prescribing Information**, including **Boxed Warning**, in pocket.
Feraheme Assist™ provides individualized support for patients

Dedicated Care Managers connect patients to support throughout their treatment journey.

Benefits Investigation Support

Each payer is unique and insurance benefits can vary. Feraheme Assist has the skill to help navigate the prescription approval process for patients who need FERAHEME therapy. A Care Manager will complete the benefits investigation, support the prior authorization process (when applicable), and offer help if you need to appeal a denial.

Reimbursement Support

After confirming coverage, a Care Manager will proactively provide plan-specific requirements for your patient and can offer information on FERAHEME coding and billing, if needed.

Patient Assistance Program

AMAG Pharmaceuticals is committed to helping patients access their FERAHEME treatment. Eligible uninsured and commercially underinsured patients may receive a single course of therapy at no cost.* To assess eligibility, complete Step 6 on the enrollment form and submit to Feraheme Assist.

To get started, connect with Feraheme Assist

Submit a completed enrollment form to Feraheme Assist via fax (877-591-2505)

- Download the enrollment form at Feraheme.com
- Include a copy of both sides of the patient insurance card(s)
  - If your patient is uninsured or commercially underinsured, complete Step 6 of the enrollment form to have your patient evaluated for participation in the Patient Assistance Program
- Ensure both you and your patient have signed the form
  - If needed, patients can sign a stand-alone authorization at www.allcareconsent.com

Upon receipt of the Feraheme Assist Enrollment Form:

- You will receive a fax confirming receipt of the enrollment form
- You will receive a detailed summary of your patient’s benefits once coverage is verified

Have questions? Connect with us.
Translation services in >250 languages to help ensure language is not a barrier.
844-635-2624 (Monday - Friday, 8 AM – 8 PM ET)
info@ferahemeassist.com

*Each patient’s eligibility is evaluated on an individual basis. To be eligible, patients must meet the FDA-approved indication for FERAHEME. In compliance with federal regulations, patients insured by a government-funded program (Medicaid, Medicare, TRICARE, etc) are not eligible. Patient must be at or below 500% federal poverty level based on residency to participate.

This program may be discontinued or modified at any time based on eligibility, state and local laws, and program availability. For full program eligibility, restrictions, and enrollment requirements, please contact Feraheme Assist.

Feraheme Important Safety Information (cont’d)

Contraindications

Feraheme is contraindicated in patients with known hypersensitivity to Feraheme or any of its components or a history of allergic reaction to any intravenous iron product.
Feraheme Assist: Working together to support patients’ FERAHEME treatment journey

Feraheme Important Safety Information (cont’d)

Warnings and Precautions

Hypersensitivity: In addition to the fatal and serious adverse reactions in the Boxed Warning, other adverse reactions associated with hypersensitivity have occurred (pruritus, rash, urticaria, and wheezing). Allergic reactions have occurred following the first dose or subsequent doses in patients in whom a previous dose was tolerated. Patients with a history of multiple drug allergies may have a greater risk of anaphylaxis with parenteral iron products. Carefully consider the potential risks and benefits before administering Feraheme to these patients. Elderly patients with multiple or serious co-morbidities who experience hypersensitivity reactions and/or hypotension following administration of Feraheme may have more severe outcomes.

Hypotension: Feraheme may cause clinically significant hypotension. Monitor patients for signs and symptoms of hypotension following each Feraheme administration.

Iron Overload: Excessive therapy with parenteral iron can lead to excess storage of iron with the possibility of iatrogenic hemosiderosis. Regularly monitor the hematologic response during parenteral iron therapy. Do not administer Feraheme to patients with iron overload.

Magnetic Resonance (MR) Imaging Test Interference: Administration of Feraheme may transiently affect the diagnostic ability of MR imaging. Alteration of MR imaging studies may persist for up to 3 months following the last Feraheme dose. Maximum alteration of vascular MR imaging is anticipated to be evident for 1 – 2 days following Feraheme administration.

Adverse Reactions

The most common adverse reactions (≥ 2%) are diarrhea, headache, nausea, dizziness, hypotension, constipation, and peripheral edema.

Please see additional Important Safety Information in this document and accompanying full Prescribing Information, including Boxed Warning, in pocket.