

# HOSPITAL OUTPATIENT BILLING AND CODING GUIDE

Medicare, Medicaid, and Commercial

### Pharmacosmos Therapeutics Inc.

120 Headquarters Plaza East Tower, 6th Floor Morristown, NJ 07960

**Updated: June 2023** 

### **INDICATIONS**

Monoferric 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
- who have non-hemodialysis dependent chronic kidney disease (NDD-CKD)

Please see Important Safety Information throughout and full Prescribing Information.

### **HOSPITAL OUTPATIENT BILLING AND CODING**



### Medicare, Medicaid, and Commercial

### Important information

The coding, coverage, and payment information contained herein is gathered from various resources, general in nature, and subject to change without notice. Third-party payment for medical products and services is affected by numerous factors. It is always the provider's responsibility to determine the appropriate healthcare setting, and to submit true and correct claims for those products and services rendered. Providers should contact third-party payers for specific information on their coding, coverage, and payment policies. Information and materials are provided to assist healthcare providers, but the responsibility to determine coverage, reimbursement, and appropriate coding for a particular patient and/or procedure remains at all times with the provider.

International Classification of Diseases, 10th Revision, Clinical Modification (ICD-10-CM) diagnosis codes
The following tables display selected diagnosis codes that may be associated with iron deficiency anemia (IDA).\*

### Primary diagnosis codes

ICD-10-CM¹ diagnosis code	Description		
D50.0	IDA secondary to blood loss (chronic)		
D50.1	Sideropenic dysphagia		
D50.8	Other IDAs		
D50.9	IDA, unspecified		
D63.0	Anemia in neoplastic disease		
D63.1	Anemia in CKD  Code CKD stage first  Confirm iron deficiency		
D63.8	Anemia in other chronic diseases classified elsewhere  Code underlying disease first  Confirm iron deficiency		
D64.81	Antineoplastic chemotherapy-induced anemia  • Confirm iron deficiency		

#### Secondary diagnosis codes

ICD-10-CM¹ diagnosis code	Description	
E83.10	Iron metabolism	
K50.0-K50.919	Crohn's disease [regional enteritis]	
K51.0-K51.919	Ulcerative colitis	
K90.0	Celiac disease	
K90.4	Malabsorption due to intolerance not elsewhere classified	
K90.9	Intestinal malabsorption unspecified	
N18.1	CKD, stage 1	
N18.2	CKD, stage 2 (mild)	
N18.3	CKD, stage 3 (moderate)	
N18.30	CKD, stage 3 unspecified	
N18.31	CKD, stage 3a	
N18.32	CKD, stage 3b	
N18.4	CKD, stage 4 (severe)	
N18.5	CKD, stage 5	
N18.6	End-stage renal disease	
N18.9	CKD, unspecified	
N92.0	Excessive and frequent menstruation with regular cycle	
N92.5	Other specified irregular menstruation	
N92.6	Irregular menstruation, unspecified	
T45.4X5A	Adverse effect of iron and its compounds, initial encounter	
T45.4X5D	Adverse effect of iron and its compounds, secondary encounter	
T45.4X5S	Adverse effect of iron and its compounds, sequela encounter	
T50.905A	Adverse effect of unspecified drugs, medicaments and biological substances, initial encounter	

<sup>\*</sup>Sample diagnosis codes for the appropriate patient prescribed Monoferric

### IMPORTANT SAFETY INFORMATION CONTRAINDICATIONS

Monoferric is contraindicated in patients with a history of serious hypersensitivity to Monoferric or any of its components. Reactions have included shock, clinically significant hypotension, loss of consciousness, and/or collapse.

## **HOSPITAL OUTPATIENT BILLING AND CODING** (cont'd)



### **Current Procedural Terminology (CPT) code**

CPT* code	Description		
96365 <sup>2</sup>	Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug)		

### Healthcare Common Procedure Coding System (HCPCS) level II codes

HCPCS code	Descriptor	Site of care	Additional information
J1437³	Injection, ferric derisomaltose, 10 mg	All sites of care	If required by the payer, include the N4 qualifier, National Drug Code (NDC), unit of measure qualifier, and amount administered to the patient in Box 43. Example: N473594931001ME1000

### Revenue codes

Revenue code	Description	Revenue code	Description	
0250	General pharmacy	0510	Clinic, general	
0260	IV therapy	0636	Pharmacy, drugs requiring detailed coding	

### **National Drug Code (NDC)**

The NDC is a unique 10-digit, 3-segment number. It is a universal product identifier for drugs in the United States present on all over-the-counter and prescription medication packages and inserts.

Many NDC numbers listed on drug packaging are in a 10-digit format. The NDC number is essential for proper claim processing when submitting claims for drugs used; however, to be recognized by payers, it must be formatted into an 11-digit, 5-4-2 sequence. This requires a 0 to be placed in a specific position to meet the 5-4-2 format requirement.<sup>4</sup> As not all NDC numbers are set up the same, the table below demonstrates how to achieve the 11-digit NDC code for Monoferric.

Please note, because many practice management systems automatically remove the hyphens, be sure they are excluded from submission on the claim.

10-digit format Trade name		Package strength NDC number		New format	NDC number for payer
5-4-1	Monoferric⁵	1000 mg iron/10 mL (100 mg/mL) single-dose vial⁵	73594-9310-1	5-4-2	73594-9310- <b>0</b> 1

### **Additional Information**

Only 1000 mg iron/10 mL (100 mg/mL) single-dose vial of Monoferric is available in the United States.

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### IMPORTANT SAFETY INFORMATION (cont'd) WARNINGS AND PRECAUTIONS

### **Hypersensitivity Reactions**

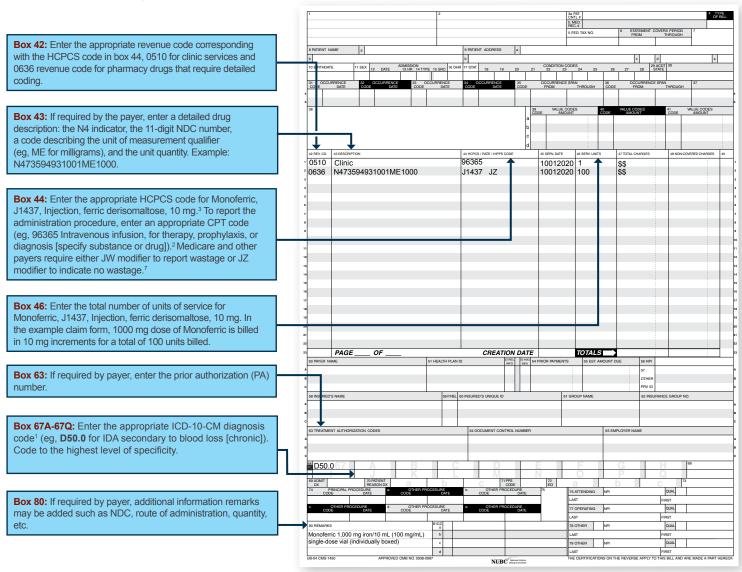
Serious hypersensitivity reactions, including anaphylactic-type reactions, some of which have been life-threatening and fatal, have been reported in patients receiving Monoferric. Patients may present with shock, clinically significant hypotension, loss of consciousness, and/or collapse. Monitor patients for signs and symptoms of hypersensitivity during and after Monoferric administration for at least 30 minutes and until clinically stable following completion of the infusion. Only administer Monoferric when personnel and therapies are immediately available for the treatment of serious hypersensitivity reactions. Monoferric is contraindicated in patients with prior serious hypersensitivity reactions to Monoferric or any of its components. In clinical trials in patients with IDA and CKD, serious or severe hypersensitivity were reported in 0.3% (6/2008) of the Monoferric treated subjects. These included 3 events of hypersensitivity in 3 patients; 2 events of infusion-related reactions in 2 patients and 1 event of asthma in one patient.

### SAMPLE UB-04 (CMS-1450) CLAIM FORM<sup>6</sup>



### Patient weight 50 kg or above: Administer 1000 mg of Monoferric as an intravenous infusion⁵

Note, only the 1000 mg iron/10 mL (100 mg/mL) single-dose vial of Monoferric is available in the United States. The UB-04 claim form (also known as CMS-1450) is the standard claim form to bill Medicare Fee-For-Service (FFS). This sample is intended to educate you on completing the form when billing for Monoferric. Although this sheet provides information that may facilitate the claims process, all coding information is for reference purposes only. Use of this sample claim form or the information in this sample claim form does not guarantee reimbursement of coverage.



Sample billing units calculation: For a 1000 mg dose of Monoferric, 100 billable units may be appropriate (1000 mg/10 mg per unit = 100)

Note: To facilitate accurate payment, report the exact dose administered.<sup>3</sup> More information on the claims process and the CMS fee schedule can be found on <a href="https://www.govinfo.gov/content/pkg/FR-2020-12-28/pdf/2020-26815.pdf">https://www.govinfo.gov/content/pkg/FR-2020-12-28/pdf/2020-26815.pdf</a>.

### IMPORTANT SAFETY INFORMATION (cont'd) WARNING AND PRECAUTIONS (cont'd)

#### Iron Overload

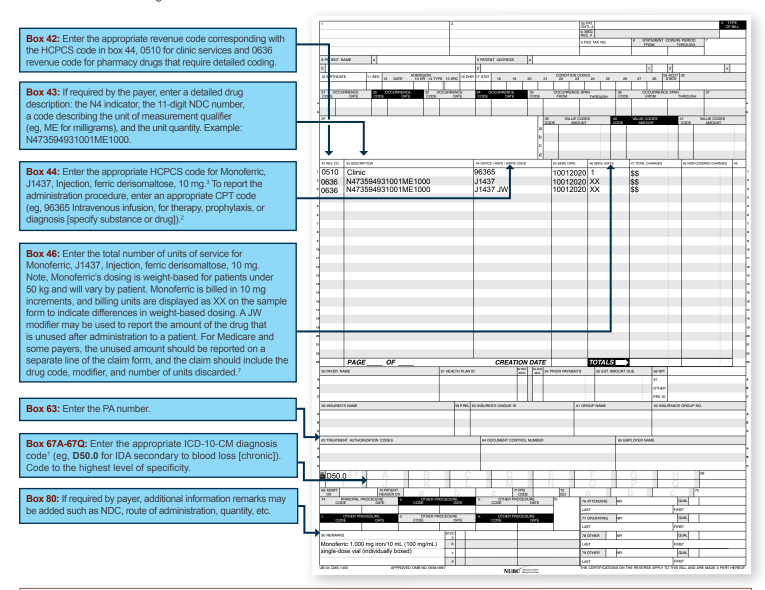
Excessive therapy with parenteral iron can lead to excess iron storage and possibly iatrogenic hemosiderosis or hemochromatosis. Monitor the hematologic response (hemoglobin and hematocrit) and iron parameters (serum ferritin and transferrin saturation) during parenteral iron therapy. Do not administer Monoferric to patients with iron overload.

### SAMPLE UB-04 (CMS-1450) CLAIM FORM<sup>6</sup>



### Patient weight less than 50 kg: Administer 20 mg/kg actual body weight as an intravenous infusion<sup>5</sup>

Note, only the 1000 mg iron/10 mL (100 mg/mL) single-dose vial of Monoferric is available in the United States. The UB-04 claim form (also known as CMS-1450) is the standard claim form to bill Medicare FFS. This sample is intended to educate you on completing the form when billing for Monoferric. Although this sheet provides information that may facilitate the claims process, all coding information is for reference purposes only. Use of this sample claim form or the information in this sample claim form does not guarantee reimbursement of coverage.



Sample billing units calculation: 20 mg/kg \* Y kg of body weight=20 \* Y mg administered. Then [20 \* Y] \* 1 billing unit/10 mg = [# Billing Units]

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Monoferric is available through the specialty pharmacy, Biologics by McKesson, if preferred by your office or required by your patient's health plan. Monoferric is also available through authorized distributors.

### IMPORTANT SAFETY INFORMATION (cont'd) ADVERSE REACTIONS

Adverse reactions were reported in 8.6% (172/2008) of patients treated with Monoferric. Adverse reactions related to treatment and reported by ≥1% of the treated patients were nausea (1.2%) and rash (1%). Adjudicated serious or severe hypersensitivity reactions were reported in 6/2008 (0.3%) patients in the Monoferric group. Hypophosphatemia (serum phosphate <2.0 mg/dL) was reported in 3.5% of Monoferric-treated patients in Trials 1 & 2.

To report adverse events, please contact Pharmacosmos at 1-888-828-0655. You may also contact the FDA at <a href="https://www.fda.gov/medwatch">www.fda.gov/medwatch</a> or call 1-800-FDA-1088.

### INDICATIONS AND IMPORTANT SAFETY INFORMATION

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References: 1. Centers for Medicare and Medicaid Services. 2023 ICD-10-CM. Accessed May 23, 2023. https://www.cms.gov/medicare/icd-10/2023-icd-10-cm 2. Find-A-Code. 96365 - CPT® Code in category: Intravenous infusion, for therapy, prophylaxis, or diagnosis (specify substance or drug). InnoviHealth Systems, Inc. Updated 2022. Accessed January 19, 2023. https://www.findacode.com/code.php?set=CPT&c=96365 3. Centers for Medicare & Medicaid Services. Healthcare Common Procedure Coding System (HCPCS) Application Summaries and Coding Decisions: Second Quarter, 2020 Coding Cycle for Drug and Biological Products. Accessed January 19, 2023. https://www.cms. gov/files/document/2020-hcpcs-application-summary-guarter-2-2020-drugs-and-biologicalsupdated-07312020.pdf 4. National Drug Code (NDC) Conversion Table. Converting NDCs from 10-digits to 11 digits. Accessed January 19, 2023. https://phpa.health.maryland.gov/OIDEOR/IMMUN/Shared%20Documents/Handout%203%20-%20NDC%20 conversion%20to%2011%20digits.pdf 5. Monoferric [Prescribing Information]. Morristown, NJ: Pharmacosmos Therapeutics Inc; 2023. 6. Centers for Medicare & Medicard Services. CMS Manual System. CMS 1450 (UB-04). Accessed January 19, 2023. https://www.cms.gov/Regulations-and-Guidance/Guidance/Manuals/Downloads/clm104c25. pdf 7. Centers for Medicare & Medicaid Services (CMS). Medicare program JW modifier: drug/biological amount discarded/not administered to any patient frequently asked questions. Accessed January 19, 2023. https://www.cms.gov/Medicare/Medicare-Fee-for-Service-Payment/HospitalOutpatientPPS/Downloads/JW-Modifier-FAQs.pdf





1-800-992-9022 | Monday-Friday, 8 AM to 8 PM ET



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