
HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use HEPARIN SODIUM INJECTION safely and effectively. See full prescribing information for HEPARIN SODIUM INJECTION.

HEPARIN SODIUM INJECTION, for intravenous or subcutaneous use Initial U.S. Approval: 1939

----- INDICATIONS AND USAGE

HEPARIN SODIUM INJECTION is an anticoagulant indicated for (1)

• Prophylaxis and treatment of venous thrombosis and pulmonary embolism

- Prevention of postoperative deep venous thrombosis and pulmonary embolism in patients undergoing major abdominothoracic surgery or who, for other reasons, are at risk of developing thromboembolic disease
- Atrial fibrillation with embolization
- Treatment of acute and chronic consumptive coagulopathies (disseminated intravascular coagulation)
- Prevention of clotting in arterial and cardiac surgery
- Prophylaxis and treatment of peripheral arterial embolism
- Use as an anticoagulant in blood transfusions, extracorporeal circulation, and dialysis procedures

----- DOSAGE AND ADMINISTRATION

Recommended Adult Dosages:

• Therapeutic Anticoagulant Effect with Full-Dose Heparin† (2.3)

Deep Subcutaneous (Intrafat) Injection Use a different site for each	Initial Dose	5,000 units by intravenous injection followed by 10,000 to 20,000 units of a concentrated solution, subcutaneously
injection	Every 8 hours or Every 12 hours	8,000 to 10,000 units of a concentrated solution 15,000 to 20,000 units of a concentrated solution
	Initial Dose	10,000 units, either undiluted or in 50 to 100 mL of 0.9% Sodium Chloride Injection, USP
Intermittent Intravenous Injection	Every 4 to 6 hours	5,000 to 10,000 units, either undiluted or in 50 to 100 mL of 0.9% Sodium Chloride Injection, USP
	Initial Dose	5,000 units by intravenous injection
Intravenous Infusion		20,000 to 40,000 units/24 hours in 1,000 mL of 0.9% Sodium Chloride Injection, USP (or in any compatible solution) for infusion
† Based on 150 lb (68 kg) pati	Continuous	n laboratory monitoring
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----- DOSAGE FORMS AND STRENGTHS ------

(porcine), preservative free	Heparin Sodium Injection, USP (porcine) contains parabens
	1 mL multiple-dose vial contains 1,000 USP units
	10 mL multiple-dose vial contains 10,000 USP units

30 mL multiple-dose vial contains 30,000 USP units
1 mL multiple-dose vial contains 5,000 USP units
1 mL multiple-dose vial contains 10,000 USP units
5 mL multiple-dose vial contains 50,000 USP units
1 mL multiple-dose vial contains 20,000 USP units

- History of heparin-induced thrombocytopenia (HIT) or heparin-induced thrombocytopenia and thrombosis (HITTS) (4)
- Known hypersensitivity to heparin or pork products (4)
- In whom suitable blood coagulation tests cannot be performed at appropriate intervals (4)
- An uncontrolled bleeding state, except when this is due to disseminated intravascular coagulation (4)

······ WARNINGS AND PRECAUTIONS ······

- Fatal Medication Errors: Confirm choice of correct strength prior to administration (5.1)
- Hemorrhage: Hemorrhage, including fatal events, has occurred in patients receiving heparin. Use caution in conditions with increased risk of hemorrhage (5.2)
- HIT and HITTS: Monitor for signs and symptoms and discontinue if indicative of HIT and HITTS (5.3)
- Benzyl Alcohol Toxicity: Use preservative-free formulation in neonates and infants
- Monitoring: Blood coagulation tests guide therapy for full-dose heparin. Periodically monitor platelet count, hematocrit, and occult blood in stool in all patients receiving heparin (5.5, 5.6)

ADVERSE REACTIONS

Most common adverse reactions are hemorrhage, thrombocytopenia, HIT and HITTS, injection site irritation, general hypersensitivity reactions, and elevations of aminotransferase levels. (6.1) **To report SUSPECTED ADVERSE REACTIONS, contact Fresenius Kabi USA, LLC at 1-800-551-7176 or FDA at 1-800-FDA-1088 or** <u>www.fda.gov/medwatch</u>.

----- DRUG INTERACTIONS ------

Drugs that interfere with coagulation, platelet aggregation or drugs that counteract coagulation may induce bleeding (7.2)

- Pregnancy: Preservative-free formulation recommended. (8.1)
- Lactation: Preservative-free formulation recommended. (8.2)
- Pediatric Use: Use preservative-free formulation in neonates and infants. (8.4)

See 17 for PATIENT COUNSELING INFORMATION.

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FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

Heparin Sodium Injection is indicated for:

- Prophylaxis and treatment of venous thrombosis and pulmonary embolism;
- Prevention of postoperative deep venous thrombosis and pulmonary embolism in patients undergoing major abdominothoracic surgery or who, for other reasons, are at risk of developing thromboembolic disease;

- Atrial fibrillation with embolization;
- Treatment of acute and chronic consumptive coagulopathies (disseminated intravascular coagulation);
- Prevention of clotting in arterial and cardiac surgery;
- Prophylaxis and treatment of peripheral arterial embolism.
- Anticoagulant use in blood transfusions, extracorporeal circulation, and dialysis procedures.

2 DOSAGE AND ADMINISTRATION

2.1 Preparation for Administration

Confirm the choice of the correct Heparin Sodium Injection vial to ensure that the 1 mL vial is not confused with a "catheter lock flush" vial or other 1 mL vial of incorrect strength [see Warnings and Precautions (5.1)]. Confirm the selection of the correct formulation and strength prior to administration of the drug.

Inspect parenteral drug products visually for particulate matter and discoloration prior to administration, whenever solution and container permit. Use only if solution is clear and the seal is intact. Do not use if solution is discolored or contains a precipitate.

When heparin is added to an infusion solution for continuous intravenous administration, the container should be inverted at least six times to ensure adequate mixing and prevent pooling of the heparin in the solution.

Administer Heparin Sodium Injection by intermittent intravenous injection, intravenous infusion, or deep subcutaneous (intrafat, i.e., above the iliac crest or abdominal fat layer) injection. The intramuscular route of administration should be avoided because of the frequent occurrence of hematoma at the injection site [see Adverse Reactions (6.1)].

2.2 Laboratory Monitoring for Efficacy and Safety

Adjust the dosage of Heparin Sodium Injection according to the patient's coagulation test results. Dosage is considered adequate when the activated partial thromboplastin time (aPTT) is 1.5 to 2 times normal or when the whole blood clotting time is elevated approximately 2.5 to 3 times the control value. When initiating treatment with Heparin Sodium Injection by continuous intravenous infusion, determine the coagulation status (aPTT, INR, platelet count) at baseline and continue to follow aPTT approximately every 4 hours and then at appropriate intervals thereafter. When the drug is administered intermittently by intravenous injection, perform coagulation tests before each injection during the initiation of treatment and at appropriate intervals thereafter. After deep subcutaneous (intrafat) injections, tests for adequacy of dosage are best performed on samples drawn 4 to 6 hours after the injection.

Periodically monitor platelet counts, hematocrit, and occult blood in stool during the entire course of heparin therapy, regardless of the route of administration.

2.3 Therapeutic Anticoagulant Effect with Full-Dose Heparin

The dosing recommendations in Table 1 are based on clinical experience. Although dosages must be adjusted for the individual patient according to the results of suitable laboratory tests, the following dosage schedules may be used as guidelines:

Table 1: Recommended Adult Full-Dose Heparin Regimens for Therapeutic Anticoagulant Effect

METHOD OF ADMINISTRATION	FREQUENCY	RECOMMENDED DOSE
Deep Subcutaneous (Intrafat) Injection Use a different site for each injection to prevent the	Initial Dose	5,000 units by intravenous injection, followed by 10,000 to 20,000 units of a concentrated solution, subcutaneously
development of hematoma	Every 8 hours or	8,000 to 10,000 units of a concentrated solution
	Every 12 hours	15,000 to 20,000 units of a concentrated solution
Intermittent Intravenous Injection	Initial Dose	10,000 units, either undiluted or in 50 to 100 mL of 0.9% Sodium Chloride Injection, USP
	Every 4 to 6 hours	5,000 to 10,000 units, either undiluted or in 50 to 100 mL of 0.9% Sodium Chloride Injection, USP
Continuous Intravenous Infusion	Initial Dose	5,000 units by intravenous injection
	Continuous	20,000 to 40,000 units/24 hours in 1,000 mL of 0.9% Sodium Chloride Injection, USP (or in any compatible solution) for infusion

*Based on 68 kg patient

2.4 Pediatric Dosing

Use preservative-free HEPARIN SODIUM INJECTION in neonates and infants [see Warnings and Precautions (5.4)].

There are no adequate and well controlled studies on heparin use in pediatric patients. Pediatric dosing recommendations are based on clinical experience. In general, the following dosage schedule may be used as a guideline in pediatric patients:

Recommended Pediatric Use	
Initial Dose	75 to 100 units/kg (IV bolus over 10 minutes)
Maintenance Dose	Infants: 25 to 30 units/kg/hour; Infants < 2 months have the highest requirements (average 28 units/kg/hour) Children > 1 year of age: 18 to 20

	units/kg/hour; Older children may require less heparin, similar to weight-adjusted adult dosage
Monitoring	Adjust heparin to maintain aPTT of 60 to 85 seconds, assuming this reflects an anti- Factor Xa level of 0.35 to 0.70

2.5 Cardiovascular Surgery

Patients undergoing total body perfusion for open-heart surgery should receive an initial dose of not less than 150 units of heparin sodium per kilogram of body weight. Frequently, a dose of 300 units per kilogram is used for procedures estimated to last less than 60 minutes, or 400 units per kilogram for those estimated to last longer than 60 minutes.

2.6 Low-Dose Prophylaxis of Postoperative Thromboembolism

The most widely used dosage has been 5,000 units 2 hours before surgery and 5,000 units every 8 to 12 hours thereafter for 7 days or until the patient is fully ambulatory, whichever is longer. Administer the heparin by deep subcutaneous (intrafat, i.e., above the iliac crest or abdominal fat layer, arm, or thigh) injection with a fine (25 to 26-gauge) needle to minimize tissue trauma.

2.7 Converting to Warfarin

To ensure continuous anticoagulation when converting from Heparin Sodium Injection to warfarin, continue full heparin therapy for several days until the INR (prothrombin time) has reached a stable therapeutic range. Heparin therapy may then be discontinued without tapering [see Drug Interactions (7.1)].

2.8 Converting to Oral Anticoagulants other than Warfarin

For patients currently receiving intravenous heparin, stop intravenous infusion of heparin sodium immediately after administering the first dose of oral anticoagulant; or for intermittent intravenous administration of heparin sodium, start oral anticoagulant 0 to 2 hours before the time that the next dose of heparin was to have been administered.

2.9 Extracorporeal Dialysis

Follow equipment manufacturers' operating directions carefully. A dose of 25 to 30 units/kg followed by an infusion rate of 1,500 to 2,000 units/hour is suggested based on pharmacodynamic data if specific manufacturers' recommendations are not available.

3 DOSAGE FORMS AND STRENGTHS

Heparin Sodium Injection, USP is available as:

Heparin Sodium Injection, USP, preservative free, is available as follows:

- 2,000 USP units per 2 mL, single-dose vial
- 5,000 USP units per 0.5 mL, single-dose vial

Heparin Sodium Injection, USP contains **benzyl alcohol** and is available as follows:

- 50,000 USP units per 10 mL, multiple-dose vial
- 40,000 USP units per 4 mL, multiple-dose vial

Heparin Sodium Injection, USP contains **parabens** and is available as follows:

- 1,000 USP units per mL, multiple-dose vial
- 10,000 USP units per 10 mL, multiple-dose vial
- 30,000 USP units per 30 mL, multiple-dose vial
- 5,000 USP units per mL, multiple-dose vial
- 10,000 USP units per mL, multiple-dose vial
- 50,000 USP units per 5 mL, multiple-dose vial
- 20,000 USP units per mL, multiple-dose vial

4 CONTRAINDICATIONS

The use of Heparin Sodium Injection is contraindicated in patients with the following conditions:

- History of heparin-induced thrombocytopenia and heparin-induced thrombocytopenia and thrombosis [see Warnings and Precautions (5.3)];
- Known hypersensitivity to heparin or pork products (e.g., anaphylactoid reactions) [see Adverse Reactions (6.1)];
- In whom suitable blood coagulation tests, e.g., the whole blood clotting time, partial thromboplastin time, etc., cannot be performed at appropriate intervals (this contraindication refers to full-dose heparin; there is usually no need to monitor coagulation parameters in patients receiving low-dose heparin);
- An uncontrolled active bleeding state [see Warnings and Precautions (5.4)], except when this is due to disseminated intravascular coagulation.

5 WARNINGS AND PRECAUTIONS

5.1 Fatal Medication Errors

Do not use Heparin Sodium Injection as a "catheter lock flush" product. Heparin Sodium Injection is supplied in vials containing various strengths of heparin, including vials that contain a highly concentrated solution of 10,000 units in 1 mL. Fatal hemorrhages have occurred in pediatric patients due to medication errors in which 1 mL Heparin Sodium Injection vials were confused with 1 mL "catheter lock flush" vials. Carefully examine all Heparin Sodium Injection vials to confirm the correct vial choice prior to administration of the drug.

5.2 Hemorrhage

Avoid using heparin in the presence of major bleeding, except when the benefits of heparin therapy outweigh the potential risks.

Hemorrhage can occur at virtually any site in patients receiving heparin. Fatal hemorrhages have occurred. Adrenal hemorrhage (with resultant acute adrenal insufficiency), ovarian hemorrhage, and retroperitoneal hemorrhage have occurred during anticoagulant therapy with heparin [see Adverse Reactions (6.1)]. A higher incidence of bleeding has been reported in patients, particularly women, over 60 years of age [see Clinical Pharmacology (12.3)]. An unexplained fall in hematocrit, fall in blood

pressure or any other unexplained symptom should lead to serious consideration of a hemorrhagic event.

Use heparin sodium with caution in disease states in which there is increased risk of hemorrhage, including:

- Cardiovascular Subacute bacterial endocarditis, severe hypertension.
- Surgical During and immediately following (a) spinal tap or spinal anesthesia or (b) major surgery, especially involving the brain, spinal cord, or eye.
- Hematologic Conditions associated with increased bleeding tendencies, such as hemophilia, thrombocytopenia and some vascular purpuras.
- Patients with hereditary antithrombin III deficiency receiving concurrent antithrombin III therapy
 - The anticoagulant effect of heparin is enhanced by concurrent treatment with antithrombin III (human) in patients with hereditary antithrombin III deficiency. To reduce the risk of bleeding, reduce the heparin dose during concomitant treatment with antithrombin III (human).
- Gastrointestinal Ulcerative lesions and continuous tube drainage of the stomach or small intestine.
- Other Menstruation, liver disease with impaired hemostasis.

5.3 Heparin-Induced Thrombocytopenia and Heparin-Induced Thrombocytopenia and Thrombosis

Heparin-induced thrombocytopenia (HIT) is a serious antibody-mediated reaction. HIT occurs in patients treated with heparin and is due to the development of antibodies to a platelet Factor 4-heparin complex that induce *in vivo* platelet aggregation. HIT may progress to the development of venous and arterial thromboses, a condition referred to as heparin-induced thrombocytopenia with thrombosis (HITT). Thrombotic events may also be the initial presentation for HITT. These serious thromboembolic events include deep vein thrombosis, pulmonary embolism, cerebral vein thrombosis, limb ischemia, stroke, myocardial infarction, mesenteric thrombosis, renal arterial thrombosis, skin necrosis, gangrene of the extremities that may lead to amputation, and possibly death. If the platelet count falls below 100,000/mm³ or if recurrent thrombosis develops, promptly discontinue heparin, evaluate for HIT and HITT, and, if necessary, administer an alternative anticoagulant.

HIT or HITT can occur up to several weeks after the discontinuation of heparin therapy. Patients presenting with thrombocytopenia or thrombosis after discontinuation of heparin sodium should be evaluated for HIT or HITT.

5.4 Risk of Serious Adverse Reactions in Infants Due to Benzyl Alcohol Preservative

Serious and fatal adverse reactions including "gasping syndrome" can occur in neonates and infants treated with benzyl alcohol-preserved drugs, including Heparin Sodium Injection multiple-dose vials. The "gasping syndrome" is characterized by central nervous system depression, metabolic acidosis, and gasping respirations.

When prescribing Heparin Sodium Injection multiple-dose vials in infants consider the combined daily metabolic load of benzyl alcohol from all sources including Heparin

Sodium Injection multiple-dose vials and other drugs containing benzyl alcohol. The minimum amount of benzyl alcohol at which toxicity may occur is not known [see Use in Specific Populations (8.4)].

5.5 Thrombocytopenia

Thrombocytopenia in patients receiving heparin has been reported at frequencies up to 30%. It can occur 2 to 20 days (average 5 to 9) following the onset of heparin therapy. Obtain platelet counts before and periodically during heparin therapy. Monitor thrombocytopenia of any degree closely. If the count falls below 100,000/mm³ or if recurrent thrombosis develops, promptly discontinue heparin, evaluate for HIT and HITT, and, if necessary, administer an alternative anticoagulant [see Warnings and Precautions (5.3)].

5.6 Coagulation Testing and Monitoring

When using a full dose heparin regimen, adjust the heparin dose based on frequent blood coagulation tests. If the coagulation test is unduly prolonged or if hemorrhage occurs, discontinue heparin promptly [see Overdosage (10)]. Periodically monitor platelet counts, hematocrit, and occult blood in stool during the entire course of heparin therapy, regardless of the route of administration [see Dosage and Administration (2.2)].

5.7 Heparin Resistance

Resistance to heparin is frequently encountered in fever, thrombosis, thrombophlebitis, infections with thrombosing tendencies, myocardial infarction, cancer, in postsurgical patients, and patients with antithrombin III deficiency. Close monitoring of coagulation tests is recommended in these cases. Adjustment of heparin doses based on anti-Factor Xa levels may be warranted.

5.8 Hypersensitivity

Patients with documented hypersensitivity to heparin should be given the drug only in clearly life-threatening situations [see Adverse Reactions (6.1)].

Because Heparin Sodium Injection is derived from animal tissue, it should be used with caution in patients with a history of allergy.

6 ADVERSE REACTIONS

The following clinically significant adverse reactions are described elsewhere in the labeling:

- Hemorrhage [see Warnings and Precautions (5.2)]
- Heparin-Induced Thrombocytopenia and Heparin-Induced Thrombocytopenia and Thrombosis [see Warnings and Precautions (5.3)]
- Risk of Serious Adverse Reactions in Infants Due to Benzyl Alcohol Preservative [see Warnings and Precautions (5.4)]
- Thrombocytopenia [see Warnings and Precautions (5.5)]
- Heparin Resistance [see Warnings and Precautions (5.7)]
- Hypersensitivity [see Warnings and Precautions (5.8)]

6.1 Postmarketing Experience

The following adverse reactions have been identified during post approval use of Heparin Sodium Injection. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

- Hemorrhage–Hemorrhage is the chief complication that may result from heparin therapy [see Warnings and Precautions (5.2)]. Gastrointestinal or urinary tract bleeding during anticoagulant therapy may indicate the presence of an underlying occult lesion. Bleeding can occur at any site but certain specific hemorrhagic complications may be difficult to detect including:
 - Adrenal hemorrhage, with resultant acute adrenal insufficiency, has occurred with heparin therapy, including fatal cases.
 - Ovarian (corpus luteum) hemorrhage developed in a number of women of reproductive age receiving short- or long-term heparin therapy.
 - Retroperitoneal hemorrhage
- HIT and HITT, including delayed onset cases [see Warnings and Precautions (5.3)].
- Local Irritation Local irritation, erythema, mild pain, hematoma or ulceration may follow deep subcutaneous (intrafat) injection of heparin sodium. Because these complications are much more common after intramuscular use, the intramuscular route is not recommended.
- Histamine-like reactions Such reactions have been observed at the site of injections. Necrosis of the skin has been reported at the site of subcutaneous injection of heparin, occasionally requiring skin grafting [see Warnings and Precautions (5.3)].
- Hypersensitivity Generalized hypersensitivity reactions have been reported, with chills, fever and urticaria as the most usual manifestations, and asthma, rhinitis, lacrimation, headache, nausea and vomiting, and anaphylactoid reactions, including shock, occurring less frequently. Itching and burning, especially on the plantar side of the feet, may occur. [see Warnings and Precautions (5.8)]
- Elevations of aminotransferases Significant elevations of aspartate aminotransferase (AST) and alanine aminotransferase (ALT) levels have occurred in patients who have received heparin.
- Others Osteoporosis following long-term administration of high doses of heparin, cutaneous necrosis after systemic administration, suppression of aldosterone synthesis, delayed transient alopecia, priapism, and rebound hyperlipemia on discontinuation of heparin sodium have also been reported.

7 DRUG INTERACTIONS

7.1 Oral Anticoagulants

Heparin sodium may prolong the one-stage prothrombin time. Therefore, when heparin sodium is given with dicumarol or warfarin sodium, a period of at least 5 hours after the last intravenous dose or 24 hours after the last subcutaneous dose should elapse before blood is drawn, if a valid prothrombin time is to be obtained.

7.2 Platelet Inhibitors

Drugs such as NSAIDS (including salicylic acid, ibuprofen, indomethacin, and celecoxib), dextran, phenylbutazone, thienopyridines, dipyridamole, hydroxychloroquine, glycoprotein IIb/IIIa antagonists (including abciximab, eptifibatide, and tirofiban), and

others that interfere with platelet-aggregation reactions (the main hemostatic defense of heparinized patients) may induce bleeding and should be used with caution in patients receiving heparin sodium. To reduce the risk of bleeding, a reduction in the dose of antiplatelet agent or heparin is recommended.

7.3 Other Interactions

Digitalis, tetracyclines, nicotine or antihistamines may partially counteract the anticoagulant action of heparin sodium. Intravenous nitroglycerin administered to heparinized patients may result in a decrease of the partial thromboplastin time with subsequent rebound effect upon discontinuation of nitroglycerin. Careful monitoring of partial thromboplastin time and adjustment of heparin dosage are recommended during coadministration of heparin and intravenous nitroglycerin.

Antithrombin III (human) – The anticoagulant effect of heparin is enhanced by concurrent treatment with antithrombin III (human) in patients with hereditary antithrombin III deficiency. To reduce the risk of bleeding, a reduced dosage of heparin is recommended during treatment with antithrombin III (human).

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

<u>Risk Summary</u>

There are no available data on heparin sodium use in pregnant women to inform a drugassociated risk of major birth defects and miscarriage. In published reports, heparin exposure during pregnancy did not show evidence of an increased risk of adverse maternal or fetal outcomes in humans. No teratogenicity, but early embryo-fetal death was observed in animal reproduction studies with administration of heparin sodium to pregnant rats and rabbits during organogenesis at doses approximately 10 times the maximum recommended human dose (MRHD) of 45,000 units/day (*see Data*). Consider the benefits and risks of Heparin Sodium Injection for the mother and possible risks to the fetus when prescribing Heparin Sodium Injection to a pregnant woman.

If available, preservative-free Heparin Sodium Injection is recommended when heparin therapy is needed during pregnancy. There are no known adverse outcomes associated with fetal exposure to the preservative benzyl alcohol through maternal drug administration; however, the preservative benzyl alcohol can cause serious adverse events and death when administered intravenously to neonates and infants [see Use in Specific Populations (8.4)].

The estimated background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2-4% and 15-20%, respectively.

<u>Data</u>

Human Data

The maternal and fetal outcomes associated with uses of heparin via various dosing

methods and administration routes during pregnancy have been investigated in numerous studies. These studies generally reported normal deliveries with no maternal or fetal bleeding and no other complications.

Animal Data

In a published study conducted in rats and rabbits, pregnant animals received heparin intravenously during organogenesis at a dose of 10,000 units/kg/day, approximately 10 times the maximum human daily dose based on body weight. The number of early resorptions increased in both species. There was no evidence of teratogenic effects.

8.2 Lactation

Risk Summary

If available, preservative-free Heparin Sodium Injection is recommended when heparin therapy is needed during lactation. Benzyl alcohol present in maternal serum is likely to cross into human milk and may be orally absorbed by a breastfed infant. There is no information regarding the presence of Heparin Sodium Injection in human milk, the effects on the breastfed infant, or the effects on milk production. Due to its large molecular weight, heparin is not likely to be excreted in human milk, and any heparin in milk would not be orally absorbed by a breastfed infant. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for Heparin Sodium Injection and any potential adverse effects on the breastfed infant from Heparin Sodium Injection or from the underlying maternal condition [see Use in Specific Populations (8.4)].

8.4 Pediatric Use

There are no adequate and well controlled studies on heparin use in pediatric patients. Pediatric dosing recommendations are based on clinical experience [see Dosage and Administration (2.4)].

Carefully examine all Heparin Sodium Injection vials to confirm choice of the correct strength prior to administration of the drug. Pediatric patients, including neonates, have died as a result of medication errors in which Heparin Sodium Injection vials have been confused with "catheter lock flush" vials [see Warnings and Precautions (5.1)].

Benzyl Alcohol Toxicity

Use preservative-free Heparin Sodium Injection in neonates and infants.

Serious adverse reactions including fatal reactions and the "gasping syndrome" occurred in premature neonates and infants in the neonatal intensive care unit who received drugs containing benzyl alcohol as a preservative. In these cases, benzyl alcohol dosages of 99 to 234 mg/kg/day produced high levels of benzyl alcohol and its metabolites in the blood and urine (blood levels of benzyl alcohol were 0.61 to 1.378 mmol/L). Additional adverse reactions included gradual neurological deterioration, seizures, intracranial hemorrhage, hematologic abnormalities, skin breakdown, hepatic and renal failure, hypotension, bradycardia, and cardiovascular collapse. Preterm, low-birth weight infants may be more likely to develop these reactions because they may be less able to metabolize benzyl alcohol.

8.5 Geriatric Use

There are limited adequate and well-controlled studies in patients 65 years and older, however, a higher incidence of bleeding has been reported in patients, particularly women, over 60 years of age [see Warnings and Precautions (5.2)]. Patients over 60 years of age may require lower doses of heparin. Lower doses of heparin may be indicated in these patients [see Clinical Pharmacology (12.3)].

10 OVERDOSAGE

Bleeding is the chief sign of heparin overdosage.

Neutralization of Heparin Effect

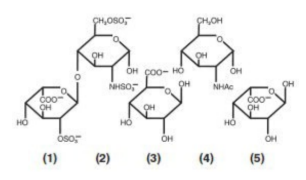
When clinical circumstances (bleeding) require reversal of the heparin effect, protamine sulfate (1% solution) by slow infusion will neutralize heparin sodium. **No more than 50 mg** should be administered, **very slowly**, in any 10-minute period. Each mg of protamine sulfate neutralizes approximately 100 USP heparin units. The amount of protamine required decreases over time as heparin is metabolized. Although the metabolism of heparin is complex, it may, for the purpose of choosing a protamine dose, be assumed to have a half-life of about 1/2 hour after intravenous injection. Because fatal reactions often resembling anaphylaxis have been reported with protamine, it should be given only when resuscitation techniques and treatment of anaphylactoid shock are readily available. For additional information consult the labeling of Protamine Sulfate Injection.

11 DESCRIPTION

Heparin is a heterogeneous group of straight-chain anionic mucopolysaccharides, called glycosaminoglycans, having anticoagulant properties. Although others may be present, the main sugars occurring in heparin are: (1) α -L-iduronic acid 2-sulfate, (2) 2-deoxy-2-sulfamino- α -D-glucose 6-sulfate, (3) β -D-glucuronic acid, (4) 2-acetamido-2-deoxy- α -D-glucose and (5) α -L-iduronic acid. These sugars are present in decreasing amounts, usually in the order (2)> (1)> (4)> (3)> (5), and are joined by glycosidic linkages, forming polymers of varying sizes. Heparin is strongly acidic because of its content of covalently linked sulfate and carboxylic acid groups. In heparin sodium, the acidic protons of the sulfate units are partially replaced by sodium ions.

Heparin Sodium Injection, USP is a sterile solution of heparin sodium derived from porcine intestinal mucosa, standardized for anticoagulant activity, in water for injection. It is to be administered by intravenous or deep subcutaneous routes. The potency is determined by a biological assay using a USP reference standard based on units of heparin activity per milligram.

Structure of Heparin Sodium (representative subunits):



Heparin Sodium Injection, USP (porcine), preservative free, is available as follows:

Each mL of the 1,000 units per mL preparation contains: 1,000 USP Heparin units (porcine); 9 mg sodium chloride; Water for Injection q.s. Made isotonic with sodium chloride. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

Each 0.5 mL of the 5,000 units per 0.5 mL preparation contains: 5,000 USP Heparin units (porcine); Water for Injection q.s. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

<u>Heparin Sodium Injection, USP (porcine), preserved with benzyl alcohol, is available as</u> <u>follows:</u>

Each mL of the 5,000 units per mL preparation contains: 5,000 USP Heparin units (porcine); 6 mg sodium chloride; 15 mg benzyl alcohol (as a preservative); Water for Injection q.s. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

Each mL of the 10,000 units per mL preparation contains: 10,000 USP Heparin units (porcine); 5 mg sodium chloride; 10.42 mg benzyl alcohol (as a preservative); Water for Injection q.s. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

Heparin Sodium Injection, USP (porcine), preserved with parabens, is available as follows:

Each mL of the 1,000 units per mL preparation contains: 1,000 USP Heparin units (porcine); 9 mg sodium chloride; 1.5 mg methylparaben; 0.15 mg propylparaben; Water for Injection q.s. Made isotonic with sodium chloride. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

Each mL of the 5,000 units per mL preparation contains: 5,000 USP Heparin units (porcine); 5 mg sodium chloride; 1.5 mg methylparaben; 0.15 mg propylparaben; Water for Injection q.s. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

Each mL of the 10,000 units per mL preparation contains: 10,000 USP Heparin units (porcine); 1.5 mg methylparaben; 0.15 mg propylparaben; Water for Injection q.s. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

Each mL of the 20,000 units per mL preparation contains: 20,000 USP Heparin units (porcine); 1.5 mg methylparaben; 0.15 mg propylparaben; Water for Injection q.s. Hydrochloric acid and/or sodium hydroxide may have been added for pH adjustment (5.0 to 7.5).

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Heparin interacts with the naturally occurring plasma protein, Antithrombin III, to induce a conformational change, which markedly enhances the serine protease activity of Antithrombin III, thereby inhibiting the activated coagulation factors involved in the clotting sequence, particularly Xa and IIa. Small amounts of heparin inhibit Factor Xa, and larger amounts inhibit thrombin (Factor IIa). Heparin also prevents the formation of a stable fibrin clot by inhibiting the activation of the fibrin stabilizing factor. Heparin does not have fibrinolytic activity; therefore, it will not lyse existing clots.

12.2 Pharmacodynamics

Various times (activated clotting time, activated partial thromboplastin time, prothrombin time, whole blood clotting time) are prolonged by full therapeutic doses of heparin; in most cases, they are not measurably affected by low doses of heparin. The bleeding time is usually unaffected by heparin.

12.3 Pharmacokinetics

Absorption

Heparin is not absorbed through the gastrointestinal tract and therefore administered via parenteral route. Peak plasma concentration and the onset of action are achieved immediately after intravenous administration.

Distribution

Heparin is highly bound to antithrombin, fibrinogens, globulins, serum proteases and lipoproteins. The volume of distribution is 0.07 L/kg.

Elimination

Metabolism

Heparin does not undergo enzymatic degradation.

Excretion

Heparin is mainly cleared from the circulation by liver and reticuloendothelial cells mediated uptake into extravascular space. Heparin undergoes biphasic clearance, a) rapid saturable clearance (zero order process due to binding to proteins, endothelial cells and macrophage) and b) slower first order elimination. The plasma half-life is dosedependent and it ranges from 0.5 to 2 h.

Specific Populations

Geriatric patients

Patients over 60 years of age, following similar doses of heparin, may have higher plasma levels of heparin and longer activated partial thromboplastin times (aPTTs) compared with patients under 60 years of age [see Use in Specific Populations (8.5)].

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

No long-term studies in animals have been performed to evaluate carcinogenic potential of heparin. Also, no reproduction studies in animals have been performed concerning mutagenesis or impairment of fertility.

16 HOW SUPPLIED/STORAGE AND HANDLING

Heparin Sodium Injection, USP (porcine), **preservative free**, is available as follows:

Product			
Code	Unit of Sale	Strength	Each
27602	NDC 63323-276-02	2,000 USP units per 2 mL	NDC 63323-276-01
	Unit of 25	(1,000 USP units per mL)	2 mL single-dose, flip-top vial
504313	NDC 63323-543-13	5,000 USP units per 0.5	NDC 63323-543-03
	Unit of 25	mL	0.5 mL fill in a 2 mL single-dose,
			flip-top vial

Use only if solution is clear and seal intact. Do not use if solution is discolored or contains a precipitate.

This container closure is not made with natural rubber latex. Discard unused portion.

Heparin Sodium Injection, USP (porcine) contains **benzyl alcohol** and is available as follows:

Product Code	Unit of Sale	Strength	Each
	NDC 63323-047-10 Unit of 25	50,000 USP units per 10 mL	NDC 63323-047-01 10 mL multiple-dose, flip-top
			vial
504514	NDC 63323-459-14	40,000 USP units per 4 mL	NDC 63323-459-04
	Unit of 25	(10,000 USP units per mL)	•
			dose, flip-top vial

Use only if solution is clear and seal intact. Do not use if solution is discolored or contains a precipitate. This container closure is not made with natural rubber latex.

Heparin Sodium Injection, USP (porcine) contains **parabens** and is available as follows:

Product Code	Unit of Sale	Strength	Each
504013	NDC 63323- 540-13 Unit of 25	1,000 USP units per mL	NDC 63323-540-03 1 mL fill in a 2 mL multiple-dose, flip-top vial
	NDC 63323- 540-15	'	NDC 63323-540-05 10 mL multiple-dose, flip-top vial

Unit of 25	(1,000 USP units per mL)	· · ·
NDC 65219- 066-10 Unit of 25	10,000 USP units per 10 mL (1,000 USP units per mL)	NDC 65219-066-01 10 mL multiple-dose, flip-top vial This product contains an RFID
NDC 63323- 540-36 Unit of 25	30,000 USP units per 30 mL (1,000 USP units per mL)	NDC 63323-540-33 30 mL multiple-dose, flip-top vial
NDC 63323- 262-06 Unit of 25	5,000 USP units per mL	NDC 63323-262-03 1 mL fill in a 2 mL multiple-dose, flip-top vial
NDC 63323- 542-13 Unit of 25	10,000 USP units per mL	NDC 63323-542-09 1 mL fill in a 2 mL multiple-dose, flip-top vial
NDC 63323- 542-14 Unit of 25	50,000 USP units per 5 mL (10,000 USP units per mL)	NDC 63323-542-04 5 mL fill in a 10 mL multiple-dose, flip-top vial
NDC 63323- 915-13 Unit of 25	20,000 USP units per mL	NDC 63323-915-03 1 mL fill in a 2 mL multiple-dose, flip-top vial

Use only if solution is clear and seal intact. Do not use if solution is discolored or contains a precipitate.

This container closure is not made with natural rubber latex.

STORAGE:

Store at 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature].

17 PATIENT COUNSELING INFORMATION

<u>Hemorrhage</u>

Inform patients that it may take them longer than usual to stop bleeding, that they may bruise and/or bleed more easily when they are treated with heparin, and that they should report any unusual bleeding or bruising to their physician. Hemorrhage can occur at virtually any site in patients receiving heparin. Fatal hemorrhages have occurred [see Warnings and Precautions (5.2)].

Prior to Surgery

Advise patients to inform physicians and dentists that they are receiving heparin before any surgery is scheduled [see Warnings and Precautions (5.2)].

Heparin-Induced Thrombocytopenia

Inform patients of the risk of heparin-induced thrombocytopenia (HIT). HIT may progress to the development of venous and arterial thromboses, a condition known as

heparin-induced thrombocytopenia and thrombosis (HITT). HIT and HITT can occur up to several weeks after the discontinuation of heparin therapy [see Warnings and *Precautions (5.3)*].

<u>Hypersensitivity</u>

Inform patients that generalized hypersensitivity reactions have been reported. Necrosis of the skin has been reported at the site of subcutaneous injection of heparin [see Warnings and Precautions (5.8), Adverse Reactions (6.1)].

Other Medications

Because of the risk of hemorrhage, advise patients to inform their physicians and dentists of all medications they are taking, including non-prescription medications, and before starting any new medication [see Drug Interactions (7.1)].

Lake Zurich, IL 60047 www.fresenius-kabi.com/us 451645B

PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Label

Not for Lock Flush

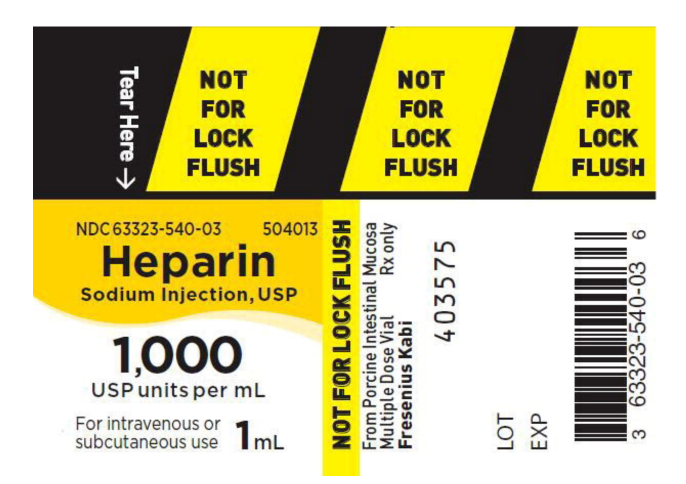
NDC 63323-540-03 504013

Heparin Sodium Injection, USP

1,000 USP units per mL

For intravenous or subcutaneous use

1 mL



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Tray Label

NDC 63323-540-13 504013

Heparin Sodium Injection, USP

1,000 USP units per mL

For intravenous or subcutaneous use

1 mL

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Label

Not for Lock Flush

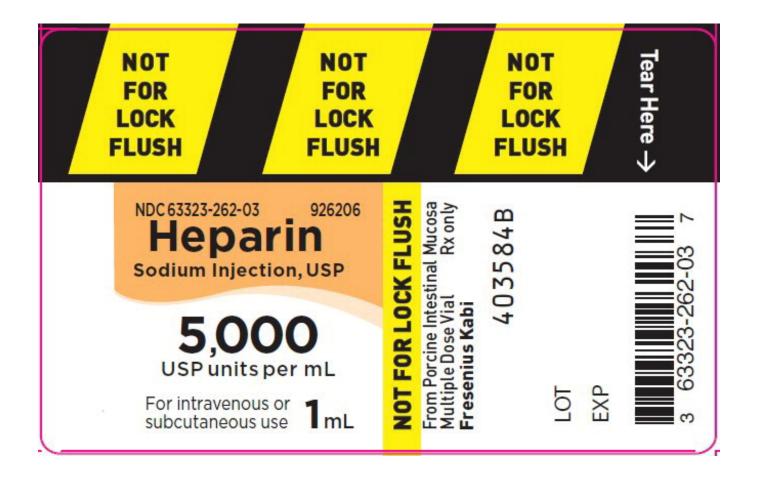
NDC 63323-262-03 926206

Heparin Sodium Injection, USP

5,000 USP units per mL

For intravenous or subcutaneous use

1 mL



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Tray Label

NDC 63323-262-06 926206

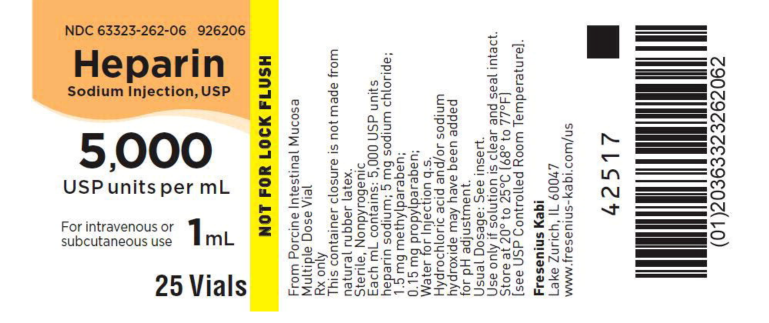
Heparin Sodium Injection, USP

5,000 USP units per mL

For intravenous or subcutaneous use

1 mL

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Label

Not for Lock Flush

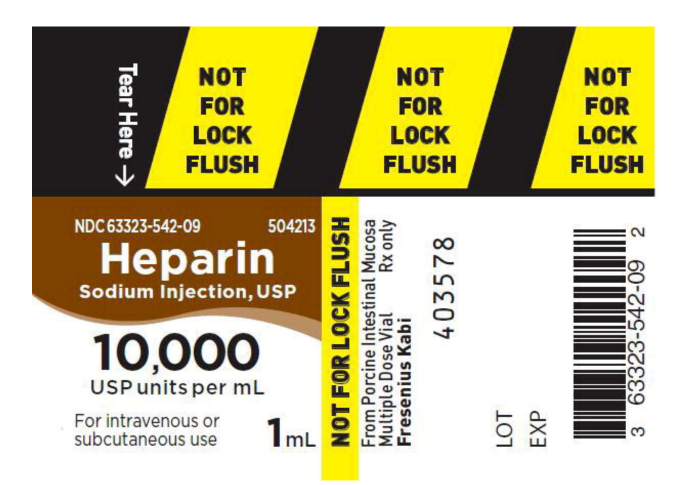
NDC 63323-542-09 504213

Heparin Sodium Injection, USP

10,000 USP units per mL

For intravenous or subcutaneous use

1 mL



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Tray Label

NDC 63323-542-13 504213

Heparin Sodium Injection, USP

10,000 USP units per mL

For intravenous or subcutaneous use

1 mL

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 4 mL Multiple Dose Vial Label

NDC 63323-459-04 504514

Heparin Sodium Injection, USP

40,000 USP units per 4 mL

(10,000 USP units per mL)

For intravenous or subcutaneous use



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 4 mL Multiple Dose Vial Tray Label

NDC 63323-459-14 504514

Heparin Sodium Injection, USP

40,000 USP units per 4 mL

(10,000 USP units per mL)

For intravenous or subcutaneous use

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Label

Not for Lock Flush

NDC 63323-915-03 915513

Heparin Sodium Injection, USP

20,000 USP units per mL

For intravenous or subcutaneous use

1 mL



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 1 mL Multiple Dose Vial Tray Label

NDC 63323-915-13 915513

Heparin Sodium Injection, USP

20,000 USP units per mL

For intravenous or subcutaneous use

1 mL

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 0.5 mL Single Dose Vial Label

Not for Lock Flush

NDC 63323-543-03 504313

Heparin Sodium Injection, USP

5,000 USP units per 0.5 mL

For IV or SC use

Preservative Free



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 0.5 mL Single Dose Vial Tray Label

NDC 63323-543-13 504313

Heparin Sodium Injection, USP

5,000 USP units per 0.5 mL

For intravenous or subcutaneous use

Preservative Free

25 Vials

NDC 63323-543-13 504313 Heparin sodium Injection, USP	sa	not made from 000 USP units r Injection q.s. sodium added ar and seal intact. to 77°FJ m TemperatureJ.	543130
5,000 USP 0.5 mL	Intestinal Mucosa	2 5 7 4	203633235
For intravenous or Preservative subcutaneous use Free	e	Vose Vi vitainer rubber vitainer vitainer sodiur sodiu vitainer vitainer soli vitainer vitainer soli vitainer vitainer soli vitainer vitainer soli vitainer soli vitainer soli vitainer soli vitainer soli vitainer soli vitainer soli vitainer soli soli vitainer soli vitainer soli soli vitainer soli vitainer soli	(01)
25 Vials	From	Single D Rx only Rx only Sterile, Piscard Fach 0.5 Fach 0.6 Hydrocic Hydrocic Hydrocic Store at Isee US Isee US	

PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 10 mL Multiple Dose Vial Label

NDC 63323-540-05 504015

Heparin Sodium Injection, USP

10,000 USP units per 10 mL

(1,000 USP units per mL)

For intravenous or subcutaneous use



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 10 mL Multiple Dose Vial Tray Label

NDC 63323-540-15 504015

Heparin Sodium Injection, USP

10,000 USP units per 10 mL

(1,000 USP units per mL)

For intravenous or subcutaneous use

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 30 mL Multiple Dose Vial Label

NDC 63323-540-33 504036

Heparin Sodium Injection, USP

30,000 USP units per 30 mL

(1,000 USP units per mL)

For intravenous or subcutaneous use



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 30 mL Multiple Dose Vial Tray Label

NDC 63323-540-36 504036

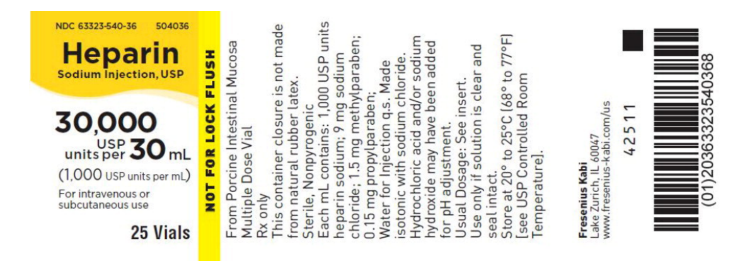
Heparin Sodium Injection, USP

30,000 USP units per 30 mL

(1,000 USP units per mL)

For intravenous or subcutaneous use

25 Vials



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 5 mL Multiple Dose Vial Label

NDC 63323-542-04 504214

Heparin Sodium Injection, USP

50,000 USP units per 5 mL

(10,000 USP units per mL)

For intravenous or subcutaneous use



PACKAGE LABEL - PRINCIPAL DISPLAY PANEL - Heparin 5 mL Multiple Dose Vial Tray Label

NDC 63323-542-14 504214

Heparin Sodium Injection, USP

50,000 USP units per 5 mL

(10,000 USP units per mL)

For intravenous or subcutaneous use

25 Vials



[see USP Controlled Room Temperature] heparin sodium; 1.5 mg methylparaben; This container closure is not made from Each mL contains: 10,000 USP units hydroxide may have been added for Store at 20° to 25°C (68° to 77°F) Hydrochloric acid and/or sodium Use only if solution is clear and Usual Dosage: See insert. Lake Zurich, IL 60047 www.fresenius-kabi.com/us 0.15 mg propylparaben Water for Injection q.s. Sterile, Nonpyrogenic natural rubber latex Multiple Dose Vial pH adjustment. Fresenius Kabi seal intact. only ž

HEPARIN SODIUM

heparin sodium injection, solution

Product Information

	oduct Type		HUMAN PRESCRIPTION DRUG		Item Code	e (Source	e)	NDC:63323-542
Ro	oute of Admini	stration	INTRAVENOUS, SUBCUTANEOUS	5				
A	ctive Ingredie	ent/Active	Moiety					
		Ingredie	nt Name	I	Basis of S	trength		Strength
HE	PARIN SODIUM	(UNII: ZZ45AB2	4CA) (HEPARIN - UNII:T2410KM04	IA) I	HEPARIN		1000	0 [USP'U] in 1 r
In	active Ingre	dients						
		Ing	redient Name				Str	ength
ME	THYLPARABEN	(UNII: A2I8C7HI	ЭТ)			1.5 mg i	n 1 m	L
PR	OPYLPARABEN (UNII: Z8IX2SC1	.OH)			0.15 mg	in 1 r	nL
	DROCHLORIC A							
	DIUM HYDROXII							
Pa	ackaging Item Code			Ν	Aarketing	Start	Ma	arketing End
Pa #	ackaging	Pac	kage Description		Date	Start	Ma	arketing End Date
Pa #	Ackaging Item Code NDC:63323-542- 13	Pac 25 in 1 TRAY	kage Description		-	Start	Ma	
Pa #	ackaging Item Code NDC:63323-542-	Pac 25 in 1 TRAY			Date	Start	Ma	
P a # 1	Ackaging Item Code NDC:63323-542- 13 NDC:63323-542-	Pac 25 in 1 TRAY 1 mL in 1 VIAL	kage Description	11/0	Date	Start	Ma	
P a # 1 2	Item Code NDC:63323-542- 13 NDC:63323-542- 09 NDC:63323-542-	Pac 25 in 1 TRAY 1 mL in 1 VIAL Product 25 in 1 TRAY	kage Description	11/0	Date 04/2019	Start	Ma	
P a # 1 2	Ackaging Item Code NDC:63323-542- 13 NDC:63323-542- 09 NDC:63323-542- 14 NDC:63323-542-	Pac 25 in 1 TRAY 1 mL in 1 VIAL Product 25 in 1 TRAY 5 mL in 1 VIAL	:kage Description ; Type 0: Not a Combination	11/0	Date 04/2019	Start	Ma	
Pa # 1 2 2	Ackaging Item Code NDC:63323-542- 13 NDC:63323-542- 09 NDC:63323-542- 14 NDC:63323-542- 04	Pac 25 in 1 TRAY 1 mL in 1 VIAL Product 25 in 1 TRAY 5 mL in 1 VIAL Product	: Kage Description ; Type 0: Not a Combination ; Type 0: Not a Combination	11/0	Date 04/2019	Start	Ma	
Pa # 1 2 2	Ackaging Item Code NDC:63323-542- 13 NDC:63323-542- 09 NDC:63323-542- 14 NDC:63323-542-	Pac 25 in 1 TRAY 1 mL in 1 VIAL Product 25 in 1 TRAY 5 mL in 1 VIAL Product	: Kage Description ; Type 0: Not a Combination ; Type 0: Not a Combination	11/0	Date 04/2019	Start	Ma	
Pa # 1 2 2	Ackaging Item Code NDC:63323-542- 13 NDC:63323-542- 09 NDC:63323-542- 14 NDC:63323-542- 04	Pac 25 in 1 TRAY 1 mL in 1 VIAL Product 25 in 1 TRAY 5 mL in 1 VIAL Product	: Kage Description ; Type 0: Not a Combination ; Type 0: Not a Combination	11/0	Date 04/2019	g Start		

HEPARIN SODIUM							
heparin sodium injection, solu	Ition						
Product Information							
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source	e) NDC:63323-915				
Route of Administration INTRAVENOUS, SUBCUTANEOUS							
Active Ingredient/Active Moiety							
Ingredie	Ingredient Name Basis of Strength Strength						
HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:T2410KM04A) HEPARIN 20000 [USP'U] in 1 mL							

In	active Ingree	dients						
			redient Name			Strength		
M	THYLPARABEN (1 mL						
PROPYLPARABEN (UNII: Z8IX2SC10H) 0.15 mg in 1 mL								
ΗY	DROCHLORIC AC	CID (UNII: QTT	17582CB)					
SC		DE (UNII: 55X04	1QC32I)					
Pa	ackaging							
#	ltem Code	Pac	ckage Description	Marketing S Date	Start	Marketing End Date		
1	NDC:63323-915- 13	25 in 1 TRAY		11/04/2019				
1	NDC:63323-915- 03	1 mL in 1 VIAL Product	; Type 0: Not a Combination					
м	arketing I	nformat	ion					
	Marketing		tion Number or Monograph	Marketin	a Start	Marketing End		
	Category	Аррпса	Citation	-	Date			
ND	-	Citation Date NDA017029 11/04/2019				Date		
	A	NDA017029			e	Date		
	A	NDA017029			e	Date		
H						Date		
	EPARIN SC	DIUM				Date		
		DIUM				Date		
ne	EPARIN SC	DIUM ijection, solu				Date		
ne P	EPARIN SC parin sodium ir roduct Inforr	DIUM ijection, solu		11/04/2019				
P P	EPARIN SC parin sodium ir roduct Inforr roduct Type	DIUM njection, solu mation	ution HUMAN PRESCRIPTION DRUG					
P P	EPARIN SC parin sodium ir roduct Inforr	DIUM njection, solu mation	ution	11/04/2019				
P P	EPARIN SC parin sodium ir roduct Inforr roduct Type	DIUM njection, solu mation	ution HUMAN PRESCRIPTION DRUG	11/04/2019				
P P R	EPARIN SC parin sodium ir roduct Inforr roduct Type	DIUM njection, solu mation	Ition HUMAN PRESCRIPTION DRUG INTRAVENOUS, SUBCUTANEOUS	11/04/2019				
P P R	EPARIN SC parin sodium ir roduct Inforr roduct Type pute of Adminis	DIUM njection, solu mation stration	Ition HUMAN PRESCRIPTION DRUG INTRAVENOUS, SUBCUTANEOUS	11/04/2019	e (Source)			
P P R A	EPARIN SC parin sodium ir roduct Inforr roduct Type oute of Adminis	DIUM njection, solu mation stration ent/Active Ingredie	Ition HUMAN PRESCRIPTION DRUG INTRAVENOUS, SUBCUTANEOUS Moiety	ltem Code	e (Source) rength	NDC:63323-543		
P P R A	EPARIN SC parin sodium ir roduct Inforr roduct Type oute of Adminis	DIUM njection, solu mation stration ent/Active Ingredie	ution HUMAN PRESCRIPTION DRUG INTRAVENOUS, SUBCUTANEOUS Moiety nt Name	ltem Code	e (Source) rength) NDC:63323-543		
P Pr Ro A	EPARIN SC parin sodium ir roduct Inforr roduct Type oute of Adminis	DIUM njection, solu mation stration ent/Active Ingredie	ution HUMAN PRESCRIPTION DRUG INTRAVENOUS, SUBCUTANEOUS Moiety nt Name	ltem Code	e (Source) rength	NDC:63323-543		

 Packaging

 #
 Item Code
 Package Description
 Marketing Start Date
 Marketing End Date

SODIUM HYDROXIDE (UNII: 55X04QC32I)

1	NDC:63323-543- 13	25 in 1 TRAY		11,	/04/2019		
1			AL; Type 0: Not a Combination				
-	03	Product					
R 4		1	!				
IV	larketing						
	Marketing Category	Applica	tion Number or Monograph Citation		ng Start ate	Marketing En Date	
NE		NDA017029			11/04/2019		
Η	EPARIN S	ODIUM					
۱e	parin sodium	injection, solu	Ition				
Ρ	roduct Info	rmation					
Ρ	roduct Type		HUMAN PRESCRIPTION DRUG		Item Coc	le (Source	e) NDC:63323-45
R	oute of Admin	istration	INTRAVENOUS, SUBCUTANEOUS				
A	ctive Ingred		-				
		-	nt Name		Basis of S	-	Strength
HE	EPARIN SODIUM	(UNII: ZZ45AB2	24CA) (HEPARIN - UNII:T2410KM04)	A)	HEPARIN		10000 [USP'U] in 1
Ir	nactive Ingre	edients					
	U		redient Name				Strength
BE	ENZYL ALCOHO	-				10.42 mg i	_
s		DE (UNII: 451W47	7IQ8X)			5 mg in 1 r	mL
SC	DDIUM HYDROX	IDE (UNII: 55X02	IQC32I)				
D	ackaging						
					Markot	ing Start	Marketing E
#	Item Code	Pa	ackage Description			ate	Date
1	NDC:63323-	25 in 1 TRAY			11/04/201	9	
	459-14 NDC:63323-	4 mL in 1 VIAI	MULTI-DOSE; Type 0: Not a				
1	459-04	Combination Pr					
Μ	larketing	Informat	ion				
	Marketing	Applica	tion Number or Monograph	1		ng Start	Marketing En
	Category		Citation		Da	ate	Date

11/04/2019

NDA

NDA017029

HEPARIN SODIUM

Product Infor	mation					
Product Type		HUMAN PRESCRIPTION DRUC	6 It	em Cod	e (Source) NDC:63323-540
Route of Admini	istration	INTRAVENOUS, SUBCUTANE	OUS			
Active Ingredi	ent/Active	Moiety				
	Ingredie	nt Name	B	asis of a	Strength	Strength
HEPARIN SODIUM	(UNII: ZZ45AB2	4CA) (HEPARIN - UNII:T2410K	M04A) HE	PARIN		1000 [USP'U] in 1 m
Inactive Ingre	dients					
	Ing	redient Name				Strength
SODIUM CHLORID	E (UNII: 451W47	IQ8X)			9 mg in 1	mL
METHYLPARABEN	(UNII: A2I8C7HI9	PT)			1.5 mg in	1 mL
PROPYLPARABEN	(UNII: Z8IX2SC1	OH)			0.15 mg i	n 1 mL
HYDROCHLORIC ACID (UNII: QTT17582CB)						
SODIUM HYDROXI						
sodium Hydroxi Packaging	DE (UNII: 55X04		Ma	nrketing Date		Marketing End Date
Packaging	DE (UNII: 55X04	QC32I)	Ma 11/04	Date		-
Packaging Htem Code NDC:63323-540- NDC:62223 540	DE (UNII: 55X04 Pac 25 in 1 TRAY	QC32I)		Date		-
Bitem Code 1 NDC:63323-540- 03 1 NDC:63323-540- 03	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL;	QC32I) kage Description	11/04	Date		-
Packaging Item Code Item Code DC:63323-540-	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL; Product 25 in 1 TRAY	QC32I) kage Description	11/04	Date /2019		-
SODIUM HYDROXI Packaging Item Code NDC:63323-540-	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL; Product 25 in 1 TRAY 10 mL in 1 VIA Product 25 in 1 TRAY 25 in 1 TRAY	QC32I) kage Description Type 0: Not a Combination L; Type 0: Not a Combinatior	11/04 11/04 11/04 11/04	Date /2019 /2019		-
Packaging Item Code Item Code NDC:63323-540- NDC:63323-540-	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL; Product 25 in 1 TRAY 10 mL in 1 VIA Product 25 in 1 TRAY 25 in 1 TRAY	QC32I) kage Description Type 0: Not a Combination	11/04 11/04 11/04 11/04	Date /2019 /2019		-
Solum Hydroxi Item Code Item Code NDC:63323-540-	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL; Product 25 in 1 TRAY 10 mL in 1 VIA Product 25 in 1 TRAY 25 in 1 TRAY 30 mL in 1 VIA	QC32I) kage Description Type 0: Not a Combination L; Type 0: Not a Combinatior	11/04 11/04 11/04 11/04	Date /2019 /2019		-
SOUUM HYDROXI Packaging # Item Code 1 NDC:63323-540- 1 NDC:63323-540- 2 NDC:63323-540- 3 NDC:63323-540- 3 NDC:63323-540- 3 NDC:63323-540-	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL; Product 25 in 1 TRAY 10 mL in 1 VIA Product 25 in 1 TRAY 30 mL in 1 VIA Product	QC32I) kage Description Type 0: Not a Combination L; Type 0: Not a Combinatior L; Type 0: Not a Combinatior	11/04 11/04 11/04 11/04	Date /2019 /2019		-
SOLIUM HYDROXI Packaging Item Code Independent NDC:63323-540-	DE (UNII: 55X04 Pac 25 in 1 TRAY 1 mL in 1 VIAL; Product 25 in 1 TRAY 25 in 1 TRAY 10 mL in 1 VIA Product 25 in 1 TRAY 30 mL in 1 VIA Product	QC32I) kage Description Type 0: Not a Combination L; Type 0: Not a Combinatior L; Type 0: Not a Combinatior	11/04 11/04 11/04	Date /2019 /2019 /2019	s s s s s s s s s s s s s s s s s s s	-

HEPARIN SODIUM

heparin sodium injection, solution

INTRAVENOUS, SUBCUTANEOUS Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:T2410KM04A) HEPARIN 5000 [USP'U] in 1 m Ingredient Name Basis of Strength 5000 [USP'U] in 1 m Inactive Ingredients Ingredient Name Strength SoDIUM CHLORIDE (UNII: 451W47/Q8X) 5 mg in 1 mL METHYLPARABEN (UNII: 281X2SC10H) 0.15 mg in 1 mL POPYLPARABEN (UNII: 281X2SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: 55X04QC32I) 0.15 mg in 1 mL Strength Portragende Marketing End Date Marketing Information		mation					
Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:T2410KM04A) HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:T2410KM04A) SODIUM CHLORIDE (UNII: 451W47IQ8X) SODIUM CHLORIDE (UNII: 451W47IQ8X) SODIUM CHLORIDE (UNII: 451W47IQ8X) SODIUM CHLORIDE (UNII: 451W47IQ8X) SODIUM CHLORIDE (UNII: 281Z5C10H) HYDROCHLORIC ACID (UNII: 281Z5C10H) HYDROCHLORIC ACID (UNII: 2717582CB) SODIUM HYDROXIDE (UNII: 55X04QC32I)	Product Type	HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:63323-262	
Ingredient Name Basis of Strength Strength HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:TZ410KM04A) HEPARIN 5000 [USP'U] in 1 m Inactive Ingredients Ingredient Name Strength Inactive Ingredients 5 mg in 1 mL SODIUM CHLORIDE (UNII: 451W47108X) 5 mg in 1 mL METHYLPARABEN (UNII: 281C27H19T) 0.15 mg in 1 mL PROPYLPARABEN (UNII: Z812SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: 2717582CB) 0.15 mg in 1 mL SODIUM HYDROXIDE (UNII: 55X04QC32I) Marketing Start MC:63323-262- 25 in 1 TRAY 03/25/2020 1 NDC:63323-262- 25 in 1 TRAY 03/25/2020 Marketing Information ImL in 1 VIAL; Type 0: Not a Combination ImL	Route of Admini	f Administration INTRAVENOUS, SUBCUTANEOUS					
Ingredient Name Basis of Strength Strength HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:TZ410KM04A) HEPARIN 5000 [USP'U] in 1 m Inactive Ingredients Ingredient Name Strength SODIUM CHLORIDE (UNII: 451W47IQ8X) 5 mg in 1 mL METHYLPARABEN (UNII: 281C97H9T) 1.5 mg in 1 mL PROPYLPARABEN (UNII: 2812SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: 2717582CB) 0.15 mg in 1 mL SODIUM HYDROXIDE (UNII: 55X04QC32I) Marketing Start MDC:63323-262- 25 in 1 TRAY 03/25/2020 1 NDC:63323-262- 25 in 1 TRAY 03/25/2020 1 NDC:63323-262- 1 mL in 1 VIAL; Type 0: Not a Combination Image: Comparison Marketing Information Image: Comparison Marketing Start Marketing End Date							
HEPARIN SODIUM (UNII: ZZ45AB24CA) (HEPARIN - UNII:T2410KM04A) HEPARIN 5000 [USP'U] in 1 m Inactive Ingredients Ingredient Name Strength SoDIUM CHLORIDE (UNII: 451W47IQ8X) 5 mg in 1 mL Magredient Name Strength SoDIUM CHLORIDE (UNII: 4218C7H19T) 5 mg in 1 mL PROPYLPARABEN (UNII: 2812SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: 0TT17582CB) SoDIUM HYDROXIDE (UNII: 55X04QC32I) Packaging # Item Code Package Description Marketing Start Marketing End Date 1 NDC:63323-262- 25 in 1 TRAY 03/25/2020 03/25/2020 ImL in 1 VIAL; Type 0: Not a Combination Image: Colspan="2">Image: Colspan="2">Marketing Colspan="2">Marketing Colspan="2">Marketing Colspan="2">Marketing Colspan="2">Marketing Colspan="2" Marketing Colspan="2" Marketing Colspan= multiper or Monograph Date Marketing Start Marketing End Date	Active Ingredi	ent/Active	Moiety				
Ingredients Ingredient Name Strength Solum CHLORIDE (UNII: 451W47IQ8X) 5 mg in 1 mL METHYLPARABEN (UNII: 4218C7HI9T) 1.5 mg in 1 mL POPYLPARABEN (UNII: 28IX2SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: 28IX2SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: 25X04QC32I) OISOLIM HYDROXIDE (UNII: 55X04QC32I) Packaging # Item Code Package Description Marketing Start Date Marketing Information 1 NDC: 63323-262- 1 mL in 1 VIAL; Type 0: Not a Combination Product O3/25/2020 Image Start Date Marketing Information		Ingredie	Basis of S	Strength	Strength		
Ingredient Name Strength Solum ChLoride (UNII: 451W47IQ8X) 5 mg in 1 mL METHYLPARABEN (UNII: 4218C7H19T) 1.5 mg in 1 mL PROPYLPARABEN (UNII: 281X2SC10H) 0.15 mg in 1 mL HydrochLoric ACID (UNII: QTT17582CB) O.15 mg in 1 mL Solum Hydroxide (UNII: 55X04QC32I) O.15 mg in 1 mL Package Description Marketing Start Date Marketing Information NDC:63323-262- 25 in 1 TRAY 03/25/2020 NDC:63323-262- 1 mL in 1 VIAL; Type 0: Not a Combination Product Marketing Information	HEPARIN SODIUM	(UNII: ZZ45AB2	4CA) (HEPARIN - UNII:T2410KM04	A) HEPARIN		5000 [USP'U] in 1 m	
Ingredient Name Strength Solum ChLoride (UNII: 451W47IQ8X) 5 mg in 1 mL METHYLPARABEN (UNII: 4218C7H19T) 1.5 mg in 1 mL PROPYLPARABEN (UNII: 281X2SC10H) 0.15 mg in 1 mL HydrochLoric ACID (UNII: QTT17582CB) O.15 mg in 1 mL Solum Hydroxide (UNII: 55X04QC32I) O.15 mg in 1 mL Package Description Marketing Start Date Marketing Information NDC:63323-262- 25 in 1 TRAY 03/25/2020 NDC:63323-262- 1 mL in 1 VIAL; Type 0: Not a Combination Product Marketing Information							
SODIUM CHLORIDE (UNII: 451W47IQ8X) 5 mg in 1 mL METHYLPARABEN (UNII: A2I8C7HI9T) 1.5 mg in 1 mL PROPYLPARABEN (UNII: 28IX2SC10H) 0.15 mg in 1 mL HYDROCHLORIC ACID (UNII: QTT17582CB) 0.15 mg in 1 mL SODIUM HYDROXIDE (UNII: 55X04QC32I) V # Item Code Package Description Marketing End Date 1 NDC:63323-262- 06 25 in 1 TRAY 03/25/2020 Marketing End Date 1 NDC:63323-262- 03 1 mL in 1 VIAL; Type 0: Not a Combination Product 03/25/2020 V V	Inactive Ingre	dients					
Marketing Category Application Number or Monograph Citation Marketing Start Date Marketing Start Marketing Start Date Marketing Start Marketing Start Date Marketing Start Marketing Start Date		Ing	redient Name			Strength	
PROPYLPARABEN (UNII: Z8IX2SC10H) 0.15 mg in 1 mL HydrochLoric ACID (UNII: QTT17582CB) 0.15 mg in 1 mL Sodium Hydroxide (UNII: 55X04QC32I) 0.15 mg in 1 mL # Item Code Package Description Marketing Date 1 NDC:63323-262- 06 25 in 1 TRAY 03/25/2020 Marketing Code 1 NDC:63323-262- 03 1 mL in 1 VIAL; Type 0: Not a Combination Product 03/25/2020 Image: Code Vertice State Marketing Code Application Number or Monograph Citation Marketing Start Date Marketing Start Date	SODIUM CHLORID	E (UNII: 451W47	IQ8X)		5 mg in 1	mL	
Hydrochloric Acid (UNII: QTT17582CB) Solum Hydroxide (UNII: 55x04QC32I) Package Description Marketing Start Date Marketing End Date 1 NDC:63323-262- 06 25 in 1 TRAY 03/25/2020 1 NDC:63323-262- 1 mL in 1 VIAL; Type 0: Not a Combination Product Herein Code Combination Product Marketing Start Date	METHYLPARABEN	(UNII: A2I8C7HI	ЭТ)		1.5 mg in	1 mL	
BOILUM HYDROXIDE (UNII: 55X04QC32I) Marketing Start Marketing End Date # Item Code Package Description Marketing Start Date Marketing End Date 1 NDC:63323-262- 06 25 in 1 TRAY 03/25/2020 03/25/2020 1 NDC:63323-262- 03 1 mL in 1 VIAL; Type 0: Not a Combination Product Image: Colspan="4">Image: Colspan="4">Marketing Start Date Marketing Category Marketing Start Date Marketing Start Date Marketing End Date	PROPYLPARABEN (UNII: Z8IX2SC10H)0.15 mg in 1 mL						
Package Description Marketing Start Date Marketing End Date 1 NDC:63323-262- 06 25 in 1 TRAY 03/25/2020 03/25/2020 1 NDC:63323-262- 03 1 mL in 1 VIAL; Type 0: Not a Combination Product							
# Item Code Package Description Marketing Start Date Marketing End Date 1 NDC:63323-262- 06 25 in 1 TRAY 03/25/2020 03/25/2020 1 NDC:63323-262- 03 1 mL in 1 VIAL; Type 0: Not a Combination Product Image: Comparison Image: Comparison Marketing Information Marketing Category Application Number or Monograph Citation Marketing Start Date Marketing End Date	Packaging						
I 06 25 III T HAT 03/25/2020 I NDC:63323-262- 1 mL in 1 VIAL; Type 0: Not a Combination Image: Comparison of the comparison of		Pac	kage Description		Start	Marketing End	
Image: Product Product Marketing Information Marketing Start Citation Number or Monograph Citation Marketing Category Application Number or Monograph Citation				Date		Date	
Marketing CategoryApplication Number or Monograph CitationMarketing Start DateMarketing End Date	1 NDC:63323-262-	25 in 1 TRAY		2410		Date	
Marketing CategoryApplication Number or Monograph CitationMarketing Start DateMarketing End Date	1 NDC:63323-262- 06 1 NDC:63323-262-	1 mL in 1 VIAL	; Type 0: Not a Combination	2410		Date	
Category Citation Date Date	1 NDC:63323-262- 06 1 NDC:63323-262-	1 mL in 1 VIAL	; Type 0: Not a Combination	2410		Date	
NDA NDA017029 03/25/2020	 NDC:63323-262- 06 NDC:63323-262- 03 	1 mL in 1 VIAL Product		2410		Date	
	 NDC:63323-262- 06 NDC:63323-262- 03 MDC:63323-262- 03 	1 mL in 1 VIAL Product	ion tion Number or Monograph	03/25/2020 Marketir		Marketing End	

Labeler - Fresenius Kabi USA, LLC (608775388)

Establishment						
Name	Address	ID/FEI	Business Operations			
Fresenius Kabi USA, LLC			ANALYSIS(63323-542, 63323-540, 63323-262, 63323-543, 63323-459, 63323-915), MANUFACTURE(63323-542, 63323-540, 63323-262, 63323-543, 63323-459, 63323- 915)			

Establishment						
Name	Address	ID/FEI	Business Operations			
Fresenius Kabi USA, LLC		023648251	ANALYSIS(63323-540), MANUFACTURE(63323-540)			