

## Important Prescribing Information

May 2, 2019

## Subject: Temporary importation of intravenous drug products to address drug shortages

Dear Healthcare Professional,

In order to address shortages of critical drug products from the aftermath of Hurricane Maria, Baxter Healthcare Corporation (Baxter) is coordinating with the U.S. Food and Drug Administration (FDA) to increase the availability of products from Baxter's manufacturing facility in the United Kingdom (UK).

Baxter has initiated temporary importation of Heparin Sodium 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusions in VIAFLEX Container. This product is manufactured by Baxter's manufacturing facility in the UK and marketed in the UK. At this time, no other entity except Baxter is authorized by the FDA to import or distribute these products in the United States. FDA has not approved Heparin Sodium BP in 0.9% w/v Sodium Chloride IV Infusions in VIAFLEX container manufactured by Baxter's manufacturing facility in the UK.

Product name and description	Size	Product code	Pack Factor	NDC
Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion in VIAFLEX container <b>(1,000 units / 500 mL)</b>	500 mL	FKB0953G	20	0338-9556-20
Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion in VIAFLEX container (2,000 units / 1,000 mL)	1,000 mL	FKB0944G	10	0338-9552-10

Effective immediately, and during this temporary period, Baxter will offer the following:

BP = British Pharmacopoeia

It is important to note the following:

- The imported products are labeled in IU/L, whereas the FDA-approved heparin products are labeled in units per mL. The imported products and FDA-approved products contain the same Heparin Sodium concentration of 2 units per mL.
- The administration port protector on the imported products contains a twist-off port protector that must be twisted off rather than pulled off. The FDA approved product includes a medication (injection) port while the imported products do not include such a port. Please refer to the image below and the product comparison chart at the end of this letter.

- The imported product's administration port system is fully compatible with IV set spike heads that meet the International Organization of Standardization (ISO) standards and with Baxter IV sets marketed in the United States.
- The imported products do not have a barcode. Institutions should manually input the product into their systems to confirm that barcode systems do not provide incorrect information when the product is scanned. Alternative procedures should be followed to assure that the correct drug product is being used and administered to individual patients. Barcodes stickers are provided with this Dear Healthcare Provider letter. Please refer to page 10 for the product barcode information.

There are some key differences in the labeling between the U.S. marketed Heparin Sodium and 0.9% Sodium Chloride Injection and the UK products. Please see the product comparison table at the end of this letter.

Please refer to the FDA-approved package insert for the full prescribing information of Heparin Sodium and 0.9% Sodium Chloride Injection drug product at:

https://www.dailymed.nlm.nih.gov/dailymed/getFile.cfm?setid=0d929726-76c3-48fc-b4e7fd06409f9fb3&type=pdf&name=0d929726-76c3-48fc-b4e7-fd06409f9fb3

If you have any questions about the information contained in this letter or the use of the imported products, please contact Baxter's Medical Information Service at 1-800-933-0303.

To place an order, please contact Baxter's Center for Service by calling 1-888-229-0001.

To report product quality issues or to replace missing barcode stickers, please contact Baxter Product Surveillance at 1-800-437-5176.

To report adverse events associated with these imported products, please call Baxter at 1-866-888-2472, or fax: 1-800-759-1801. Adverse events or quality problems experienced with the use of this product may also be reported to the FDA's MedWatch Adverse Event Reporting program either online, by regular mail or by fax:

- Complete and submit the report **Online**: www.fda.gov/medwatch/report.htm
- Regular mail or Fax: Download form <a href="http://www.fda.gov/MedWatch/getforms.htm">http://www.fda.gov/MedWatch/getforms.htm</a> or call 1-800-332-1088 to request a reporting form, then complete and return to the address on the pre-addressed form, or submit by fax to 1-800-FDA-0178.

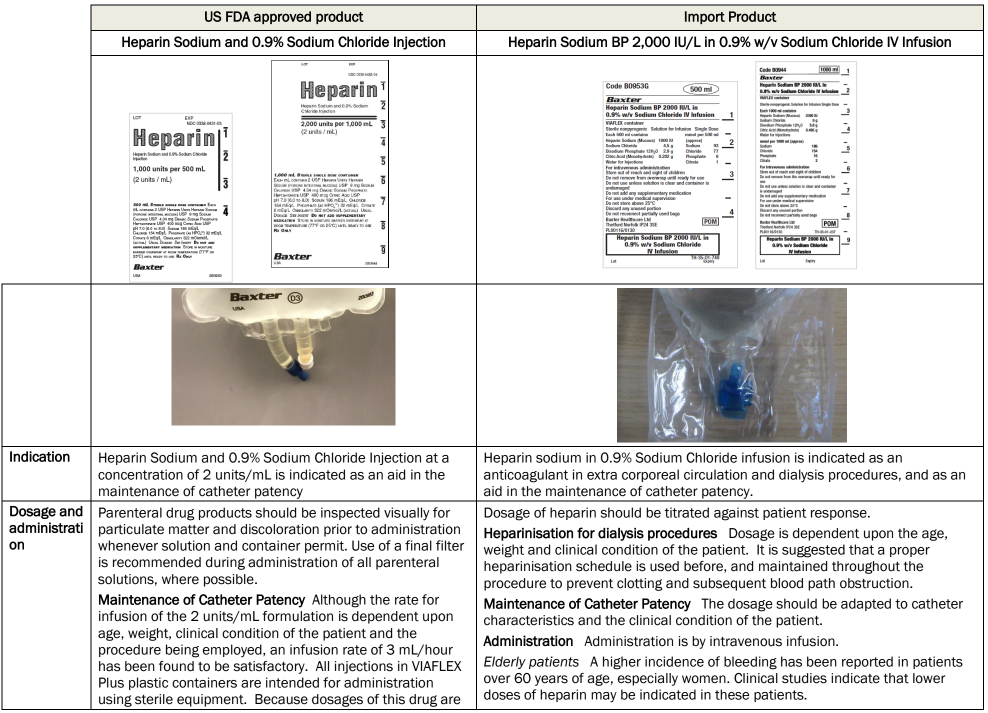
Sincerely,

Dan Varfu

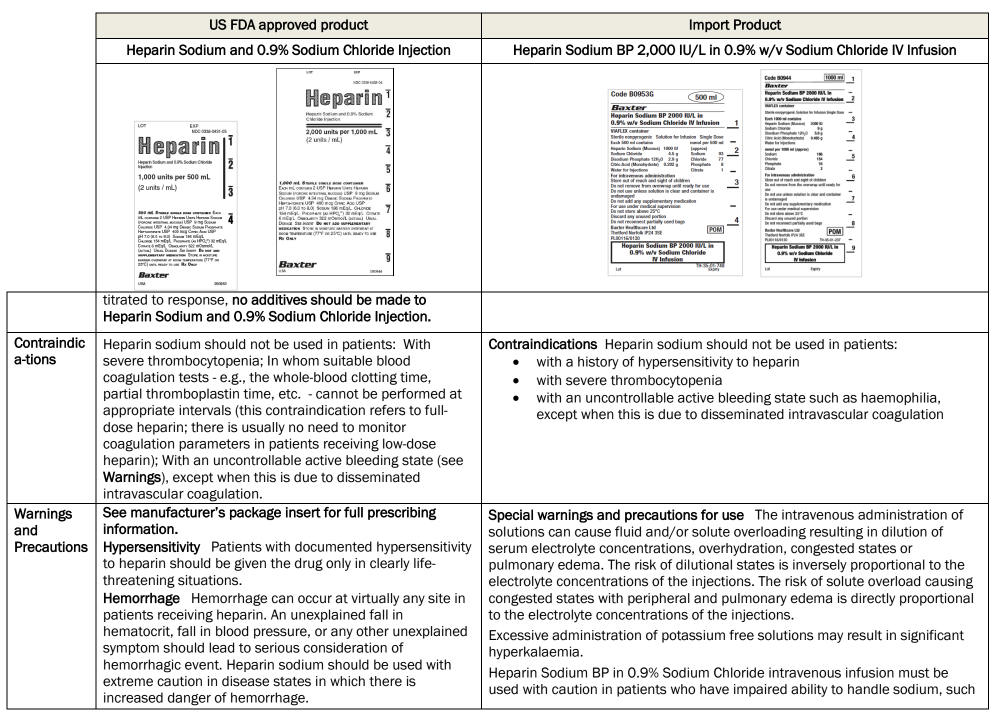
Dennis Vaughn Vice President, Marketing Operations Baxter Healthcare Corporation

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	US FDA approved product	Import Product Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion	
	Heparin Sodium and 0.9% Sodium Chloride Injection		
	LOT EXP NDC 0338-041-05 Herebo and 0.9% Sodum Claudin Hyperbo and 0.9% Sodum Claudin Department of the sodum Claudin (2 units / mL) Sodum Sodum Sodum Claudin Sodum Sodum	Code B09536       S00 ml         Heparin Sodium BP 2000 IU/L in 0.9% w/v Sodium Chloride IV Infusion       1 1         NAFLE contains       The provide the status of the st	
Ingredients	Each 500 mL contains 1,000 units Heparin Sodium (porcine Intestinal Mucosa) USP, 4.5g Sodium Chloride USP, 2.17 g Dibasic Sodium phosphate Heptahydrate USP, 0.2 g Citric Acid USP Each 1000 mL contains 2,000 units Heparin Sodium (porcine Intestinal Mucosa) USP, 9g Sodium Chloride USP, 4.34 g Dibasic Sodium phosphate Heptahydrate USP, 0.4 g Citric Acid USP	Each 500 mL contains 1,000IU Heparin Sodium (Mucous), 4.5 g Sodium Chloride, 2.9 g Disodium Phosphate 12H <sub>2</sub> O, 0.202 g Citric Acid (Monohydrate), Water for Injection Each 1,000 mL contains 2,000 IU Heparin Sodium (Mucous), 9.0 g Sodium Chloride, 5.8 g Disodium Phosphate 12H <sub>2</sub> O, 0.405 g Citric Acid (Monohydrate), Water for Injection	
Additional Information	Each 500 mL and 1000 mL container contains: Sodium 186 mEq/L; Chloride 154 mEq/L; Phosphate (as HPO <sub>4</sub> =) 32 mEq/L; Citrate 6 mEq/L pH 7.0 (6.0 to 8.0); Osmolarity 322 mOsmol/L	Mmol per 500 mL (approx.)Mmol per1000 mL (approx.)SodiumSodium93Chloride186Chloride154Phosphate8Citrate116Citrate2	
Description	Heparin Sodium and 0.9% Sodium Chloride Injection is a buffered, sterile, nonpyrogenic solution of Heparin Sodium, USP derived from porcine intestinal mucosa, standardized for anticoagulant activity supplied in single dose containers for vascular administration. It contains no antimicrobial agents. The potency is determined by a biological assay using a USP reference standard based on units of heparin activity per milligram.	Sterile non pyrogenic aqueous solution intended for intravenous administration.	
Administrati on ports	Medication port with rubber closure PLUS Administration port with pull off port protector	Administration port with Twist off Protector (No medication port)	



USMP/G74/17-0001(2) 05/19



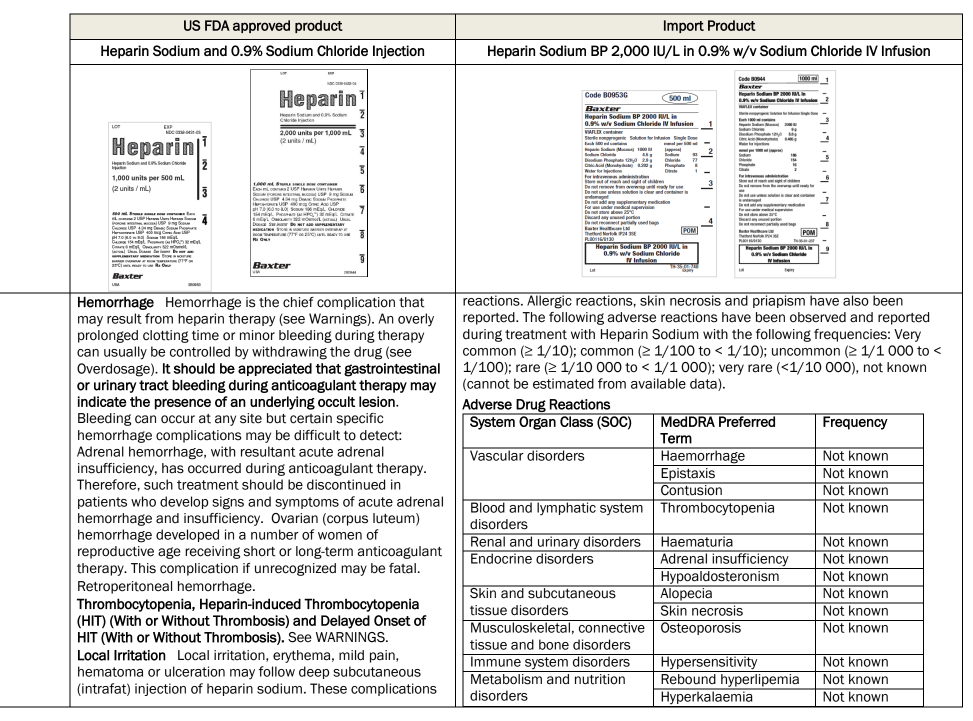
US FDA approved product	Import Product
Heparin Sodium and 0.9% Sodium Chloride Injection	Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion
$ \int dt = $	Code B09536     500 ml       Hearts Galaxies     1       Hoppins Galaxies     1 <tr< th=""></tr<>
<ul> <li>Coagulation Testing When heparin sodium is administered in therapeutic amounts, its dosage should be regulated by frequent blood coagulation tests. If the coagulation test is unduly prolonged or if hemorrhage occurs, heparin sodium should be discontinued promptly (see Overdosage).</li> <li>Thrombocytopenia Thrombocytopenia has been reported to occur in patients receiving heparin with a reported incidence of up to 30%. Platelet counts should be obtained at baseline and periodically during heparin administration. Mild thrombocytopenia (count greater than 100,000/mm3) may remain stable or reverse even if heparin is continued. However, thrombocytopenia of any degree should be monitored closely. If the count falls below 100,000/mm3 or if recurrent thrombosis develops (see Heparin-induced Thrombocytopenia (HIT) With or Without Thrombosis), the heparin product should be discontinued and, if necessary, an alternative anticoagulant administered.</li> <li>Heparin-induced Thrombocytopenia (HIT) (With or Without Thrombosis) HIT is a serious immune-mediated reaction resulting from irreversible aggregation of platelets. HIT may progress to the development of venous and arterial thromboses, a condition referred to as HIT with thrombosis. Thrombotic events may also be the initial presentation for HIT. Once HIT (with or without thrombosis) is diagnosed or</li> </ul>	as renal insufficiency and congestive heart failure, and in clinical states in which there exists oedema with sodium retention. Do not use unless solution is clear and container undamaged. Heparin sodium BP in 0.9% w/v sodium chloride intravenous infusion should not be administered orally. Heparin should be used with extreme care in patients suffering from conditions in which there is an increased danger of haemorrhage. Haemorrhage can occur at virtually any site in patients receiving heparin. An unexplained fall in haematocrit, fall in blood pressure, or any other unexplained symptom should lead to serious consideration of haemorrhagic event. Heparin sodium should be used with extreme caution in disease states in which there is increased danger of haemorrhage. Some of the conditions in which increased danger of haemorrhage exists are: 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. Haematologic - Conditions associated with increased bleeding tendencies, such as haemophilia, thrombocytopenia, and some vascular purpuras. Gastrointestinal - Ulcerative lesions and continuous tube drainage of the stomach or small intestine. Other - Menstruation, liver disease with impaired haemostasis. Periodic hematocrit tests, and tests for occult blood in stool are recommended during the entire course of heparin therapy, regardless of the route of administration.

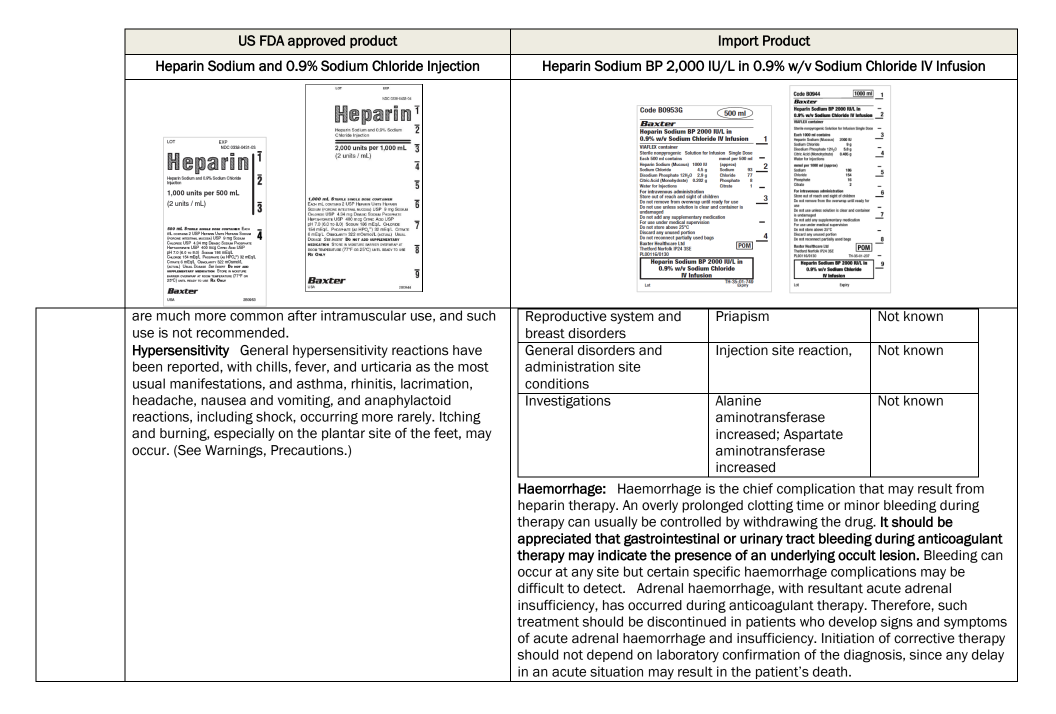
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US FDA approved product	Import Product
Heparin Sodium and 0.9% Sodium Chloride Injection	Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion
International constructionInternational constructionInternationInternational constructionInternational constructionInternational constructionInternational constructionInternationInternational constructionInternational constructionInternational constructionInternational constructionInternationInternational constructionInternational constructionInternational constructionInternational constructionInternationInternational constructionInternational construction	Code B09536     Son m       Heartin Solution BP 2000 IU/L in Sparkin Solution BP 2000 IU/L in Sparkin Solution BP 2000 IU/L in Sparkin Solution Microany 100 II (upper)     Imparities Solution for Influences       VAREE container     1       Basedim Phosphate 12Hg 2.3.9     Choird 9.7       Bosdim Phosphate 12Hg 2.3.9     Choird 9.7       Disodim Phosphate 12Hg 2.3.9     Choird 9.7
<ul> <li>heparin flushes) should be discontinued and an alternative anticoagulant used. Future use of heparin sodium, especially within 3 to 6 months following the diagnosis of HIT (with or without thrombosis), and while patients test positive for HIT antibodies, should be avoided. Delayed Onset of HIT (With or Without Thrombosis) Heparin-induced thrombocytopenia (with or without thrombosis) 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 (with or without thrombosis).</li> <li>PRECAUTIONS General Thrombocytopenia, Heparin-induced Thrombocytopenia (HIT) (With or Without Thrombosis) and Delayed Onset of HIT (With or Without Thrombosis).</li> <li>PRECAUTIONS General Thrombocytopenia, Heparin-induced Thrombocytopenia (HIT) (With or Without Thrombosis). Heparin Resistance: Increased resistance to heparin is frequently encountered in fever, thrombosis, thrombophlebitis, infections with thrombosing tendencies, myocardial infarction, cancer and in postsurgical patients. Increased Risk in Older Patients, Especially Women: A higher incidence of bleeding has been reported in patients, particularly women, over 60 years of age. Solutions Containing Sodium: These solutions should be used with caution in patients receiving corticosteroids or corticotropin. Laboratory Tests Periodic platelet counts, hematocrits, and tests for occult blood in stool are recommended during the</li> </ul>	Heparin can suppress adrenal secretion of aldosterone leading to hyperkalaemia, particularly in patients such as those with diabetes mellitus, chronic renal failure, pre-existing metabolic acidosis, a raised plasma potassium, or taking potassium sparing drugs. The risk of hyperkalaemia appears to increase with duration of therapy but is usually reversible. Plasma potassium should be measured in patients at risk before starting heparin therapy and in all patients treated for more than 7 days. Thrombocytopenia is commonly seen in patients receiving heparin. Platelet counts should be obtained at baseline and periodically during heparin administration. Mild thrombocytopenia (count greater than 100,000/mm <sup>3</sup> ) may remain stable or reverse even if heparin is continued. However, thrombocytopenia of any degree should be monitored closely. If the count falls below 100,000/mm <sup>3</sup> or if recurrent thrombosis develops, the heparin product should be discontinued and, if necessary, an alternative anticoagulant administered. HIT is a serious immune-mediated disorder resulting from irreversible aggregation of platelets. HIT may progress to the development of venous and arterial thromboses, a condition referred to as HIT with thrombosis. Thrombotic events may also be the initial presentation for HIT. 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 fatal outcomes. Once HIT (with or without thrombosis) is diagnosed or strongly suspected, heparin

US FDA approved product	Import Product
Heparin Sodium and 0.9% Sodium Chloride Injection	Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion
<text><text><text></text></text></text>	Code B0953G     500 ml       Haparin Sodium BP 2000 II/L in 0.9% w/v Sodium Chloride IV Infusion     1       VAREE Container     1       Varei de comprogenic Scherie nongrogenic Check Add (Moodenhan)     2       Discidium Thosphats 12H, 0.2 g.g.     Check add
entire course of heparin therapy, regardless of the route of administration (see Dosage and Administration). <b>Carcinogenesis, Mutagenesis, Impairment of Fertility</b> 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. <b>Pregnancy</b> Teratogenic Effects - Pregnancy Category C: Animal reproduction studies have not been conducted with heparin sodium. It is not known whether heparin sodium can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. Heparin sodium should be given to a pregnant woman only if clearly needed. Nonteratogenic Effects: Heparin does not cross the placental barrier. <b>Nursing Mothers</b> Heparin is not excreted in human milk. <b>Pediatric Use</b> Safety and effectiveness in pediatric patients have not been established. See Dosage and Administration. <b>Geriatric Use</b> A higher incidence of bleeding has been reported in patients over 60 years of age, especially women (see Precautions, General). Clinical studies indicate that lower doses of heparin may be indicated in these patients (see Precautions, General and Clinical Pharmacology).	<ul> <li>sodium (including heparin flushes) should be discontinued and an alternative anticoagulant used. Future use of heparin sodium, especially within 3 to 6 months following the diagnosis of HIT (with or without thrombosis), and while patients test positive for HIT antibodies, should be avoided.</li> <li>Elevations of aminotransferase (SGOT [S-AST] and SGPT [S-ALT]) levels have been commonly seen in patients (and healthy subjects) who have received heparin. Since aminotransferase determinations are important in the differential diagnosis of myocardial infarction, liver disease, and pulmonary emboli, rises that might be caused by drugs (like heparin) should be interpreted with caution.</li> <li>Resistance to heparin has been noted in fever, thrombosis, thrombophlebitis, infections with thrombosing tendencies, myocardial infarction, cancer and in postsurgical patients.</li> <li>These solutions should be used with caution in patients receiving corticosteroids or corticotropin.</li> <li>Pregnancy: The safety of heparin sodium in 0.9% w/v Sodium Chloride intravenous infusion has not been demonstrated in pregnant women. There are no or limited amount of data from the use of Heparin Sodium in pregnant women. Animal studies are insufficient with respect to reproductive toxicity. Heparin Sodium is not recommended during pregnancy.</li> <li>Breast-feeding: Heparin does not pass the placental barrier; it is not excreted in human milk. Heparin Sodium can be used during breast-feeding.</li> </ul>

	US FDA approved product	Import Product           Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion	
	Heparin Sodium and 0.9% Sodium Chloride Injection		
	INT EXC   INT EXC   INT EXC   INT EXC   INT EXC   INT INT	Code B0953G     500 ml       Heartin Solition BP 2000 IU/L in 0.9% w/v Solitium Chlorido IV Infusion Solition Robotion for Infusion Single Dose Each 500 et contains     1       Heartin Solition Merzous 1000 II (Lin Solition Robotion for Infusion Single Dose Each 500 et contains     1       Solition Robotion for Infusion Single Dose Each 500 et contains     1       Solition Robotion for Infusion Single Dose Each 500 et contains     1       Disclose Merzous 1000 II (Lin Solition Robotion for Infusion Single Dose Each 500 et contains     3       Disclose Merzous 1000 II (Lin Solition Robotion for Infusion Single Dose Each 500 et contains     3       Disclose Merzous 1000 II (Lin Solition Robotion for Infusion Single Dose Each 500 et contains     3       Disclose Merzous 1000 II (Lin Solition Robotion for Infusion Single Dose Each 500 et contains     3       Disclose Merzous 1000 II (Lin Solition Robotion for Infusion Single Dose Torus 1144 and solition for Infusion Single Dose Each 500 et contains     3       Do not researce and solition for Infusion Single Dose Torus 1144 and solition for Infusion Single Dose Each 500 et contains     4       Do not researce and solition for Infusion Single Dose Torus 1144 and solition for Infusion Single Dose Each 500 et contains     4       Do not researce and solition for Infusion Do not researce and solition for Infusion Each 500 et contains     6       Do not researce and solition for Infusion Do no	
Drug Interactions	See manufacturer's package insert for full prescribing information. 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. Platelet inhibitors: Drugs such as acetylsalicylic acid, dextran, phenylbutazone, ibuprofen, indomethacin, dipyridamole, hydroxychloroquine 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. Other interactions; Digitalis, tetracyclines, nicotine, or antihistamines may partially counteract the anticoagulant action of heparin sodium.	<ul> <li>Interaction with other medicinal products and other forms of interaction</li> <li>Heparin may prolong the one stage prothrombin time. Accordingly, when</li> <li>Heparin is given with dicoumarol or warfarin sodium, a period of at least 5</li> <li>hours after the last intravenous dose of heparin should elapse before blood is</li> <li>drawn, if a valid prothrombin time is to be obtained.</li> <li>Drugs such as acetylsalicylic acid, dextran, phenylbutazone, ibuprofen,</li> <li>indomethacin, dipyridamole, hydroxychloroquine and others which interfere</li> <li>with platelet aggregation (the main haemostatic defense of heparinized</li> <li>patients) may induce bleeding and should be used with caution in patients on</li> <li>heparin therapy.</li> <li>The use of ACE inhibitors and angiotensin-II antagonists in conjunction with</li> <li>heparin increase the risk of hyperkalaemia.</li> <li>Incompatibilities Do not add other drugs to Heparin Sodium in 0.9% Sodium</li> <li>Chloride Intravenous Infusion.</li> </ul>	
	Drug/Laboratory Tests Interactions Hyperaminotransferasemia Significant elevations of aminotransferase (SGOT [S-AST] and SGPT [S-ALT]) levels have occurred in a high percentage of patients (and healthy subjects) who have received heparin.		
Adverse Events	See manufacturer's package insert for full prescribing information.	The most frequently reported undesirable effects are bleeding events, reversible increase in liver enzymes, thrombocytopenia and various skin	





	US FDA approved product	Import Product
	Heparin Sodium and 0.9% Sodium Chloride Injection	Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion
	Image:	Code B09536     500 ml       Bacter     Boom       Heparin Sodium BP 2000 IV/L in Solidin Container     1       Hostini Sodium BP 2000 IV/L in Solidin Container     1       Solidin Chorido IV Infusion     1       Solidin Chorido IV Infusion     1       Solidin Chorido IV Infusion     3       Dotted Solidin For Infusion Single Oscie     1       Solidin Chorido IV Infusion     3       Solidin Chorido IV Infusion     3       Solidin Chorido IV Infusion     3       Discolutin Provphate 1724/0 2.8 g     Chorido 7       Discolutin Threadout 174     10       Solidin Chorido IV Infusion     3       Discolutin Solidin I Solidi
		Ovarian (corpus luteum) haemorrhage developed in a number of women of reproductive age receiving short or long-term anticoagulant therapy. This complication if unrecognized may be fatal.
Overdose and Treatment	Symptoms Bleeding is the chief sign of heparin overdosage. Nosebleeds, blood in urine or tarry stools may be noted as the first sign of bleeding. Easy bruising or petechial formations may precede frank bleeding. Treatment Neutralization of heparin effect. When clinical circumstances (bleeding) require reversal of heparinization, 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. Administration of protamine sulfate can cause severe hypotensive and anaphylactoid reactions. Because fatal reactions often resembling anaphylaxis have been reported, the drug should be given only when resuscitation techniques and treatment of anaphylactoid shock are readily available.	Overdose Bleeding is the chief sign of heparin overdosage. Protamine Sulphate (1% w/v solution) by slow intravenous infusion will neutralise heparin. No more than 50 mg should be given very slowly in any 10 minute period. Each mg of protamine sulphate neutralises approximately 100 units of heparin (or 1 to 1.5 mg neutralises approximately 1 mg of heparin). Heparins derived from various animal sources require different amounts of protamine sulphate for neutralisation. Decreasing amounts of protamine are required as time from the last heparin injection increases. Thirty minutes after a dose of heparin, approximately 0.5 mg of protamine is sufficient to neutralise each 100 units of heparin. Blood or plasma transfusions may be necessary; these dilute but do not neutralise heparin.

	US FDA approved product	Import Product           Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion	
	Heparin Sodium and 0.9% Sodium Chloride Injection		
	I I   I I <th>Code B09536     500 ml       Hearin Sodium RP 2000 IU/L in 0.9% w/x Sodium Chloride IV Infusion     1 1       WHX So diama Chloride IV Infusion     1 1       Number of Medicing Processing     1 1       Sodium Chloride IV Infusion     1 1       Sodium Chloride IV Infusion     1 1       Number of Medicing Processing     1 1       Sodium Chloride IV Infusion     1 1        Sodium Chlorid</th>	Code B09536     500 ml       Hearin Sodium RP 2000 IU/L in 0.9% w/x Sodium Chloride IV Infusion     1 1       WHX So diama Chloride IV Infusion     1 1       Number of Medicing Processing     1 1       Sodium Chloride IV Infusion     1 1       Sodium Chloride IV Infusion     1 1       Number of Medicing Processing     1 1       Sodium Chloride IV Infusion     1 1        Sodium Chlorid	
	For additional information the labeling of Protamine Sulfate Injection, USP products should be consulted.		
Storage Conditions	<ul> <li>Exposure of pharmaceutical products to heat should be minimized. Avoid excessive heat.</li> <li>It is recommended the product be stored at room temperature (25°C); brief exposure up to 40°C does not adversely affect the product.</li> </ul>	<ul> <li>Keep out of the sight and reach of children.</li> <li>Do not store above 25 °C.</li> <li>Heparin Sodium Solution must not be used if the container is damaged or the solution is not clear.</li> </ul>	
Directions for Use	<b>Warning:</b> Do not use plastic containers in series connections. Such use could result in air embolism due to residual air being drawn from the primary container before administration of the fluid from the secondary container is completed.	For use under medical supervision: The solution should only be used once. Any left over solution should be discarded. Do not use unless solution is clear and the container is undamaged. Discard any unused portion. Do not reconnect partially used bags.	
How Supplied	Heparin Sodium and 0.9% Sodium Chloride Injection in VIAFLEX Plus plastic container is supplied in 500 mL and 1,000 mL bags of Heparin and 0.9% Sodium Chloride Injection as follows: <b>2B0953</b> Heparin Sodium 1,000 units in 0.9% Sodium Chloride (500 mL) NDC 0338-0431-03 <b>2B0944</b> Heparin Sodium 2,000 units in 0.9% Sodium Chloride (1,000 mL) NDC 0338-0433-04	It is supplied as a clear solution for infusion (slow injection) in a VIAFLEX plastic bag with a plastic overpouch. Do not remove from overpouch until ready for use. 500 mL bag: FKB0953G Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion ( <b>1,000 IU/500 mL</b> ) NDC 0338-9556-20 1,000 mL bag: FKB0944G Heparin Sodium BP 2,000 IU/L in 0.9% w/v Sodium Chloride IV Infusion ( <b>2000 IU/1,000 mL</b> ) NDC 0338-9552-10	



## Barcode stickers for imported Heparin Sodium BP 2,000 IU/L\* in 0.9% w/v Sodium Chloride IV Infusion in VIAFLEX container

## Barcode stickers – Instructions for use:

To replace missing barcode stickers, please contact Baxter Product Surveillance at 1-800-437-5176.

- 1. Confirm receipt of the Dear Healthcare Provider (DHCP) letter and barcode sticker sheet consisting of product barcodes.
- 2. Review and confirm that the DHCP letter, the product received, and the barcodes match.
- 3. Affix barcode sticker onto the overwrap.
- 4. Scan the barcode on the product overwrap at the time of use.



1,000 units in 500 mL\* Product code: FKB0953G NDC 0338-9556-20 **FKB0953G** Heparin Sodium BP 2 units per mL in 0.9% w/v Sodium Chloride IV Infusion in VIAFLEX container **(1,000 units / 500 mL)** 



\* The imported products and FDA-approved products contain the same Heparin Sodium concentration of 2 units per mL.