POTASSIUM CHLORIDE IN DEXTROSE AND SODIUM CHLORIDE- dextrose monohydrate, sodium chloride, and potassium chloride injection, solution Fresenius Kabi USA, LLC

POTASSIUM CHLORIDE IN DEXTROSE AND SODIUM CHLORIDE

free flex®

 R_x only

Potassium Chloride in 5% Dextrose and 0.9% Sodium Chloride Injection, USP

Flexible Plastic Container

DESCRIPTION

Intravenous solutions with potassium chloride (I.V. solutions with KCl) are sterile and nonpyrogenic solutions in water for injection. They are for administration by intravenous infusion only.

See table below for summary of content and characteristics of these solutions.

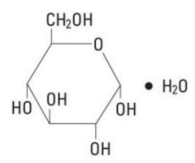
Table 1		COMPOSITION (g/L)				Approx. Ionic Concentrations (mEq/L)				is (mEq/L)
Potassium										-
Chloric	le									
in 5% Dextrose										
and 0.9	and 0.9%									
	Sodium									
Chloric	-									
Injection,	1								I	
mEq					Calculated	рΗ				Approximate
Potassium	(mL)	Hydrous	Chloride		Osmolarity		(Na+)	(K+)	(Cŀ)	Kcal/L
					(mOsmol/L)					
20 mEq	1000	50	9	1.49	600	4.2		20	174	170
						(3.5				
						to				
						6.5)				
40 mEq	1000	50	9	2.98	640	4.2	154	40	194	170
						(3.5				
						to				
						6.5)				

May contain HCl for pH adjustment.

The solutions contain no bacteriostat, antimicrobial agent or added buffer and each is intended only for use as a single-dose injection. When smaller doses are required the unused portion should be discarded.

These solutions are parenteral fluid, nutrient and/or electrolyte replenishers.

Dextrose, USP is chemically designated D-glucose, monohydrate ($C_6H_{12}O_6 \cdot H_2O$), a hexose sugar freely soluble in water. It has the following structural formula:



Potassium Chloride, USP is chemically designated KCl, a white granular powder freely soluble in water.

Sodium Chloride, USP is chemically designated NaCl, a white crystalline powder freely soluble in water.

Water for Injection, USP is chemically designated H_2O .

The flexible plastic container is fabricated from a specially formulated non-plasticized, film containing polypropylene and thermoplastic elastomers (**free***flex*[®] bag). Water can permeate from inside the container into the overwrap but not in amounts sufficient to affect the solution significantly. Solutions in contact with the plastic container may leach out certain chemical components from the plastic in very small amounts; however, biological testing was supportive of the safety of the plastic container materials. Exposure to temperatures above 25°C/77°F during transport and storage will lead to minor losses in the moisture content. Higher temperatures lead to greater losses. It is unlikely that these minor losses will lead to clinically significant changes within the expiration period.

CLINICAL PHARMACOLOGY

When administered intravenously, these solutions provide a source of water and potassium chloride with carbohydrate (dextrose) and sodium chloride. See **HOW SUPPLIED** section for specific concentrations of these various solutions.

Solutions containing carbohydrate in the form of dextrose restore blood glucose levels and provide calories. Carbohydrate in the form of dextrose may aid in minimizing liver glycogen depletion and exerts a protein-sparing action. Dextrose injected parenterally undergoes oxidation to carbon dioxide and water.

Intravenous solutions containing potassium chloride are particularly intended to provide needed potassium cation (K⁺). Potassium is the chief cation of body cells (160 mEq/liter of intracellular water). It is found in low concentration in plasma and extracellular fluids (3.5 to 5.0 mEq/liter in a healthy adult). Potassium plays an important role in electrolyte balance. Normally about 80 to 90% of the potassium intake is excreted in the urine; the remainder in the stools and to a small extent, in the perspiration. The kidney does not conserve potassium well so that during fasting or in patients on a potassium-free diet, potassium loss from the body continues resulting in potassium depletion. A deficiency of either potassium or chloride will lead to a deficit of the other.

Sodium chloride in water dissociates to provide sodium (Na⁺) and chloride (Cl⁻) ions. Sodium (Na⁺) is the principal cation of the extracellular fluid and plays a large part in the therapy of fluid and electrolyte disturbances. Chloride (Cl⁻) has an integral role in buffering action when oxygen and carbon dioxide exchange occurs in the red blood cells. The distribution and excretion of sodium (Na⁺) and chloride (Cl⁻) are largely under the control of the kidney which maintains a balance between intake and output.

Water is an essential constituent of all body tissues and accounts for approximately 70% of total body weight. Average normal adult daily requirement ranges from two to

three liters (1.0 to 1.5 liters each for insensible water loss by perspiration and urine production).

Water balance is maintained by various regulatory mechanisms. Water distribution depends primarily on the concentration of electrolytes in the body compartments and sodium (Na⁺) plays a major role in maintaining physiologic equilibrium.

INDICATIONS AND USAGE

These solutions are indicated in patients requiring parenteral administration of potassium chloride with minimal carbohydrate calories and sodium chloride.

CONTRAINDICATIONS

Solutions containing potassium chloride are contraindicated in diseases where high potassium levels may be encountered.

WARNINGS

Solutions which contain potassium ions should be used with great care, if at all, in patients with hyperkalemia, severe renal failure and in conditions in which potassium retention is present.

To avoid potassium intoxication, do not infuse these solutions rapidly. In patients with severe renal insufficiency or adrenal insufficiency, administration of potassium chloride may cause potassium intoxication.

Solutions containing sodium ions should be used with great care, if at all, in patients with congestive heart failure, severe renal insufficiency and in clinical states in which there exists edema with sodium retention.

In patients with diminished renal function, administration of solutions containing sodium or potassium ions may result in sodium or potassium retention.

The intravenous administration of these solutions can cause fluid and/or solute overloading resulting in dilution of serum electrolyte concentrations, overhydration, congested states or pulmonary edema.

The risk of dilutional states is inversely proportional to the electrolyte concentration of administered parenteral solutions. The risk of solute overload causing congested states with peripheral and pulmonary edema is directly proportional to the electrolyte concentrations of such solutions.

PRECAUTIONS

Clinical evaluation and periodic laboratory determinations are necessary to monitor changes in fluid balance, electrolyte concentrations and acid-base balance during prolonged parenteral therapy or whenever the condition of the patient warrants such evaluation.

Solutions containing dextrose should be used with caution in patients with known subclinical or overt diabetes mellitus.

Caution must be exercised in the administration of parenteral fluids, especially those containing sodium ions, to patients receiving corticosteroids or corticotropin.

Potassium replacement therapy should be guided primarily by serial electrocardiograms. Plasma potassium levels are not necessarily indicative of tissue potassium levels. High plasma concentrations of potassium may cause death through cardiac depression, arrhythmias or arrest.

Potassium-containing solutions should be used with caution in the presence of cardiac disease, particularly in digitalized patients or in the presence of renal disease.

Care should be exercised to insure that the needle (or catheter) is well within the lumen of the vein and that extravasation does not occur.

Do not administer unless solution is clear and container is undamaged. Discard unused portion.

Carcinogenesis, Mutagenesis, Impairment of Fertility:

Studies with solutions from flexible plastic containers have not been performed to evaluate carcinogenic potential, mutagenic potential or effects on fertility.

Pregnancy Animal reproduction studies have not been conducted with dextrose, potassium chloride or sodium chloride. It is also not known whether dextrose, potassium chloride or sodium chloride can cause fetal harm when administered to a pregnant woman or can affect reproduction capacity. Dextrose, potassium chloride or sodium chloride or a pregnant woman only if clearly needed.

Nursing Mothers:

Caution should be exercised when solutions from flexible plastic containers are administered to a nursing mother.

Pediatric Use:

The safety and effectiveness in the pediatric population are based on the similarity of the clinical conditions of the pediatric and adult populations. In neonates or very small infants the volume of fluid may affect fluid and electrolyte balance.

Frequent monitoring of serum glucose concentrations is required when dextrose is prescribed to pediatric patients, particularly neonates and low birth weight infants.

In very low birth weight infants, excessive or rapid administration of dextrose injection may result in increased serum osmolality and possible intracerebral hemorrhage.

Geriatric Use:

An evaluation of current literature revealed no clinical experience identifying differences in response between elderly and younger patients. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy. Sodium and potassium ions are known to be substantially excreted by the kidney, and the risk of toxic reactions may be greater in patients with impaired renal function. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selection, and it may be useful to monitor renal function.

ADVERSE REACTIONS

Reactions which may occur because of the solutions or technique of administration include febrile response, infection at the site of injection, venous thrombosis or phlebitis extending from the site of injection, extravasation and hypervolemia.

If an adverse reaction does occur, discontinue the infusion, evaluate the patient, institute appropriate therapeutic countermeasures and save the remainder of the fluid

for examination if deemed necessary.

Nausea, vomiting, abdominal pain and diarrhea have been reported with potassium therapy. The signs and symptoms of potassium intoxication include paresthesias of the extremities, flaccid paralysis, listlessness, mental confusion, weakness and heaviness of the legs, hypotension, cardiac arrhythmias, heart block, electrocardiographic abnormalities such as disappearance of P waves, spreading and slurring of the QRS complex with development of a biphasic curve and cardiac arrest.

Potassium-containing solutions are intrinsically irritating to tissues. Therefore, extreme care should be taken to avoid perivascular infiltration. Local tissue necrosis and subsequent sloughing may result if extravasation occurs. Chemical phlebitis and venospasm have also been reported.

Should perivascular infiltration occur, I.V. administration at that site should be discontinued at once. Local infiltration of the affected area with procaine hydrochloride, 1%, to which hyaluronidase may be added, will often reduce venospasm and dilute the potassium remaining in the tissues locally. Local application of heat may also be helpful.

To report SUSPECTED ADVERSE REACTIONS, contact Fresenius Kabi USA, LLC at 1-800-551-7176 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

OVERDOSAGE

In the event of potassium overdosage, discontinue the infusion immediately and institute intensive corrective therapy to reduce serum potassium levels. See **WARNINGS** and **PRECAUTIONS**.

DOSAGE AND ADMINISTRATION

These solutions should be administered only by intravenous infusion and as directed by the physician. The dose and rate of injection are dependent upon the age, weight and clinical condition of the patient. If the serum potassium level is greater than 2.5 mEq/liter, potassium should be given at a rate not to exceed 10 mEq/hour in a concentration less than 30 mEq/liter. Somewhat faster rates and greater concentrations (usually up to 40 mEq/liter) of potassium may be indicated in patients with more severe potassium deficiency. The total 24- hour dose should not generally exceed 200 mEq of potassium.

As reported in the literature, the dosage and constant infusion rate of intravenous dextrose must be selected with caution in pediatric patients, particularly neonates and low birth weight infants, because of the increased risk of hyperglycemia/hypoglycemia.

Drug Interactions

Additives may be incompatible. Consult with pharmacist, if available. When introducing additives, use aseptic technique, mix thoroughly and do not store.

Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit. See **PRECAUTIONS**.

HOW SUPPLIED

Potassium Chloride in 5% Dextrose and 0.9% Sodium Chloride Injection, USP is supplied in single-dose **free***flex*[®] plastic containers as follows:

Product Code Unit of Use	Strength	Unit of Sale
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219118	NDC 65219-118-01 One 1000 mL free <i>flex</i> [®] bag	20 mEq Potassium	NDC 65219-118-10 Package of 10 free flex [®]
		Chloride	bags
219119	NDC 65219-119-01	40 mEq	NDC 65219-119-10
	One 1000 mL free flex [®] bag	Potassium	Package of 10 free <i>flex</i> ®
		Chloride	bags

Avoid excessive heat. Store at 20 to 25°C (68 to 77°F). [See USP Controlled Room Temperature.] Protect from freezing.

The container closure is not made with natural rubber latex. Non-PVC, Non-DEHP, Sterile.

INSTRUCTIONS FOR USE

Check flexible container solution composition, lot number, and expiry date.

Do not remove solution container from its overwrap until immediately before use. Use sterile equipment and aseptic technique.

Flexible Plastic Container (**free***flex*[®] bag)

<u>To Open</u>

- 1. Turn solution container over so that the text is face down. Using the pre-cut corner tabs, peel open the overwrap and remove solution container.
- 2. Check the solution container for leaks by squeezing firmly. If leaks are found, or if the seal is not intact, discard the solution.
- 3. Do not use if the solution is cloudy or a precipitate is present.

To Add Medication

- 1. Identify WHITE Additive Port with arrow pointing toward container.
- 2. Immediately before injecting additives, break off WHITE Additive Port Cap with the arrow pointing toward container.
- 3. Hold base of WHITE Additive Port horizontally.
- 4. Insert needle horizontally through the center of WHITE Additive Port's septum and inject additives.
- 5. Mix container contents thoroughly.

Preparation for Administration

- 1. Immediately before inserting the infusion set, break off BLUE Infusion Port Cap with the arrow pointing away from container.
- 2. Use a non-vented infusion set or close the air-inlet on a vented set.
- 3. Close the roller clamp of the infusion set.
- 4. Hold the base of BLUE Infusion Port.
- 5. Insert spike through BLUE Infusion Port by rotating wrist slightly until the spike is inserted. **NOTE:** See full directions accompanying administration set.

WARNING: Do not use flexible container in series connections.

Manufactured for:



Lake Zurich, Illinois 60047

Made in Norway

Issued: June 2022

PACKAGE LABEL - PRINCIPAL DISPLAY - Potassium Chloride in 5% Dextrose and 0.9% Sodium Chloride Injection, USP Bag Label

20 mEq POTASSIUM

1000 mL NDC 65219-**118**-01 **freeflex**[®] 20 mEq POTASSIUM CHLORIDE

in 5% Dextrose and

0.9% Sodium Chloride Injection, USP

Rx Only

100	20 mEq POTASSIUM
	1000 mL NDC 65219- 118 -01 free <i>flex</i> [®]
200	20 mEq in 5% Dextrose and
300	0.9% Sodium Chloride Injection, USP
400	Each 100 mL contains: Potassium Chloride 149 mg; Sodium Chloride 900 mg; Dextrose, Hydrous 5 g in Water for Injection. May contain HCI for pH adjustment. Electrolytes per 1000 mL (not including ions for pH adjustment): Potassium 20 mEq; Sodium 154 mEq; Chloride 174 mEq. 600 mOsmol/L (Calc.) pH 4.2 (3.5 to 6.5)
	ADDITIVES MAY BE INCOMPATIBLE. CONSULT WITH PHARMACIST, IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC TECHNIQUE, MIX THOROUGHLY AND DO NOT STORE.
500	IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC
500	IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC TECHNIQUE, MIX THOROUGHLY AND DO NOT STORE. Single-Dose Container. For Intravenous Use.
500	IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC TECHNIQUE, MIX THOROUGHLY AND DO NOT STORE. Single-Dose Container. For Intravenous Use. Discard Unused Portion. Usual dosage: See package insert. The overwrap is a moisture barrier.
	IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC TECHNIQUE, MIX THOROUGHLY AND DO NOT STORE. Single-Dose Container. For Intravenous Use. Discard Unused Portion. Usual dosage: See package insert. The overwrap is a moisture barrier. Use immediately once removed from overwrap. STORE AT: 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature]. Avoid excessive heat.
	IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC TECHNIQUE, MIX THOROUGHLY AND DO NOT STORE. Single-Dose Container. For Intravenous Use. Discard Unused Portion. Usual dosage: See package insert. The overwrap is a moisture barrier. Use immediately once removed from overwrap. STORE AT: 20° to 25°C (68° to 77°F) [see USP Controlled Room Temperature]. Avoid excessive heat. Protect from freezing. 403883 The container closure is not made with natural FUH 3354 01-62-16-020

Made in Norway www.fresenius-kabi.com/us

LOT NO

EXP DATE

PACKAGE LABEL - PRINCIPAL DISPLAY - Potassium Chloride in 5% Dextrose and 0.9% Sodium Chloride Injection, USP Case Label

NDC 65219-118-10

20 mEq Potassium Chloride in 5% Dextrose and

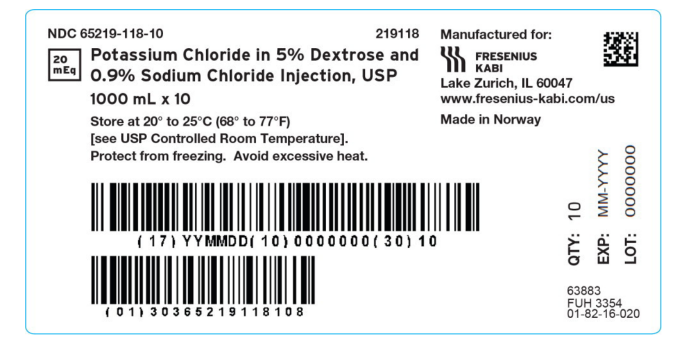
0.9% Sodium Chloride Injection, USP

1000 mL x 10

Store at 20° to 25°C (68° to 77°F)

[see USP Controlled Room Temperature].

Protect from freezing. Avoid excessive heat.



PACKAGE LABEL - PRINCIPAL DISPLAY - Potassium Chloride in 5% Dextrose and 0.9% Sodium Chloride Injection, USP Bag Label

40 mEq POTASSIUM

1000 mL NDC 65219-**119**-01 **free***flex*[®] **40 mEq POTASSIUM CHLORIDE**

in 5% Dextrose and

0.9% Sodium Chloride Injection, USP

Rx Only

100	40 mEq POTASSIUM
	1000 mL NDC 65219- 119 -01 free <i>flex</i> [®]
200	40 mEq FOTASSIUM CHLORIDE
300	in 5% Dextrose and 0.9% Sodium Chloride Injection, USP
400	Each 100 mL contains: Potassium Chloride 298 mg; Sodium Chloride 900 mg; Dextrose, Hydrous 5 g in Water for Injection. May contain HCl for pH adjustment. Electrolytes per 1000 mL (not including ions for pH adjustment): Potassium 40 mEq; Sodium 154 mEq; Chloride 194 mEq. 640 mOsmol/L (Calc.) pH 4.2 (3.5 to 6.5)
	ADDITIVES MAY BE INCOMPATIBLE. CONSULT WITH PHARMACIST, IF AVAILABLE. WHEN INTRODUCING ADDITIVES, USE ASEPTIC TECHNIQUE, MIX THOROUGHLY AND DO NOT STORE.
500	Single-Dose Container. For Intravenous Use. Discard Unused Portion. Usual dosage: See package insert. The overwrap is a moisture barrier. Use immediately once removed from overwrap.
600	STORE AT: 20° to 25°C (68° to 77°F) [see USPControlled Room Temperature]. Avoid excessive heat.Protect from freezing.403884The container closure is not made with naturalFUH 3364rubber latex.Non-PVC, Non-DEHP, Sterile.
700	Manufactured for: (01)00365219119012 (01)00365219119012 Lake Zurich, IL 60047 Made in Norway

PACKAGE LABEL - PRINCIPAL DISPLAY - Potassium Chloride in 5% Dextrose and 0.9% Sodium Chloride Injection, USP Case Label

NDC 65219-119-10

40 mEq Potassium Chloride in 5% Dextrose and

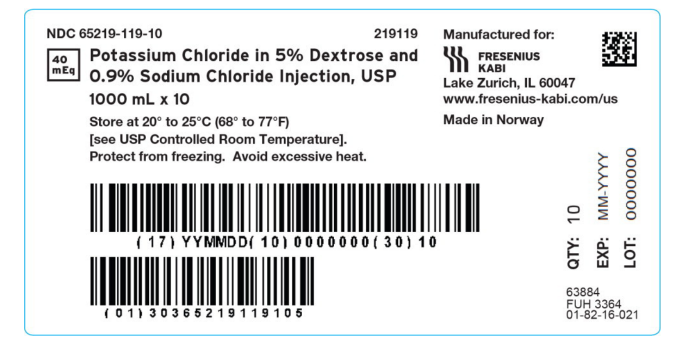
0.9% Sodium Chloride Injection, USP

1000 mL x 10

Store at 20° to 25°C (68° to 77°F)

[see USP Controlled Room Temperature].

Protect from freezing. Avoid excessive heat.



POTASSIUM CHLORIDE IN DEXTROSE AND SODIUM CHLORIDE					
dextrose monohydrate, sodium chloride, and potassium chloride injection, solution					
Product Information					
Product Type	ct Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:65219-118			5219-118	
Route of Administration INTRAVENOUS					
Active Increasiont/Active	Maiah				
Active Ingredient/Active	Molety				
Ingredient Name Basis of Strength Strength					
DEXTROSE MONOHYDRATE (UNII: LX22YL083G) (ANHYDROUS DEXTROSE - DEXTROSE 50 g					

UN	III:5SL0G7R0C)K)			MONO	HYDRATE	in 1000 mL		
SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37, CHLORIDE ION - UNII:Q32ZN48698)					SODIU	М	9 g in 1000 mL		
PC	TASSIUM CI	HLORIDE (UNII: 66	DYQ98110) (POTASSIUM CATION - UNII:Q32Z N48698)			SIUM	1.49 g in 1000 mL		
In	active Ing	gredients							
Ingredient Name							Strength		
W	ATER (UNII: 0	59QF0KO0R)							
HJ	DROCHLOR	IC ACID (UNII: QTT	17582CB)						
Pa	ackaging								
#	ltem Code		Package Description		Marketi Start Da		Marketing End Date		
1	NDC:65219- 118-10	10 in 1 CASE			08/31/2022				
1	NDC:65219- 118-01	1000 mL in 1 BAG Device/System (s	i; Type 2: Prefilled Drug Delivery yringe, patch, etc.)						
Μ	larketin	g Informat	ion						
			J 1		keting Start M Date		arketing End Date		
AN	IDA	ANDA21344	5 08/31/2022		2022				
-			DE IN DEXTROSE AN				RIDE		
ae	xtrose mor	ionydrate, sodil	Im chloride, and potassium cl	nioriae i	njection, so	lution			
Р	roduct In	formation							
	roduct Type		HUMAN PRESCRIPTION DRUG		ode (Source	e) (e	NDC:65219-119		
		ministration	INTRAVENOUS						

Active Ingredient/Active Moiety Basis of Strength **Ingredient Name** Strength DEXTROSE MONOHYDRATE (UNII: LX22YL083G) (ANHYDROUS DEXTROSE -DEXTROSE 50 g in 1000 mL UNII:5SL0G7R0OK) MONOHYDRATE SODIUM CHLORIDE (UNII: 451W47IQ8X) (SODIUM CATION - UNII:LYR4M0NH37, SODIUM 9 g in 1000 mL CHLORIDE ION - UNII:Q32ZN48698) CHLORIDE POTASSIUM CHLORIDE (UNII: 660YQ98I10) (POTASSIUM CATION -POTASSIUM 2.98 g

UNII:295053K152, CHLORIDE ION - UNII:Q32ZN48698)

 Inactive Ingredients

 Ingredient Name
 Strength

 WATER (UNII: 059QF0K00R)

 HYDROCHLORIC ACID (UNII: QTT17582CB)

 Packaging

in 1000 mL

CHLORIDE

#	ltem Code		Package Description			Marketing End Date
1	NDC:65219- 119-10	10 in 1 C	ASE	08/31/2022		
1	NDC:65219- 119-01		000 mL in 1 BAG; Type 2: Prefilled Drug Delivery Device/System (syringe, patch, etc.)			
M	larketin	g Info	ormation			
	Marketing Category		Application Number or Monograph Citation	Marketing Start Date		Marketing End Date
AN	ANDA		DA213445	08/31/2	2022	

Labeler - Fresenius Kabi USA, LLC (013547657)

Establishment

Name	Address	ID/FEI	Business Operations
HP Halden Pharma AS		347747373	MANUFACTURE(65219-118, 65219-119) , ANALYSIS(65219-118, 65219-119) , PACK(65219-119)

Revised: 5/2024

Fresenius Kabi USA, LLC