KOGENATE FS- antihemophilic factor (recombinant) Bayer HealthCare LLC

HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use KOGENATE FS safely and effectively. See full prescribing information for KOGENATE FS. KOGENATE FS (Antihemophilic Factor [Recombinant], Formulated with Sucrose)

For Intravenous Use, Lyophilized Powder for Reconstitution with Vial Adapter Initial U.S. Approval: 1993

- On-demand treatment and control of bleeding episodes in adults and children with hemophilia A.
- Perioperative management of bleeding in adults and children with hemophilia A.
- Routine prophylaxis to reduce the frequency of bleeding episodes in children with hemophilia A and to reduce the risk of joint damage in children without pre-existing joint damage.
- Routine prophylaxis to reduce the frequency of bleeding episodes in adults with hemophilia A.

Kogenate FS is not indicated for the treatment of von Willebrand disease (1).

DOSAGE AND ADMINISTRATION

For intravenous use only.

• Each vial of Kogenate FS contains the labeled amount of recombinant factor VIII in international units (IU, unit).

Control of bleeding episodes and perioperative management (2.1):

- Dose (units) = body weight (kg) x desired factor VIII rise (IU/dL or % of normal) x 0.5 (IU/kg per IU/dL).
- Titrate doses to patient's clinical response.
- Determine treatment frequency based on type of bleeding episode.

For routine prophylaxis in adults: 25 units per kg three times a week (2.1). For routine prophylaxis in children: 25 units per kg every other day (2.1).

Available as lyophilized powder in single use vials containing nominally 250, 500, 1000, 2000, and 3000 IU (3). Kogenate FS is provided with a sterile vial adapter with 15-micrometer filter and a prefilled diluent

syringe, which together serve as an alternative needleless reconstitution system.

Do not use in patients who have life-threatening hypersensitivity reactions, including anaphylaxis to mouse or hamster protein or other constituents of the product (4).

------ WARNINGS AND PRECAUTIONS ------

- Hypersensitivity reactions, including anaphylaxis, are possible. Should symptoms occur, discontinue treatment with Kogenate FS and administer appropriate treatment (5.1).
- Development of activity-neutralizing antibodies can occur in patients receiving factor VIII-containing products, including Kogenate FS. If expected plasma factor VIII activity levels are not attained, or if bleeding is not controlled with an expected dose, perform an assay that measures factor VIII inhibitor concentration (5.2).
- When clotting is normalized by treatment with factor VIII, development of cardiovascular risk factors may be the same as the risk for non-hemophilic patients (5.3).
- Monitor plasma factor VIII levels during infusions when indicated (5.4).

ADVERSE REACTIONS The most common adverse reactions (\geq 4%) in clinical trials are inhibitor formation (neutralizing antibodies) in previously untreated and minimally treated patients (PUPs and MTPs), skin-associated hypersensitivity reactions (e.g., rash, pruritus, urticaria), infusion site reactions (e.g., inflammation, pain), and central venous access device (CVAD) associated infections (6).

To report SUSPECTED ADVERSE REACTIONS, contact Bayer HealthCare at 1-888-842-2937 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

USE IN SPECIFIC POPULATIONS

• Pediatric Use: Higher factor VIII clearance may occur in children (4.4-16 years). Dose adjustment may be needed (8.4).

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling. Revised: 12/2019

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1 INDICATIONS AND USAGE

Kogenate[®] FS is a recombinant antihemophilic factor indicated for:

- On-demand treatment and control of bleeding episodes in adults and children with hemophilia A.
- Perioperative management of bleeding in adults and children with hemophilia A.
- Routine prophylaxis to reduce the frequency of bleeding episodes in children with hemophilia A and to reduce the risk of joint damage in children without pre-existing joint damage.
- Routine prophylaxis to reduce the frequency of bleeding episodes in adults with hemophilia A.

Kogenate FS is not indicated for the treatment of von Willebrand disease.

2 DOSAGE AND ADMINISTRATION

For intravenous use after reconstitution only.

2.1 Dose

- Dosage and duration of treatment depend on the severity of the factor VIII deficiency, the location and extent of bleeding, and the patient's clinical condition.¹ Careful control of replacement therapy is especially important in cases of major surgery or life-threatening bleeding episodes.
- Each vial of Kogenate FS has the recombinant factor VIII (rFVIII) potency in international units (IU, unit) stated on the label. One IU (unit), as defined by the World Health Organization standard for blood coagulation factor VIII, human, is approximately equal to the level of factor VIII activity found in 1 mL of fresh pooled human plasma.
- The expected *in vivo* peak increase in factor VIII level expressed as IU/dL (or % normal) can be estimated using the following formulas:

Dosage (units) = body weight (kg) x desired factor VIII rise (IU/dL or % of normal) x 0.5 (IU/kg per IU/dL)

or

IU/dL (or % normal) = Total Dose (IU)/body weight (kg) x 2 [IU/dL]/[IU/kg]

• Titrate dose to the patient's clinical response. Patients may vary in their pharmacokinetic (e.g., half-life, *in vivo* recovery) and clinical responses to Kogenate FS.^{2,3,4} Although the dose can be estimated by the calculations above, it is highly recommended that appropriate laboratory tests, including serial factor VIII activity assays, are performed [see Warnings and Precautions (5.3) and Clinical Pharmacology (12.3)].

On-Demand Treatment andControl of Bleeding Episodes

A guide for dosing Kogenate FS for on-demand treatment and control of bleeding episodes is provided in Table 1. The goal of treatment is to maintain a plasma factor VIII

activity level at or above the plasma levels (in % of normal or in IU/dL) outlined in Table 1.

Table 1 Dosing for On-Demand Treatment and Control of Bleeding Episodes

Type of Bleeding Episodes	Factor VIII Level Required (IU/dL or % of normal)	Dose (IU/kg)	Frequency of Doses (hours)	Duration of Therapy (days)
Minor Early hemarthrosis, minor muscle or oral bleeds.	20 - 40	10 - 20	Repeat dose if there is evidence of further bleeding.	Until bleeding is resolved
Moderate Bleeding into muscles, bleeding into the oral cavity, definite hemarthroses, and known trauma.	30 - 60	15 - 30	12 - 24	Until bleeding is resolved
Major Gastrointestinal bleeding. Intracranial, intra- abdominal or intrathoracic bleeding, central nervous system bleeding, bleeding in the retropharyngeal or retroperitoneal spaces, or iliopsoas sheath. Fractures. Head trauma.	80 - 100	Initial: 40 – 50 Repeat: 20 – 25	8 - 12	Until bleeding is resolved

Perioperative Management of Bleeding

A guide for dosing Kogenate FS during surgery (perioperative management of bleeding) is provided in Table 2. The goal of treatment is to maintain a plasma factor VIII activity level at or above the plasma level (in % of normal or in IU/dL) outlined in Table 2.

Factor VIIILevelType of SurgeryRequired		Frequency of Doses	Duration of Therapy
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	(IU/dL or % of normal)	(IU/KY)	(hours)	(days)
Minor Including tooth extraction	30 - 60	15 - 30	12 - 24	Until bleeding is resolved.
Major Examples include tonsillectomy, inguinal herniotomy, synovectomy, total knee replacement, craniotomy, osteosynthesis, trauma.	100	50 Pre-operatively to achieve 100% activity.	6 – 12 to keep FVIII activity in desired range	Until healing is complete.

Routine Prophylaxis in Adults

The recommended dose for routine prophylaxis is 25 units per kg of body weight three times per week.

Routine Prophylaxis in Children

The recommended dose for routine prophylaxis is 25 units per kg of body weight every other day.⁵

2.2 Preparation and Reconstitution

Kogenate FS is administered by intravenous injection after reconstitution. Patients should follow the specific reconstitution and administration procedures provided by their physicians.

Reconstitute and administer Kogenate FS with the components provided with each package. If any component of the package is opened or damaged, do not use this component.

Product reconstitution, administration, and handling of the administration set and needles must be done with caution because percutaneous puncture with a needle contaminated with blood can transmit infectious viruses, including HIV (AIDS) and hepatitis. Place needles in a sharps container after single use. Discard all equipment, including any reconstituted Kogenate FS product, in an appropriate container. Obtain immediate medical attention if injury occurs.

The procedures below are provided as general guidelines for the reconstitution of Kogenate FS provided with a sterile vial adapter with 15-micrometer filter and a prefilled diluent syringe, which together serve as an alternative needleless reconstitution system.

- Work on a clean surface and wash hands thoroughly using soap and warm water before performing the procedures.
- Reconstitute Kogenate FS with the components provided with each package. If any component of the package is opened or damaged, do not use this component.

• Filter the reconstituted product prior to administration to remove potential particulate matter in the solution. Filtering can be achieved by using the vial adapter.

Vacuum Transfer and Reconstitution

- Prepare the product under aseptic conditions.
- Warm both unopened vial and syringe in your hands to a comfortable temperature (do not exceed 37°C or 99°F).
- Remove protective cap from the vial (A). Aseptically cleanse the rubber stopper with alcohol, being careful not to handle the rubber stopper.
- Place product vial on a firm, non-skid surface. Peel off the paper cover on the vial adapter plastic housing. Do not remove the adapter from the plastic housing. Holding the adapter housing, place over the product vial and firmly press down (B). The adapter will snap over the vial cap. Do not remove the adapter housing at this step.
- Holding the syringe by the barrel, snap the syringe cap off the tip (C). Do not touch the syringe tip with your hand or any surface. Set the syringe aside for further use.
- Now remove and discard the adapter housing (D).
- Attach the prefilled syringe to the threaded vial adapter by turning clockwise (E).
- Grasp the plunger rod by the top plate and remove from carton. Avoid touching the sides and threads of the plunger rod. Immediately attach the plunger rod by turning it firmly clockwise into the threaded syringe rubber stopper (F).
- Inject the diluent by slowly pushing down on the plunger rod (G).
- Swirl vial gently until all material is dissolved (H). Do not shake vial. Be sure that the powder is completely dissolved. Do not use solutions containing visible particles or that are cloudy.
- Withdraw solution into the syringe by holding the vial on end above the vial adapter and syringe (I) then draw the plunger rod out slowly and smoothly. Ensure that the entire content of the vial is drawn into the syringe.
- With the plunger rod in place, remove the syringe from the vial adapter (the latter should remain attached to the vial). Attach the syringe to the administration set provided and inject intravenously (J).
- If the same patient is to receive more than one bottle, reconstitute each bottle with the diluent syringe provided then combine solutions in a larger syringe (not provided) and administer as usual.

2.3 Administration

For intravenous use after reconstitution only.

- Inspect Kogenate FS visually for particulate matter and discoloration prior to administration, wherever solution and container permit. Do not use Kogenate FS if you notice any particulates or turbidity in the solution.
- Store the reconstituted Kogenate FS at room temperature prior to administration, but administer it within 3 hours.
- Administer Kogenate FS using the administration set provided over a period of 1 to 15 minutes. Adapt the rate of administration to the response of each individual patient. Determine the pulse rate before and during administration of Kogenate FS. If there is a significant increase in pulse rate, reduce the rate of administration or temporarily halt the infusion allowing the symptoms to disappear promptly.

3 DOSAGE FORMS AND STRENGTHS

Kogenate FS is available as a lyophilized powder in single use glass vials containing nominally 250, 500, 1000, 2000, and 3000 International Units (IU, unit).

Each vial of Kogenate FS is labeled with the recombinant antihemophilic factor activity expressed in International Units per vial. This potency assignment employs a factor VIII concentrate standard that is referenced to a WHO International Standard for factor VIII concentrates, and is evaluated by appropriate methodology to ensure accuracy of the results.

4 CONTRAINDICATIONS

Kogenate FS is contraindicated in patients who have life-threatening hypersensitivity reactions, including anaphylaxis to mouse or hamster protein or other constituents of the product (sucrose, glycine, histidine, sodium, calcium chloride, polysorbate 80, imidazole, tri-n-butyl phosphate, and copper).

5 WARNINGS AND PRECAUTIONS

5.1 Hypersensitivity Reactions

Hypersensitivity reactions, including anaphylaxis have been reported with Kogenate FS. Reported symptoms included facial swelling, flushing, hives, decrease in blood pressure, nausea, rash, restlessness, shortness of breath, tachycardia, tightness of the chest, tingling, urticaria, and vomiting.

Kogenate FS contains trace amounts of mouse immunoglobulin G (MuIgG) and hamster (BHK) proteins *[see Description (11)]*. Patients treated with this product may develop hypersensitivity to these non-human mammalian proteins.

Discontinue Kogenate FS if symptoms occur and seek immediate emergency treatment.

5.2 Neutralizing Antibodies

Neutralizing antibodies (inhibitors) have been reported following administration of Kogenate FS, predominantly in previously untreated patients (PUPs) [see Adverse Reactions (6)]. Carefully monitor patients for the development of factor VIII inhibitors, using appropriate clinical observations and laboratory tests.⁶ If expected plasma factor VIII activity levels are not attained, or if bleeding is not controlled with an expected dose, perform an assay that measures factor VIII inhibitor concentration [see Warnings and Precautions (5.4)].

5.3 Cardiovascular Risk Factors

Hemophilic patients with cardiovascular risk factors or diseases may be at the same risk to develop cardiovascular events as non-hemophilic patients when clotting has been normalized by treatment with factor VIII.

5.4 Monitoring Laboratory Tests

- Monitor plasma factor VIII activity levels by the one-stage clotting assay to confirm the adequate factor VIII levels have been achieved and maintained, when clinically indicated [see Dosage and Administration (2)].
- Monitor for development of factor VIII inhibitors. Perform assay to determine if factor VIII inhibitor is present. If expected factor VIII activity plasma levels are not attained or if bleeding is not controlled with the expected dose of Kogenate FS, use Bethesda Units (BU) to titer inhibitors.
- If the inhibitor is less than 10 BU per mL, the administration of additional Kogenate FS concentrate may neutralize the inhibitor and may permit an appropriate hemostatic response.
- If inhibitor titers are above 10 BU per mL, adequate hemostasis may not be achieved. The inhibitor titer may rise following Kogenate FS infusion as a result of an anamnestic response to factor VIII. The on-demand treatment and control of bleeding in such patients requires the use of alternative therapeutic approaches and agents.

6 ADVERSE REACTIONS

Serious adverse reactions seen with Kogenate FS are systemic hypersensitivity reactions, including bronchospastic reactions and/or hypotension and anaphylaxis, and the development of high-titer inhibitors necessitating alternative treatments to factor VIII.

The most common adverse reactions (\geq 4%) observed in clinical trials were inhibitor formation in previously untreated patients (PUPs) and minimally treated patients (MTPs), skin-related hypersensitivity reactions (e.g., rash, pruritus), infusion site reactions (e.g., inflammation, pain), and central venous access device (CVAD) associated infections.

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in clinical trials of another drug and may not reflect the rates observed in clinical practice.

Previously Treated Patients (PTPs)

During the open-label clinical studies conducted in 73 PTPs, there were 24 adverse reactions reported in the course of 24,936 infusions.

Adverse reactions reported by \geq 4% of the patients are listed in Table 3 below.

Table 3 Adverse Reactions (AR) in Previously Treated Patients with Frequency of ≥ 4% (Age Range 12-59 years)

MedDRA Primary SOC	Preferred Term	N = 73 AR (%)
Skin and Subcutaneous Tissue Disorders	Rash, pruritus	6 (8.2%)
General Disorders and Administration Site Conditions	Infusion site reactions	3 (4.1%)
SOC = System Organ Class		

Previously Untreated Patients (PUPs) and Minimally Treated Patients (MTPs)

In clinical studies with pediatric PUPs and MTPs, there were 29 adverse reactions reported in the course of 9,389 infusions.

Adverse reactions reported by \geq 4% of the patients are listed in Table 4 below.

Table 4 Adverse Reactions (AR) in Previously Untreated Patients and Minimally Treated Patients with Frequency of $\geq 4\%$ (Age Range 2-27 months)

MedDRA Primary SOC	Preferred Term	N = 61 AR (%)
Skin and Subcutaneous Tissue Disorders	Rash, pruritus, urticaria	10 (16%)
Blood and Lymphatic System Disorders	Factor VIII inhibition (neutralizing antibodies)	9 (15%)*
General Disorders and Administration Site Conditions	Infusion site reactions	4 (7%)
SOC = System Organ Class	· · · · · · · · · · · · · · · · · · ·	

* Denominator for *de novo* inhibitors is N = 60, since one patient had a pre-existing inhibitor.

Minimally Treated Patients (MTPs) in the Joint Outcome Study

In the Joint Outcome Study with pediatric MTPs treated with routine prophylaxis or episodic enhanced treatment for 5.5 years, 46 of the 65 randomized patients experienced adverse events over the study duration.

Table 5 Adverse Reactions in Minimally Treated Patients in the Joint OutcomeStudy (Age Range 0-6 years)

MedDRA Primary SOC	Preferred Term	Prophylaxis Arm	Enhanced Episodic Arm
		N = 32	N = 33
		AR (%)	AR (%)

Surgical and Medical Procedures	Central venous catheterization, Catheter removal	19 (59%)	18 (55%)*
Infections and Infestations	Central line infection	6 (19%)	6 (18%)
General Disorders and Administration Site Conditions	Pyrexia	1 (3%)	4 (12%)
SOC = System Organ	Class		

* Three patients from the enhanced episodic arm had catheter removal.

Immunogenicity

In clinical studies with 73 PTPs (defined as having more than 100 exposure days), one patient had a pre-existing inhibitor. In the other 72 patients, followed over 4 years, no *de novo* inhibitors were observed.

In clinical studies with pediatric PUPs and MTPs, inhibitor development was observed in 9 out of 60 patients (15%), 6 were high titer¹ (> 5 BU) and 3 were low-titer inhibitors. Inhibitors were detected at a median number of 7 exposure days (range 2 to 16 exposure days).

In the Joint Outcome Study with Kogenate FS,⁵ *de novo* inhibitor development was observed in 8 of 64 patients with negative baseline values (12.5%), 2 patients developed high titer¹ (> 5 BU) and were withdrawn from the study. Six patients developed low-titer inhibitors. Inhibitors were detected at a median number of 44 exposure days (range 5 to 151 exposure days).

Inhibitor data in PUPs have been collected in several postmarketing registries [see *Postmarketing Experience (6.2)*].

The detection of antibody formation is dependent on the sensitivity and specificity of the assay. Additionally, the observed incidence of antibody (including neutralizing antibody) positivity in an assay may be influenced by several factors including assay methodology, sample handling, timing of sample collection, concomitant medications, and underlying disease. For these reasons, comparison of the incidence of antibodies to Kogenate FS with the incidence of antibodies to other products may be misleading.

6.2 Postmarketing Experience

Because adverse 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. The following adverse reaction has been identified during post approval use of Kogenate FS:

Sensory System – Dysgeusia

Immunogenicity - Postmarketing Registries

Data from the Research of Determinants of Inhibitor Development (RODIN) study⁷, French National Registry (FranceCoag)⁸ and United Kingdom Haemophilia Centre Doctors' Organisation (UKHCDO)⁹ registry reported an inhibitor development rate in PUPs for Kogenate FS of 38%, 50% and 35%, respectively, which is comparable to previously-reported inhibitor rates¹⁰ for FVIII products. These registry studies show a trend towards an increased risk of inhibitor development in PUPs, as compared to the reference rFVIII product. A survey of Canadian hemophilia centers¹¹ (2005 to 2012) and available data from the European Haemophilia Safety Surveillance (EUHASS)¹² registry from 2009 to 2013, reported an inhibitor development rate in PUPs for Kogenate FS of 42% and 31%, respectively, with no statistically significant differences observed across FVIII products.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

There are no data with Kogenate FS use in pregnant women to inform on drugassociated risk. Animal reproduction studies have not been conducted with Kogenate FS. It is also not known whether Kogenate FS can cause fetal harm when administered to a pregnant woman or affect reproductive capacity. 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.

8.2 Lactation

Risk Summary

There is no information regarding the presence of Kogenate FS in human milk, the effects on the breastfed infant, or the effects on milk production. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for Kogenate FS and any potential adverse effects on the breastfed child from Kogenate FS or from the underlying maternal condition.

8.4 Pediatric Use

Safety and efficacy studies have been performed in previously untreated and minimally treated pediatric patients. Children, in comparison to adults, present higher factor VIII clearance values and, thus, lower half-life and recovery of factor VIII. This may be due to differences in body composition.¹³ Account for this difference in clearance when dosing or following factor VIII levels in the pediatric population [see Clinical Pharmacology (12.3)].

Routine prophylactic treatment in children ages 0–2.5 years with no pre-existing joint damage has been shown to reduce spontaneous joint bleeding and the risk of joint damage. This data can be extrapolated to ages >2.5–16 years for children who have no existing joint damage [see Clinical Studies (14)].

8.5 Geriatric Use

Clinical studies with Kogenate FS did not include patients aged 65 and over. Dose selection for an elderly patient should be individualized.

11 DESCRIPTION

Kogenate FS Antihemophilic Factor (Recombinant) is a coagulation factor VIII produced by recombinant DNA technology. It is produced by Baby Hamster Kidney (BHK) cells into which the human factor VIII gene has been introduced.¹⁴ The cell culture medium contains Human Plasma Protein Solution (HPPS) and recombinant insulin, but does not contain any proteins derived from animal sources. Kogenate FS is a purified glycoprotein consisting of multiple peptides including an 80 kD and various extensions of the 90 kD subunit. It has the same biological activity as factor VIII derived from human plasma. No human or animal proteins, such as albumin, are added during the purification and formulation processes of Kogenate FS.

The purification process includes a solvent/detergent virus inactivation step in addition to the use of the purification methods of ion exchange chromatography, monoclonal antibody immunoaffinity chromatography, along with other chromatographic steps designed to purify recombinant factor VIII and remove contaminating substances.

Additionally, the manufacturing process was investigated for its capacity to decrease the infectivity of an experimental agent of transmissible spongiform encephalopathy (TSE), considered as a model for the vCJD and CJD agents.¹⁵⁻¹⁹ Several of the individual production and raw material preparation steps in the Kogenate FS manufacturing process have been shown to decrease TSE infectivity of that experimental model agent. These TSE reduction steps include the Fraction II+III separation step for HPPS (6.0 log10) and an anion exchange chromatography step (3.6 log10).

Kogenate FS is formulated with the following as stabilizers (see Table 6) in the final container and is then lyophilized. The final product does not contain any preservative. It is a sterile, nonpyrogenic, powder preparation for intravenous injection. Intravenous administration of sucrose contained in Kogenate FS will not affect blood glucose levels.

Stabilizer	250 IU, 500 IU, 1000 IU Nominal Vial Sizes	2000 IU, 3000 IU Nominal Vial Sizes	
Sucrose	0.9-1.3%	0.9-1.2%	
Glycine	21-25 mg/mL	20–24 mg/mL	
Histidine	18-23 mmol/L	17-22 mmol/L	

Table 6 Stabilizers Contained in Kogenate FS Final Container

Table 7 lists the inactive ingredients/excipients also contained in the final product.

Inactive Ingredient/Excipient	250 IU, 500 IU, 1000 IU	2000 IU, 3000 IU
Sodium	27-36 mEq/L	26–34 mEq/L
Calcium	2.0-3.0 mmol/L	1.9–2.9 mmol/L
Chloride	32-40 mEq/L	31-38 mEq/L
Polysorbate 80	64–96 μg/mL	64–96 μg/mL
Sucrose	28 mg/vial	52 mg/vial
Imidazole, tri-n-butyl phosphate, and copper	Trace amounts	Trace amounts

Table 7 Inactive Ingredients/Excipients

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Kogenate FS temporarily replaces the missing clotting factor VIII that is needed for effective hemostasis.

12.2 Pharmacodynamics

The activated partial thromboplastin time (aPTT) is prolonged in patients with hemophilia. Determination of aPTT is a conventional *in vitro* assay for biological activity of factor VIII. Treatment with Kogenate FS normalizes the aPTT over the effective dosing period.

12.3 Pharmacokinetics

The pharmacokinetic properties of Kogenate FS were investigated in two separate studies in adult and pediatric previously treated patients (PTPs).

Pharmacokinetic studies with Kogenate FS were conducted in 20 PTPs (ages 12 to 33 years) with severe hemophilia A. The pharmacokinetic parameters for Kogenate FS were measured in a randomized, crossover clinical trial with the predecessor KOGENATE product using a single dose administration of 50 IU per kg. After 24 weeks, the same dose of Kogenate FS was administered to the same patients. The recovery and half-life data for Kogenate FS were unchanged after 24 weeks of continued treatment with sustained efficacy and no evidence of factor VIII inhibition (see Table 8).

Table 8 Pharmacokinetic Parameters for Kogenate FS Compared toKOGENATE

Parameter	Kogen	KOGENATE	
	Initial PK (Mean±SD)	PK at week 24 (Mean±SD)	Reference (Mean±SD)
AUC (IU•h/dL)	1588.05 ± 344.32	1487.08 ± 381.73	1879.02 ± 412.32
Cmax (IU/dL)	114.95 ± 20.19	109.42 ± 20.09	127.40 ± 33.21
Half-life (hr)	13.74 ± 1.82	14.60 ± 4.38	14.07 ± 2.62
In Vivo Recovery (IU/dL / IU/kg)	2.20 ± 0.34	2.11 ± 0.37	2.43 ± 0.60

The pharmacokinetics of Kogenate FS were investigated in pediatric PTPs (4.4–18.1 years of age, average age 12).¹³ The pharmacokinetic parameters in children compared to adults show differences in higher clearance, lower incremental *in vivo* factor VIII recovery and a shorter factor VIII half-life. The pharmacokinetic parameters are depicted in Table 9.

Parameter	Mean (range)
AUC (IU•h/dL)	1320.0
Clearance (mL/h•kg)	4.1
Half-life (hr)	10.7 (7.8–15.3)

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

No studies have been conducted with Kogenate FS to assess its mutagenic or carcinogenic potential and impairment of fertility. By inference, the predecessor KOGENATE product and Kogenate FS would be expected to have equivalent mutagenic and carcinogenic potential.

The predecessor product did not demonstrate reverse mutation or chromosomal aberrations at doses substantially greater than the maximum expected clinical dose. *In vivo* evaluation with the predecessor product in animals using doses ranging between 10 and 40 times the expected clinical maximum also indicated that the predecessor product did not possess a mutagenic potential. Long-term investigations of carcinogenic potential in animals have not been performed due to the immune response to heterologous proteins in all non-human mammalian species.

13.2 Animal Toxicology and/or Pharmacology

Preclinical studies evaluating Kogenate FS in hemophilia A with mice, rats, rabbits, and dogs demonstrated safe and effective restoration of hemostasis. Doses several fold higher than the recommended clinical dose (related to body weight) did not demonstrate any acute or subacute toxic effect for Kogenate FS in laboratory animals.

Kogenate FS has been shown to be comparable to the predecessor KOGENATE product with respect to its biochemical and physiochemical properties, as well as its non-clinical in vivo pharmacology and toxicology.

14 CLINICAL STUDIES

Previously Treated Patients (PTPs) Clinical Studies

A total of 73 patients with severe ($\leq 2\%$ FVIII) hemophilia A, ages 12–59, who had been previously treated with other recombinant or with plasma-derived AHF products, were treated up to 54 months in open label studies with Kogenate FS. A total of 5,684 bleeding episodes were treated during the studies; 92.7% of the bleeds were treated with one (79.7%) or two (13.0%) infusions. Patients could be treated with on-demand or prophylaxis. Regularly scheduled prophylaxis treatment represented 76% of all infusions (treatment regimens of 2–3 infusions per week).

A total of 30 patients received Kogenate FS for 41 surgical procedures during the PTP studies. There were both minor and major surgery types, 16 and 25 respectively. Efficacy was measured by the attending surgeon based on a comparison of estimated blood loss from experience with non-hemophilic patients undergoing similar procedures. The surgeon or treating physician assigned a rating to the hemostatic outcome according to 4 categories: "excellent (blood loss less than expected)," "good (blood loss as expected)," "moderate (blood loss more than expected)," or "none (uncontrolled bleeding)." Hemostasis was rated as satisfactory ("excellent" or "good") in all cases.

Previously Untreated Patients (PUPs) and Minimally Treated Patients (MTPs)

Clinical Study

Kogenate FS has been used in the treatment of bleeding episodes in pediatric PUPs and MTPs with severe (<2% FVIII) hemophilia A. There were 37 PUPs and 24 MTPs (defined as having equal to or less than 4 exposure days) treated with a total of 9,419 infusions of Kogenate FS for a follow up duration up to 3.1 years. A total of 1047 bleeding episodes were treated; the bleeds were treated with one (73%) or two (15%) infusions.

A total of 27 surgical procedures were performed in 22 patients during the PUPs and MTPs study. There were both minor and major surgery types, 21 and 6 respectively. The attending surgeon measured efficacy and assigned a rating to the hemostatic outcome according to 4 categories as described above for PTPs. Hemostasis was rated as satisfactory ("excellent" or "good") in all cases.

Adult Prophylaxis for Bleeding Frequency Reduction

A 3-year, multicenter, open-label, parallel-group, prospective, randomized, controlled clinical study of the effect of routine prophylaxis with Kogenate FS versus on-demand use on bleeding frequency in adults and adolescents included 84 PTPs with severe Hemophilia A (FVIII level < 1 IU/dL), age 15 to 50 years. Patients were matched at baseline on demographic and disease characteristics. The median number of bleeds in the year before enrollment was 18.

Patients were randomized 1:1 to prophylaxis (25 units per kg three times a week) or ondemand use of Kogenate FS. Escalation of the prophylaxis dose by 5 units per kg/infusion after years 1 and 2, up to a maximum of 35 units per kg/infusion, was allowed.

Bleeding frequency was analyzed in the intent-to-treat population after a median followup period of 1.4 years. Patients who received prophylaxis experienced statistically significantly fewer bleeds (p<0.0001) compared to patients treated on-demand regardless of baseline subgroups examined including age, bleeding history, and presence or absence of target joints. The ratio of the mean bleeding frequency was 15.2 (95% CI: 8.5, 27.2; p<0.0001) for on-demand versus prophylaxis, indicating that patients who received on-demand treatment experienced on average 15.2 times as many bleeds compared to patients treated with prophylaxis. The mean annualized bleed rates (bleeds/subject/year) were 37 in the on-demand group versus 2 in the prophylaxis group. The median annualized bleed rate (bleeds/subject/year) in the on-demand group was 33 versus zero in the prophylaxis group. Most of the bleeding occurred in joints: the median joint bleed rate (joint bleeds/subject/year) was 24 in the on-demand group versus zero in the prophylaxis group. The mean annualized joint bleed rate was 29 in the on-demand group versus 2 in the prophylaxis group.

Twenty-two of 42 (52%) prophylaxis subjects experienced no bleeding, and 12 of 42 (29%) prophylaxis subjects experienced only 1–2 bleeds during the follow-up period.

Among prophylaxis patients the mean number of infusions/week was 2.8, and the median dose per prophylaxis infusion was 26 units per kg.

Pediatric Prophylaxis for Joint Damage Risk Reduction

A total of 65 boys less than 30 months of age with severe hemophilia A (FVIII level ≤ 2 IU/dL) and with ≤ 2 bleeds into each index joint and normal baseline joint imaging, were observed for up to 5.5 years in a multicenter, open-label, prospective, randomized, controlled clinical study.⁵ Patients received either 25 IU per kg every other day (primary

prophylaxis; n = 32) or at least 3 doses totaling a minimum of 80 IU per kg at the time of a bleeding episode (enhanced episodic; n = 33). Joint damage was evaluated by magnetic resonance imaging (MRI) or radiography, as well as the frequency of bleeding episodes. Joint damage detected by MRI or radiography in the ankles, knees, and elbows (i.e., index joints) was statistically significantly lower (p = 0.002) for subjects receiving prophylactic therapy (7%) than for subjects receiving episodic therapy (42%). This corresponds to a 6.29-fold relative risk of joint damage for subjects treated with enhanced episodic therapy compared to prophylaxis. The mean rate of index joint hemorrhages for subjects on episodic therapy was 4.89 bleeds per year, versus 0.63 bleeds per year observed in the prophylaxis arm. Three of 33 (9.1%) subjects in the episodic arm experienced recurrent life threatening bleeds (intracranial, gastrointestinal) compared to no subjects in the prophylaxis arm. On a per joint basis, joints in the regular prophylaxis arm were 8-fold more likely to remain damage-free than those in the episodic arm. Joint damage was most frequently observed in ankle joints and was detected at higher rates by MRI than by radiography. Ankles were also the index joint that demonstrated the highest frequency of bleeding events in this study (left ankle, mean 2.7 hemorrhages; right ankle, mean 2.6 hemorrhages).

As shown in Table 10 below, the incidence of joint damage was statistically significantly lower in the prophylactic group as compared to the episodic treatment group when assessed by MRI, or either MRI or radiography, using predefined criteria (described below) for establishing joint damage. However, there was no statistically significant difference between the two groups when joint damage was assessed by radiography alone.

To evaluate joint damage, MRIs were scored using a scale developed by Nuss et al.,²⁰ and X-rays were scored using the method of Pettersson et al.²¹ Both scales have been validated in various clinical trials and are routinely used for joint damage evaluation in hemophiliacs. Joint damage was defined as bone and/or cartilage damage including subchondral cysts, erosions and cartilage loss with narrowing of joint space. This corresponded to a total MRI score of \geq 7 or an X-ray score of \geq 1 in any of the following categories: subchondral cysts, erosions of joint surfaces or narrowing of joint spaces. Images were read separately by two independent radiologists centrally. Any discrepant reading was read by an independent third radiologist who was not aware of the initial reading results. The concordant reading of two out of three readers was used for analysis purposes.

Endpoint	Proph	nylaxis	Episodic	p-value	
Assessment	Incidence (%)	Relative Risk (95% Cl)	Incidence (%)	Relative Risk (95% Cl)	
MRI	2/27 (7%)	0.17 (0.04, 0.67)	13/29 (45%)	6.05 (1.50, 24.38)	0.002
Radiography	1/28 (4%)	0.19 (0.02, 1.55)	5/27 (19%)	5.19 (0.65, 41.54)	0.101
MRI or Radiography	2/30 (7%)	0.16 (0.04, 0.65)	13/31 (42%)	6.29 (1.55, 25.55)	0.002

Table 10 Subjects with Joint Damage (Subjects with Available Baseline and Endpoint Data)

Relative Risk is the risk of damage to one or more index joints on the given therapy as compared to the other therapy.

P-value is from the 2-sided Fisher Exact Test comparing the incidence of joint damage between treatment groups.

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16 HOW SUPPLIED/STORAGE AND HANDLING

How Supplied

Kogenate FS with vial adapter, with 15-micrometer filter and a prefilled diluent syringe, which together serve as an alternative needleless reconstitution system, is supplied in the following single use glass vial sizes. A prefilled diluent syringe containing Sterile Water for Injection, USP, a sterile vial adapter for reconstitution, and an administration set are also provided.

Kit NDC Number	Approximate FVIII Activity (IU)	Diluent (mL)
0026-3782-25	250	2.5
0026-3783-35	500	2.5
0026-3785-55	1000	2.5
0026-3786-65	2000	5.0
0026-3787-75	3000	5.0

Actual factor VIII activity in IU is stated on the label of each Kogenate FS Vial. Use the

actual potency as indicated on the vial label to calculate the dose.

Storage and Handling

The product vial and diluent syringe are not made with natural rubber latex.

Product as Packaged for Sale

- Store Kogenate FS at +2°C to +8°C (36°F to 46°F) for up to 30 months from the date of manufacture. Within this period, Kogenate FS may be stored for a period of up to 12 months at temperatures up to +25°C or 77°F.
- Record the starting date of room temperature storage on the unopened product carton. Once stored at room temperature, do not return the product to the refrigerator. The shelf-life then expires after storage at room temperature, or after the expiration date on the product vial, whichever is earlier.
- Do not use Kogenate FS after the expiration date indicated on the vial.
- Do not freeze.
- Protect from extreme exposure to light and store the lyophilized powder in the carton prior to use.

Product After Reconstitution

- After reconstitution, store the Kogenate FS solution at room temperature and administer within 3 hours.
- Use the administration set provided.

17 PATIENT COUNSELING INFORMATION

- Advise the patient to read the FDA-approved patient labeling (Patient Information and Instructions for Use).
- Advise patients to report any adverse reactions or problems following Kogenate FS administration to their physician or healthcare provider.
- Allergic-type hypersensitivity reactions have been reported with Kogenate FS. Warn
 patients of the early signs of hypersensitivity reactions [including hives (rash with
 itching), generalized urticaria, tightness of the chest, wheezing, hypotension] and
 anaphylaxis. Advise patients to discontinue use of the product if these symptoms
 occur and seek immediate emergency treatment with resuscitative measures such
 as the administration of epinephrine and oxygen.
- Inhibitor formation may occur at any time in the treatment of a patient with hemophilia A. Advise patients to contact their physician or treatment center for further treatment and/or assessment, if they experience a lack of clinical response to factor VIII replacement therapy, as this may be a manifestation of an inhibitor.
- Advise patients to consult with their healthcare provider prior to travel. While traveling advise patients to bring an adequate supply of Kogenate FS based on their current regimen of treatment.

FDA-Approved Patient Labeling

Patient Information

Kogenate FS (kō-jen-ate)

Antihemophilic Factor (Recombinant)

Formulated with Sucrose

This leaflet summarizes important information about Kogenate FS with vial adapter. Please read it carefully before using this medicine. This information does not take the place of talking with your healthcare provider, and it does not include all of the important information about Kogenate FS. If you have any questions after reading this, ask your healthcare provider.

Do not attempt to self-infuse unless you have been taught how by your healthcare provider or hemophilia center.

What is Kogenate FS?

Kogenate FS is a medicine used to replace clotting factor (factor VIII or antihemophilic factor) that is missing in people with hemophilia A (also called "classic" hemophilia). Hemophilia A is an inherited bleeding disorder that prevents blood from clotting normally.

Kogenate FS is used to treat and control bleeding in adults and children with hemophilia A. Your healthcare provider may give you Kogenate FS when you have surgery. Kogenate FS can reduce the number of bleeding episodes when used regularly (prophylaxis). Kogenate FS can reduce the risk of joint damage in children.

Kogenate FS is not used to treat von Willebrand Disease.

Who should not use Kogenate FS?

You should not use Kogenate FS if you

- are allergic to rodents (like mice and hamsters).
- are allergic to any ingredients in Kogenate FS.

What should I tell my healthcare provider before I use Kogenate FS?

Tell your healthcare provider about all of your medical conditions.

Tell your healthcare provider and pharmacist about all of the medicines you take, including all prescription and non-prescription medicines, such as over-the-counter medicines, supplements, or herbal remedies.

Tell your healthcare provider if you have been told you have heart disease or are at risk for heart disease.

Tell your healthcare provider if you have been told that you have inhibitors to factor VIII (because Kogenate FS may not work for you).

Tell your healthcare provider if you are pregnant or planning to become pregnant. It is not known if Kogenate FS may harm your unborn baby.

Tell your healthcare provider if you are breastfeeding. It is not known if Kogenate FS passes into the milk.

What are the possible side effects of Kogenate FS?

You could have an allergic reaction to Kogenate FS. Call your healthcare provider right away and stop treatment if you get

• rash or hives

- itching
- tightness of the chest or throat
- difficulty breathing
- light-headed, dizziness
- nausea
- decrease in blood pressure

Your body can also make antibodies, called "inhibitors," against Kogenate FS, which may stop Kogenate FS from working properly. Consult with your healthcare provider to make sure you are carefully monitored with blood tests for the development of inhibitors to factor VIII.

Other common side effects of Kogenate FS are

- Local injection site reactions (pain, swelling, irritation at infusion site)
- Infections from implanted injection device

Tell your healthcare provider about any side effect that bothers you or that does not go away.

Finding veins for injections may be difficult in young children. When frequent injections are required your child's healthcare provider may propose to have a device surgically placed under the skin to facilitate access to the bloodstream. These devices may result in infections.

These are not all the possible side effects with Kogenate FS. You can ask your healthcare provider for information that is written for healthcare professionals.

How do I store Kogenate FS?

Do not freeze Kogenate FS.

Store Kogenate FS at +2°C to +8°C (36°F to 46°F) for up to 30 months from the date of manufacture. Within this period, Kogenate FS may be stored for a period of up to 12 months at temperatures up to +25°C or 77°F.

Record the starting date of room temperature storage clearly on the unopened product carton. Once stored at room temperature, do not return the product to the refrigerator. The product then expires after storage at room temperature, or after the expiration date on the product vial, whichever is earlier. Store vials in their original carton and protect them from extreme exposure to light.

Reconstituted product (after mixing dry products with wet diluent) must be used within 3 hours and cannot be stored.

Throw away any unused Kogenate FS after the expiration date.

Do not use reconstituted Kogenate FS if it is not clear to slightly cloudy and colorless.

What else should I know about Kogenate FS and hemophilia A?

Medicines are sometimes prescribed for purposes other than those listed here. Do not use Kogenate FS for a condition for which it is not prescribed. Do not share Kogenate FS with other people, even if they have the same symptoms that you have.

This leaflet summarizes the most important information about Kogenate FS. If you would like more information, talk to your healthcare provider. You can ask your healthcare

provider or pharmacist for information about Kogenate FS that was written for healthcare professionals.

Instructions for use

How should I take Kogenate FS?

Do not attempt to self-infuse unless you have been taught how by your healthcare provider or hemophilia center.

See the step-by-step instructions for reconstituting Kogenate FS at the end of this leaflet and the specific infusion instruction leaflet provided.

You should always follow the specific instructions given by your healthcare provider. The steps listed below are general guidelines for using Kogenate FS. If you are unsure of the procedures, please call your healthcare provider before using.

Call your healthcare provider right away if bleeding is not controlled after using Kogenate FS.

Your healthcare provider will prescribe the dose that you should take.

Your healthcare provider may need to take blood tests from time to time.

Talk to your healthcare provider before traveling. You should plan to bring enough Kogenate FS for your treatment during this time.

Carefully handle Kogenate FS. Dispose of all materials, including any leftover reconstituted Kogenate FS product, in an appropriate container.

Reconstitution and use of Kogenate FS

Always work on a clean surface and wash your hands before performing the following procedure. Use only the components for reconstitution and administration that are provided with each package of Kogenate FS. If a package is opened or damaged, do not use this component. If these components cannot be used, please contact your healthcare provider. If you have any questions about Kogenate FS contact Bayer at 1-888-84-BAYER (1-888-842-2937).

1. Warm the unopened diluent and the concentrate to a temperature not to exceed 37°C or 99°F.

2. Remove protective cap from the vial (A). Aseptically cleanse the rubber stopper with alcohol, being careful not to handle the rubber stopper.



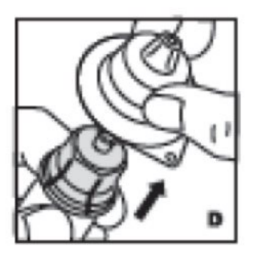
3. Place product vial on a firm, non-skid surface. Peel off the paper cover on the vial adapter plastic housing. **Do not remove the adapter from the plastic housing.** Holding the adapter housing, place over the product vial and firmly press down (B). The adapter will snap over the vial cap. **Do not remove the adapter housing at this step.**



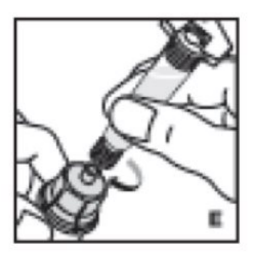
4. Holding the syringe by the barrel, snap the syringe cap off the tip (C). **Do not touch the syringe tip with your hand or any surface.** Set the syringe aside for further use.



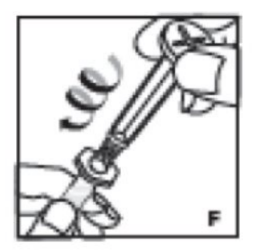
5. Now remove and discard the adapter housing (D).



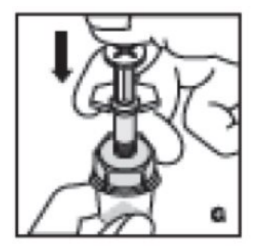
6. Attach the prefilled syringe to the threaded vial adapter by turning clockwise (E).



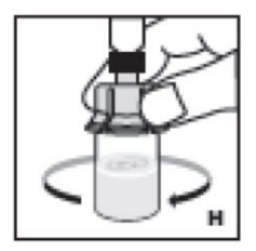
the plunger rod. Immediately attach the plunger rod by turning it firmly clockwise into the threaded syringe rubber stopper (F).



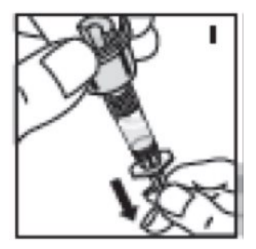
8. Inject the diluent by slowly pushing down on the plunger rod (G).



9. Swirl vial gently until all material is dissolved (H). **Do not shake vial.** Be sure that the powder is completely dissolved. **Do not use solutions containing visible particles or that are cloudy.**



10. Withdraw solution into the syringe by holding the vial on end above the vial adapter and syringe (I) then draw the plunger rod out slowly and smoothly. Ensure that the entire content of the vial is drawn into the syringe.



11. With the plunger rod in place, remove the syringe from the vial adapter (the latter should remain attached to the vial). Attach the syringe to the administration set provided and inject intravenously (J). NOTE: follow instructions for infusion set provided.



12. If the same patient is to receive more than one bottle, reconstitute each bottle with the diluent syringe provided then combine solutions in a larger syringe (not provided) and administer as usual.

Rate of administration

The entire dose of Kogenate FS can usually be infused within 1 to 15 minutes. However, your healthcare provider will determine the rate of administration that is best for you.

Resources at Bayer available to the patient:

For Adverse Reaction Reporting contact:

Bayer Medical Communications 1-888-84-BAYER (1-888-842-2937)

Contact Bayer to receive more product information:

Kogenate FS Customer Service 1-888-606-3780

Bayer Reimbursement HELPline 1-800-288-8374

For more information, visit www.kogenatefs.com

Bayer HealthCare LLC Whippany, NJ 07981 USA

U.S. License No. 8 (License Holder: Bayer Corporation)

http://www.kogenatefs.com/

The following are representative examples of Kogenate FS labeling. See the "How Supplied" section for a complete listing of all components.

Kogenate FS with Vial Adapter Carton 250 IU

NDC 0026-3782-25

Kogenate FS

Antihemophilic Factor (Recombinant)

Formulated with Sucrose

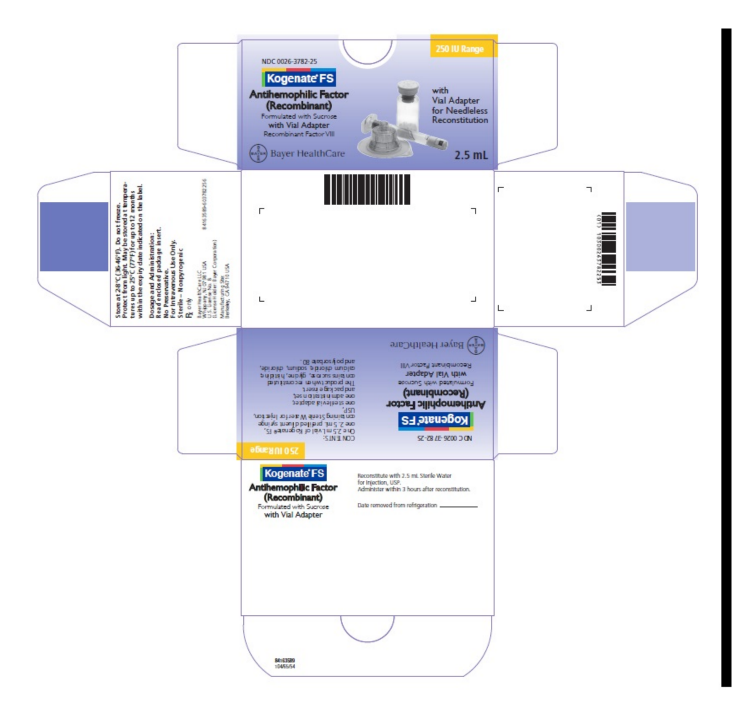
with Vial Adapter

Recombinant Factor VIII

With Vial Adapter for Needleless Reconstitution

2.5 mL

Bayer HealthCare



Kogenate FS with Vial Adapter Carton 500 IU

NDC 0026-3783-35

Kogenate FS

Antihemophilic Factor

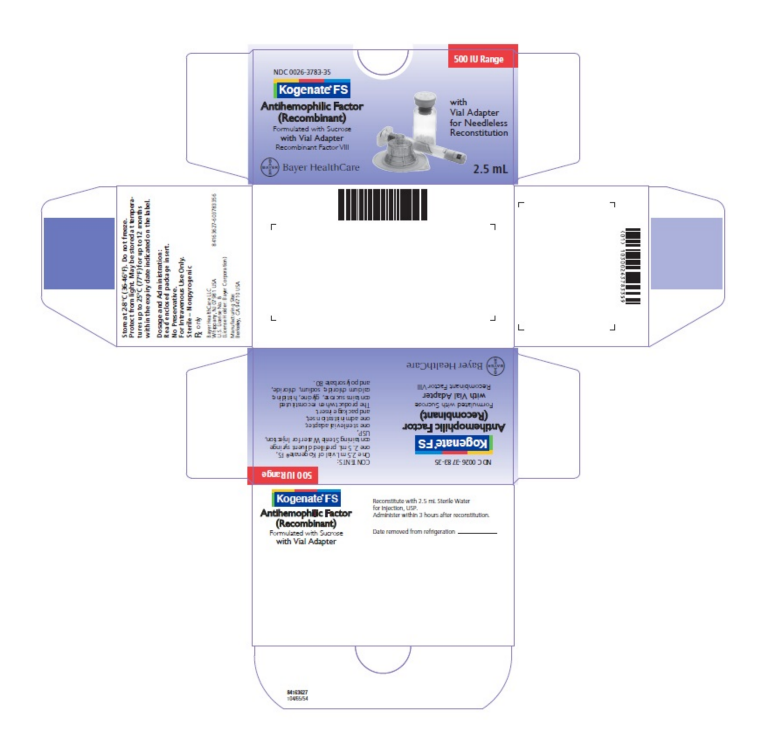
(Recombinant)

Formulated with Sucrose with Vial Adapter Recombinant Factor VIII

With Vial Adapter for Needleless Reconstitution

2.5 mL

Bayer HealthCare



Kogenate FS with Vial Adapter Carton 1000 IU

NDC 0026-3785-55

Kogenate FS

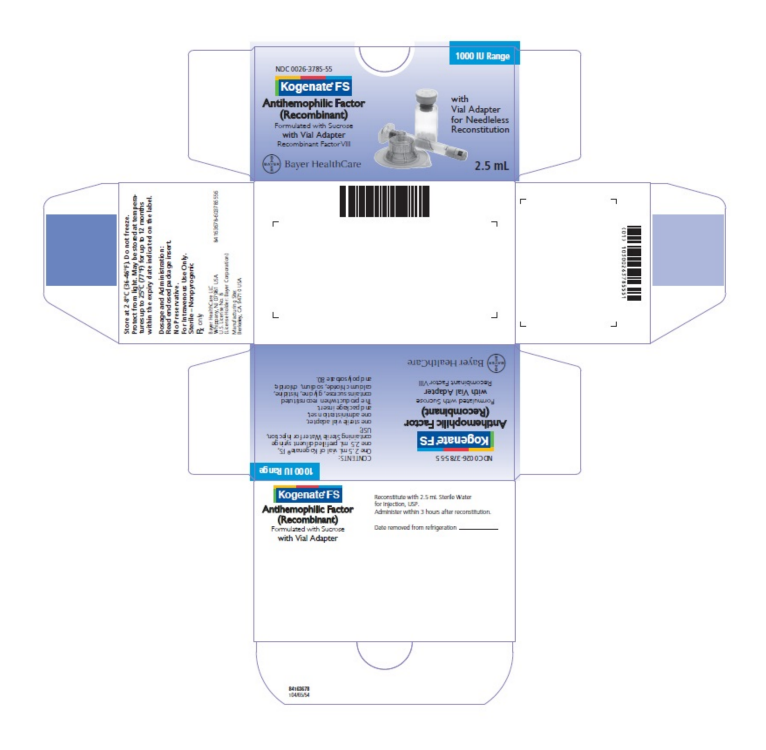
Antihemophilic Factor (Recombinant)

Formulated with Sucrose with Vial Adapter Recombinant Factor VIII

With Vial Adapter for Needleless Reconstitution

2.5 mL

Bayer HealthCare



Kogenate FS with Vial Adapter Carton 2000 IU

NDC 0026-3786-65

Kogenate FS

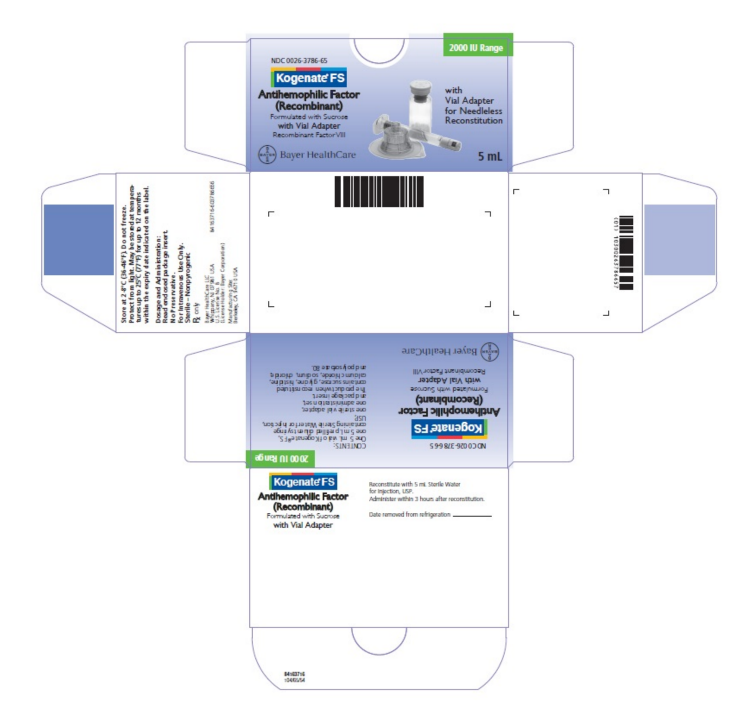
Antihemophilic Factor (Recombinant)

Formulated with Sucrose with Vial Adapter Recombinant Factor VIII

With Vial Adapter for Needleless Reconstitution

5 mL

Bayer HealthCare



Kogenate FS with Vial Adapter Carton 3000 IU

NDC 0026-3787-75

Kogenate FS

Antihemophilic Factor (Recombinant)

Formulated with Sucrose with Vial Adapter Recombinant Factor VIII

With Vial Adapter for Needleless Reconstitution

5 mL

Bayer HealthCare



KOGENATE FS antihemophilic factor	5 or (recombinant) kit		
Product Inform	ation		
Product Type	PLAS MA DERIVATIVE	Item Code (Source)	NDC:0026-3785
Packaging			

	ltem Code		Package Descrip	tion	Marketing Start Date	-
	DC:0026-		Other Type of Part 3	Combination Product		
3	785-55	(e.g., Drug/Device/	Biological Product)			
Qu	antity o	f Parts				
Par	t #	Package C	Juantity	Tota	l Product Quar	ntity
Part	: 1 VIAL	, SINGLE-USE		2.5 mL		
Part	2 1 SYR	INGE		2.5 mL		
Pa	rt 1 of	⁻ 2				
КС	JGENA	TE FS				
anti	ihemoph	lic factor (recom	binant) injection, p	owder, lyophilized,	for solution	
Pro	oduct In	formation				
lter	n Code (S	Source)	NDC:0026-4785			
Rou	te of Ad	ministration	INTRAVENOUS			
Δct	ive Ina	edient/Active	Moiety			
	ive mg		dient Name		Basis of Stre	ength Strengt
ANT	ІНЕМОРН	•		JNII: P89DR4NY54)	ANTIHEMOPHILIC F	-
			RECOMBINANT - UNII:P		HUMAN RECOMBIN	
Ina	ctive In	gredients				
			Ingredient Nam	ne		Strength
CAL	CIUM (UNI	: SY7Q814VUP)	_			
CHL	ORIDE IOI	I (UNII: Q32ZN4869	8)			
СОР	PER (UNII:	789U1901C5)				
GLY	CINE (UNII	TE7660XO1C)				
HIST	TIDINE (UN	II: 4QD397987E)				
POL	YSORBAT	E 80 (UNII: 60ZP39)	ZG8H)			
		METAPHOSPHATE	(UNII: P1BM4ZH95L)			
SOD	ROSE (UN	I. CISIH6M354)				
SOD SUC		PHOSPHATE (UNII:	95UAS8YAF5)			
SOD SUC			95UAS 8YAF5)			
SOD SUC TRI-			95UAS8YAF5)			
SOD SUC TRI- Pac	N-BUTYL C kaging Item			ption	Marketi	
SOD SUC TRI-	N-BUTYL Ckaging Item Code	PHOSPHATE (UNII:	Package Descri		Marketi Start Da	
SOD SUC TRI- Pac #	N-BUTYL ckaging Item Code DC:0026-	PHOSPHATE (UNII: 2.5 mL in 1 VIAL, SI		er Type of Part 3		

Marketin		tion			
Category		tion Number or Monograph Citation	Marketir Da		Marketing En Date
BLA	BLA103332		06/26/2013		
Part 2 of	2				
DILUENT water solutio					
Product In	formation				
ltem Code (S	ource)	NDC:0026-0426			
Route of Adr	ninistration	INTRAVENOUS			
Inactive In	-				
	-	redient Name		9	Strength
WATER (UNII: 0	35010100010				
ltom		Package Description		Marketi Start Da	
# Item Code		Package Description E; Type 9: Other Type of Part 3 Con /Device/Biological Product)	nbination		-
# Item Code		GE; Type 9: Other Type of Part 3 Con	nbination		-
Item Code 1 NDC:0026- 0426-02		GE; Type 9: Other Type of Part 3 Con /Device/Biological Product)	nbination		-
Item Code 1 NDC:0026- 0426-02	Product (e.g., Drug g Informat g Applica	GE; Type 9: Other Type of Part 3 Con /Device/Biological Product)	nbination Marketir Da	Start Da	-
 Code NDC:0026- 0426-02 Marketin Marketin 	Product (e.g., Drug g Informat g Applica	SE; Type 9: Other Type of Part 3 Con /Device/Biological Product)	Marketir	Start Da	ate End Dat Marketing En
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 # Item Code NDC:0026- 0426-02 Marketin Category BLA 	g Informat g Applica	E; Type 9: Other Type of Part 3 Con /Device/Biological Product)	Marketir Da	Start Da	ate End Dat Marketing En
 # Item Code NDC:0026- 0426-02 Marketin Category BLA 	Product (e.g., Drug g Informat g Applica BLA103332 g Informat g Applica	E; Type 9: Other Type of Part 3 Con /Device/Biological Product)	Marketir Da	Start Da ng Start te	ate End Dat Marketing En

KOGENATE FS

antihemophilic factor (recombinant) kit

Product Information

_			-					
Pro	oduct Type		PLASMA	DERIVATIVE	Item Code (Sour	ce)	NDC:0026-3	3783
Pa	ckaging							
# I	tem Code		Р	ackage Descrip	tion	Marketing Start Date		eting End Date
	NDC:0026- 3783-35			3: Prefilled Biologic I ringe, patch, etc.)	Delivery			
			<u> </u>	3 -, pr - , ,				
Ou	antity of	Parts						
-	rt #		kage O	uantity	Tota	al Product Qu	antity	
Par	t 1 1 VIAL, 1	SINGLE-U	-		2.5 mL		- -	
Par	t2 1 SYRIN	GE			2.5 mL			
Pa	art 1 of	2						
	DGENAT		/ 1					
ant	ihemophilic	factor	(recomb	pinant) injection,	powder, lyophilized	, for solution		
Pr	oduct Inf	ormati	on					
lte	m Code (So	ource)		NDC:0026-4783				
Ro	ute of Adm	inistrati	ion	INTRAVENOUS				
Act	tive Ingre	dient//	Active	Moiety				
			Ingre	dient Name		Basis of St	rength	Strengt
					(UNII: P89DR4NY54)	ANTIHEMOPHILIC		500 [iU]
(AN⊺	TIHEMOPHILIC	FACTOR,	HUMAN F	Recombinant - Unii:	P89DR4NY54)	HUMAN RECOME	SINANT	in 2.5 mL
Ina	active Ing	redien	ts					
				Ingredient Na	me		Sti	rength
CAL	CIUM (UNII: :	SY7Q814	VUP)					
СНГ	ORIDE ION	(UNII: Q32	2ZN48698	3)				
со	PPER (UNII: 7	89U19010	C5)					
GLY	CINE (UNII: T	E7660XO	1C)					
	TIDINE (UNII:							
	YSORBATE							
				(UNII: P1BM4ZH95L)				
	CROSE (UNII:							
TRI	-N-BUTYL PH	IOSPHAT	re (Unii: 9	95UAS8YAF5)				
Da	ckaging							

# Item			Package Description		Market Start Da		Marketing End Date
1 NDC:002 4783-01			IGLE-USE; Type 9: Other Type of Pa t (e.g., Drug/Device/Biological Produ				
Marke	ting In	format	ion				
Mark		Applicat	tion Number or Monograph Citation	Marketin		Mar	keting End
Cate BLA	gory	BLA103332	Citation	Dat 06/26/2013	e		Date
Part 2	of 2						
DILUE	NT						
water sol							
Product	: Inform	ation					
	e (Source		NDC:0026-0426				
	Administ		INTRAVENOUS				
Inactive	Ingredi	ionts					
mactive	mgreu						
		Ingi	redient Name		5	Streng	th
WATER (U	NII: 059QF0	-	redient Name		3	Streng	lth
WATER (U	NII: 059QF0	-	redient Name		5	streng	ith
		-	redient Name		5	Streng	lth
Packagi " Item	ng	-	Package Description		Marketii	ng	Marketing
Packagi # Item Code	ng 2 26- 2.5 mL	KOOR) . in 1 SYRINGI	Package Description E; Type 9: Other Type of Part 3 Con	nbination		ng	
Packagi # Item Code	ng 2 26- 2.5 mL	KOOR) . in 1 SYRINGI	Package Description	nbination	Marketii	ng	Marketing
Packagi # Item Code	ng 2 26- 2.5 mL	KOOR) . in 1 SYRINGI	Package Description E; Type 9: Other Type of Part 3 Con	nbination	Marketii	ng	Marketing
Packagi # Item Code 1 NDC:002 0426-02	26- Produc	KOOR) . in 1 SYRINGI	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product)	nbination	Marketii	ng	Marketing
Packagi # Item Code 1 NDC:002 0426-02 Marke	ing 26- 2.5 mL Produc ting In eting	KOOR) . in 1 SYRINGI t (e.g., Drug/	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product)	nbination Marketing Dat	Marketin Start Da	ng ite	Marketing
Packagi # Item Code 1 NDC:002 0426-02 Marke Cate	ing 26- 2.5 mL Produc ting In eting	KOOR) . in 1 SYRINGI t (e.g., Drug/	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product) ion	Marketing	Marketin Start Da	ng ite	Marketing End Date keting End
Packagi # Item Code 1 NDC:002 0426-02 Marke Cate	ing 26- 2.5 mL Produc ting In eting	KOOR) in 1 SYRINGI it (e.g., Drug/ format Applicat	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product) ion	Marketin Dat	Marketin Start Da	ng ite	Marketing End Date keting End
Packagi # Item Code 1 NDC:002 0426-02 Marke Mark Cate	ing 26- 2.5 mL Produc ting In eting gory	koor) . in 1 SYRINGI t (e.g., Drug/ format Applicat BLA103332	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product) ion tion Number or Monograph Citation	Marketin Dat	Marketin Start Da	ng ite	Marketing End Date keting End
Packagi # Item Code 1 NDC:002 0426-02 Marke BLA Marke Marke	ing 26- 2.5 mL Produc ting In eting gory	koor) in 1 SYRINGF t (e.g., Drug/ format BLA103332	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product) ion tion Number or Monograph Citation	Marketin Dat	Marketin Start Da	ng ite Marl	Marketing End Date keting End
 # Item Code NDC:002 NDC:002	ing 26- 2.5 mL Produc ting In eting gory	koor) in 1 SYRINGF t (e.g., Drug/ format BLA103332	Package Description E; Type 9: Other Type of Part 3 Con Device/Biological Product) ion tion Number or Monograph Citation	Marketing Dat 06/26/2013 Marketing	Marketin Start Da	ng ite Marl	Marketir End Dat keting En Date

KOGENATE FS

		ic factor (I	recomb	inant) kit				
ununer	noprii		ccomb					
Produ	uct In	formati	on					
Produ	ct Typ	e	PLAS MA	DERIVATIVE	Item Code (Source	e) NC	DC:0026-3	3782
Packa	aging							
I	em ode			Package Descrip	cription Marketing Market Start Date End D			
1 NDC: 3782	0026- -25			Other Type of Part 3 Biological Product)	Combination Product			
Quan	tity o	f Parts						
Part #	÷	Pac	kage Q	uantity	Tota	l Product Qua	ntity	
Part 1		, SINGLE-US	SE		2.5 mL			
Part 2	1 SYR	INGE			2.5 mL			
Part	1 of	2						
Produ	uct In	formati	on					
	-	Source)	011	NDC:0026-4782				
		ministrati	ion	INTRAVENOUS				
Noute	UT AU	ininistiati	UT	INTRAVENCOS				
Active	e Ingr	edient/A	Active	Moiety				
			Ingre	dient Name		Basis of Str	ength	Strengt
				AN RECOMBINANT (RECOMBINANT - UNII:P		ANTIHEMOPHILIC HUMAN RECOMBII		250 [iU] in 2.5 mL
Inacti	ive In	gredient	ts					
				Ingredient Nan	ne		St	rength
CALCIU	M (UNI	: SY7Q814\	/UP)					
CHLOR	IDE IO	I (UNII: Q32	ZN48698	3)				
COPPE	R (UNII:	789U19010	25)					
GLYCIN	ie (Unii:	TE7660XO	1C)					
HISTID	INE (UN	II: 4QD3979	987E)					
POLYS	ORBAT	e 80 (UNII:	60ZP392	ZG8H)				
SODIU	M POLY	МЕТАРНО	SPHATE	(UNII: P1BM4ZH95L)				

SUCROSE (UNII: C151H8M554)

	DUOCD						
TRI-N-BUTYI	L PHUSP	HATE (UNII:	950A58TAF5)				
Packagin	a						
ltom	9				Marke	tina	Marketing
# Code			Package Description		Start D		End Date
1 NDC:0026-4782-01			NGLE-USE; Type 9: Other Type of Pa t (e.g., Drug/Device/Biological Produ				
4702-01	Combin						
Marketi	ina In	format	ion				
Market	-		tion Number or Monograph	Marketin	a Start	Ма	rketing End
Catego			Citation	Dat			Date
BLA		BLA103332		06/26/2013			
D							
Part 2 c	of 2						
DILUEN	Т						
water solut	ion						
Product I	nform	ation					
ltem Code	(Source	<u>a)</u>	NDC:0026-0426				
Route of A			INTRAVENOUS				
Noute of A							
Inactive I	ngredi	ients					
		Ing	redient Name			Stren	gth
WATER (UNII	: 059QF0	KO0R)					
Packagin	g						
# Item Code			Package Description		Market		Marketing End Date
1 NDC:0026-			E; Type 9: Other Type of Part 3 Com	bination			
• 0426-02	Produc	t (e.g., Drug/	Device/Biological Product)				
		£					
Marketi	-						
Market Catego		Applica	tion Number or Monograph Citation	Marketin Dat		Ма	rketing End Date
BLA		BLA103332		06/26/2013			
Marketi	ing In	format	ion				

Marketing Category Application Number or Monograph Citation Marketing Start Marketing End Date BLA BLA103332 06/26/2013 06/26/2013 KOGENATE FS antihemophilic factor (recombinant) kit 06/26/2013 NDC:0026-3786 Product Information Product Type PLASMA DERIVATIVE Item Code (Source) NDC:0026-3786 Packaging # Item Code (Source) NDC:0026-3786 End Date Packaging # Item Code (Source) NDC:0026-3786 1 NDC:0026-1786 End Date End Date 2 NDC:0026-1786 End Date End Date 3786-026 Itele and Date Start Date End Date 2 NDC:0026-1786 End Date End Date 3786-027 Itele and Date Start Date End Date 2 1 NDC:0026-1786 End Date End Date 2 Itele and Date SmL SmL End Date 2 1 SYRINGE S mL SmL End Date Part 1 of 2 KOGENATE FS NDC:0026-4786 </th <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th> <th></th>										
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antihemophilic factor (recombinant) kit Product Information Product Type PLASMA DERIVATIVE Item Code (Source) MDC:0026-3786 Package Package Description Marketing Start Date Part 2 NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 I in 1 BOX; Type 9: Other Type of Part 3 Combination Product I NDC:0026 Part # Package Quantity Fart 1 I VAL, SINGLE-USE S mL Part 1 I VAL, SINGLE-USE S mL Part 1 I S YRINGE Part 1 Of 2 KOGENATE FS antihemophilic factor (recombinant) injection, powder, lyophilized, for solution Product Information Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Molety Ingredient Name AntiHemophilic FACTOR, HUMAN RECOMBINANT - UNIE PS9DRANYS4) ANTIHEMOPHILC FACTOR, HUMAN RECOMBINANT - UNIE PS9DRANYS4) Inactive IngredientS Ingredient Name Ingredient Name CALCIUM (UNIE SYT0814VUP) CHLORIDE ION (UNIE 032ZIM8698)	BLA		BLA103332	2		06/26/201	3			
antihemophilic factor (recombinant) kit Product Information Product Type PLASMA DERIVATIVE Item Code (Source) MDC:0026-3786 Packaging										
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Product Type PLASMA DERIVATIVE Item Code (Source) NDC:0026-3786 Package Package Description Marketing Start Date Marketing End Date # Rem Code Package Description Marketing End Date Marketing End Date 1 NDC:0026-3786 Lin 1 BOX; Type 9: Other Type of Part 3 Combination Product Marketing End Date Marketing End Date Quantity of Parts Package Quantity Total Product Quantity Value Smc Part 1 1 VAL, SINOLE-USE 5 mL Smc Smc Part 2 1 SYRINGE 5 mL Smc Smc Part 1 1 VAL, SINOLE-USE 5 mL Smc Smc Part 2 1 of 2 KOGENATE FS Smc Smc Smc Product Information INTRAVENOUS INTRAVENOUS Smc Smc Smc Active Ingredient/Active Molecty Ingredient Name Basis of Strength Smc Smc Inactive Ingredients Ingredient Name Strength Codo [U] Smc Smc Instrument Ingredient Name Basis of Strength Strength Catcum (UMII: SY70	antihen	nophilic fa	ctor (recom	binant) kit						
Product Type PLASMA DERIVATIVE Item Code (Source) NDC:0026-3786 Packaging Marketing Start Date Marketing End Date # Rem Code (Source) Marketing End Date Marketing End Date 1 NDC:0026-3786 Lin 1 BOX; Type 9: Other Type of Part 3 Combination Product Marketing End Date Quantity of Parts Package Quantity Total Product Quantity Marketing End Date Part 1 1 VAL, SINOLE-USE 5 mL SmL SmP Part 1 1 VAL, SINOLE-USE 5 mL SmL SmP Part 1 1 SYRINGE 5 mL SmL SmP Part 1 1 SYRINGE SmC:0026-4786 Smoute of Administration INTRAVENOUS Route of Administration INTRAVENOUS ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UMI: P89DRAMY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UMI: P89DRAMY54) ANTIHEMOPHILIC FACTOR, 2000 [U] Inactive IngredientS Ingredient Name Basis of Strength Strength CALCUM (UMI: SY70814YUP Ingredient Name Strength CALCUM (UMI: SY20814YUP Ingredient Name Strength CALCUM (UMI: SY20814YUP Ingredient Name Strength <td>Duadu</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>	Duadu									
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Item Code Package Description Marketing Start Date Marketing End Date INDC:0026 I in 1 80X; Type 9; Other Type of Part 3 Combination Product Image: Combination Product Produc	Produc	t Type	PLAS MA	A DERIVATIVE	ltem Code	(Source)		NDC:0026-3	3786	
Item Code Package Description Marketing Start Date Marketing End Date INDC:0026-1 in 1 80X; Type 9; Other Type of Part 3 Combination Product Imarketing Start Date Marketing End Date Quantity of Parts Package Quantity Total Product Quantity Part 4 Package Quantity S mL Part 1 I VAL, SINGLE-USE S mL Part 2 I SYRINGE S mL Part 1 Of 2 KOGENATE FS antihemophilic factor (recombinant) injection, powder, lyophilized, for solution NDC:0026-4786 Route of Administration INTRAVENOUS Strength Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII: P89DRANY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII: P89DRANY54) ANTIHEMOPHILIC FACTOR, STRENGTH Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SYZ0814/UP) Ingredient Name Strength										
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A 3786-65 (e.g., Drug/Device/Biological Product) Quantity of Parts Total Product Quantity Part # Package Quantity S mL Part 1 1 VIAL, SINGLE-USE S mL Part 2 1 SYRINGE S mL Part 3 1 SYRINGE S mL Part 1 1 VIAL, SINGLE-USE S mL Part 2 1 SYRINGE S mL Part 1 of 2 KOGENATE FS Solution Route of Administration Ingredient, Active Roues Solution Product Information INTRAVENOUS AntiHemophilic Factor, HUMAN RECOMBINANT (UNII: P89DRANY54) Basis of Strength Strength Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength Inactive Ingredients Ingredient Name Strength Strength Inactive Ingredients Ingredient Name Strength Inactive Ingredients Ingredient Name Strength Calcum (UNII: SY70814VUP) Ingredient Name Strength Calcum (UNII: SY70814VUP) Chloribe Ion (UNII: Gazz N48698) Strength	Co									
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Part # Package Quantity Total Product Quantity Part 1 1 VIAL, SINGLE-USE 5 mL Part 2 1 SYRINGE 5 mL Part 2 1 SYRINGE 5 mL Part 1 of 2 KOGENATE FS antihemophilic factor (recombinant) injection, powder, lyophilized, for solution Product Information Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS VERNOUS Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) Ingredient Name Ingredient Name Ingredient Name Ingredient Name Ingredient Name Strength Ingredient Name Ingredient Name Ingredient Name Ingredient Name Ingredient Name Ingredient Name Ingredient Name <td></td> <td></td> <td>J</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>			J							
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KOGENATE FS antihemophilic factor (recombinant) injection, powder, lyophilized, for solution Product Information Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Ingredient Name Basis of Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) Inactive Ingredients Ingredient Name CALCIUM (UNII: SY70814VUP) Ingredient Name CALCIUM (UNII: SY70814VUP) Ingredient Name	rait 2	I SHANGE	•		JIIL					
KOGENATE FS antihemophilic factor (recombinant) injection, powder, lyophilized, for solution Product Information Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Ingredient Name Basis of Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) Inactive Ingredients Ingredient Name CALCIUM (UNII: SY70814VUP) Ingredient Name CALCIUM (UNII: SY70814VUP) Ingredient Name										
KOGENATE FS antihemophilic factor (recombinant) injection, powder, lyophilized, for solution Product Information Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Ingredient Name Basis of Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) Inactive Ingredients Ingredient Name CALCIUM (UNII: SY70814VUP) Ingredient Name CALCIUM (UNII: SY70814VUP) Ingredient Name	Dart	1 of 2								
antihemophilic factor (recombinant) injection, powder, lyophilized, for solution Product Information NDC:0026-4786 Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Strength AntiHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) Inactive Ingredients Ingredient Name CALCIUM (UNII: SY7Q814VUP) Ingredient Name CALCIUM (UNII: SY7Q814VUP) Ingredient Name CALCIUM (UNII: SY7Q814VUP) CALCIUM (UNII: SY7Q814VUP)										
Product Information NDC:0026-4786 Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 2000 [iU] 2000 [iU] Inactive Ingredients Ingredient Name Strength Inactive Ingredients Strength CALCIUM (UNII: SY7Q814YUP) Ingredient Name Strength CALCIUM (UNII: SY3Q814YUP) CHLORIDE ION (UNII: Q32ZN48698) Image Strength			-							
Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Strength Active Ingredient/Active Moiety Basis of Strength AntiHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 2000 [iU] In active Ingredients Strength Inactive Ingredients Strength CALCIUM (UNII: SY7Q814VUP) Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CALCIUM (UNII: SY7Q814VUP) Strength	antiher	nophilic f	actor (recon	nbinant) injection, p	owder, lyop	hilized, fo	rsolution			
Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Strength Active Ingredient/Active Moiety Basis of Strength Strength AntiHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 1000 [iU] 2000 [iU] Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CALCIUM (UNII: SY7Q814VUP) Ingredient Name										
Item Code (Source) NDC:0026-4786 Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Basis of Strength Active Ingredient/Active Moiety Basis of Strength AntiHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 2000 [iU] Inactive Ingredients Ingredient Name Strength Strength CALCIUM (UNII: SY7Q814VUP) Ingredient Name CALCIUM (UNII: SY7Q814VUP) Strength CHLORIDE ION (UNII: Q32ZN48698) Strength										
Route of Administration INTRAVENOUS Active Ingredient/Active Moiety Basis of Strength Ingredient Name Basis of Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) Ingredients 2000 [iU] Instrine Ingredients 5 mL Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) Ingredient Name CHLORIDE ION (UNII: Q32Z N48698) Strength	Produ	ict Infor	mation							
Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT 2000 [iU] (ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT 2000 [iU] Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CALCIUM (UNII: Q32ZN48698) Imactive Ingredients	ltem C	ode (Sou	rce)	NDC:0026-4786						
Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 1000 [i0] 2000 [i0] (ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) ANTIHEMOPHILIC FACTOR, 100 [i0] in 5 mL Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) Strength Strength CHLORIDE ION (UNII: Q32ZN48698) Ingredient Name Ingredient Name	Route	of Admini	stration	INTRAVENOUS						
Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 1000 [i0] 2000 [i0] (ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) ANTIHEMOPHILIC FACTOR, 100 [i0] in 5 mL Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) Strength Strength CHLORIDE ION (UNII: Q32ZN48698) Ingredient Name Ingredient Name										
Ingredient Name Basis of Strength Strength ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, 1000 [i0] 2000 [i0] (ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) ANTIHEMOPHILIC FACTOR, 100 [i0] in 5 mL Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) Strength Strength CHLORIDE ION (UNII: Q32ZN48698) Ingredient Name Ingredient Name	Active	Ingredi	ent/Active	Mojety						
ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT (UNII: P89DR4NY54) ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT 2000 [iU] 2000 [iU] Inactive Ingredients Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CHLORIDE ION (UNII: Q32ZN48698) I		ingredi					Basis of St	trenath	Strength	
(ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT - UNII:P89DR4NY54) HUMAN RECOMBINANT - In 5 mL Inactive Ingredients Ingredient Name CALCIUM (UNII: SY7Q814VUP) Strength CHLORIDE ION (UNII: Q32ZN48698) I	ANTIHE	MOPHILIC	-		UNII: P89DR4N			-	-	
Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CHLORIDE ION (UNII: Q32ZN48698)							JMAN RECOMI	BINANT	in 5 mL	
Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CHLORIDE ION (UNII: Q32ZN48698)										
Ingredient Name Strength CALCIUM (UNII: SY7Q814VUP) CHLORIDE ION (UNII: Q32ZN48698)	Inacti	ve Inare	dients							
CALCIUM (UNII: SY7Q814VUP) CHLORIDE ION (UNII: Q32ZN48698)	mact	ie ingre		Ingredient Nan	ne			Stu	renath	
CHLORIDE ION (UNII: Q32ZN48698)	CALCIU	M (UNII: SY	7Q814VUP)	ingreatent nam				50	Singth	
COPPER (UNII: 789U1901C5)				98)						
	COPPER	R (UNII: 789	U1901C5)							

GLYCINE (UNII: TE766	0X01C)				
HISTIDINE (UNII: 4QD3	•				
POLYSORBATE 80 (U		(G8H)			
SODIUM POLYMETAR	PHOSPHATE	(UNII: P1BM4ZH95L)			
SUCROSE (UNII: C151	H8M554)				
TRI-N-BUTYL PHOSP	HATE (UNII: 9	95UAS 8YAF5)			
Packaging					
# Item Code		Package Description		Marketi Start Da	
1 NDC:0026- 5 mL in		LE-USE; Type 9: Other Type of Part			
4786-01 Combin	ation Product	: (e.g., Drug/Device/Biological Produ	ct)		
Marketing In	formati	ion			
Marketing Category	Applicat	ion Number or Monograph Citation	Marketin Dat		Marketing End Date
BLA	BLA103332		06/26/2013		
Part 2 of 2					
DILUENT					
water solution					
Product Inform					
Item Code (Source	e)	NDC:0026-0426			
Route of Administ	ration	INTRAVENOUS			
Inactive Ingredi	ents				
3		edient Name		S	trength
WATER (UNII: 059QF0	-				-
Dealessie					
Packaging				Marilant	Manler Hurr
# Item Code		Package Description		Marketin Start Dat	
		Type 9: Other Type of Part 3 Comb Device/Biological Product)	ination		
	-				
Marketing In	formati	ion			
Marketing		ion Number or Monograph	Marketin		Marketing End
Category		Citation	Dat	e	Date

BLA			BL	A103332			06/26/	2013		
Ma	arke	ting	l Info	rmation						
	Marke Cate			Application I	Number or Mo Citation	onograph	Ман	keting Start Date	ſ	Marketing End Date
BLA			BL	A103332			06/26/	2013		
KO	GEN		EFS							
-	-		_	recombinant	:) kit					
Pro	oduct	Info	ormati	ion						
Pro	duct 1	Гуре		PLAS MA DERIV	ATIVE	ltem Code	(Sour	ce)	NDC	0026-3787
Pac	ckagi	ng								
# It	tem C	ode		Packa	age Descript	ion		Marketin Start Dat	g e	Marketing End Date
	DC:002 787-75			DX; Type 3: Pre ystem (syringe	filled Biologic De , patch, etc.)	elivery				
		y of I	Parts							
Par				kage Quant	tity		Tot	al Product Q	uan	tity
Part		VIAL, S SYRINO	SINGLE-U	SE		5 mL 5 mL				
	1					5 ME				
Pa	rt 1	of 2	2							
			E FS factor	(recombinan	t) injection, p	owder. Ivon	hilized	l. for solution		
anu										

Product Information					
Item Code (Source)	NDC:0026-4787				
Route of Administration	ute of Administration INTRAVENOUS				
Active Ingredient/Active Moiety					
Ingre	Basis of Strength	Strength			
ANTIHEMOPHILIC FACTOR, HUM (ANTIHEMOPHILIC FACTOR, HUMAN	ANTIHEMOPHILIC FACTOR, HUMAN RECOMBINANT	3000 [iU] in 5 mL			

	active In	gredients	5						
				Ingredient Name				Strength	
CAI	LCIUM (UNI	I: SY7Q814VI	JP)					j	
		N (UNII: Q32Z)					
		789U1901C							
		: TE7660XO1							
HISTIDINE (UNII: 4QD397987E)									
POLYSORBATE 80 (UNII: 60ZP39ZG8H)									
SODIUM POLYMETAPHOSPHATE (UNII: P1BM4ZH95L)									
		II: C151H8M5							
		PHOSPHATE		5UAS 8YAF5)					
Pa	ckaging								
#	ltem Code			Package Description		Market Start D		Marketing End Date	
	NDC:0026- 4787-01	5 mL in 1 VIA Combination	L, SINGI Product	E-USE; Type 9: Other Type of Part (e.g., Drug/Device/Biological Produ	3 uct)				
M	arketir	ng Infor	mati	on					
			pplicat	ion Number or Monograph	Marketing Start Date		Ma	Marketing End Date	
	catege.	У		Citation	Dat	te		Date	
BLA	-	-	103332	Citation	Da1 06/26/2013	te		Date	
Pa	art 2 of	BLA1	103332	Citation		te		Date	
Pa DI	art 2 of	вцај f 2	103332	Citation				Date	
Pa DI wa	art 2 of LUENT ter solutio	BLAI							
Pa DI wa	art 2 of LUENT ter solutio	вцај f 2							
Pa DI wa Pr	art 2 of LUENT ter solutio	BLAI	n	Citation					
Pa DI wa Pr	art 2 of LUENT ter solutio oduct Ir m Code (2	BLAI	'n						
Pa DI wa Pr Ite	art 2 of LUENT ter solutio oduct Ir m Code (2	BLAI	'n	NDC:0026-0426					
Pa Di wa Pr Ite Ro	art 2 of LUENT ter solutio oduct Ir m Code (i ute of Ad	BLAI	on S	NDC:0026-0426 INTRAVENOUS					
Pa Di wa Pr Ite Ro	art 2 of LUENT ter solutio oduct Ir m Code (ute of Ad active In	BLAI	on S Ingr	NDC:0026-0426			Strer		
Pa Di wa Pr Ite Ro	art 2 of LUENT ter solutio oduct Ir m Code (ute of Ad active In	BLAI	on S Ingr	NDC:0026-0426 INTRAVENOUS			Strer		
DI wa Pr Ite Ro	art 2 of LUENT ter solutio oduct Ir m Code (ute of Ad active In	BLAI	on S Ingr	NDC:0026-0426 INTRAVENOUS			Strer		
Pa DI wa Pr Ite Ro	art 2 of LUENT ter solutio oduct Ir m Code () ute of Ad active Ir	BLAI	on S Ingr	NDC:0026-0426 INTRAVENOUS			ing		

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
BLA	BLA103332	06/26/2013				
Marketing	nformation					
Marketing Marketing Category	nformation Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			

Labeler - Bayer HealthCare LLC (127769128)

Establishment

Name	Address	ID/FEI	Business Operations
Bayer HealthCare LLC		127769128	MANUFACTURE(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426), API MANUFACTURE(0026-3785, 0026-4785, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-0426, 0026-3785, 0026-0426, 0026-3785, 0026-0426, 0026-3785, 0026-0426, 0026-3785, 0026-0426, 0026-3785, 0026-0426, 0026-3783, 0026-0426, 0026-3782, 0026-0426, 0026-3782, 0026-0426, 0026-3785, 0026-0426, 0026-3785, 0026-0426, 0026-3783, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
Bayer Healthcare Manufacturing Srl		630762938	ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426), LABEL(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
BioReliance Corporation			ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
Nelson Laboratories, LLC			ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426)

Establishment					
Name	Address	ID/FEI	Business Operations		

MANUFACTURE(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-0426), ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-0426), 0026-3782, 0026-4782, 0026-0426, 0026-3787, 0026-4783, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426, 0026-3787, 0026-4787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
Vetter Pharma Fertigung GmbH & Co. KG (Ravensburg Mooswiesen)		312670654	ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426), MANUFACTURE(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3785, 0026-0426, 0026-3787, 0026-0426, 0026-3785, 0026-0426, 0026-3783, 0026-0426, 0026-3782, 0026-0426, 0026-3783, 0026-0426, 0026-3782, 0026-0426, 0026-3782, 0026-0426, 0026-3785, 0026-0426, 0026-3782, 0026-0426, 0026-3785, 0026-0426, 0026-3787, 0026-0426, 0026-3783, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426, 0026-3787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
Bayer AG		315097875	PACK(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026- 3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026- 4787, 0026-0426), LABEL(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426), ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026-0426), MANUFACTURE(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-0426, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026-0426, 0026-4782, 0026-0426, 0026-4782, 0026-0426, 0026-4786, 0026-0426, 0026-4782, 0026-0426, 0026-4782, 0026-0426, 0026-4786, 0026-0426, 0026-4782, 0026-0426, 0026-4782, 0026-0426, 0026-4782, 0026-0426, 0026-4786, 0026-0426, 0026-4782, 0026-0426, 0026-4782, 0026-0426, 0026-4786, 0026-0426, 0026-4782, 0026-0426, 0026-4786, 0026-0426, 0026-4782, 0026-0426, 0026-4786, 0026-0426, 0026-4787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
Vetter Pharma Fertigung GmbH & Co. KG (Langenargen Eisenbahnstrasse)		344217323	ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026-4783, 0026- 0426, 0026-3782, 0026-4782, 0026-0426, 0026-3786, 0026-4786, 0026- 0426, 0026-3787, 0026-4787, 0026-0426), MANUFACTURE(0026-3785, 0026- 4785, 0026-0426, 0026-3783, 0026-4783, 0026-0426, 0026-3782, 0026- 4782, 0026-0426, 0026-3786, 0026-4786, 0026-0426, 0026-3787, 0026- 4787, 0026-0426)

Establishment

Name	Address	ID/FEI	Business Operations
Vetter Pharma Fertigung GmbH & Co. KG (Ravensburg Helmut- Vetter-Strasse)		341629292	ANALYSIS(0026-3785, 0026-4785, 0026-0426, 0026-3783, 0026- 4783, 0026-0426, 0026-3782, 0026-4782, 0026-0426, 0026- 3786, 0026-4786, 0026-0426, 0026-3787, 0026-4787, 0026- 0426)

Revised: 12/2019

Bayer HealthCare LLC