HEMGENIX- etranacogene dezaparvovec CSL Behring

HIGHLIGHTS OF PRESCRIBING INFORMATION

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

HEMGENIX (etranacogene dezaparvovec-drlb) suspension, for intravenous infusion Initial U.S. Approval: 2022

------INDICATIONS AND USAGE

HEMGENIX is an adeno-associated virus vector-based gene therapy indicated for the treatment of adults with Hemophilia B (congenital Factor IX deficiency) who: (1)

- Currently use Factor IX prophylaxis therapy, or
- Have current or historical life-threatening hemorrhage, or
- Have repeated, serious spontaneous bleeding episodes.

------DOSAGE AND ADMINISTRATION ------

For single-use intravenous infusion only. (2)

- Perform baseline testing to select patients, including testing for Factor IX inhibitor presence and liver health tests. (2.1)
- The recommended dose of HEMGENIX is 2×10^{13} genome copies (gc) per kg of body weight. (2.1)
- Administer HEMGENIX as an intravenous infusion after dilution with 0.9% normal saline at a constant infusion rate of 500 ml/hour (8 mL/min). (2.1)

----- DOSAGE FORMS AND STRENGTHS -----

HEMGENIX is a suspension for intravenous infusion. (3)

HEMGENIX is provided in kits containing 10 to 48 single-use vials, each kit constituting a dosage unit based on the patient's body weight. (3)

HEMGENIX has a nominal concentration of 1×10^{13} gc/mL, and each vial contains an extractable volume of not less than 10 mL (3)

------CONTRAINDICATIONS -----

None. (4)

------ WARNINGS AND PRECAUTIONS

- Infusion reactions: Monitor during administration and for at least 3 hours after end of infusion. If symptoms occur, slow or interrupt administration. Re-start administration at a slower infusion once resolved. (2.3, 5.1)
- Hepatotoxicity: Closely monitor transaminase levels once per week for 3 months after HEMGENIX
 administration to mitigate the risk of potential hepatotoxicity. Continue to monitor transaminases in all
 patients who developed liver enzyme elevations until liver enzymes return to baseline. Consider
 corticosteroid treatment should elevations occur. (5.2)
- Hepatocellular carcinogenicity: For patients with preexisting risk factors (e.g., cirrhosis, advanced hepatic fibrosis, hepatitis B or C, non-alcoholic fatty liver disease (NAFLD), chronic alcohol consumption, non-alcoholic steatohepatitis (NASH), and advanced age), perform regular (e.g., annual) liver ultrasound and alpha-fetoprotein testing following administration. (5.4)
- Monitoring Laboratory tests: Monitor for Factor IX activity and Factor IX inhibitors. (5.5)

----- ADVERSE REACTIONS

The most common adverse reactions (incidence ≥5%) were elevated ALT, headache, blood creatine kinase elevations, flu-like symptoms, infusion-related reactions, fatigue, malaise and elevated AST. (6)

To report SUSPECTED ADVERSE REACTIONS, contact CSL Behring at 1-866-915-6958 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

------USE IN SPECIFIC POPULATIONS ------

No dose adjustment is required in geriatric, hepatic, or renal impaired patients. (8.5, 8.6, 8.7)

See 17 for PATIENT COUNSELING INFORMATION.

Revised: 12/2022

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1 INDICATIONS AND USAGE

HEMGENIX is an adeno-associated virus vector-based gene therapy indicated for treatment of adults with Hemophilia B (congenital Factor IX deficiency) who:

- Currently use Factor IX prophylaxis therapy, or
- Have current or historical life-threatening hemorrhage, or
- Have repeated, serious spontaneous bleeding episodes.

2 DOSAGE AND ADMINISTRATION

For single-use intravenous infusion only.

For patient selection:

- Perform Factor IX inhibitor titer testing.
 In case of a positive test result for human Factor IX inhibitors, perform a re-test within approximately 2 weeks. If both the initial test and re-test results are positive, do not administer HEMGENIX to this patient.
- Perform liver health assessments, including:
 - Enzyme testing [alanine aminotransferase (ALT), aspartate aminotransferase (AST), alkaline phosphatase (ALP) and total bilirubin)],
 - Hepatic ultrasound and elastography.

In case of radiological liver abnormalities and/or sustained liver enzyme elevations, consider a consultation with hepatologist to assess eligibility for HEMGENIX [see Warnings and Precautions (5.2)].

2.1 Dose

The recommended dose of HEMGENIX is 2×10^{13} genome copies (gc) per kilogram (kg) of body weight (or 2 mL/kg body weight) administered as an intravenous infusion after dilution with 0.9% sodium chloride solution (normal saline) [see Dosage and Administration (2.2)].

Calculate the dose as follows:

HEMGENIX dose (in mL) = patient body weight (in kilogram) \times 2

The multiplication factor 2 represents the per kilogram dose (2 \times 10¹³ gc/kg) divided by the amount of genome copies per mL of the HEMGENIX solution (1 \times 10¹³ gc/mL).

Number of HEMGENIX vials needed = HEMGENIX dose (in mL) divided by 10 (round up to next whole number of vials).

The division factor 10 represents the extractable volume of HEMGENIX from each vial (10 mL).

The total volume of the patient's HEMGENIX dose to be diluted may be less than the total volume of vials needed.

Example calculation for 72 kg patient

Weight	(body weight multiplied by 2)	dose (mL) divided by 10, then rounded up]
72 kg	144 mL	15

HEMGENIX can be administered only once.

2.2 Preparation

The vials are for single-dose only.

General precautions

- Prepare HEMGENIX using sterile technique under aseptic conditions, proper engineering controls (e.g., biological safety cabinet or isolator) and according to institutional policies.
- Do not expose HEMGENIX to the light of an ultraviolet radiation disinfection lamp.
- Confirm that the patient's identity matches with the patient-specific identifier number on the outer carton.
- Verify the required dose of HEMGENIX based on the patient's body weight.
- Confirm that the carton contains sufficient number of vials to prepare the diluted HEMGENIX patient-specific infusion bag.
- Parenteral drug products should be inspected visually for particulate matter and discoloration prior to administration, whenever solution and container permit.

Required supplies and materials:

Normal saline infusion bag(s)* of 500 mL (1 to 2 bags based on patient's body weight) Labels** for the infusion bag(s) of 500 mL

IV Infusion line/drip chamber* primed with 0.9% normal saline

Infusion bag connector(s)

20 mL or larger Luer-lock syringes*

20 G Needles* or vial adaptors*

70% isopropyl alcohol

Sharps disposal container

The following Table shows the supplies and materials compatible with HEMGENIX:

Component*	Material of Construction	
	PE/PP copolymer (PVC-free)	
Normal saline infusion bag	(Stability after dilution was established	
(0.9% normal saline)	using PE/PP copolymer, PVC-free	
	infusion bags with 0.9% normal saline.)	
20 G Needle	Stainless Steel	
Vial adapter	PP, Silicone; PP, stainless; MABS, acrylic	
viai adaptei	silicone; ABS	
Luer-lock syringe	PP, Silicone	
IV Infusion line/drip	PVC/TOTM, PP/styrene-ethylene-	
chamber	butylene-styrene	

MABS = Methyl methacrylate acrylonitrile butadiene styrene; PE = Polyethylene; PP = Polypropylene; PVC = Polyvinyl chloride; TOTM = Trioctyltrimellitate, Acrylonitrile butadiene styrene (ABS)

**Information to be included on the infusion bag label:

- Product name: Diluted Hemgenix
- Patient identifier
- Expiration date/time (24 h from the vial removal from refrigerator)
- Storage condition: Room Temperature [15-25 °C (59-77 °F] protected from light.
- Contains genetically modified organisms
- Number of infusion bag: 1 of 2 bags / 2 of 2 bags

Preparation of 0.9% normal saline infusion bags

- 1. Prior to dilution, spike the infusion bag(s) of 0.9% normal saline solution with applicable connector.
- 2. Connect a luer-lock syringe at the mixing adapter site of the applicable connector.
- 3. Withdraw the volume equal to the calculated HEMGENIX dose (in mL) from the 500 mL infusion bag(s) of 0.9% normal saline solution. The volume to be withdrawn and number of infusion bag(s) needed will vary based on the patient body weight:

Patient body weight	Number of 500 mL 0.9% normal saline infusion bag(s) required	Volume of saline solution to withdraw
Less than 120 kg body weight	One	Equal to the total HEMGENIX dose (in mL) from one bag
Equal to or more than 120 kg body weight	Two	Equal to the total HEMGENIX dose (in mL). Remove half of the dose equivalent volume from each of the two bags.

HEMGENIX injection to the 0.9% normal saline infusion bags

- Dilute HEMGENIX with 0.9% normal saline solution only prior to administration.
- 4. Prior to dilution, inspect each of the HEMGENIX single-dose vials.
 - If particulates, cloudiness, or discoloration is visible, DO NOT use the vial(s).
- 5. Gently swirl the vials 3 times (about 10 seconds) to homogenize the HEMGENIX suspension.
 - To avoid foaming, DO NOT shake the HEMGENIX vial(s).
- 6. Remove the plastic flip-off cap from the vials and disinfect the rubber stopper with a sterilizing agent (for example sterile 70% isopropyl alcohol).
- 7. Withdraw HEMGENIX from each vial using a 20 G needle/vial adapter and syringe.
 - Use recommended 20 mL luer-lock or larger syringe that is suitable for volume measuring and a needle.
 - DO NOT use filter needles during preparation of HEMGENIX.
 - Use a new needle/vial adapter and syringe for each HEMGENIX vial.
 - Dispose of the needle and syringe in an appropriate container.

- 8. Slowly add the required HEMGENIX dose from the syringe(s) directly to the 0.9% normal saline solution in the infusion bag(s) (from step #3) to bring the total volume in each infusion bag back to 500 mL.
 - DO NOT add HEMGENIX into the airspace of the bag to avoid foaming throughout this process.
- 9. Repeat steps 7 and 8 with additional needles/vial adaptors and syringes to inject the total calculated HEMGENIX volume to the infusion bag(s) required for the patient dose.
- 10.Gently invert the infusion bag(s) at least 3 times (about 10 seconds) to mix the solution and ensure even distribution of the diluted product.
 - To avoid foaming, DO NOT shake the diluted HEMGENIX infusion bag(s).
- 11.Label the infusion bag(s).
- 12.Connect the infusion bag(s) to an infusion tube pre-filled with sterile 0.9% normal saline solution to reduce the risk of spillage and/or aerosol formation.
- 13. Transport the diluted HEMGENIX infusion bag(s) in the transport container/bag protected from light to the administration site, avoiding any shaking or excessive agitation.

2.3 Administration

Required supplies and materials for administration:

Winged intravenous needle or catheter set*
Infusion pump
0.2 mcm in-line filter*
Antiseptic skin preps
70% isopropyl alcohol wipes
Gauze and tape, or transparent dressing
Sharps disposal container
Virucidal agent to treat spill/spill kit

The following Table shows the supplies and materials compatible for infusion of HEMGENIX:

Component*	Material of Construction
Winged IV needle or catheter set	PVC/TOTM, MABS
0.2 mcm in-line filter	PES
Catheter	PVC/DEHT, Stainless steel

DEHP = Di(2-ethylhexyl)phthalate; DEHT = Di(2-

ethylhexyl)terephthalate; MABS = Methyl methacrylate acrylonitrile

butadiene styrene; PES = Polyether sulfone; PVC = Polyvinyl

chloride

Administer HEMGENIX as a single-dose intravenous infusion through a peripheral venous catheter:

1. Visually inspect diluted HEMGENIX prior to administration. The diluted HEMGENIX should be clear and colorless.

- DO NOT use if particulates, cloudiness, or discoloration are visible.
- Use the diluted HEMGENIX within 24 hours after the dose preparation [see How supplied/Storage and Handling (16.2)].
- 2. Use an integrated (in-line) 0.2 mcm filter made out of PES.
- 3. Subsequently, connect the pre-filled IV infusion line/drip chamber to the main intravenous line which has been primed with sterile 0.9% normal saline solution prior to use.
- 4. Infuse diluted HEMGENIX at a constant infusion rate of 500 mL/hour (8 mL/min).
 - DO NOT administer HEMGENIX as an intravenous push or bolus.
 - DO NOT infuse the diluted HEMGENIX solution in the same intravenous line with any other products.
 - DO NOT use a central line or port.

In the event of an infusion reaction during administration [see Warnings and Precautions (5.1)]:

- The rate of infusion may be reduced or stopped, to manage the infusion reaction.
 If the infusion is stopped, restart at a slower rate when the infusion reaction is resolved.
- If the infusion rate needs to be reduced, or stopped and restarted, HEMGENIX should be infused within 24 hours after the dose preparation [see How supplied/Storage and handling (16.2)].
- 5. After the entire content of the bag(s) is infused, flush the IV infusion line/drip chamber at the same infusion rate with 0.9% normal saline solution to ensure all HEMGENIX is delivered.
 - Treat spills of HEMGENIX with a virucidal agent with proven activity against nonenveloped viruses.
 - Dispose of unused product and disposable materials that may have come in contact with HEMGENIX in accordance with local biosafety guidelines applicable for handling and disposal of the pharmaceutical waste.

Monitoring Post-Administration

Conduct the following tests after HEMGENIX administration [see Warnings and Precautions (5.2, 5.3, 5.4)]:

- Perform regular liver enzyme testing to monitor for liver enzyme elevations which may indicate immune-mediated hepatotoxicity:
 - Monitor ALT and AST (transaminase) levels by testing weekly for 3 months following administration of HEMGENIX. Continue to monitor transaminases in all patients who developed liver enzyme elevations until liver enzymes return to baseline.
 - In the event of ALT increase to above normal limits or to twice the patient`s
 baseline in the first 3 months post-dose, consider implementing a course of
 corticosteroids. For patients with clinically relevant ALT increases who need
 corticosteroid treatment, administer the recommended starting dose of 60
 mg/day of oral prednisolone or prednisone, with a subsequent taper in response
 to normalization of the ALT levels (see Table 1):

Table 1. Prednisolone Treatment Applied in Clinical Studies With HEMGENIX:

Timeline	Dose (mg/day)	
Week 1	60	
Week 2	40	
Week 3	30	
Week 4	30	
Maintenance dose until ALT level returns to baseline level	20	
Taper dose after ALT baseline level has been reached	Reduce daily dose by 5 mg/week	

^{*} Medications equivalent to prednisolone may also be used. A combined immunosuppressant regimen or the use of other products can be considered in case of prednisolone treatment failure or contraindication.

In the clinical studies, the mean duration of corticosteroid use for elevated transaminases was 81.4 days [Standard Deviation (SD) 28.6] and ranged from 51 to 130 days [see Warnings and Precautions (5.2)].

- Monitor Factor IX activity (e.g., weekly for 3 months).
 - Monitor patients regularly for their Factor IX activity, in particular when exogenous Factor IX is administered. It may take several weeks before improved hemostatic control becomes apparent after HEMGENIX infusion; therefore, continued hemostatic support with exogenous human Factor IX may be needed during the first weeks after HEMGENIX infusion [see Clinical Pharmacology (12.3)].
 - The use of different assays may impact the test results; therefore, use the same assay and reagents to monitor patients over time, if feasible [see Monitoring Laboratory Tests (5.5)].
 - Use of exogenous Factor IX concentrates before and after HEMGENIX administration may impede assessment of endogenous, HEMGENIX-derived Factor IX activity.
- Perform regular alpha-fetoprotein (AFP) level testing and abdominal ultrasound (e.g., annually) in patients with preexisting risk factors for hepatocellular carcinoma (e.g., in patients with cirrhosis, advanced hepatic fibrosis, hepatitis B or C, non-alcoholic fatty liver disease (NAFLD), chronic alcohol consumption, non-alcoholic steatohepatitis (NASH), and advanced age).
- Monitor patients for human Factor IX inhibitors. Post-dose inhibitor testing should be performed if bleeding is not controlled, or plasma Factor IX activity levels decrease [see Warnings and Precautions (5.5)].

3 DOSAGE FORMS AND STRENGTHS

HEMGENIX is a clear and colorless suspension for intravenous infusion.

HEMGENIX is provided in a kit containing 10 to 48 vials. Each kit constitutes a dosage unit based on the patient's body weight.

HEMGENIX has a nominal concentration of 1×10^{13} gc/mL, and each vial contains an

extractable volume of not less than 10 mL.

4 CONTRAINDICATIONS

None.

5 WARNINGS AND PRECAUTIONS

5.1 Infusion Reactions

Infusion reactions, including hypersensitivity reactions and anaphylaxis, may occur. Symptoms may include chest tightness, headaches, abdominal pain, lightheadedness, flu-like symptoms, shivering, flushing, rash, and hypertension. Closely monitor patients for signs or symptoms of an infusion reaction throughout the infusion period and for at least 3 hours after end of infusion. Do not infuse the product faster than 500 mL/hour [see Adverse Reactions (6)].

In the event of an infusion reaction during administration, the infusion may be slowed or stopped. If the infusion is stopped, restart at a slower rate when the infusion reaction has resolved. Consider treatment with a corticosteroid or antihistamine for management of an infusion reaction [see Dosage and Administration (2.1)].

5.2 Hepatotoxicity

Intravenous administration of a liver-directed AAV vector could potentially lead to liver transaminase elevations (transaminitis). Transaminitis, particularly when observed in the first 3 months after HEMGENIX administration, is presumed to occur due to immunemediated injury of transduced hepatocytes and may reduce the therapeutic efficacy of the AAV-vector based gene therapy.

In clinical studies with HEMGENIX, most subjects had asymptomatic, and predominantly mild elevations in transaminases. Elevated ALT levels occurred most often in the first 4 months after HEMGENIX administration. There were some subjects who had a late onset of elevated ALT levels between Months 6-24 (range = 42 IU/L-193 IU/L); however, all of these ALT values were <2× ULN with the exception of one subject. Three additional subjects had AST elevations with onset and resolution between Months 6 and 12 (range = 41 IU/L - 96 IU/L).

In one subject, an ALT elevation $>5 \times$ ULN occurred 24 days after HEMGENIX administration and resolved by 51 days post-HEMGENIX administration. There was one subject who had an AST elevation $> 5 \times$ ULN that occurred 11 months post-HEMGENIX administration and resolved to $<2 \times$ ULN eight days later.

The majority of the elevated ALT values returned to baseline, however 9 subjects' ALT values never resolved to normal (range at 2-year follow up = 48 IU/L - 193 IU/L) [see Adverse Reactions (6)].

Closely monitor transaminase levels once per week for 3 months after HEMGENIX administration to mitigate the risk of potential hepatotoxicity. Continue to monitor transaminases in all patients who developed liver enzyme elevations until liver enzymes return to baseline [see Dosage and Administration (2.3)].

In case of increased ALT levels above the upper limit of normal or double baseline levels,

consider implementing a course of corticosteroid, along with human Factor IX activity monitoring [see Dosage and Administration (2.3)].

5.3 Immune-mediated neutralization of the AAV5 vector capsid

In AAV-vector based gene therapies, preexisting neutralizing anti-AAV antibodies may impede transgene expression at desired therapeutic levels. Following treatment with HEMGENIX all subjects developed neutralizing anti-AAV antibodies. Currently, there is no validated neutralizing anti-AAV5 antibody assay.

In the clinical studies with HEMGENIX, an unvalidated clinical trial assay was utilized to assess preexisting neutralizing anti-AAV5 antibodies. The subject sub-group with detectable preexisting neutralizing anti-AAV5 antibodies up to titers of 1:678 showed mean Factor IX activity that was numerically lower compared to that subject sub-group without detectable preexisting neutralizing anti-AAV5 antibodies. Subjects, with and without preexisting neutralizing anti-AAV5 antibodies, demonstrated hemostatic protection. In one subject with a preexisting neutralizing anti-AAV5 antibody titer of 1:3212, no human Factor IX expression was observed, and restart of the exogenous Factor IX prophylaxis was needed for bleeding events. [see Clinical Studies (14)].

Anti-AAV5 Antibody Study

Patients who intend to receive treatment with HEMGENIX are encouraged to enroll in a study to measure pre-existing anti-AAV5 neutralizing antibodies by calling CSL Behring at 1-800-504-5434. The study evaluates the effect of pre-existing anti-AAV5 neutralizing antibodies on the risk of bleeding.

5.4 Hepatocellular carcinogenicity

The integration of liver-targeting AAV vector DNA into the genome may carry the theoretical risk of hepatocellular carcinoma development.

HEMGENIX is composed of a non-replicating AAV5 vector whose DNA persists largely in episomal form. Random integration of HEMGENIX vector DNA to the human DNA at low frequency is possible. No HEMGENIX-associated clonal expansion or carcinogenicity was observed in clinical studies [see Clinical Studies (14)]. One subject with preexisting risk factors for developing hepatic cancer developed a hepatocellular carcinoma, which was assessed as not likely related to HEMGENIX treatment based on vector integration site analyses and whole genome sequencing.

Patients with preexisting risk factors for hepatocellular carcinoma (e.g., patients with cirrhosis, advanced hepatic fibrosis, hepatitis C or B, non-alcoholic fatty liver disease (NAFLD), chronic alcohol consumption, non-alcoholic steatohepatitis (NASH), and advanced age) should receive abdominal ultrasound screenings and be monitored regularly (e.g., annually) for alpha-fetoprotein (AFP) elevations in the 5 years following administration [see Dosage and Administration (2.3)].

5.5 Monitoring Laboratory Tests

After HEMGENIX administration, regularly monitor patient's Factor IX activity levels.

When using an in vitro activated partial thromboplastin time (aPTT)-based one-stage clotting assay (OSA) for determining Factor IX activity, plasma Factor IX activity results can be affected by both the type of aPTT reagent and the reference standard used in

the assay. This is important to consider particularly when changing the laboratory and/or reagents used in the assay. Therefore, the same assay and reagents are recommended to be used to monitor Factor IX activity over time.

The results of Factor IX activity tests are lower if measured with chromogenic substrate assay (CSA) compared to OSA.

In the clinical efficacy study with HEMGENIX, the post-dose Factor IX activity measured with CSA returned lower values with the mean CSA to OSA Factor IX activity ratio ranging from 0.41 to 0.55.

Monitor patients through appropriate clinical observations and laboratory tests for the development of inhibitors to Factor IX after HEMGENIX administration. Perform an assay that detects Factor IX inhibitors if bleeding is not controlled, or plasma Factor IX activity levels decrease.

6 ADVERSE REACTIONS

The most common adverse reactions (incidence ≥5%) reported in clinical studies were ALT elevations, headache, blood creatine kinase elevations, flu-like symptoms, infusion-related reactions, fatique, malaise, and AST elevations.

The following adverse reactions are discussed in greater detail in other sections of the label:

- Infusion related reactions [see Warnings and Precautions (5.1)].
- Hepatotoxicity [see Warnings and Precautions (5.2)].
- Immune-mediated neutralization of the AAV5 vector capsid [see Warnings and Precautions (5.3)].

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 the clinical trials of another drug and may not reflect the rates observed in clinical practice.

The safety of HEMGENIX was evaluated in two clinical studies (the first study enrolled 3 subjects and the second study 54 subjects). Both studies enrolled adult male subjects with moderately severe or severe Hemophilia B (N = 57), who received a single intravenous dose of 2×10^{13} gc/kg body weight of HEMGENIX. All subjects entered a follow-up period of 5 years.

No serious adverse reactions were reported [see Clinical Studies (14)]. The most common adverse reactions observed in $\geq 5\%$ of subjects post-dose are listed in Table 2:

Table 2. Adverse Reactions (Incidence ≥5%) Following Treatment with HEMGENIX

Adverse Reactions ≥5%	Subjects (%) (N = 57)
Alanine aminotransferase increased	24 (42%)
Headache	10 (18%)
Blood creatine kinase increased	24 (42%)
Flu-like symptoms	8 (14%)

Infusion-related reactions* (see below)	19* (33%)
Hypersensitivity	2 [†] (4%)
Fatigue	7 (12%)
Aspartate aminotransferase increased	24 (42%)
Nausea	4 (7%)
Malaise	7 (12%)

- * Infusion-related reaction: In 7 subjects symptoms occurred during infusion, in 12 subjects after infusion. Symptoms occurring in ≥ 5% of subjects were: Dizziness, Flu-like symptoms and Headache. Symptoms occurring in < 5% of subjects were: Abdominal pain, Abdominal discomfort, Chest discomfort, Chills, Eye pruritus, Fever (Pyrexia), Flushing, Hives (Urticaria), Infusion site reaction, and Tachycardia. Eleven subjects recovered on the day or day one after infusion. Eight subjects recovered within 8 days after infusion.
- † 1of 2 hypersensitivity reactions 12 minutes after initiation of administration of HEMGENIX, the patient experienced high blood pressure, red eyes, feeling warm, dizziness, coughing, dyspnea, elevated heart rate, shivering, and leg cramps. Infusion was stopped and not restarted. Only 10% of the HEMGENIX dose was administered. The patient recovered on the same day after treatment with intravenous diphenhydramine and intramuscular epinephrine.

 2 of 2 hypersensitivity reactions 10 minutes after initiation of administration of HEMGENIX, the patient experienced itching, tightness of throat, and swelling of the right side of the neck. The HEMGENIX dose was not interrupted and administered in full. All symptoms resolved on the same day without treatment.

Infusion-related reactions were observed in 19 subjects. Infusions were temporarily interrupted in 3 subjects and resumed at a slower infusion rate after treatment with antihistamines and/or corticosteroids. In one subject, infusion was stopped and not resumed (see footnote of Table 2).

There were 24 subjects who had elevated ALT values from Day 8 to 731 postadministration.

Five subjects had ALT elevations $>2-3\times$ ULN (range = 89 IU/L - 130 IU/L), one subject had an ALT elevation $> 3-5\times$ ULN (193 IU/L) and one subject had an ALT elevation $> 5\times$ ULN (275 IU/L). The subject who had the ALT elevation $> 5\times$ ULN occurred 3 weeks after HEMGENIX administration.

Five subjects had AST elevations $> 2-3 \times$ ULN (range = 71 IU/L - 118 IU/L), three subjects had AST elevations $> 3-5 \times$ ULN (range = 127 IU/L - 163 IU/L) and one subject had an AST elevation $> 5 \times$ ULN (327 IU/L). The subject who had the AST elevation $> 5 \times$ ULN occurred 11 months post-HEMGENIX administration.

Seventeen subjects had elevations in ALT levels within the first 4 months after HEMGENIX infusion (range = 41 IU/L - 275 IU/L), eleven of these subjects' ALT levels resolved within 4 months post-infusion (range = 41 IU/L - 275 IU/L) and five of these subjects' ALT levels never normalized as of last follow-up (range of values at 2-year follow-up = 48 IU/L - 110 IU/L). Seven additional subjects had ALT elevations with onset between Months 6 to 24 (range = 42 IU/L - 193 IU/L), five of these subjects had additional risk factors for having elevated transaminase levels including Hepatitis C and Human Immunodeficiency Virus (HIV). ALT levels never normalized as of last follow-up (range of values at 2-year follow-up = (59 IU/L - 193 IU/L) in three of the subjects with ALT

elevations with onset between Months 6 to 24.

Nineteen subjects had elevations in AST levels within 3 months after HEMGENIX infusion (range = 32 IU/L - 163 IU/L). Nine of these subjects' AST elevations resolved within 4 months post-infusion (range = 35 IU/L - 163 IU/L), three resolved within 7 to 13 months post-infusion (range = 35 IU/L - 62 IU/L), and seven of these subjects' AST levels never normalized as of last follow-up (range of values at 2-year follow-up = 36 IU/L - 327 IU/L). The remaining 5 subjects with AST elevation had onset of between 6 months and 2 years post-infusion (range = 36 IU/L - 127 IU/L), and AST levels had not normalized as of the last follow-up for one subject (AST at 2-year follow-up = 127 IU/L) who had additional risk factors for having elevated transaminase levels.

Nine subjects with ALT elevations received a tapered course of corticosteroids. The mean duration of corticosteroid treatment for the elevated ALT was 81.4 days. Nineteen of the 24 subjects with ALT elevations also had a related AST elevation. Twenty-one subjects had elevated transaminase levels and were not treated with corticosteroids. [see Clinical Studies (14)].

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

HEMGENIX is not intended for administration in women. No adverse effects on mating rate and fertility indices or fetal weights were observed in healthy naïve female mice mated with healthy male mice that were intravenously administered a predecessor of HEMGENIX product 6 days prior to mating. Vector DNA was not detected in the uterus, placenta, or fetus.

In the United States general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2 to 4% and 15 to 20%, respectively.

8.2 Lactation

Risk Summary

HEMGENIX is not intended for administration in women.

8.3 Females and Males of Reproductive Potential

Risk Summary

No clinical studies have been performed to evaluate the effects of HEMGENIX on fertility in humans. Twenty days after intravenous administration of a predecessor of HEMGENIX product in healthy male mice, vector DNA was detected in all reproductive tissues examined (epididymis, seminal vesicles, testes, and sperm). However, no differences were observed in mating rates and fertility indices in healthy naïve female mice following mating with the dosed males.

8.4 Pediatric Use

The safety and efficacy of HEMGENIX in pediatric patients have not been established.

8.5 Geriatric Use

The clinical studies included a total of 6 geriatric subjects with Hemophilia B, aged 68 to 75 years at time of enrollment. No meaningful differences in the safety and efficacy profile were observed in these subjects compared to subjects aged 18 to 65 years, and no dose adjustment was made [see Clinical Studies (14)].

8.6 Hepatic Impairment

Limited clinical data in subjects with liver impairment indicate numerically lower FIX activity as compared to subjects without hepatic impairment [see Clinical Pharmacology (12.3)]. In the clinical studies, no dose adjustment was made in subjects with hepatic pathologies. The safety and efficacy in subjects with advanced hepatic impairment, including cirrhosis, advanced liver fibrosis, or uncontrolled Hepatitis B and C, have not been studied.

8.7 Renal Impairment

Limited clinical data are available in subjects with mild and moderate renal impairment [see Clinical Pharmacology (12.3)]. In the clinical studies, no dose adjustment was made in these subjects. The safety and efficacy in subjects with severe renal impairment and end-stage renal disease have not been studied.

11 DESCRIPTION

HEMGENIX (etranacogene dezaparvovec-drlb) is an adeno-associated viral vector-based gene therapy for intravenous infusion after dilution. HEMGENIX is a non-replicating recombinant AAV5 containing a codon-optimized DNA sequence of the gain-of-function Padua variant of human Factor IX (variant R338L), under control of a liver-specific promotor 1 (LP1).

HEMGENIX has a nominal concentration of 1×10^{13} gc/mL. Each vial contains an extractable volume of no less than 10 mL of HEMGENIX and the following excipients: sucrose (50 mg/mL), polysorbate-20 (0.22 mg/mL), potassium chloride (0.2 mg/mL), potassium phosphate (0.2 mg/mL), sodium chloride (8 mg/mL), and sodium phosphate (1.2 mg/mL).

HEMGENIX is sterile, clear and colorless suspension, and contains no preservative. After dilution, HEMGENIX should be clear and colorless suspension.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

HEMGENIX is an adeno-associated virus serotype 5 (AAV5) based gene therapy designed to deliver a copy of a gene encoding the Padua variant of human coagulation Factor IX (hFIX-Padua). Single intravenous infusion of HEMGENIX results in cell transduction and increase in circulating Factor IX activity in patients with Hemophilia B.

12.2 Pharmacodynamics

Factor IX activity

The mean Factor IX activity levels over time, as measured by one-stage [activated Partial Thromboplastin Time (aPTT)-based] assay are summarized in Table 3. Subjects achieved a mean (\pm SD) uncontaminated (i.e., excluding measurements within five half-lives of Factor IX replacement therapy) Factor IX activity levels of 39% (\pm 18.7), 41.5% (\pm 21.7), 36.9% (\pm 21.4) and 36.7 (\pm 19.0) of normal, respectively, at 6, 12, 18 and 24 months. The time to onset of Factor IX protein expression post-dose was detectable by first uncontaminated measurement at Week 3 in the clinical efficacy study (N = 54) [see Clinical Studies (14)].

Table 3: Summary of Uncontaminated Factor IX Activity Over Time Following Administration of 2×10^{13} gc/kg of HEMGENIX [FAS; One-Stage (aPTT-Based) Assay]

	Factor IX Activity in % (One-stage)			
	Subject Number (*n)	Median (Min, Max)	Mean (SD)	
Week 3	43	23.7 (4.9, 56.7)	26.8 (12.7)	
Month 3	51	33.8 (7.6, 91.0)	36.8 (18.2)	
Month 6	51	37.3 (8.2, 97.1)	39.0 (18.7)	
Month 12	50	39.9 (5.9, 113.0)	41.5 (21.7)	
Month 18	50	33.6 (4.5, 122.9)	36.9 (21.4)	
Month 24	50	33.9 (4.7, 99.2)	36.7 (19.0)	

Abbreviations: SD = Standard Deviation; FAS = Full Analysis Set including all 54 subjects dosed; Min = Minimum; Max = Maximum. Uncontaminated Factor IX activity values exclude measurements within five half-lives of Factor IX replacement therapy.

Pharmacodynamics in specific populations

<u>Age</u>

Limited data (N = 7) from 60 -75 years subgroup showed that the mean Factor IX activity levels were approximately up to 2-fold higher in this subgroup compared to 18 to < 40 years age subgroup (N = 31), but comparable to 40 to <60 years age subgroup (N = 15).

Hepatic Impairment

In the clinical efficacy study, subjects with varying degree of baseline liver pathology, specifically the degree of hepatic steatosis with the Controlled Attenuation Parameter (CAP) score of \geq S2 (\geq 260 decibels/m; range: 262 to 400; n = 12) versus <S2 (<260 decibels/m; range: 100 to 259; n = 28;) and missing score (n = 14) were compared [see Clinical Studies (14)]. The mean (\pm SD) uncontaminated Factor IX activity for <S2 versus \geq S2 subgroups at Months 6, 12, 18, and 24 post dose were 40.8 (\pm 20.1) versus 34.5 (\pm 13.7), 46.4 (\pm 24.1) versus 32.6 (\pm 18.6), 41.6 (\pm 25.7) versus 29.2 (\pm 13.7), and 40.2 (\pm 19.8) versus 28.4 (\pm 13.1), respectively.

^{*} Contaminated and missing values are not shown here. Specifically, the number of subjects excluded for contamination with Factor IX replacement therapy at Week 3, Month 3, Month 6, Month 12, Month 18, and Month 24, were 10, 3, 3, 3, 3, 2, respectively

Subjects with advanced liver impairment and advanced fibrosis (elastography of e.g., \geq 9 kPA, or suggestive of or equal to METAVIR Stage 3 disease), were not studied.

Renal Impairment

In the clinical efficacy study, subjects with mild renal impairment (creatinine clearance (CLcr) = 60 to 89 mL/min defined by Cockcroft-Gault equation, n = 7) had about 37% higher Factor IX activity relative to those with normal renal function (CLcr \geq 90 mL/min; n = 45) following HEMGENIX administration. One subject with moderate renal impairment (CLcr = 30 to 59 mL/min) had similar Factor IX activity as subjects with normal renal function.

HEMGENIX was not studied in subjects with severe renal impairment (CLcr = 15 to 29 mL/min) or end-stage renal disease (CLcr < 15 mL/min).

12.3 Pharmacokinetics

<u>Vector Biodistribution (within the body) and Vector Shedding (excretion/secretion)</u>

Nonclinical data

Biodistribution of HEMGENIX was evaluated after intravenous administration in healthy male mice and non-human primates (NHPs). The highest levels of vector DNA were detected in the liver and adrenal glands in both species. Vector DNA was also detected in all reproductive tissues examined (epididymis, seminal vesicles, and testes). In a mating study evaluating a predecessor of HEMGENIX, transmission of vector DNA to naïve female mice following mating with dosed males was not observed [see Nonclinical Toxicology (13.2)].

Clinical data

Following administration of the predecessor of HEMGENIX at doses of 5×10^{12} (N = 5) and 2×10^{13} gc/kg (N = 5) in a clinical study, the pharmacokinetics of vector DNA in blood and viral shedding in saliva, nasal secretions, semen, urine, and feces were characterized. Clearance of vector DNA as confirmed by 3 subsequent measurements below limit of detection (LOD), was achieved in all subjects at both dose levels from all the matrices except for semen, where clearance was achieved in 9/10 subjects. One subject was unable to produce semen due to a historical medical condition and, therefore, shedding from semen could not be assessed. The maximum time to clearance of vector DNA was 22 weeks for urine, 26 weeks for saliva and nasal secretions, 40 weeks for feces, 52 weeks for semen, and 159 weeks for blood.

Subsequently, the pharmacokinetics of vector DNA in blood, and viral shedding in semen following HEMGENIX administration was characterized in 2 clinical studies.

In an initial clinical study (N = 3), clearance of vector DNA from semen and blood (i.e., confirmed with 3 subsequent measurements below LOD of vector DNA) was achieved in 2/3 subjects, and in all subjects, respectively, after 3 years post-administration. One subject did not return the required number of semen samples to assess the shedding status as per the definition of 3 subsequent measurements below LOD of vector DNA.

In the clinical efficacy study (N = 54), a total of 56% (30/54) of subjects achieved absence of vector DNA from blood and 69% (37/54) from semen by Month 24. Several subjects did not return the required number of blood and semen samples to assess the shedding status as per the definition of 3 subsequent measurements below LOD of

vector DNA. Considering results obtained from 2 available consecutive samples below LOD, a total of 40/54 (74%) and 47/54 (87%) subjects were identified to have reached absence of vector DNA from blood and semen, respectively, at 24 months postadministration.

12.6 Immunogenicity

In clinical studies sustained humoral immune response to infused AAV5 capsid was observed in all subjects following treatment with HEMGENIX. The neutralizing anti-AAV5 antibody levels raised above the upper limit of quantification by week 3 post-administration and remained elevated, as measured at month 24 post-dose. Readministration of HEMGENIX in the presence of high anti-AAV5 antibody titer has not been evaluated. Currently, there is no validated neutralizing anti-AAV5 antibody assay.

13 NONCLINICAL TOXICOLOGY

Nonclinical studies were initiated with a predecessor of HEMGENIX product, rAAV5 expressing the wild type human coagulation factor IX (rAAV5-hFIX). HEMGENIX was developed by introducing a 2-nucleotide change in the transgene for hFIX, generating the naturally occurring Padua variant of Factor IX (rAAV5-hFIX-Padua).

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

No traditional nonclinical carcinogenicity or mutagenicity studies were conducted with HEMGENIX; such studies were not indicated. No adverse effects were observed in mating rates and fertility indices in healthy naïve female mice following mating with males that were administered the predecessor of HEMGENIX [see Use in specific populations (8.3)]. To evaluate vector integration, host genomic DNA was isolated from liver tissue obtained from healthy mice and NHPs following intravenous administration of the predecessor of HEMGENIX. For both species, the identified rAAV5-hFIX vector DNA sequences represented episomal forms that were not integrated into the host DNA. A low level of integrated rAAV5-hFIX DNA was distributed throughout the host genome with no predilection to specific integration sites, including in genes associated with malignant transformation in humans.

13.2 Animal Toxicology and/or Pharmacology

A pharmacology study was conducted in a murine model of Hemophilia B ($B6.129P2-F9^{tm1Dws}$). Intravenous administration of the predecessor of HEMGENIX at dose levels ranging from 5×10^{11} to 2.3×10^{14} gc/kg, resulted in dose-dependent increases in plasma hFIX protein levels, plasma hFIX clotting activity, and vector transduction in the liver at 4 weeks post-dose.

Intravenous administration of HEMGENIX resulted in a no-observed-adverse-effect-level of 5×10^{13} gc/kg (the maximum dose level administered) in healthy mice and 9×10^{13} gc/kg in NHPs. Vector biodistribution to the liver and hFIX protein levels in the plasma occurred in a dose-dependent manner in both species. Anti-hFIX antibodies developed in 5 out of 12 NHPs administered HEMGENIX, which correlated with a decline in circulating hFIX protein levels beginning at 13 weeks post-dose.

One out of 10 healthy mice administered 5 \times 10¹³ gc/kg of HEMGENIX or the predecessor of HEMGENIX developed pulmonary thrombi at 13 weeks post-dose. This

dose level is 2.5-fold higher than the recommended dose level for HEMGENIX. Compared to concurrent controls, prolonged prothrombin time, decreased activated partial thromboplastin time and decreased heart rates were observed in NHPs administered 9 \times 10¹³ gc/kg of HEMGENIX during the 26-week study. This dose level is 4.5-fold higher than the recommended dose level for HEMGENIX.

14 CLINICAL STUDIES

The efficacy of HEMGENIX was evaluated in a prospective, open-label, single-dose, single-arm, multi-national study (N = 54). The study enrolled adult male subjects aged 19 to 75 years, with severe or moderately severe Hemophilia B, who received a single intravenous dose of 2 \times 10¹³ gc/kg body weight of HEMGENIX and entered a follow-up period of 5 years. The study is on-going.

The 54 subjects prospectively completed a lead-in period of at least six months with the intent to receive standard of care routine Factor IX prophylaxis. These 54 subjects then received the indicated single intravenous dose of HEMGENIX. Subjects were then followed up monthly until Month 12, and then at 6-month intervals until Year 5. For the efficacy evaluation, data up to 18 months post-treatment were used. Of the 54 subjects, 53 subjects completed at least 18 months of follow-up in the ongoing study. One subject with numerous cardiovascular and urologic risk factors, aged 75 years at screening, died of urosepsis and cardiogenic shock at Month 15 post-dose (at age 77 years) unrelated to treatment. Another subject received around 10% of the intended dose of HEMGENIX due to an infusion-related hypersensitivity reaction.

The main efficacy outcome was a non-inferiority test of annualized bleeding rate (ABR) during Months 7 to 18 after HEMGENIX treatment compared with ABR during the lead-in period. All bleeding episodes, regardless of investigator assessment, were counted. Subjects were allowed to continue prophylaxis during Months 0 to 6. The estimated mean ABR during Months 7 to 18 after HEMGENIX treatment was 1.9 bleeds/year with a 95% confidence interval (CI) of (1.0, 3.4), compared with an estimated mean ABR of 4.1 [95% CI: 3.2, 5.4] during the lead-in period. The ABR ratio (Months 7 to 18 post-treatment / lead-in) was 0.46 [95% CI: 0.26, 0.81], demonstrating non-inferiority of ABR during Months 7 to 18 compared to the lead-in period.

Two subjects were not able to stop routine prophylaxis after HEMGENIX treatment. During Months 7 to 18, an additional subject received prophylaxis from Days 396-534 [approximately 20 weeks].

Table 4. Total Bleeding Events and ABRs (Full Analysis Set: N=54)

	Lead-in Period*	Months 7 to 18 [†] after HEMGENIX treatment
All Bleeds	136	96 [‡]
Follow-up time (Person-Year)	33	52
Mean Adjusted ABR (95% CI)§	4.1 (3.2, 5.4)	1.9 (1.0, 3.4)

Subjects with Bleeds	40 (74%)	20 (37%)
Subjects with Zero Bleeds	14 (26%)	34 (63%)
Observed Spontaneous Bleed Count (Proportion of total bleeds)¶	50 (37%)	14 (26%)
Observed Joint Bleed Count (Proportion of total bleeds)¶	77 (57%)	19 (35%)

Abbreviations: ABR = Annualized Bleeding Rate; CI = Confidence Interval

- * During the observational lead-in period subjects used their individualized approach to Factor IX prophylaxis derived prior to enrollment in the study, rather than a standardized approach to Factor IX prophylaxis. Not all subjects complied with their prescribed prophylaxis regimen during the lead-in period.
- † Efficacy evaluation started from Month 7 after HEMGENIX treatment, to allow Factor IX expression to reach a steady state.
- ‡ An ABR of 20 was imputed for the period when three subjects were on continuous prophylaxis.
- § Non-inferiority comparison and mean ABR estimates were based on a repeated measures generalized estimating equations negative binomial regression model.
- ¶ For spontaneous and joint bleed counts, no imputation was done for the three subjects receiving continuous prophylaxis during Months 7 to 18.

After a single-dose of HEMGENIX, increases in Factor IX activity were observed [see Pharmacokinetics (12.3)].

16 HOW SUPPLIED/STORAGE AND HANDLING

16.1 How Supplied

HEMGENIX is supplied as sterile, preservative-free, clear, and colorless suspension. HEMGENIX has a nominal concentration of 1×10^{13} gc/mL.

HEMGENIX is provided as a customized kit to meet dosing requirements for each patient [see Dosage and Administration (2.1)], with each kit containing 10 (ten) to 48 (forty-eight) single-use vials (NDC 0053-0099-01), each with an extractable volume of no less than 10 mL of HEMGENIX (see5). The total number of vials in each kit corresponds to the dosing requirement for the individual patient depending on the patient`s body weight [see Dosage and Administration (2.1)]. The customized kit is accompanied with patient`s specific identifier number (Lot) on the outer carton. Each HEMGENIX kit may contain different drug product lots.

Kit sizes and National Drug Codes (NDC) are provided in Table 5:

Table 5. HEMGENIX Multi-Vial Kits

Total Number Dotions Books Total			
Total Number of Vials per Kit	Patient Body Weight (kg)	Volume per Kit (mL)	NDC Number
10	46-50	100	0053-0100-10
11	51-55	110	0053-0110-11
12	56-60	120	0053-0120-12
13	61-65	130	0053-0130-13
14	66-70	140	0053-0140-14
15	71-75	150	0053-0150-15
16	76-80	160	0053-0160-16
17	81-85	170	0053-0170-17
18	86-90	180	0053-0180-18
19	91-95	190	0053-0190-19
20	96-100	200	0053-0200-20
21	101-105	210	0053-0210-21
22	106-110	220	0053-0220-22
23	111-115	230	0053-0230-23
24	116-120	240	0053-0240-24
25	121-125	250	0053-0250-25
26	126-130	260	0053-0260-26
27	131-135	270	0053-0270-27
28	136-140	280	0053-0280-28
29	141-145	290	0053-0290-29
30	146-150	300	0053-0300-30
31	151-155	310	0053-0310-31
32	156-160	320	0053-0320-32
33	161-165	330	0053-0330-33
34	166-170	340	0053-0340-34
35	171-175	350	0053-0350-35
36	176-180	360	0053-0360-36
37	181-185	370	0053-0370-37
38	186-190	380	0053-0380-38
39	191-195	390	0053-0390-39
40	196-200	400	0053-0400-40
41	201-205	410	0053-0410-41
42	206-210	420	0053-0420-42
43	211-215	430	0053-0430-43
44	216-220	440	0053-0440-44
45	221-225	450	0053-0450-45
46	226-230	460	0053-0460-46
47	231-235	470	0053-0470-47
48	236-240	480	0053-0480-48

16.2 Storage and Handling

- HEMGENIX is shipped at 2°C to 8°C (36°F to 46°F).
- Upon receipt, store HEMGENIX vials in a refrigerator at 2°C to 8°C (36°F to 46°F).
- Store HEMGENIX in the original carton until use.
- Protect HEMGENIX from light until time of dilution and administration.
- Do NOT FREEZE.

After dilution

- Once diluted, store HEMGENIX in the infusion bag protected from light.
- Store diluted HEMGENIX in the infusion bag at 15°C to 25°C (59°F to 77°F).
- Infuse the diluted product within 24 hours after the dose preparation [see Dosage and Administration (2.2)].

17 PATIENT COUNSELING INFORMATION

<u>Inform patients that:</u>

- Pre-infusion blood tests will be necessary to look for Factor IX inhibitors. If these
 exist, the patient may not be a good candidate for HEMGENIX [see Dosage and
 Administration (2)].
- Prior to HEMGENIX treatment, a liver ultrasound and elastography will be performed. Patients found to have pre-existing risk factors for hepatocellular carcinoma will be monitored annually in the 5 years following infusion [see Warnings and Precautions (5.4)].
- Infusion reactions can occur. Patients will be monitored during and for at least 3 hours following administration. If a reaction occurs, the infusion rate may be slowed or interrupted, then started at a slower rate [see Warnings and Precautions (5.1)].
- HEMGENIX can elevate certain liver enzymes. Weekly blood tests will be required to monitor for this for 3 months after treatment. Corticosteroid treatment may be necessary if this occurs [see Warnings and Precautions (5.2)].
- If post-infusion bleeding is not controlled or if bleeding returns, then blood tests will be performed for Factor IX activity and neutralizing Factor IX inhibitors [see Warnings and Precautions (5.5)].
- Vector distribution in blood (within the body), and vector shedding in semen and other excreta and secreta can occur post-infusion. It is not known how long this will continue. Patients should not donate blood, organs, tissues, or cells for transplantation [see Pharmacokinetics (12.3)].

Manufactured by:

uniQure, Inc.

113 Hartwell Avenue Lexington, MA 02421 USA

Manufactured for:

CSL Behring LLC

King of Prussia, PA 19406, USA US License No. 1767

Distributed by:

CSL Behring LLC

Kankakee, IL 60901 USA

For Patent information: www.cslbehring.com/products/patents (in-licensed from uniQure)

PRINCIPAL DISPLAY PANEL - 10 ml Vial Label

NDC 0053-0099-01 10 ml

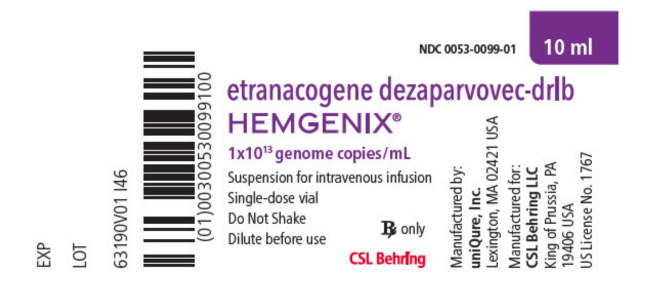
etranacogene dezaparvovec-drlb HEMGENIX®

1x10¹³ genome copies/mL

Suspension for intravenous infusion Single-dose vial Do Not Shake Dilute before use

Rx only

CSL Behring



PRINCIPAL DISPLAY PANEL - 10 ml Vial Carton

NDC 0053-0099-01

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

1x10¹³ genome copies/mL Suspension for intravenous infusion

Single-dose vial 1 x 10¹³ genome copies/mL. No US standard of potency.

Upon receipt store refrigerated at

2°C to 8°C (36°F to 46°F)

Do Not Freeze

Protect from Light

Do Not Shake

Dilute before use

See enclosed prescribing information for dosage and directions for use

Rx only CSL Behring



PRINCIPAL DISPLAY PANEL - Kit Carton

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

1x10¹³ genome copies/mL Suspension for intravenous infusion

Single-dose vial Rx only

Upon receipt store refrigerated at 2°C to 8°C (36°F to 46°F)

Do Not Freeze

Protect from Light

Do Not Shake

Dilute before use

Manufactured by: uniQure, Inc. Lexington, MA 02421 USA

Manufactured for: CSL Behring LLC King of Prussia, PA 19406 USA US License No. 1767

Distributed by: CSL Behring LLC Kankakee, IL 60901 USA

CSL Behring

etranacogene dezaparvovec-drlb HEMGENIX®

1 x 10¹³ genome copies/mL

Suspension for intravenous infusion

Single-dose vial

Upon receipt store refrigerated at 2°C to 8°C (36°F to 46°F)

Do Not Freeze

Dilute before use

B only

10 mL vial x





GTN: 12345678901234 SN: 12345678901234 EXP: MMM.DD,YYYY LOT: P123456789

ΧХ

REG-5717-01

С2Г Верцид

For Patent information:www.cslbehring.com/products/patents (in-licensed from uniQure)

Áuo 在

Contains no preservatives. Discard any unused portion.

See enclosed prescribing information for dosage and directions for use

1 x 10 to for intravenous infusion Suspension for intravenous infusion

HEMGENIX_e ețtauscodeue qezabarvovec-qup

etranacogene dezaparvovec-drlb **HEMGENIX®**

1 x 1013 genome copies/mL Suspension for intravenous infusion

Single-dose vial

Be only

Upon receipt store refrigerated at 2°C to 8°C (36°F to 46°F)

Do Not Freeze

Protect from Light

Do Not Shake

Dilute before use

Manufactured by:

uniQure, Inc. Lexington, MA 02421 USA

Manufactured for:

CSL Behring LLC

King of Prussia, PA 19406 USA

US License No. 1767

Distributed by:

CSL Behring LLC

Kankakee, IL 60901 USA

CSL Behring

Contains no preservatives.

Prot ect from Light

Discard any unused portion

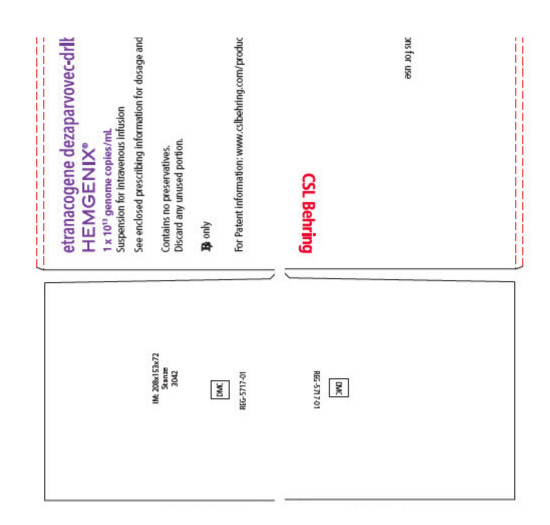
See en closed prescribing information for dosage and direction

1 x 1013 genome copies/m L No US standard of potency.

ts/patents (in-licensed from uniQure)

directions for use

1 x 10¹³ genome copies /mL Suspension for intravenous infusion etranacogene dezaparvovec-drlb HEMGENIX®



PRINCIPAL DISPLAY PANEL - NDC: 0053-0100-10_ 46-50 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\mathbb{R}}$

Patient Weight: 46-50 kg





10 mL vial x

vial x 10

PRINCIPAL DISPLAY PANEL - NDC: 0053-0110-11_ 51-55 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 51-55 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0120-12_ 56-60 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 56-60 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0130-13_ 61-65 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 61-65 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0140-14_ 66-70 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 66-70 kg



GT SN EX LO

SN: EXP:

10 mL vial x

14

PRINCIPAL DISPLAY PANEL - NDC: 0053-0150-15_ 71-75 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\mathbb{B}}$

Patient Weight: 71-75 kg





SN:

LOT:

10 mL vial x

15

PRINCIPAL DISPLAY PANEL - NDC: 0053-0160-16_ 76-80 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 76-80 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0170-17_ 81-85 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 81-85 kg





10 mL vial x

17

PRINCIPAL DISPLAY PANEL - NDC: 0053-0180-18_ 86-90 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 86-90 kg



10 mL vial x

L vial x 18

PRINCIPAL DISPLAY PANEL - NDC: 0053-0190-19_ 91-95 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 91-95 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0200-20_ 96-100 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 96-100 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0210-21_ 101-105 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 101-105 kg





x 21

10 mL vial x

PRINCIPAL DISPLAY PANEL - NDC: 0053-0220-22_ 106-110 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 106-110 kg





10 mL vial x

22

PRINCIPAL DISPLAY PANEL - NDC: 0053-0230-23_ 111-115 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\otimes}$

Patient Weight: 111-115 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0240-24_ 116-120 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 116-120 kg





10 mL vial x

24

PRINCIPAL DISPLAY PANEL - NDC: 0053-0250-25_ 121-125 KG KIT VARIABLE **LABEL**

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 121-125 kg

NDC 0053-0250-25



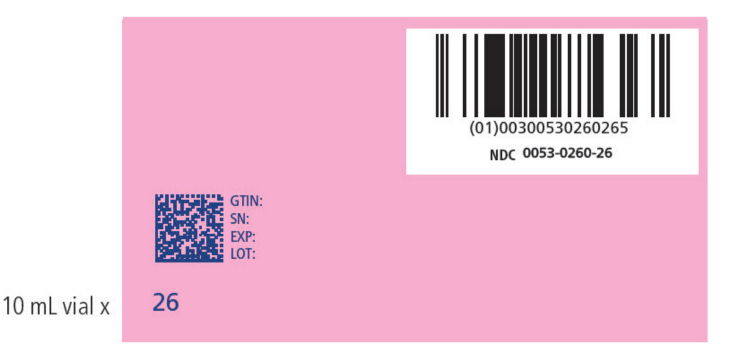
10 mL vial x

25

PRINCIPAL DISPLAY PANEL - NDC: 0053-0260-26_ 126-130 KG KIT VARIABLE **LABEL**

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\mathbb{B}}$

Patient Weight: 126-130 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0270-27_ 131-135 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 131-135 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0280-28_ 136-140 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 136-140 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0290-29_ 140-145 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 140-145 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0300-30_ 146-150 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 146-150 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0310-31_ 151-155 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 151-155 kg





10 mL vial x

31

PRINCIPAL DISPLAY PANEL - NDC: 0053-0320-32_ 156-160 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 156-160 kg





SN: EXP:

32

10 mL vial x

PRINCIPAL DISPLAY PANEL - NDC: 0053-0330-33_ 161-165 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

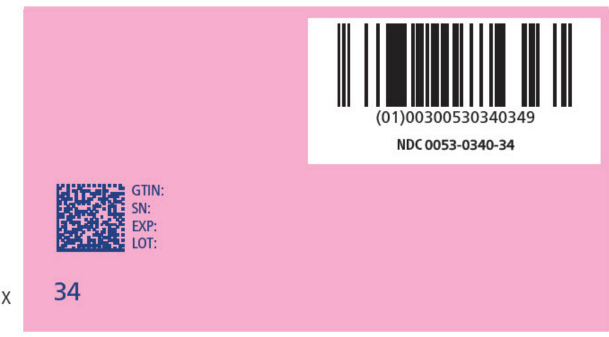
Patient Weight: 161-165 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0340-34_ 166-170 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 166-170 kg



10 mL vial x

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 171-175 kg



10 mL vial x

PRINCIPAL DISPLAY PANEL - NDC: 0053-0360-36_ 176-180 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 176-180 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0370-37_ 181-185 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 181-185 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0380-38_ 186-190 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 186-190 kg





10 mL vial x

38

PRINCIPAL DISPLAY PANEL - NDC: 0053-0390-39_ 191-195 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 191-195 kg



10 mL vial x

PRINCIPAL DISPLAY PANEL - NDC: 0053-0400-40_ 196-200 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\texttt{®}}$

Patient Weight: 196-200 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0410-41_ 201-205 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 201-205 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0420-42_ 206-210 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 206-210 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0430-43_ 211-215 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 211-215 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0440-44_ 216-220 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 216-220 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0450-45_ 221-225 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 221-225 kg





10 mL vial x

45

PRINCIPAL DISPLAY PANEL - NDC: 0053-0460-46_ 226-230 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb HEMGENIX®

Patient Weight: 226-230 kg





46

10 mL vial x

PRINCIPAL DISPLAY PANEL - NDC: 0053-0470-47_ 231-235 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 231-235 kg



PRINCIPAL DISPLAY PANEL - NDC: 0053-0480-48_ 236-240 KG KIT VARIABLE LABEL

etranacogene dezaparvovec-drlb $\mathsf{HEMGENIX}^{\$}$

Patient Weight: 236-240 kg



10 mL vial x

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0100

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
	NDC:0053-0100- 10	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022			

Quantity of Parts					
Part #	Package Quantity	Total Product Quantity			
Part 1	10 VIAL, GLASS	10			

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

 Item Code (Source)
 NDC:0053-0099

 Route of Administration
 INTRAVENOUS

Active Ingredient/Active Moiety

Active Ingredient/Active Molety				
Ingredient Name	Basis of Strength	Strength		
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}		

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0110

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0110- 11	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	11 VIAL, GLASS	11

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety Ingredient Name Basis of Strength Strength

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene dezaparvovec UNII:Z5XCD5Q9RL)

etranacogene dezaparvovec 1013 {GC}

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	

1 NDC:0053-0099- 1 in 1 VIAL, GLASS; Type 0: Not a Combination Product

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BI A	BI A125772	11/22/2022	

Marketing Information

· · · · · · · · · · · · · · · · · · ·				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		
BLA	BLAIZ3772	11/22/2022		

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0120

Packaging

# Item Code	Package Description	Marketing Start Date	Marketing End Date
1 NDC:0053-0120-	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	12 VIAL, GLASS	12

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety Ingredient Name Basis of Strength etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL) 1013 {GC}

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

HEMGENIX

etranacogene dezaparvovec kit

Product Information Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0130

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0130-	1 in 1 CARTON; Type 0: Not a Combination	11/22/2022	

Quantity of Parts					
Part #	Package Quantity	Total Product Quantity			
Part 1	13 VIAL, GLASS	13			

Part 1 of 1 HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source) NDC:0053-0099

Route of Administration INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene - UNII:Z5XCD5Q9RL)

dezaparvovec

1013 {GC}

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
_	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

Marketing Information

Marketing information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0140

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π	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0140- 14	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts					
Part #	Package Quantity	Total Product Quantity			
Part 1	14 VIAL GLASS	14			

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information			
Item Code (Source)	NDC:0053-0099		
Route of Administration	INTRAVENOUS		

l	Active Ingredient/Active Moiety					
l	Ingredient Name	Basis of Strength	Strength			
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}			

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0053-0150
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Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0150- 15	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quant	Quantity of Parts					
Part #	Package Quantity	Total Product Quantity				
Part 1	15 VIAL, GLASS	15				

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information			
Item Code (Source)	NDC:0053-0099		
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvove - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}			

l	P	Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
	1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information			
Marketing	Application Number or Monograph	Marketing Start	Marketing End

Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0160

Packaging

l		· uchaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
		NDC:0053-0160- 16	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022		

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	16 VIAL, GLASS	16

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}

ı	P	Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
	1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product				

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0170

F	Packaging				
4	# Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0170-	1 in 1 CARTON; Type 0: Not a Combination	11/22/2022		

Quant	Quantity of Parts		
Part #	Package Quantity	Total Product Quantity	
Part 1	17 VIAL, GLASS	17	

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0180

Packaging # Item Code Package Description Marketing Start Date Marketing End Date 1 NDC:0053-0180-18 1 in 1 CARTON; Type 0: Not a Combination Product 11/22/2022

Quanti	Quantity of Parts				
Part #	Package Quantity	Total Product Quantity			
Part 1 18 VIAL, GLASS 1		18			

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source) NDC:0053-0099

INTRAVENOUS Route of Administration

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene - UNII:Z5XCD5Q9RL)

dezaparvovec

1013 {GC}

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

Marketing Information

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Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

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etranacogene dezaparvovec kit

Product Information

HUMAN PRESCRIPTION DRUG NDC:0053-0190 **Product Type Item Code (Source)**

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0190- 19	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	19 VIAL, GLASS	19

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)NDC:0053-0099Route of AdministrationINTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name
Basis of Strength
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec
- UNII:Z5XCD5Q9RL)

Basis of Strength
etranacogene
dezaparvovec

etranacogene
dezaparvovec

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information					
Marketing Application Number or Monograph Marke Category Citation I			Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

HEMGENIX

etranacogene dezaparvovec kit

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Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0200

Packaging

	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
l		NDC:0053-0200- 20	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts				
Part #	Package Quantity	Total Product Quantity		
Part 1	20 VIAL, GLASS	20		

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information			
Item Code (Source)	NDC:0053-0099		
Route of Administration	INTRAVENOUS		

Active Ingredient/Active Moiety				
Ingred	dient Name	Basis of Strength	Strength	
etranacogene dezaparvovec (UNII: - UNII:Z5XCD5Q9RL)	Z5XCD5Q9RL) (etranacogene dezaparvovec	etranacogene dezaparvovec	1013 {GC}	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Application Number or Monograph Marketing Start Marketing En Category Citation Date Date				
BLA	BLA125772	11/22/2022		

Marketing Information				
Marketing Application Number or Monograph Marketing Start Marketing Category Citation Date				
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0210

F	Packaging					
#	tem Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:0053-0210-	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022			

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	21 VIAL, GLASS	21

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety Ingredient Name Basis of Strength etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL) Basis of Strength etranacogene dezaparvovec 1013 {GC}

P	Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0220

l	P	Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
	1	NDC:0053-0220- 22	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022			

Quantity of Parts						
Part #	Package Quantity	Total Product Quantity				
Part 1	22 VIAL, GLASS	22				

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength Strengt			
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}		

Packaging		

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing In	iformation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information				
Marketing Application Number or Monograph Category Citation		Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0230

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0230-	1 in 1 CARTON; Type 0: Not a Combination	11/22/2022		

Quantity of Parts

Part #	Package Quantity	Total Product Quantity	
Part 1	23 VIAL, GLASS	23	

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII: Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}			

P	Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0240

Packaging			
# Item Code	Package Description	Marketing Start Date	Marketing End Date
NDC:0053-0240- 24	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quant	Quantity of Parts					
Part # Package Quantity		Total Product Quantity				
Part 1	24 VIAL, GLASS	24				

Part 1 of 1

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source) NDC:0053-0099 **Route of Administration INTRAVENOUS**

Active Ingredient/Active Moiety

Ingredient Name Basis of Strength Strength etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene 1013 {GC} - UNII:Z5XCD5Q9RL) dezaparvovec

P	Packaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Product Information

HUMAN PRESCRIPTION DRUG NDC:0053-0250 **Product Type Item Code (Source)**

P	Packaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0250- 25	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity

25

Part 1 25 VIAL, GLASS

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source) NDC:0053-0099

Route of Administration INTRAVENOUS

Active Ingredient/Active Moiety

l	Dania of Chuomath	Churanantha
Ingredient Name	Basis of Strength	Strength

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene - UNII:Z5XCD5Q9RL) dezaparvovec

1013 {GC}

Packaging

ı				
	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:0053-0099-	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
RIΔ	ΒΙ Δ125772	11/22/2022	

Marketing Information

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Product	Informat	ion
PIUUULL	IIII OI III a	.IUI

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0260

l	Packaging			
	# Item Code	Package Description	Marketing Start Date	Marketing End Date
П				

1 NDC:0053-0260- 1 in 1 CARTON; Type 0: Not a Combination Product 11/22/2022

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	26 VIAL, GLASS	26

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

l	Active Ingredient/Active Moiety			
l	Ingredient Name	Basis of Strength	Strength	
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0270

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0270- 27	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	27 VIAL, GLASS	27

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety Ingredient Name Basis of Strength etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL) tranacogene dezaparvovec

Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information			
Marketing Application Number or Monograph Category Citation		Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0280

l	Packaging				
	# Item Code	Package Description	Marketing Start Date	Marketing End Date	
	1 NDC:0053-0280-	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022		

Quantity of Parts

Part # Package Quantity Total Product Quantity
Part 1 28 VIAL, GLASS 28

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information Item Code (Source) NDC:0053-0099 Route of Administration INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec	etranacogene	1013 {GC}
- UNII:Z5XCD5Q9RL)	dezaparvovec	1013 (00)

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0290

Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0290- 29	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts					
Part #	Package Quantity	Total Product Quantity			
Part 1	29 VIAL, GLASS	29			

Part 1 of 1

HEMGENIX

Product Information

Item Code (Source) NDC:0053-0099

Route of Administration INTRAVENOUS

Active Ingredient/Active Moiety

Active Ingredient/Active Molety			
Ingredient Name	Basis of Strength	Strength	
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0300

ı	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0300- 30	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	30 VIAL, GLASS	30

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

 Item Code (Source)
 NDC:0053-0099

 Route of Administration
 INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name
Basis of Strength
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec
- UNII:Z5XCD5Q9RL)

Basis of Strength
etranacogene
dezaparvovec

1013 {GC}

Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product				

Marketing In	formation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing I	nformation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Prod	luct	Inform	ation

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0310

P	Packaging						
#	Item Code	Package Description	Marketing Start Date	Marketing End Date			
1	NDC:0053-0310- 31	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022				

Quant	ity of Parts	
Part #	Package Quantity	Total Product Quantity
Part 1	31 VIAL, GLASS	31

HEMGENIX

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety					
Ingredient Name	Basis of Strength	Strength			
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}			

Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Ir	nformation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing In	nformation		
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0320

l	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:0053-0320- 32	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	32 VIAL, GLASS	32

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

l	Active Ingredient/Active Moiety			
l	Ingredient Name	Basis of Strength Stren		
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0330

ı	Packaging			
7	# Item Code	Package Description	Marketing Start Date	Marketing End Date
:	NDC:0053-0330-	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts					
Part #	Package Quantity	Total Product Quantity			
Part 1	33 VIAL. GLASS	33			

Part 1 of 1

HEMGENIX

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category		Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
	BLA	BLA125772	11/22/2022	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0340

l	P	Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date		
		NDC:0053-0340- 34	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022			

Quantity of Parts Part # Package Quantity Total Product Quantity Part 1 34 VIAL, GLASS 34

Part 1 of 1

HEMGENIX

Product Information	
Item Code (Source)	NDC:0053-0099

Rout	- വ	· 24	minis	strati	On.

INTRAVENOUS

Active Ingredient/Active Moiety

Basis of Strength Strength

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene - UNII:Z5XCD5Q9RL)

dezaparvovec

1013 {GC}

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

Marketing Information

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0350

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0350- 35	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

ı	Part #	Package Quantity	Total Product Quantity
п			

Part 1 35 VIAL, GLASS

35

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source) NDC:0053-0099

Route of Administration INTRAVENOUS

Active Ingredient/Active Moiety

Product

Basis of Strength Strength Ingredient Name

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene - UNII:Z5XCD5Q9RL)

dezaparvovec

1013 {GC}

ı	Packaging					
	# Item Code	Package Description	Marketing Start Date	Marketing End Date		
	NDC:0053-0099-	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product				

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
BLA	BLA125772	11/22/2022				

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0360

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Marketing Start Marketing End Package Description Item Code Date Date

		NDC:0053-0360- 36	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	
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Quantity of Parts				
Part #	Package Quantity	Total Product Quantity		
Part 1	36 VIAL, GLASS	36		

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information		
Item Code (Source)	NDC:0053-0099	
Route of Administration	INTRAVENOUS	

l	Active Ingredient/Active Moiety				
l	Ingredient Name	Basis of Strength	Strength		
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

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etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0370

ı	P	ackaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:0053-0370- 37	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	37 VIAL, GLASS	37

Part 1 of 1

HEMGENIX

Product Information			
Item Code (Source)	NDC:0053-0099		
Route of Administration	INTRAVENOUS		

l	Active Ingredient/Active Moiety			
l	Ingredient Name	Basis of Strength	Strength	
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0380

Packaging				
#	tem Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0380-	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts					
Part #	Package Quantity	Total Product Quantity			
Part 1	38 VIAL, GLASS	38			

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information Item Code (Source) NDC:0053-0099 Route of Administration INTRAVENOUS

I	Active Ingredient/Active Moiety				
l	Ingredient Name	Basis of Strength	Strength		
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}		

F	Packaging			
#	tem Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0099-	1 in 1 VIAL, GLASS; Type 0: Not a Combination		

Marketing	Application Number or Monograph	Marketing Start	Marketing End
Category	Citation	Date	Date
BLA	BLA125772	11/22/2022	

Marketing Information

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Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0390

Packaging

Ш	• '	ackaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0390- 39	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	39 VIAL, GLASS	39

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information					
Marketing Application Number or Monograph Marketing Start Category Citation Date			Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0400

 Packaging

 #
 Item Code
 Package Description
 Marketing Start Date
 Marketing End Date

 1
 NDC:0053-0400-40
 1 in 1 CARTON; Type 0: Not a Combination Product
 11/22/2022

Quant	Quantity of Parts				
Part #	art # Package Quantity Total Product Quantity				
Part 1	40 VIAL, GLASS	40			

Part 1 of 1

HEMGENIX

Product Information		
Item Code (Source)	NDC:0053-0099	
Route of Administration	INTRAVENOUS	

Active Ingredient/Active Moiety				
Ingredient Name	Basis of Strength	Strength		
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}		

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information					
Marketing Category	Marketing End Date				
BLA	BLA125772	11/22/2022			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0053-0410

F	Packaging			
#	tem Code	Package Description	Marketing Start Date	Marketing End Date
3	NDC:0053-0410- 41	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	41 VIAL, GLASS	41

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etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety

	Ingredient Name	Basis of Strength	Strength
l	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}

l	Packaging			
	# Item Code	Package Description	Marketing Start Date	Marketing End Date
	1 NDC:0053-0099-	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0053-0420

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0420- 42	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts		
Part #	Package Quantity	Total Product Quantity
Part 1	42 VIAL, GLASS	42

HEMGENIX

Product Information	
Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

	Active Ingredient/Active Moiety				
l	Ingredient Name	Basis of Strength	Strength		
	$\begin{tabular}{ll} \textbf{etranacogene dezaparvovec} & \textbf{(UNII: Z5XCD5Q9RL)} & \textbf{(etranacogene dezaparvovec - UNII: Z5XCD5Q9RL)} \\ \end{tabular}$	etranacogene dezaparvovec	1013 {GC}		

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing In			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
BLA	BLA125772	11/22/2022		

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0430

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0430- 43	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Part #	Package Quantity	Total Product Quantity
Part 1	43 VIAL, GLASS	43

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}

Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing In			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	
BLA	BLAIZSTTZ	11/22/2022	

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0440

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	NDC:0053-0440- 44	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts

Qualit	Quantity of Farts					
Part #	Package Quantity	Total Product Quantity				
Part 1	44 VIAL, GLASS	44				

Part 1 of 1

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source)	NDC:0053-0099
Route of Administration	INTRAVENOUS

Active Ingredient/Active Moiety				
	Ingredient Name	Basis of Strength	Strength	
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

Packaging					
#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information						
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date			
BLA	BLA125772	11/22/2022				

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0450

l	Packaging					
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date	
		NDC:0053-0450- 45	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022		

Quantity of Parts Part # Package Quantity Total Product Quantity Part 1 45 VIAL, GLASS 45

Part 1 of 1

HEMGENIX

Product Information	
Item Code (Source)	NDC:0053-0099

Route			

INTRAVENOUS

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
(1011) 75/(0050001) / 1		

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene - UNII:Z5XCD5Q9RL)

dezaparvovec

1013 {GC}

l	Packaging				
	# Item Code	Package Description	Marketing Start Date	Marketing End Date	
	1 NDC:0053-0099-	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product			

Marketing Information					
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date		
BLA	BLA125772	11/22/2022			

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0460

l	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:0053-0460- 46	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts			
Part #	Package Quantity	Total Product Quantity	
Part 1	46 VIAL, GLASS	46	

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information

Item Code (Source) NDC:0053-0099

Route of Administration INTRAVENOUS

Active Ingredient/Active Moiety

Product

Ingredient Name Basis of Strength Strength

etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec etranacogene dezaparvovec UNII:Z5XCD5Q9RL)

dezaparvovec 1013 {GC}

Item Code Package Description Marketing Start Date Marketing End Date

NDC:0053-0099- 1 in 1 VIAL, GLASS; Type 0: Not a Combination

Marketing InformationMarketing CategoryApplication Number or Monograph CitationMarketing Start DateMarketing End DateBLABLA12577211/22/2022

Marketing InformationMarketing CategoryApplication Number or Monograph CitationMarketing Start DateMarketing End DateBLABLA12577211/22/2022

HEMGENIX

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etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0470

Packaging

Item Code Package Description Marketing Start Date Marketing End

Quantity of Parts		
Part #	Package Quantity	Total Product Quantity
Part 1	47 VIAL, GLASS	47

HEMGENIX

etranacogene dezaparvovec injection, suspension

Product Information		
Item Code (Source)	NDC:0053-0099	
Route of Administration	INTRAVENOUS	

	Active Ingredient/Active Moiety			
ı	Ingredient Name	Basis of Strength	Strength	
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

l	Packaging				
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
		NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

HEMGENIX

etranacogene dezaparvovec kit

Product Information

Product Type HUMAN PRESCRIPTION DRUG Item Code (Source) NDC:0053-0480

l	P	Packaging			
	#	Item Code	Package Description	Marketing Start Date	Marketing End Date
	1	NDC:0053-0480- 48	1 in 1 CARTON; Type 0: Not a Combination Product	11/22/2022	

Quantity of Parts		
Part #	Package Quantity	Total Product Quantity
Part 1	48 VIAL, GLASS	48

Part 1 of 1

HEMGENIX

Product Information		
Item Code (Source)	NDC:0053-0099	
Route of Administration	INTRAVENOUS	

l	Active Ingredient/Active Moiety			
l	Ingredient Name	Basis of Strength	Strength	
	etranacogene dezaparvovec (UNII: Z5XCD5Q9RL) (etranacogene dezaparvovec - UNII:Z5XCD5Q9RL)	etranacogene dezaparvovec	1013 {GC}	

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0053-0099- 01	1 in 1 VIAL, GLASS; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
BLA	BLA125772	11/22/2022	

Labeler - CSL Behring (058268293)

Establishment			
Name	Address	ID/FEI	Business Operations
CSL Behring LLC		058268293	MANUFACTURE(0053-0100, 0053-0110, 0053-0120, 0053-0130, 0053-0140, 0053-0150, 0053-0160, 0053-0170, 0053-0180, 0053-0190, 0053-0200, 0053-0210, 0053-0220, 0053-0230, 0053-0240, 0053-0250, 0053-0260, 0053-0270, 0053-0280, 0053-0290, 0053-0300, 0053-0310, 0053-0320, 0053-0330, 0053-0340, 0053-0350, 0053-0360, 0053-0370, 0053-0380, 0053-0390, 0053-0400, 0053-0410, 0053-0420, 0053-0430, 0053-0440, 0053-0450, 0053-0460, 0053-0470, 0053-0480)

Revised: 5/2023 CSL Behring