MOZOBIL- plerixafor injection, solution sanofi-aventis U.S. LLC

HIGHLIGHTS	OF	PRESCRIBING	INFORMATION
------------	----	-------------	-------------

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

MOZOBIL® (plerixafor) injection, for subcutaneous	s use
Initial U.S. Approval: 2008	

------INDICATIONS AND USAGE

Mozobil, a hematopoietic stem cell mobilizer, is indicated in combination with filgrastim to mobilize hematopoietic stem cells (HSCs) to the peripheral blood for collection and subsequent autologous transplantation in patients with non-Hodgkin's lymphoma or multiple myeloma. (1)

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

- Initiate Mozobil treatment after the patient has received filgrastim once daily for 4 days. (2.1)
- Repeat Mozobil dose up to 4 consecutive days. (2.1)
- Dose based on patient weight
 - Less than or equal to 83 kg: 20 mg dose or select dose based on 0.24 mg/kg actual body weight.
 (2.1)
 - greater than 83 kg: select dose based on 0.24 mg/kg actual body weight. (2.1)
- Administer by subcutaneous injection approximately 11 hours prior to initiation of apheresis. (2.1)
- Renal impairment: If creatinine clearance is ≤50 mL/min, decrease dose by one-third to 0.16 mg/kg.
 (2.3)

----- DOSAGE FORMS AND STRENGTHS

• Injection: 24 mg/1.2 mL (20 mg/mL) in a single-dose vial. (3)

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

• History of hypersensitivity to Mozobil. (4)

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

- Anaphylactic Shock and Serious Hypersensitivity Reactions have occurred. Monitor patients during and after completion of Mozobil administration. (5.1)
- Tumor Cell Mobilization in Leukemia Patients: Mozobil may mobilize leukemic cells and should not be used in leukemia patients. (5.2)
- Hematologic Effects: Increased circulating leukocytes and decreased platelet counts have been observed. Monitor blood cell counts and platelet counts during Mozobil use. (5.3)
- Potential for Tumor Cell Mobilization: Tumor cells may be released from marrow during HSC mobilization with Mozobil and filgrastim. Effect of reinfusion of tumor cells is unknown. (5.4)
- Splenic Rupture: Evaluate patients who report left upper abdominal and/or scapular or shoulder pain. (5.5)
- Embryo-Fetal Toxicity: Can cause fetal harm. Advise women not to become pregnant when taking Mozobil. (5.6, 8.1)

..... ADVERSE REACTIONS

Most common adverse reactions (≥10%): diarrhea, nausea, fatigue, injection site reactions, headache, arthralgia, dizziness, and vomiting. (6)

To report SUSPECTED ADVERSE REACTIONS, contact Genzyme Corporation at 1-877-4MOZOBIL or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

------USE IN SPECIFIC POPULATIONS

Lactation: Advise not to breastfeed. (8.2)

See 17 for PATIENT COUNSELING INFORMATION.

Revised: 9/2023

FULL PRESCRIBING INFORMATION: CONTENTS*

- 1 INDICATIONS AND USAGE
- 2 DOSAGE AND ADMINISTRATION
 - 2.1 Recommended Dosage and Administration
 - 2.2 Recommended Concomitant Medications
 - 2.3 Dose Modifications in Renal Impairment
- **3 DOSAGE FORMS AND STRENGTHS**
- **4 CONTRAINDICATIONS**
- 5 WARNINGS AND PRECAUTIONS
 - 5.1 Anaphylactic Shock and Hypersensitivity Reactions
 - 5.2 Tumor Cell Mobilization in Leukemia Patients
 - 5.3 Hematologic Effects
 - 5.4 Potential for Tumor Cell Mobilization
 - 5.5 Splenic Enlargement and Rupture
 - 5.6 Embryo-Fetal Toxicity

6 ADVERSE REACTIONS

- 6.1 Clinical Trials Experience
- 6.2 Postmarketing Experience

8 USE IN SPECIFIC POPULATIONS

- 8.1 Pregnancy
- 8.2 Lactation
- 8.3 Females and Males of Reproductive Potential
- 8.4 Pediatric Use
- 8.5 Geriatric Use
- 8.6 Renal Impairment

10 OVERDOSAGE

11 DESCRIPTION

12 CLINICAL PHARMACOLOGY

- 12.1 Mechanism of Action
- 12.2 Pharmacodynamics
- 12.3 Pharmacokinetics

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

14 CLINICAL STUDIES

16 HOW SUPPLIED/STORAGE AND HANDLING

17 PATIENT COUNSELING INFORMATION

* Sections or subsections omitted from the full prescribing information are not listed.

FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

Mozobil is indicated in combination with filgrastim to mobilize hematopoietic stem cells (HSCs) to the peripheral blood for collection and subsequent autologous transplantation

in patients with non-Hodgkin's lymphoma (NHL) or multiple myeloma (MM).

2 DOSAGE AND ADMINISTRATION

2.1 Recommended Dosage and Administration

Begin treatment with Mozobil after the patient has received filgrastim once daily for 4 days [see Dosage and Administration (2.2)]. Administer Mozobil approximately 11 hours prior to initiation of each apheresis for up to 4 consecutive days.

The recommended dose of Mozobil by subcutaneous injection is based on body weight:

- 20 mg fixed dose or 0.24 mg/kg of body weight for patients weighing less than or equal to 83 kg. [see Clinical Pharmacology (12.3)]
- 0.24 mg/kg of body weight for patients weighing greater than 83 kg

Use the patient's actual body weight to calculate the volume of Mozobil to be administered. Each vial delivers 1.2 mL of 20 mg/mL solution, and the volume to be administered to patients should be calculated from the following equation:

 $0.012 \times \text{patient's actual body weight (in kg)} = \text{volume to be administered (in mL)}$

In clinical studies, Mozobil dose has been calculated based on actual body weight in patients up to 175% of ideal body weight. Mozobil dose and treatment of patients weighing more than 175% of ideal body weight have not been investigated.

Based on increasing exposure with increasing body weight, the Mozobil dose should not exceed 40 mg/day [see Clinical Pharmacology (12.3)].

Vials should be inspected visually for particulate matter and discoloration prior to administration and should not be used if there is particulate matter or if the solution is discolored.

Discard unused portion.

2.2 Recommended Concomitant Medications

Administer daily morning doses of filgrastim 10 mcg/kg for 4 days prior to the first evening dose of Mozobil and on each day prior to apheresis [see Clinical Studies (14)].

2.3 Dose Modifications in Renal Impairment

In patients with moderate and severe renal impairment (estimated creatinine clearance (CL_{CR}) less than or equal to 50 mL/min), reduce the dose of Mozobil by one-third based on body weight category as shown in Table 1. If CL_{CR} is less than or equal to 50 mL/min the dose should not exceed 27 mg/day, as the mg/kg-based dosage results in increased plerixafor exposure with increasing body weight [see Clinical Pharmacology (12.3)]. Similar systemic exposure is predicted if the dose is reduced by one-third in patients with moderate and severe renal impairment compared with subjects with normal renal function [see Clinical Pharmacology (12.3)].

Table 1: Recommended Dosage of Mozobil in Patients with Renal Impairment

Estimated Creatinine Clearance (mL/min)	Dose		
	Body Weight less than or equal to 83 kg	Body Weight greater than 83 kg and less than 160 kg	
greater than 50	20 mg or 0.24 mg/kg once daily	0.24 mg/kg once daily (not to exceed 40 mg/day)	
less than or equal to 50	13 mg or 0.16 mg/kg once daily	0.16 mg/kg once daily (not to exceed 27 mg/day)	

The following (Cockcroft-Gault) formula may be used to estimate CL_{CR}:

Males:

Creatinine clearance (mL/min) = $\frac{\text{weight (kg)} \times (140 - \text{age in years)}}{72 \times \text{serum creatinine (mg/dL)}}$

Females:

Creatinine clearance (mL/min) = $0.85 \times \text{value}$ calculated for males

There is insufficient information to make dosage recommendations in patients on hemodialysis.

3 DOSAGE FORMS AND STRENGTHS

Injection: 24 mg/1.2 mL (20 mg/mL) sterile, clear, colorless to pale-yellow solution in a single-dose vial.

4 CONTRAINDICATIONS

Mozobil is contraindicated in patients with a history of hypersensitivity to plerixafor [see Warnings and Precautions (5.1)]. Anaphylactic shock has occurred with use of Mozobil.

5 WARNINGS AND PRECAUTIONS

5.1 Anaphylactic Shock and Hypersensitivity Reactions

Serious hypersensitivity reactions, including anaphylactic-type reactions, some of which have been life-threatening with clinically significant hypotension and shock have occurred in patients receiving Mozobil [see Adverse Reactions (6.2)]. Observe patients for signs and symptoms of hypersensitivity during and after Mozobil administration for at least 30 minutes and until clinically stable following completion of each administration. Only administer Mozobil when personnel and therapies are immediately available for the treatment of anaphylaxis and other hypersensitivity reactions.

In clinical studies, mild or moderate allergic reactions occurred within approximately 30

minutes after Mozobil administration in less than 1% of patients [see Adverse Reactions (6.1)].

5.2 Tumor Cell Mobilization in Leukemia Patients

For the purpose of HSC mobilization, Mozobil may cause mobilization of leukemic cells and subsequent contamination of the apheresis product. Therefore, Mozobil is not intended for HSC mobilization and harvest in patients with leukemia.

5.3 Hematologic Effects

<u>Leukocytosis</u>

Administration of Mozobil in conjunction with filgrastim increases circulating leukocytes as well as HSC populations. Monitor white blood cell counts during Mozobil use [see Adverse Reactions (6.1)].

<u>Thrombocytopenia</u>

Thrombocytopenia has been observed in patients receiving Mozobil. Monitor platelet counts in all patients who receive Mozobil and then undergo apheresis.

5.4 Potential for Tumor Cell Mobilization

When Mozobil is used in combination with filgrastim for HSC mobilization, tumor cells may be released from the marrow and subsequently collected in the leukapheresis product. The effect of potential reinfusion of tumor cells has not been well-studied.

5.5 Splenic Enlargement and Rupture

Higher absolute and relative spleen weights associated with extramedullary hematopoiesis were observed following prolonged (2 to 4 weeks) daily plerixafor SC administration in rats at doses approximately 4-fold higher than the recommended human dose based on body surface area. The effect of Mozobil on spleen size in patients was not specifically evaluated in clinical studies. Cases of splenic enlargement and/or rupture have been reported following the administration of Mozobil in conjunction with filgrastim. Evaluate individuals receiving Mozobil in combination with filgrastim who report left upper abdominal pain and/or scapular or shoulder pain for splenic integrity.

5.6 Embryo-Fetal Toxicity

Based on findings from animal reproduction studies, Mozobil can cause fetal harm when administered to a pregnant woman. Plerixafor administration to pregnant rats during organogenesis resulted in embryo-fetal mortality, structural abnormalities, and alterations to growth at exposures approximately 10 times the exposure at the recommended human dose.

Advise pregnant women of the potential risk to the fetus. Advise females of reproductive potential to use an effective form of contraception during treatment with Mozobil and for one week after the final dose [see Use in Specific Populations (8.1)].

6 ADVERSE REACTIONS

The following clinically significant adverse reactions are discussed elsewhere in the

labeling:

- Anaphylactic shock and hypersensitivity reactions [see Warnings and Precautions (5.1)]
- Potential for tumor cell mobilization in leukemia patients [see Warnings and Precautions (5.2)]
- Increased circulating leukocytes and decreased platelet counts [see Warnings and Precautions (5.3)]
- Potential for tumor cell mobilization [see Warnings and Precautions (5.4)]
- Splenic enlargement [see Warnings and Precautions (5.5)]

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 practice.

The most common adverse reactions (≥10%) reported in patients who received Mozobil in conjunction with filgrastim regardless of causality and more frequent with Mozobil than placebo during HSC mobilization and apheresis were diarrhea, nausea, fatigue, injection site reactions, headache, arthralgia, dizziness, and vomiting.

Safety data for Mozobil in combination with filgrastim were obtained from two randomized placebo-controlled studies (301 patients) and 10 uncontrolled studies (242 patients). Patients were primarily treated with Mozobil at daily doses of 0.24 mg/kg SC. Median exposure to Mozobil in these studies was 2 days (range 1 to 7 days).

In the two randomized studies in patients with NHL and MM, a total of 301 patients were treated in the Mozobil and filgrastim group and 292 patients were treated in the placebo and filgrastim group. Patients received daily morning doses of filgrastim 10 mcg/kg for 4 days prior to the first dose of Mozobil 0.24 mg/kg SC or placebo and on each morning prior to apheresis. The adverse reactions that occurred in \geq 5% of the patients who received Mozobil regardless of causality and were more frequent with Mozobil than placebo during HSC mobilization and apheresis are shown in Table 2.

Table 2: Adverse Reactions in ≥5% of Non-Hodgkin's Lymphoma and Multiple Myeloma Patients Receiving Mozobil and More Frequent than Placebo during HSC Mobilization and Apheresis

		Percent of Patients (%)				
	Mozobi	Mozobil and Filgrastim (n=301)			and Filo (n=292)	
	All Grades*	Grade 3	Grade 4	All Grades	Grade 3	Grade 4
Gastrointestinal disorders						
Diarrhea	37	<1	0	17	0	0
Nausea	34	1	0	22	0	0
Vomiting	10	<1	0	6	0	0
Flatulence	7	0	0	3	0	0

General disorders and administration site conditions						
Injection site reactions	34	0	0	10	0	0
Fatigue	27	0	0	25	0	0
Musculoskeletal and connective tissue disorders						
Arthralgia	13	0	0	12	0	0
Nervous system disorders						
Headache	22	<1	0	21	1	0
Dizziness	11	0	0	6	0	0
Psychiatric disorders						
Insomnia	7	0	0	5	0	0

^{*} Grades based on criteria from the World Health Organization (WHO)

In the randomized studies, 34% of patients with NHL or MM had mild to moderate injection site reactions at the site of subcutaneous administration of Mozobil. These included erythema, hematoma, hemorrhage, induration, inflammation, irritation, pain, paresthesia, pruritus, rash, swelling, and urticaria.

Mild to moderate allergic reactions were observed in less than 1% of patients within approximately 30 min after Mozobil administration, including one or more of the following: urticaria (n=2), periorbital swelling (n=2), dyspnea (n=1) or hypoxia (n=1). Symptoms generally responded to treatments (e.g., antihistamines, corticosteroids, hydration or supplemental oxygen) or resolved spontaneously.

Vasovagal reactions, orthostatic hypotension, and/or syncope can occur following subcutaneous injections. In Mozobil oncology and healthy volunteer clinical studies, less than 1% of subjects experienced vasovagal reactions following subcutaneous administration of Mozobil doses ≤ 0.24 mg/kg. The majority of these events occurred within 1 hour of Mozobil administration. Because of the potential for these reactions, appropriate precautions should be taken.

Other adverse reactions in the randomized studies that occurred in <5% of patients but were reported as related to Mozobil during HSC mobilization and apheresis included abdominal pain, hyperhidrosis, abdominal distention, dry mouth, erythema, stomach discomfort, malaise, hypoesthesia oral, constipation, dyspepsia, and musculoskeletal pain.

Hyperleukocytosis: In clinical trials, white blood cell counts of 100,000/mcL or greater were observed, on the day prior to or any day of apheresis, in 7% of patients receiving Mozobil and in 1% of patients receiving placebo. No complications or clinical symptoms of leukostasis were observed.

6.2 Postmarketing Experience

In addition to adverse reactions reported from clinical trials, the following adverse

reactions have been reported from postmarketing experience with Mozobil. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

Blood and Lymphatic System: Splenomegaly and splenic rupture

Immune System Disorders: Anaphylactic reactions, including anaphylactic shock

Psychiatric Disorders: Abnormal dreams and nightmares

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

Limited available data with Mozobil use in pregnant women are insufficient to inform a drug-associated risk of adverse developmental outcomes. In animal reproduction studies, subcutaneous administration of plerixafor to pregnant rats during organogenesis at doses 10-times the maximum recommended human doses resulted in embryo-fetal mortality, structural abnormalities, and alterations to growth [see Data].

Advise pregnant women of the potential risk to the fetus.

Advise women of reproductive potential to avoid becoming pregnant while receiving treatment with Mozobil [see Warnings and Precautions (5.6)].

In the U.S. 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. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. The background risk of major birth defects and miscarriages for the indicated population is unknown.

Data

Animal data

Plerixafor administered to pregnant rats induced embryo-fetal toxicity, including fetal death, increased resorptions and postimplantation loss, decreased fetal weights, anophthalmia, shortened digits, cardiac interventricular septal defect, ringed aorta, globular heart, hydrocephaly, dilatation of olfactory ventricles, and retarded skeletal development. Embryo-fetal toxicities occurred mainly at a dose of 90 mg/m² (approximately 10 times the recommended human dose of 0.24 mg/kg when compared on a mg/m² basis).

8.2 Lactation

Risk Summary

There are no data on the presence of plerixafor in human milk, the effect on the breastfed child, or the effect on milk production. Because of the potential serious adverse reactions in the breastfed child, advise females that breastfeeding is not recommended during treatment with Mozobil and for one week after the final dose.

8.3 Females and Males of Reproductive Potential

Pregnancy Testing

Verify pregnancy status in females of reproductive potential prior to initiating Mozobil [see Use in Specific Populations (8.1)].

Contraception

Females

Mozobil can cause embryo-fetal harm when administered to pregnant women [see Use in Specific Populations (8.1)]. Advise females of reproductive potential to use effective contraception during treatment with Mozobil and for one week after the final dose.

Males

Males treated with Mozobil should use effective contraception during treatment and for one week after cessation of treatment.

8.4 Pediatric Use

The safety and effectiveness of Mozobil have not been established in pediatric patients. Effectiveness was not demonstrated in a single phase 1/2 randomized, open-label, comparative study of Mozobil plus standard regimens for mobilization of hematopoietic stem cells to the peripheral blood for collection and subsequent autologous transplantation. Forty-five pediatric patients ages 1 to <17 years with solid tumors or lymphoma were randomized 2:1 to Mozobil in addition to standard mobilization regimens (N=30) or standard mobilization regimens alone (N=15) (comparator arm). No new safety signals were observed in pediatric patients in this trial.

The day prior to the first apheresis, median peripheral blood CD34+ counts were 35 \times 10^6 cells/L in the comparator arm and 15 \times 10^6 cells/L in the Mozobil arm. On the day of the first apheresis, median peripheral blood CD34+ counts were 64 \times 10^6 cells/L in the comparator arm and 77 \times 10^6 cells/L in the Mozobil arm.

Plerixafor exposure, shown as AUC, in pediatric patients aged 12 to 18 years was within the range of values previously observed in adults, while the AUC of plerixafor in pediatric patients aged 2 to 12 years was 67% to 87% of that observed in adults, given the same adult dose (0.24 mg/kg).

8.5 Geriatric Use

Of the total number of subjects in controlled clinical studies of Mozobil, 24% were 65 and over, while 0.8% were 75 and over. No overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients, but greater sensitivity of some older individuals cannot be ruled out.

Since plerixafor is mainly excreted by the kidney, no dose modifications are necessary in elderly individuals with normal renal function. In general, care should be taken in dose selection for elderly patients due to the greater frequency of decreased renal function with advanced age. Dosage adjustment in elderly patients with CL_{CR} less than or equal to 50 mL/min is recommended [see Dosage and Administration (2.3) and Clinical Pharmacology (12.3)].

8.6 Renal Impairment

In patients with moderate and severe renal impairment (CL_{CR} less than or equal to 50 mL/min), reduce the dose of Mozobil by one-third to 0.16 mg/kg [see Dosage and Administration (2.3) and Clinical Pharmacology (12.3)].

10 OVERDOSAGE

Based on limited data at doses above the recommended dose of 0.24 mg/kg SC, the frequency of gastrointestinal disorders, vasovagal reactions, orthostatic hypotension, and/or syncope may be higher.

11 DESCRIPTION

Mozobil (plerixafor) injection is a sterile, preservative-free, clear, colorless to pale-yellow, isotonic solution for subcutaneous injection. Each mL of the sterile solution contains 20 mg of plerixafor. Each single-dose vial is filled to deliver 1.2 mL of the sterile solution that contains 24 mg of plerixafor and 5.9 mg of sodium chloride in Water for Injection adjusted to a pH of 6.0 to 7.5 with hydrochloric acid and with sodium hydroxide, if required.

Plerixafor is a hematopoietic stem cell mobilizer with a chemical name 1,4-Bis((1,4,8,11-tetraazacyclotetradecan-1-yl)methyl)benzene. It has the molecular formula $C_{28}H_{54}N_8$. The molecular weight of plerixafor is 502.79 g/mol. The structural formula is provided in Figure 1.

Figure 1: Structural Formula

Plerixafor is a white to off-white crystalline solid. It is hygroscopic. Plerixafor has a typical melting point of 131.5° C. The partition coefficient of plerixafor between 1-octanol and pH 7 aqueous buffer is <0.1.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Plerixafor is an inhibitor of the CXCR4 chemokine receptor and blocks binding of its cognate ligand, stromal cell-derived factor- 1α (SDF- 1α). SDF- 1α and CXCR4 are recognized to play a role in the trafficking and homing of human hematopoietic stem cells (HSCs) to the marrow compartment. Once in the marrow, stem cell CXCR4 can act to help anchor these cells to the marrow matrix, either directly via SDF- 1α or through the induction of other adhesion molecules. Treatment with plerixafor resulted in leukocytosis and elevations in circulating hematopoietic progenitor cells in mice, dogs

and humans. CD34+ cells mobilized by plerixafor were capable of engraftment with long-term repopulating capacity up to one year in canine transplantation models.

12.2 Pharmacodynamics

Data on the fold increase in peripheral blood CD34+ cell count (cells/mcL) by apheresis day were evaluated in two placebo-controlled clinical studies in patients with NHL and MM (Study 1 and Study 2, respectively). The fold increase in CD34+ cell count (cells/mcL) over the 24-hour period starting from the day prior to the first apheresis and ending the next morning just before the first apheresis is summarized in Table 3. During this 24-hour period, a single dose of Mozobil or placebo was administered 10 to 11 hours prior to apheresis.

Table 3: Fold Increase in Peripheral Blood CD34+ Cell Count Following Pretreatment with Filgrastim and Administration of Plerixafor

	Mozobil and Filgrastim			cebo and grastim
Study	Median	Mean (SD)	Median	Mean (SD)
Study 1	5.0	6.1 (5.4)	1.4	1.9 (1.5)
Study 2	4.8	6.4 (6.8)	1.7	2.4 (7.3)

In pharmacodynamic studies of Mozobil in healthy volunteers, peak mobilization of CD34+ cells was observed between 6 and 9 hours after administration. In pharmacodynamic studies of Mozobil in conjunction with filgrastim in healthy volunteers, a sustained elevation in the peripheral blood CD34+ count was observed from 4 to 18 hours after plerixafor administration with a peak CD34+ count between 10 and 14 hours.

QT/QTc Prolongation

There is no indication of a QT/QTc prolonging effect of Mozobil in single doses up to 0.40 mg/kg. In a randomized, double-blind, crossover study, 48 healthy subjects were administered a single subcutaneous dose of plerixafor (0.24 mg/kg and 0.40 mg/kg) and placebo. Peak concentrations for 0.40 mg/kg Mozobil were approximately 1.8-fold higher than the peak concentrations following the 0.24 mg/kg single subcutaneous dose.

12.3 Pharmacokinetics

The single-dose pharmacokinetics of plerixafor 0.24 mg/kg were evaluated in patients with NHL and MM following pretreatment with filgrastim (10 mcg/kg once daily for 4 consecutive days). Plerixafor exhibits linear kinetics between the 0.04 mg/kg to 0.24 mg/kg dose range. The pharmacokinetics of plerixafor was similar across clinical studies in healthy subjects who received plerixafor alone and NHL and MM patients who received plerixafor in combination with filgrastim.

A population pharmacokinetic analysis incorporated plerixafor data from 63 subjects (NHL patients, MM patients, subjects with varying degrees of renal impairment, and healthy subjects) who received a single SC dose (0.04 mg/kg to 0.24 mg/kg) of

plerixafor. A two-compartment disposition model with first order absorption and elimination was found to adequately describe the plerixafor concentration-time profile. Significant relationships between clearance and creatinine clearance (CL_{CR}), as well as between central volume of distribution and body weight were observed. The distribution half-life ($t_{1/2\alpha}$) was estimated to be 0.3 hours and the terminal population half-life ($t_{1/2\beta}$) was 5.3 hours in patients with normal renal function.

The population pharmacokinetic analysis showed that the mg/kg-based dosage results in an increased plerixafor exposure (AUC_{0-24h}) with increasing body weight. In order to compare the pharmacokinetics and pharmacodynamics of plerixafor following 0.24 mg/kg-based and fixed (20 mg) doses, a follow-up trial was conducted in patients with NHL (N=61) who were treated with 0.24 mg/kg or 20 mg of plerixafor. The trial was conducted in patients weighing 70 kg or less. The fixed 20 mg dose showed 1.43-fold higher exposure (AUC_{0-10h}) than the 0.24 mg/kg dose (Table 4). The fixed 20 mg dose also showed numerically higher response rate (5.2% [60.0% vs 54.8%] based on the local lab data and 11.7% [63.3% vs 51.6%] based on the central lab data) in attaining the target of \geq 5 \times 10 6 CD34+ cells/kg than the mg/kg-based dose. However, the median time to reach \geq 5 \times 10 6 CD34+ cells/kg was 3 days for both treatment groups, and the safety profile between the groups was similar. Based on these results, further analysis was conducted by FDA reviewers and a body weight of 83 kg was selected as an appropriate cut-off point to transition patients from fixed to weight based dosing.

Table 4: Systemic Exposure (AUC_{0-10h}) Comparisons of Fixed and Weight-Based Regimens

Regimen	Geometric Mean AUC
Fixed 20 mg (n=30)	3991.2
0.24 mg/kg (n=31)	2792.7
Ratio (90% CI)	1.43 (1.32,1.54)

There is limited experience with the 0.24 mg/kg dose of plerixafor in patients weighing above 160 kg. Therefore, the dose should not exceed that of a 160 kg patient (i.e., 40 mg/day if CL_{CR} is greater than 50 mL/min and 27 mg/day if CL_{CR} is less than or equal to 50 mL/min) [see Dosage and Administration (2.1, 2.3)].

Absorption

Peak plasma concentrations occurred at approximately 30 to 60 minutes after a SC dose.

Distribution

Plerixafor is bound to human plasma proteins up to 58%. The apparent volume of distribution of plerixafor in humans is 0.3 L/kg demonstrating that plerixafor is largely confined to, but not limited to, the extravascular fluid space.

Metabolism

The metabolism of plerixafor was evaluated with *in vitro* assays. Plerixafor is not metabolized as shown in assays using human liver microsomes or human primary hepatocytes and does not exhibit inhibitory activity *in vitro* towards the major drug metabolizing cytochrome P450 enzymes (1A2, 2A6, 2B6, 2C8, 2C9, 2C19, 2D6, 2E1 and

3A4/5). In *in vitro* studies with human hepatocytes, plerixafor does not induce CYP1A2, CYP2B6, or CYP3A4 enzymes. These findings suggest that plerixafor has a low potential for involvement in cytochrome P450-dependent drug-drug interactions.

Elimination

The major route of elimination of plerixafor is urinary. Following a 0.24 mg/kg dose in healthy volunteers with normal renal function, approximately 70% of the dose was excreted in the urine as the parent drug during the first 24 hours following administration. In studies with healthy subjects and patients, the terminal half-life in plasma ranges between 3 and 5 hours. At concentrations similar to what are seen clinically, plerixafor did not act as a substrate or inhibitor of P-glycoprotein in an *in vitro* study with MDCKII and MDCKII-MDR1 cell models.

Special Populations

Renal impairment

Following a single 0.24 mg/kg SC dose, plerixafor clearance was reduced in subjects with varying degrees of renal impairment and was positively correlated with CL_{CR} . The mean AUC_{0-24h} of plerixafor in subjects with mild $(CL_{CR} 51-80 \text{ mL/min})$, moderate $(CL_{CR} 31-50 \text{ mL/min})$, and severe $(CL_{CR} < 31 \text{ mL/min})$ renal impairment was 7%, 32%, and 39% higher than healthy subjects with normal renal function, respectively. Renal impairment had no effect on C_{max} . A population pharmacokinetic analysis indicated an increased exposure (AUC_{0-24h}) in patients with moderate and severe renal impairment compared to patients with $CL_{CR} > 50 \text{ mL/min}$. These results support a dose reduction of one-third in patients with moderate to severe renal impairment $(CL_{CR} \le 50 \text{ mL/min})$ in order to match the exposure in patients with normal renal function. The population pharmacokinetic analysis showed that the mg/kg-based dosage results in an increased plerixafor exposure (AUC_{0-24h}) with increasing body weight; therefore, if CL_{CR} is $\le 50 \text{ mL/min}$ the dose should not exceed 27 mg/day [see Dosage and Administration (2.3)].

Since plerixafor is primarily eliminated by the kidneys, coadministration of plerixafor with drugs that reduce renal function or compete for active tubular secretion may increase serum concentrations of plerixafor or the coadministered drug. The effects of coadministration of plerixafor with other drugs that are renally eliminated or are known to affect renal function have not been evaluated.

Race

Clinical data show similar plerixafor pharmacokinetics for Caucasians and African Americans, and the effect of other racial/ethnic groups has not been studied.

Gender

Clinical data show no effect of gender on plerixafor pharmacokinetics.

Age

Clinical data show no effect of age on plerixafor pharmacokinetics.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

Carcinogenicity studies with plerixafor have not been conducted.

Plerixafor was not genotoxic in an *in vitro* bacterial mutation assay (Ames test in *Salmonella*), an *in vitro* chromosomal aberration test using V79 Chinese hamster cells, or an *in vivo* bone marrow micronucleus test in rats after subcutaneous doses up to 25 mg/kg (150 mg/m²).

The effect of plerixafor on human fertility is unknown. The effect of plerixafor on male fertility was not studied in designated reproductive toxicology studies. The staging of spermatogenesis measured in a 28-day repeated dose toxicity study in rats revealed no abnormalities considered to be related to plerixafor. No histopathological evidence of toxicity to male or female reproductive organs was observed in 28-day repeated dose toxicity studies.

No adverse effects on estrus or reproductive indices were observed in an investigative fertility study in female rats administered plerixafor at doses up to 90 mg/m 2 (15mg/kg/day) or approximately 10 times the recommended human dose of 0.24 mg/kg when compared on a mg/m 2 basis.

14 CLINICAL STUDIES

The efficacy and safety of Mozobil in conjunction with filgrastim in non-Hodgkin's lymphoma (NHL) Study AMD 3100-3101 (referred to as study 1) (NCT00103610) and multiple myeloma (MM) Study AMD 3100-3102 (referred to as study 2) (NCT00103662) were evaluated in two placebo-controlled studies (Studies 1 and 2). Patients were randomized to receive either Mozobil 0.24 mg/kg or placebo on each evening prior to apheresis. Patients received daily morning doses of filgrastim 10 mcg/kg for 4 days prior to the first dose of Mozobil or placebo and on each morning prior to apheresis. Two hundred and ninety-eight (298) NHL patients were included in the primary efficacy analyses for Study 1. The mean age was 55 years (range 29–75) and 58 years (range 22–75) in the Mozobil and placebo groups, respectively, and 93% of subjects were Caucasian. In study 2, 302 patients with MM were included in the primary efficacy analyses. The mean age (58 years) and age range (28–75) were similar in the Mozobil and placebo groups, and 81% of subjects were Caucasian.

In Study 1, 59% of NHL patients who were mobilized with Mozobil and filgrastim collected \geq 5 × 10⁶ CD34+ cells/kg from the peripheral blood in four or fewer apheresis sessions, compared with 20% of patients who were mobilized with placebo and filgrastim (p <0.001). Other CD34+ cell mobilization outcomes showed similar findings (Table 5).

Table 5: Study 1 Efficacy Results - CD34+ Cell Mobilization in NHL Patients

Efficacy Endpoint	Mozobil and Filgrastim (n=150)	Placebo and Filgrastim (n=148)	p-value*
Patients achieving ≥5 × 10 ⁶ cells/kg in ≤4 apheresis days	89 (59%)	29 (20%)	<0.001

Patients achieving $\geq 2 \times 10^6$	130 (87%)	70 (47%)	-0.001
cells/kg in ≤4 apheresis days	130 (87 /8)	70 (47 /0)	~0.001

^{*} p-value calculated using Pearson's Chi-Squared test

The median number of days to reach $\geq 5 \times 10^6$ CD34+ cells/kg was 3 days for the Mozobil group and not evaluable for the placebo group. Table 6 presents the proportion of patients who achieved $\geq 5 \times 10^6$ CD34+ cells/kg by apheresis day.

Table 6: Study 1 Efficacy Results - Proportion of Patients Who Achieved ≥5 × 10⁶ CD34+ cells/kg by Apheresis Day in NHL Patients

Days	Proportion* in Mozobil and Filgrastim (n=147 [†])	Proportion* in Placebo and Filgrastim (n=142 [†])
1	27.9%	4.2%
2	49.1%	14.2%
3	57.7%	21.6%
4	65.6%	24.2%

^{*} Percents determined by Kaplan Meier method

In Study 2, 72% of MM patients who were mobilized with Mozobil and filgrastim collected \geq 6 × 10⁶ CD34+ cells/kg from the peripheral blood in two or fewer apheresis sessions, compared with 34% of patients who were mobilized with placebo and filgrastim (p <0.001). Other CD34+ cell mobilization outcomes showed similar findings (Table 7).

Table 7: Study 2 Efficacy Results - CD34+ Cell Mobilization in Multiple Myeloma Patients

Efficacy Endpoint	Mozobil and Filgrastim (n=148)	Placebo and Filgrastim (n=154)	p-value*
Patients achieving $\geq 6 \times 10^6$ cells/kg in ≤ 2 apheresis days	106 (72%)	53 (34%)	<0.001
Patients achieving \geq 6 × 10 ⁶ cells/kg in \leq 4 apheresis days	112 (76%)	79 (51%)	<0.001
Patients achieving ≥2 × 10 ⁶ cells/kg in ≤4 apheresis days	141 (95%)	136 (88%)	0.028

^{*} p-value calculated using Pearson's Chi-Squared test

The median number of days to reach $\geq 6 \times 10^6$ CD34+ cells/kg was 1 day for the Mozobil group and 4 days for the placebo group. Table 8 presents the proportion of patients who achieved $\geq 6 \times 10^6$ CD34+ cells/kg by apheresis day.

Table 8: Study 2 - Proportion of Patients Who Achieved

[†] n includes all patients who received at least one day of apheresis

≥6 × 10⁶ CD34+ cells/kg by Apheresis Day in MM Patients

Days	Proportion*in Mozobil and Filgrastim (n=144 [†])	Proportion* in Placebo and Filgrastim (n=150 [†])
1	54.2%	17.3%
2	77.9%	35.3%
3	86.8%	48.9%
4	86.8%	55.9%

^{*} Percents determined by Kaplan Meier method

Multiple factors can influence time to engraftment and graft durability following stem cell transplantation. For transplanted patients in the Phase 3 studies, time to neutrophil and platelet engraftment and graft durability were similar across the treatment groups.

16 HOW SUPPLIED/STORAGE AND HANDLING

Mozobil® (plerixafor) injection 24 mg/1.2 mL (20 mg/mL) is a sterile, preservative-free, clear, colorless to pale-yellow solution supplied in a 2 mL clear glass single-dose vial.

NDC Number: 0024-5862-01

Store at 25°C (77°F); excursions permitted to 15°C-30°C (59°F-86°F) [see USP Controlled Room Temperature].

17 PATIENT COUNSELING INFORMATION

Advise patients of the potential for anaphylactic reactions, including signs and symptoms such as urticaria, periorbital swelling, dyspnea, or hypoxia during and following Mozobil injection and to report these symptoms immediately to a healthcare professional [see Adverse Reactions (6.1, 6.2)].

Advise patients to contact healthcare professional immediately if they experience left upper abdominal pain and/or scapular or shoulder pain [see Adverse Reactions (6.1,6.2)].

Advise patients to inform a healthcare professional immediately if symptoms of vasovagal reactions such as orthostatic hypotension or syncope occur during or shortly after their Mozobil injection [see Adverse Reactions (6.1)].

Advise patients who experience itching, rash, or reaction at the site of injection to notify a healthcare professional, as these symptoms have been treated with over-the-counter medications during clinical trials [see Adverse Reactions (6.1)].

Advise patients that Mozobil may cause gastrointestinal disorders, including diarrhea, nausea, vomiting, flatulence, and abdominal pain. Patients should be told how to manage specific gastrointestinal disorders and to inform their healthcare professional if severe events occur following Mozobil injection [see Adverse Reactions (6.1)].

Advise females of reproductive potential of the potential risk to a fetus. Advise females

[†] n includes all patients who received at least one day of apheresis

to contact their healthcare provider if they become pregnant, or if pregnancy is suspected, during treatment with Mozobil [see Warnings and Precautions (5.6), Use in Specific Populations (8.1)].

Advise females and males of reproductive potential to use effective contraceptive methods during Mozobil use and for 1 week following cessation of treatment [see Warnings and Precautions (5.6), Use in Specific Populations (8.1)].

Advise women not to breastfeed during treatment with Mozobil and for 1 week following the last dose [Use in Specific Populations (8.2)].

Genzyme Corporation Cambridge, MA 02141 A SANOFI COMPANY

© 2023 Genzyme Corporation. All rights reserved. Mozobil is a registered trademark of Genzyme Corporation.

PRINCIPAL DISPLAY PANEL - 24 mg/1.2 mL Vial Carton

NDC 0024-5862-01

Mozobil[®] (plerixafor) injection

24 mg/1.2 mL (20 mg/mL)

For subcutaneous injection only

One single-dose vial.

Discard unused portion.

Rx only

SANOFI GENZYME



MOZOBIL

plerixafor injection, solution

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:0024-5862
Route of Administration	SUBCUTANEOUS		

Active Ingredient/Active Moiety		
Ingredient Name	Basis of Strength	Strength
PLERIXAFOR (UNII: S915P5499N) (PLERIXAFOR - UNII:S915P5499N)	PLERIXAFOR	24 mg in 1.2 mL

Inactive Ingredients				
Ingredient Name	Strength			
SODIUM CHLORIDE (UNII: 451W47IQ8X)	5.9 mg in 1.2 mL			
HYDROCHLORIC ACID (UNII: QTT17582CB)				
WATER (UNII: 059QF0KO0R)				
SODIUM HYDROXIDE (UNII: 55X04QC32I)				

P	Packaging			
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:0024- 5862-01	1 in 1 CARTON	09/01/2013	
1		1.2 mL in 1 VIAL, SINGLE-USE; Type 0: Not a Combination Product		

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
NDA	NDA022311	09/01/2013	

Labeler - sanofi-aventis U.S. LLC (824676584)

Establishment			
Name	Address	ID/FEI	Business Operations
Aptuit (Oxford) Ltd		217370698	analysis(0024-5862), api manufacture(0024-5862)

Establishment			
Name	Address	ID/FEI	Business Operations
Genzyme Corporation		098066215	analysis(0024-5862), manufacture(0024-5862), pack(0024-5862), label(0024-5862)

Revised: 10/2023 sanofi-aventis U.S. LLC