

PRODUCT INFORMATION

INDICATION

MONJUVI (tafasitamab-cxix), in combination with lenalidomide, is indicated for the treatment of adult patients with relapsed or refractory diffuse large B-cell lymphoma (DLBCL) not otherwise specified, including DLBCL arising from low grade lymphoma, and who are not eligible for autologous stem cell transplant (ASCT).

This indication is approved under accelerated approval based on overall response rate. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

IMPORTANT SAFETY INFORMATION

Contraindications: None.

Warnings and Precautions:

- Infusion-Related Reactions (IRRs). MONJUVI can cause IRRs, including fever, chills, rash, flushing, dyspnea, and hypertension. Premedicate patients and monitor frequently during infusion. Based on the severity of the IRR, interrupt or discontinue MONJUVI and institute appropriate medical management.
- Myelosuppression. MONJUVI can cause serious or severe myelosuppression, including neutropenia, thrombocytopenia, and anemia. Monitor complete blood counts (CBC) prior to administration of each treatment cycle and throughout treatment. Monitor patients with neutropenia for signs of infection.
 Consider granulocyte colony-stimulating factor administration. Withhold MONJUVI based on the severity of the adverse reaction. Refer to the lenalidomide prescribing information for dosage modifications.

(continued on page 6)



INDICATION

MONJUVI (tafasitamab-cxix), in combination with lenalidomide, is indicated for the treatment of adult patients with relapsed or refractory diffuse large B-cell lymphoma (DLBCL) not otherwise specified, including DLBCL arising from low grade lymphoma, and who are not eligible for autologous stem cell transplant (ASCT).

This indication is approved under accelerated approval based on overall response rate. Continued approval for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s).

Manufactured by	<i>MorphoSys US Inc.</i> 844-MOR-1992
Marketed by	MorphoSys US Inc. www.MorphoSys-US.com Incyte Corporation www.Incyte.com
Product Name	MONJUVI
Established Name	tafasitamab-cxix
Product Website	www.MONJUVIHCP.com





Not actual size



National Comprehensive Cancer Network® (NCCN®) Preferred Treatment Option

NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) recommend tafasitamab-cxix (MONJUVI) in combination with lenalidomide as a preferred second-line or subsequent therapy option (if not previously used) for DLBCL in patients who are not candidates for transplant (Category 2A)¹¹

*It is unclear if tafasitamab or loncastuximab tesirine or if any other CD-19 directed therapy would have a negative impact on the efficacy of subsequent anti-CD19 CAR T-cell therapy.

NCCN makes no warranties of any kind whatsoever regarding their content, use, or application and disclaims any responsibility for their application or use in any way.

PRODUCT INFORMATION	
Description	200 mg single-dose vial as lyophilized powder for reconstitution
Sales Unit	One single-dose vial
Units Per Case	One vial per carton
NDC	10-Digit - 73535-208-01 11-Digit - 73535- 0 208-01
Global Trade Identification Numbers	00373535208013 30373535208014 (case)

HCPCS CODING

J9349 (Injection, tafasitamab-cxix, 2mg) Effective April 1, 2021 Wholesale Acquisition Cost (WAC) \$1,328.28 / Vial Established January 2024

PRICING

Please see Important Safety Information on pages 1 and 6 and full Prescribing Information.



PRODUCT AVAILABILITY

MONJUVI is available through a limited number of Specialty Distributors authorized to distribute and/or dispense the medication. For prescribers who do not wish to use buy-and-bill, distribution through a select Specialty Pharmacy is available

DISTRIBUTION

▶ All orders must be placed with one of the Specialty Distributors listed below:

AmerisourceBergen Specialty Distribution

AmerisourceBergen

5025 Plano Parkway, Carrollton, TX 75010 Phone: 800-746-6273 | Fax: 800-547-9413

Service@asdhealthcare.com www.asdhealthcare.com MONJUVI Item # 58057

Cardinal Health Specialty Pharmaceutical Distribution

233 Mason Road, LaVergne, TN 37086

Phone: 855-855-0708 | Phone: 877-453-3972

Fax: 877-274-9897

GMB-SPD-Specialty@cardinalhealth.com

GMB-SPDOncology Sales Team@cardinalhealth.com

MONJUVI Item # 5653530

McKesson Plasma & Biologics

6535 N State Highway 161, Irving, TX 75039 Phone: 877-625-2566 | Fax: 888-752-7276

MPBOrders@mckesson.com connect.mckesson.com MONJUVI Item # 1559434

Oncology Supply

AmerisourceBergen

2801 Horace Shepard Drive, Dothan, AL 36303

Phone: 800-633-7555

Service@oncologysupply.com www.oncologysupply.com MONJUVI Item # 58057

CuraScript SD

255 Technology Park, Lake Mary, FL 32746 Phone: 877-599-7748 | Fax: 800-862-6208

Customer.Service@curascript.com

www.curascriptsd.com MONJUVI Item # 413402

McKesson Specialty Care Distribution

6535 N State Highway 161, Irving, TX 75039

Phone: 800-482-6700 mscs.mckesson.com MONJUVI Item # 5010390

▶ For Specialty Pharmacy services, customers can contact:

Biologics by McKesson

13000 Weston Parkway, Suite 105, Cary, NC 27513

Phone: 800-850-4306 | Fax: 800-823-4506 | Email: MyCareTeam@mckesson.com

biologics.mckesson.com

DISPENSING PACK DIMENSIONS (APPROXIMATE)

Depth: 33.5 mm Height: 76 mm Width: 37.5 mm

STORAGE INFORMATION

- ▶ MONJUVI for injection is a sterile, preservative-free, white to slightly yellowish lyophilized powder for reconstitution supplied as a 200 mg single-dose vial. Each 200 mg vial is individually packaged in a carton
- ➤ Store refrigerated at 36°F to 46°F (2°C to 8°C) in the original carton to protect from light. Do not shake.

 Do not freeze
- ▶ Use the reconstituted MONJUVI solution immediately. If needed, store the reconstituted solution in the vial for a maximum of 12 hours either refrigerated at 36°F to 46°F (2°C to 8°C) or room temperature at 68°F to 77°F (20°C to 25°C) before dilution. Protect from light during storage. Do not freeze or shake
- ▶ If not used immediately, store the diluted MONJUVI infusion solution refrigerated for up to 18 hours at 36°F to 46°F (2°C to 8°C) and/or at room temperature for up to 12 hours at 68°F to 77°F (20°C to 25°C). The room temperature storage includes time for infusion. Protect from light during storage. Do not freeze or shake

PRODUCT EXPIRATION

Expiration date printed on both single-dose vial and carton

Please see Important Safety Information on pages 1 and 6 and full Prescribing Information.



DOSAGE AND ADMINISTRATION HIGHLIGHTS

- ► The recommended dose of MONJUVI is 12 mg/kg based on actual body weight administered as an intravenous infusion according to the dosing schedule shown below
- ▶ Administer MONJUVI in combination with lenalidomide 25 mg orally on Days 1 to 21 of each 28-day cycle for a maximum of 12 cycles, then continue MONJUVI as monotherapy until disease progression or unacceptable toxicity. Refer to the lenalidomide prescribing information for lenalidomide dosage recommendations
- MONJUVI Dosing Schedule:
 - Cycle 1 Days 1, 4, 8, 15 and 22
 - Cycles 2 and 3 Days 1, 8, 15 and 22
 - Cycle 4 and beyond Days 1 and 15
- ▶ MONJUVI should be administered by a healthcare professional with immediate access to emergency equipment and appropriate medical support to manage infusion-related reactions (IRRs)

See full <u>Prescribing Information</u> for additional details on dosing and administration including preparing the infusion, prophylaxis for infusion related reactions, and dose modifications for adverse reactions.

RECONSTITUTION

- ▶ Calculate the dose (mg) and determine the number of vials needed
- ▶ Reconstitute each 200 mg MONJUVI vial with 5 mL Sterile Water for Injection, USP with the stream directed toward the wall of each vial to obtain a final concentration of 40 mg/mL tafasitamab-cxix
- ▶ Gently swirl the vial(s) until completely dissolved. Do not shake or swirl vigorously. Complete dissolution may take up to 5 minutes
- ▶ Visually inspect the reconstituted solution for particulate matter or discoloration. The reconstituted solution should appear as a colorless to slightly yellow solution. Discard the vial(s) if the solution is cloudy, discolored, or contains visible particles
- ▶ Use the reconstituted MONJUVI solution immediately. If needed, store the reconstituted solution in the vial for a maximum of 12 hours either refrigerated at 36°F to 46°F (2°C to 8°C) or room temperature at 68°F to 77°F (20°C to 25°C) before dilution. Protect from light during storage

DILUTION

- ▶ Determine the volume (mL) of the 40 mg/mL reconstituted MONJUVI solution needed based on the required dose
- ► Remove a volume equal to the required MONJUVI solution from a 250 mL 0.9% Sodium Chloride Injection, USP infusion bag and discard it
- ▶ Withdraw the necessary amount of MONJUVI and slowly dilute in the infusion bag that contains the 0.9% Sodium Chloride Injection, USP to a final concentration of 2 mg/mL to 8 mg/mL. Discard any unused portion of MONJUVI remaining in the vial
- ▶ Gently mix the intravenous bag by slowly inverting the bag. <u>Do not shake</u>. Visually inspect the infusion bag with the diluted MONJUVI infusion solution for particulate matter and discoloration prior to administration
- ▶ If not used immediately, store the diluted MONJUVI infusion solution refrigerated for up to 18 hours at 36°F to 46°F (2°C to 8°C) and/or at room temperature for up to 12 hours at 68°F to 77°F (20°C to 25°C). The room temperature storage includes time for infusion. Protect from light during storage

Do not shake or freeze the reconstituted or diluted infusion solutions

Please see Important Safety Information on pages 1 and 6 and full Prescribing Information.



PRODUCT ORDER AND RETURN INFORMATION

For information on ordering, please contact ICS:

Fax: 833-958-2078 Email: MorphoSysCS@icsconnect.com

Credit for returns is subject to MorphoSys' current Return Goods Policy

Please contact ICS for more information:

Phone: 866-765-0919 Email: MorphoSysReturns@icsconnect.com

ADVERSE EVENT REPORTING

▶ Contact MorphoSys US Inc. or the FDA to report an adverse event:

MorphoSys US Inc. FDA

Phone: 844-667-1992 Phone: 1-800-FDA-1088

Email: MedInfo@MorphoSys.com Web: www.fda.gov/medwatch

PRODUCT COMPLAINTS

▶ For product complaints other than adverse event reporting, please contact MorphoSys US Inc. at:

Phone: 844-667-1992 Email: MedInfo@MorphoSys.com

US MEDICAL INFORMATION INQUIRES

MY MISSION SUPPORT

- ▶ My MISSION Support offers robust patient access and support services to eligible patients and caregivers
- ▶ My MISSION Support's services include help understanding health insurance coverage requirements, reimbursement support, and financial assistance options for eligible patients* such as copay assistance for commercially insured patients† and free product for eligible patients through the My MISSION Support Patient Assistance Program
- ► For personalized support from a My MISSION Support Program Specialist, call 855-421-6172, Monday Friday 8 AM to 8 PM ET, or visit www.MyMISSIONSupport.com to learn more

^{*} Other terms and conditions apply. Visit www.MyMISSIONSupport.com for full eligibility criteria.

[†] Program is not valid for prescriptions that are eligible to be reimbursed, in whole or in part, by Medicaid, Medicare, or other federal or state healthcare programs (including any state prescription drug assistance programs and the Government Health Insurance Plan available in Puerto Rico [formerly known as "La Reforma de Salud"]).

IMPORTANT SAFETY INFORMATION (continued)

Warnings and Precautions (continued):

- Infections. Fatal and serious infections, including opportunistic infections, occurred in patients during treatment with MONJUVI and following the last dose. 73% of the 81 patients developed an infection. The most frequent infections were respiratory tract infection, urinary tract infection, bronchitis, nasopharyngitis and pneumonia. Grade 3 or higher infection occurred (30% of 81 patients). The most frequent grade 3 or higher infection was pneumonia. Infection-related deaths were reported (2.5% of 81 patients). Monitor patients for signs and symptoms of infection and manage infections as appropriate.
- Embryo-Fetal Toxicity. Based on its mechanism of action, MONJUVI may cause fetal B-cell depletion when administered to a pregnant woman. Advise pregnant women of the potential risk to a fetus and women of reproductive potential to use effective contraception during treatment with MONJUVI and for at least 3 months after the last dose. The combination of MONJUVI with lenalidomide is contraindicated in pregnant women. Refer to the lenalidomide prescribing information on use during pregnancy.

Adverse Reactions:

The most common adverse reactions (≥20%) were neutropenia (51%), fatigue (38%), anemia (36%), diarrhea (36%), thrombocytopenia (31%), cough (26%), pyrexia (24%), peripheral edema (24%), respiratory tract infection (24%), and decreased appetite (22%).

You may report side effects to the FDA at (800) FDA-1088 or www.fda.gov/medwatch. You may also report side effects to MORPHOSYS US INC. at (844) 667-1992.

Please see the full <u>Prescribing Information</u> for additional Important Safety Information.

REFERENCES: 1. Referenced with permission from the NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines®) for B-Cell Lymphomas V.5.2023. © National Comprehensive Cancer Network, Inc. 2023. All rights reserved. Accessed July 7, 2023. To view the most recent and complete version of the guideline, go online to NCCN.org.





MONJUVI and the MONJUVI logo are registered trademarks of MorphoSys AG.
© 2023 August 2023 RC-US-TAF-01823
Distributed and marketed by MorphoSys US Inc. and marketed by Incyte.
MorphoSys is a registered trademark of MorphoSys AG.
Incyte and the Incyte logo are registered trademarks of Incyte.
All other trademarks are the property of their respective owners.