TN100 Covey Drive Suite 307 Franklin, TN 37067





Office: 212-803-3339 Fax: 646-768-8600

OMVOH (mirikizumab-mrkz)

Provider Order Form

PATIENT INFORMATION				
Name:	DOB:		SEX: M □ F □	
ICD-10 code (required):	ICD-10 descript	ICD-10 description:		
□NKDA Allergies:			Weight lbs/kg:	
REFERRAL STATUS				
REFERENCE STATOS				
□New Referral □Referral Renewal □Medicat	tion/Order Change 🗆 Benefit	ts Verification Only	\square Discontinuation Order	
PHYSICIAN INFORMATION				
Referral Coordinator Name:	Referral Coordin	Referral Coordinator Email:		
Ordering Provider:	Provider NPI:	Provider NPI:		
Referring Practice Name:	Phone:	Phone: Fax:		
Practice Address:	City:	State:	Zip Code:	

INDICATIONS AND USAGE

OMVOHTM is an interleukin-23 antagonist indicated for the treatment of moderately to severely active ulcerative colitis in adults (1)

DOSAGE AND ADMINISTRATION

Prior to Treatment Initiation

- Evaluate patients for tuberculosis (TB) infection. (2.1, 5.3)
- Obtain liver enzymes and bilirubin levels. (2.1, 5.4)
- · Complete all age-appropriate vaccinations according to current Recommended Dosage
- The recommended induction dosage is 300 mg administered by intravenous infusion over at least 30 minutes at Weeks 0, 4, and 8.
- The recommended maintenance dosage is 200 mg administered by subcutaneous injection (given as two consecutive injections of 100 mg each) at Week 12, and every 4 weeks thereafter. (2.2)

Preparation and Administration

See the full prescribing information for preparation, administration and storage information for intravenous infusion and subcutaneous injection. (2.3, 2.4)

DOSAGE FORMS AND STRENGTH

Intravenous Infusion (3)

- Injection: 300 mg/15 mL (20 mg/mL) solution in a single-dose vial Subcutaneous Injection (3):
- Injection: 100 mg/mL solution in a single-dose prefilled pen

CONTRAINDICATIONS

History of serious hypersensitivity reaction to mirikizumab-mrkz or any of the excipients. (4, 5.1).

WARNINGS AND PRECAUTIONS

- Hypersensitivity Reactions: Serious hypersensitivity reactions, including anaphylaxis and infusion-related reactions, have been reported. If a severe hypersensitivity reaction occurs, discontinue and initiate appropriate treatment. (5.1)
- Infections: OMVOH may increase the risk of infection. Do not initiate treatment with OMVOH in patients with a clinically important active infection until the infection resolves or is adequately treated. If a serious infection develops, do not administer OMVOH until the infection resolves, (5.2)
- Tuberculosis: Do not administer OMVOH to patients with active TB infection. Monitor patients receiving OMVOH for signs and symptoms of active TB during and after treatment. (5.3)
- · Hepatotoxicity: Drug-induced liver injury has been reported. Monitor liver enzymes and bilirubin levels at baseline and for at

least 24 weeks of treatment and thereafter according to routine patient management. Interrupt treatment if drug-induced liver injury is suspected, until this diagnosis is excluded. (5.4)

Immunizations: Avoid use of live vaccines. (5.5)

ADVERSE REACTIONS

Most common adverse reactions (≥2%) are:

Date:

- Induction: upper respiratory tract infections and arthralgia. (6.1)
- Maintenance: upper respiratory tract infections, injection site reactions, arthralgia, rash, headache, and herpes viral infection. (6.1)

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