

Updated: 02/2025 DMMA Approved: 02/2025

Request for Prior Authorization for Enspryng (satralizumab-mwge) and Uplizna (Inebilizumab-cdon)

Website Form — www.highmarkhealthoptions.com
Submit request via: Fax - 1-855-476-4158

All requests for Enspryng (satralizumab-mwge) and Uplizna (Inebilizumab-cdon) require a Prior Authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

Enspryng (satralizumab-mwge) and Uplizna (Inebilizumab-cdon) Prior Authorization Criteria:

For all requests for Enspryng (satralizumab-mwge) and Uplizna (Inebilizumab-cdon) all of the following criteria must be met:

- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines
- Is age-appropriate according to FDA-approved package labeling, nationally recognized compendia, or peer-reviewed medical literature
- Member must not have an active hepatitis B infection or active or untreated latent tuberculosis

Coverage may be provided with a <u>diagnosis</u> of Neuromyelitis Optica Spectrum Disorder (NMOSD) and the following criteria are met:

- Medication is prescribed by, or in consultation with a neurologist
- Documentation of a positive test for AQP4-IgG antibodies
- The prescriber submits documentation of baseline number of relapse(s), which occurred over the last year.
- Documentation of an Expanded Disability Status Scale (EDSS) score of ≤ 6.5 for Enspryng or a score of ≤ 8 for Uplizna (Inebilizumab-cdon)
- Must have documentation of inadequate response, contraindication or intolerance to one (1) immunosuppressant (e.g., mycophenolate mofetil, azathioprine, methotrexate) or an inadequate response, contraindication or intolerance to rituximab or any of its biosimilars
- **Initial Duration of Approval:** 12 months
- Reauthorization criteria
 - Documentation the member has experienced a decrease from baseline in the number of NMOSD relapse(s).
- Reauthorization Duration of Approval: 12 months

Coverage may be provided for any non-FDA labeled indication if it is determined that the use is a medically accepted indication supported by nationally recognized pharmacy compendia or peer-reviewed medical literature for treatment of the diagnosis(es) for which it is prescribed. These requests will be reviewed on a case by case basis to determine medical necessity.

Drugs are authorized in generic form unless the branded product is on the preferred drug list or the prescriber has indicated in writing that the branded product is medically necessary. If only the



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HEALTH OPTIONSDMMA Approved: 02/2025 branded product is on the preferred drug list, the generic form will be considered non-preferred and shall not require the prescriber to indicate in writing that the branded product is medically necessary.



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ENSPRYNG (SATRALIZUMAB-MWGE) AND UPLIZNA (INEBILIZUMAB-CDON) PRIOR AUTHORIZATION FORM

Please complete and fax all requested information below including any progress notes, laboratory test results, or chart documentation as applicable to Highmark Health Options Pharmacy Services. FAX: (855) 476-4158

If needed, you may call to spe	ak to a Pharmacy Services Represe	entative. PHONE : (844) 325	5-6251 Mon- Fri 8:00am to 7:00pm
	PROVIDER IN	FORMATION	
Requesting Provider:		NPI:	
Provider Specialty:		Office Contact:	
Office Address:		Office Phone:	
		Office Fax:	
	MEMBER INF	ORMATION	
Member Name:		DOB:	
Member ID:		Member weight:	Height:
	REQUESTED DRUG	G INFORMATION	
Medication:		Strength:	
Directions:		Quantity:	Refills:
Is the member currently received	ing requested medication? Yes		
Is this medication being used for	or a chronic or long-term condition		may be necessary for the life of
the patient? Yes No			
	Billing Inf		
This medication will be billed:		cally, JCODE:	
Place of Service: Hospital		ber's home Other	
	Place of Service		
Name:		NPI:	
Address:		Phone:	
	MEDICAL HISTORY (Co		
Is documentation of a positive what is the member's Expande How many relapse(s) have occur		or untreated latent tuberculor vided? Yes No Score? mmunosuppressant or ritux	
	CURRENT or PRE	VIOUS THERAPY	
Medication Name	Strength/ Frequency	Dates of Therapy	Status (Discontinued & Why / Current)
Use the member eveninged a	REAUTHO decrease in the number of NMOS		omit documentation. No
	UPPORTING INFORMATION		
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Prescribing Pro	vider Signature		Date



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