

Kaiser Permanente Health Plan of Mid-Atlantic States, Inc.
REYVOW (lasmiditan succinate) Prior Authorization (PA)
Pharmacy Benefits Prior Authorization Help Desk
Length of Authorizations: Initial- 4 months; Continuation- 12 months

Instructions:

This form is used by Kaiser Permanente and/or participating providers for coverage **REYVOW** (lasmiditan succinate). Please complete all sections, incomplete forms will delay processing. Fax this form back to Kaiser Permanente within 24 hours fax: 1-866-331-2104. If you have any questions or concerns, please call 1-866-331-2103. **Requests will not be considered unless all sections are complete.**

KP-MAS Formulary can be found at: Pharmacy | Community Provider Portal | Kaiser Permanente

1 - Patient Information				
Patient Name:	Kaiser Medical ID#:	Date of Birth:		
2 – Prescriber Information				
Is the prescriber a Neurologist or Pain Management Specialist with expertise in diagnosis/treating headaches? □ No □ Yes				
If consulted with a specialist, specialist name and specialty:				
Prescriber Name:	Specialty:	NPI:		
Prescriber Address:				
Prescriber Phone #:	Prescriber Fax #:			
Do you have an approved provider referral number from Kaiser Permanente? □ Yes – please provide your provider referral number here:				
3 – Pharmacy Information				
Pharmacy Name:	Pharmacy NPI:			
Pharmacy Phone #	Pharmacy Fax #:			
4 – Drug Therapy Requested				
Sig:				
Drug 2: Name/Strength/Formulation:				
Sig:				

5- Diagnosis/Clinical Criteria

		J- Diagnosis/ Cillical	citeria		
1.	Is this request for initial or co	ntinuing therapy?			
	□ Initial therapy	□ Continuing therapy, start da	nte:		
2.	2. Indicate the Member's diagnosis for the requested medication:				
	Clinical Criteria:				
1.	 Is the medication being prescribed for the treatment of acute migraine? 				
	□ No □ Yes				
2.	2. Does the patient have documented trial (≥ 2 months) with treatment failure, or inadequate response, to at least 3				
	generic oral triptan agents at maximally tolerated doses?				
	□ No □ Yes				
2	Has the nationt failed or has	contraindication to Uhroby (uhrogor	ant\2		
٥.	3. Has the patient failed or has contraindication to Ubrelvy (ubrogepant)?□ No □ Yes				
	□ NO □ Tes				
Fo	r Continuation of Therapy. Ple	ase Respond to Additional Question	s Below:		
Does the patient meet all the initial criteria for coverage?					
	□ No □ Yes				
2. After 3 months of treatment, does the patient have evidence of positive clinical response?					
□ No □ Yes					
6 – Prescriber Sign-Off					
	Iditional Information –		and and backle to this assured		
		nedical records for the patient that	· ·		
2	•		nale/explanation and any additional supporting		
information that should be taken into consideration for the requested medication:					
I certify that the information provided is accurate. Supporting documentation is available for State audits.					
Р	rescriber Signature:		Date:		
_	Note This deep control of the	and the form of the standard and the sta	The state of the s		
	Please Note: This document contains confidential information, including protected health information, intended for a specific individual and purpose. The information is private and legally protected by law, including HIPAA. If you are not the intended recipient, you are hereby notified that any disclosure, copying, distribution or taking of				
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