

### State of Oklahoma SoonerSelect > 4aetna SoonerCare





## Nurtec® ODT (Rimegepant) Prior Authorization Form

Pharmacy billing (NDC:) Start date (or date of next dose):	Member I	Name:	_ Date of Birth:	Member ID#:
Billing Provider Information   Provider NPI:			Drug Information	on
Provider NPI:	Pharmac	y billing (NDC:	) Start date (	or date of next dose):
Provider NPI:	Dose:	Regimen:		Fill Quantity/Day Supply:
Prescriber NPI:			ing Provider Infor	
Prescriber NPI:	Provider	NPI:	Provider Nam	e:
Prescriber NPI:	Provider	Phone:	Provider Fax	c:
Prescriber Phone:			Prescriber Inform	ation
All information must be provided and SoonerCare may verify through further requested documentation. The member's medication history will be reviewed prior to approval.  *Page 1 of 2—Please complete and return all pages. Failure to complete all pages will result in processing delays  For Initial Authorization:  1. What is the member's diagnosis?    Acute Treatment of Migraine in Adults     Preventive Treatment of Episodic Migraines in Adults     Other, please list:   Other, please list:   Whill the member take Nurtec ODT concurrently with an injectable prophylactic calcitonin gene-related peptide (CGRP) inhibitor (e.g., Emgality®, Ajovy®, Aimovig®, Vyepti®)? Yes   No   If yes, please list:   Medication   Date Span   Dosing     Medication   Date Span   Dosing     C. If the member has no triptan trials, please provide a patient-specific, clinically significant reason why a triptan appropriate for the member:   3. If diagnosis is Preventive Treatment of Episodic Migraines in Adults, please provide the following (initial approvals be for 3 months):   a. Does the member have documented:   Episodic Migraine Headaches     b. Date of member's episodic migraine diagnosis?     c. Number of episodic migraines per day, on average, for the past 3 months?     d. Have the following medical conditions known to cause or exacerbate migraines been ruled out/treated?     i. Increased intracranial pressure (e.g., post-lumbar puncture headache, dural tear after trauma)? Yes   No   Iii. Decreased intracranial pressure (e.g., post-lumbar puncture headache, dural tear after trauma)? Yes   No   Iii. Obstructive sleep apnea? Yes   No   Iiii. Obstructive sleep apnea? Yes   No   Iiiii. Obstructive sleep apnea? Yes   No   Iiiiii. Obstructive sleep apnea? Yes   No   Iiiiiiiiiiiiiiiiiiiiiiiiiiiiiiiiiiii	Prescribe	er NPI:	Prescriber Name:	
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III. Medication Date Span Dosing	f.	(antihypertensives, anticonvulsants i. Medicationii. Medicationiii. Medication	s, antidepressants, etc.)? Date S Date S Date S	Yes No If yes, please list:  Span Dosing  Dosing  Dosing  Dosing
g. If the trial duration for the medication(s) listed above is not at least 8 weeks, please document the reason(s): Medication(s)	g.	Medication(s)	on(s) listed above is not a	at least 8 weeks, please document the reason(s):

Fax completed prior authorization request form to 888-601-8461 or submit Electronic Prior Authorization through CoverMyMeds® or SureScripts. All requested data must be provided. Incomplete forms or forms without the chart notes will be returned. Pharmacy

Coverage Guidelines are available at AetnaBetterHealth.com/Oklahoma.

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# State of Oklahoma SoonerCare



# Nurtec® ODT (Rimegepant) Prior Authorization Form

Метрег Name:	Date of Birtn:	Wember ID#:	
	Criteria		
The member's drug history will b	and SoonerCare may verify througe e reviewed prior to approval. nd return <u>all</u> pages. <i>Failure to comp</i>	•	
h. Is the member taking any of the absence of intractable condition i. Decongestants (alone or ir ii. Combination analgesics or iii. Opioid-containing medicat iv. Analgesic medications included v. Ergotamine-containing medications. Triptans? Yes No Label	eventive Treatment of Episodice following medications known to cause known to cause known to cause known to cause chronic pain? In combination products)? Yes Notationing caffeine and/or butalbital? Young acetaminophen or non-steroidadications? Yes Notation Notations? Yes Notation how the tractable conditions known to cause of ation(s) listed in Question h, please list	use medication overuse or rebound holds for the local section of the local section overuse of the local section overuse of the local section over section over section of the local section over section	neadaches in the
	ation(s) listed in Question h, please p ued use of medication(s) known to car		
k. Has the member been evaluate recommended as treatment? You i. If yes, please include named ii. Will member use Nurtec® ODT calcitonin gene-related peptide m. If applicable, are other aggrava being treated (e.g., smoking)? You. Please provide a patient-specific	cations that are <b>likely</b> to be the caused within the last 6 months by a neuroles \( \sum \text{No} \) No \( \sum \text{loop} \) e of neurologist recommending Nurte concurrently with botulinum toxin for (CGRP) inhibitor? Yes \( \sum \text{No} \) No \( \sum \text{loop} \) ting factors that contribute to the deverges \( \sum \text{No} \) No \( \sum \text{No} \) Not Applicable \( \sum \text{loop} \) ic, clinically significant reason why the open of the contribute of the contribu	ologist for episodic migraines and wa ec <sup>®</sup> ODT treatment_ the prevention of migraine or with ar elopment of episodic/chronic migrain ]	n alternative ne headaches
continued approval):  1. Has the member been complian 2. Has the member responded we 3. Please provide the member's c	compliance and information regent with Nurtec <sup>®</sup> ODT (rimegepant) treall to treatment with Nurtec <sup>®</sup> ODT (rimurrent number of migraine days per n	natment? Yes No No negepant) ? Yes No No nonth:	d for
Prescriber Signature:		_ Date:	
	is medically necessary and all informa		
	cific information will be requested if neces		-

Please complete and return <u>all</u> pages. Failure to complete all pages will result in processing delays. Page 2 of 2

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Pharm – 193 2/10/2023