

Prior Authorization Form



Note: Please provide as much information as possible on this form. Missing data may cause processing delays for requested authorization(s). Attach additional sheets to this form if necessary.

Please fax the completed PA form and any additional informational sheets to Nirvanahealth at the following fax number:
+1(866) 871-8565

Patient Information	Prescriber Information
Patient Name: _____	Prescriber Name: _____
Health Plan Name: _____	Prescriber Address: _____
Patient Insurance Id: _____	_____
Patient Date of Birth: _____	Prescriber Phone: () _____
Patient Phone: _____	Prescriber Fax: () _____
	Prescriber Specialty: _____
	Prescriber DEA: _____
	Prescriber NPI: _____

Medication & Medical Information	
Requested Drug(s) & Strength(s):	<input type="checkbox"/> Aimovig Autoinjector 140 mg/mL subcutaneous auto-injector <input type="checkbox"/> Aimovig Autoinjector 70 mg/mL subcutaneous auto-injector
Requested Daily Quantity Limit – Amount:	
Requested Daily Quantity Limit – Days:	
Expected Length of Therapy:	
Directions:	
Diagnosis and Diagnosis Codes (ICD-10 Standard Codes):	
List drugs used previously to treat the same condition:	
Additional clinical information or history. Please include any relevant test results and/or medical record notes:	

Questionnaire

Q1: I, as the provider or designated representative of the provider, certify and attest that the information provided is complete and accurate and that, upon request, I shall provide any information to RxAdvance that RxAdvance determines is reasonably necessary to verify my responses. (Check only one that apply)

Yes

No

Q2: Is the member currently treated with this medication? (Check only one that apply)

Yes (please list start date of therapy (month/day/year)) _____
(*Required)

No

Q3: What is the member's diagnosis? (Check only one that apply)

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Episodic Migraines (EM)

Chronic Migraines (CM)

Other (please specify the member's diagnosis and provide clinical rationale for the request)
_____ (*Required)

Q4: Has the member continues to be monitored for medication overuse headache? (Check only one that apply)

Yes

No (please provide medical justification for continuation of therapy)
_____ (*Required)

Q5: Does the medication prescribed by or in consultation with neurologist, headache specialist, or pain specialist? (Check only one that apply)

Yes (please specify prescriber specialty) _____ (*Required)

No (please provide medical justification for continuation of therapy)
_____ (*Required)

Q6: Does the member has a positive clinical response to therapy, demonstrated by a reduction in headache frequency and, or intensity? (Check only one that apply)

Yes (please specify the positive clinical response) _____ (*Required)

No (please provide medical justification for continuation of therapy)
_____ (*Required)

Q7: Is the use of acute migraine medications (e.g., non-steroidal anti-inflammatory drugs [NSAIDs] [e.g., ibuprofen, naproxen], triptans [e.g., eletriptan, rizatriptan, sumatriptan]) decreased since the start of calcitonin gene-related peptide (CGRP) therapy? (Check only one that apply)

Yes (please specify name of drug(s) and details of decrease use)
_____ (*Required)

No (please provide medical justification for continuation of therapy)
_____ (*Required)

Q8: Will the request be used in combination with another oral, calcitonin gene-related peptide (CGRP) inhibitor for the preventive treatment of migraines? (Check only one that apply)

Yes (please provide clinical rationale for the request) _____ (*Required)

No

Q9: What is the member's diagnosis? (Check only one that apply)

Episodic Migraines (EM)

Chronic Migraines (CM)

Other (please specify the member's diagnosis and provide clinical rationale for the request)
_____ (*Required)

Q10: Has the member have at least 4 but not more than 14 headache days per month? (Check only one that apply)

Yes (please specify number of headache days per month) _____ (*Required)

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No (please specify number of headache days per month and please provide clinical rationale for the request) _____ (*Required)

Q11: Has member's medication overuse headache been considered and potentially offending medication(s) have been discontinued? (Check only one that apply)

Yes (please specify name of drug(s) and date of discontinuation (month/year)) _____ (*Required)

No (please provide clinical rationale for the request) _____ (*Required)

Q12: Has the member had greater than or equal to 15 headache days per month, of which at least 8 must be migraine days for at least 3 months? (Check only one that apply)

Yes (please specify number of headache days per month) _____ (*Required)

No (please specify number of headache days per month and please provide clinical rationale for the request) _____ (*Required)

Q13: Has the member had an inadequate response or intolerance to at least 2-month trial of Elavil (amitriptyline) or Effexor (venlafaxine)? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q14: Does the member have any contraindication(s) to Elavil (amitriptyline)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q15: Does the member have any contraindication(s) to Effexor (venlafaxine)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q16: Has the member had an inadequate response or intolerance to at least 2-month trial of Depakote/Depakote ER (divalproex sodium) or Topamax (topiramate)? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q17: Does the member have any contraindication(s) to Depakote/Depakote ER (divalproex sodium)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q18: Does the member have any contraindication(s) to Topamax (topiramate)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

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No

Q19: Has the member had an inadequate response or intolerance to at least 2-month trial of one of the following beta blockers: atenolol, propranolol, nadolol, timolol, or metoprolol? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q20: Does the member have any contraindication(s) to atenolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q21: Does the member have any contraindication(s) to propranolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q22: Does the member have any contraindication(s) to nadolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q23: Does the member have any contraindication(s) to timolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q24: Does the member have any contraindication(s) to metoprolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No

Q25: Has the member had an inadequate response, intolerance or experienced contraindication(s) to Atacand (candesartan)? (Check only one that apply)

Yes (please specify corresponding contraindication(s) or intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q26: Has the member had an inadequate response or intolerance to at least 2-month trial of Depakote/Depakote ER (divalproex sodium) or Topamax (topiramate)? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q27: Does the member have any contraindication(s) to Depakote/Depakote ER (divalproex sodium)? (Check only one that apply)

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Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q28: Does the member have any contraindication(s) to Topamax (topiramate)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q29: Has the member had an inadequate response or intolerance to at least 2-month trial of one of the following beta blockers: atenolol, propranolol, nadolol, timolol, or metoprolol? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q30: Does the member have any contraindication(s) to atenolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q31: Does the member have any contraindication(s) to propranolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q32: Does the member have any contraindication(s) to nadolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q33: Does the member have any contraindication(s) to timolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q34: Does the member have any contraindication(s) to metoprolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No

Q35: Has the member had an inadequate response, intolerance or experienced contraindication(s) to Atacand (candesartan)? (Check only one that apply)

Yes (please specify corresponding contraindication(s) or intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

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Q36: Has the member had an inadequate response or intolerance to at least 2-month trial of one of the following beta blockers: atenolol, propranolol, nadolol, timolol, or metoprolol? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q37: Does the member have any contraindication(s) to atenolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No (please provide clinical rationale for the request) _____ (*Required)

Q38: Does the member have any contraindication(s) to propranolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No (please provide clinical rationale for the request) _____ (*Required)

Q39: Does the member have any contraindication(s) to nadolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No (please provide clinical rationale for the request) _____ (*Required)

Q40: Does the member have any contraindication(s) to timolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No (please provide clinical rationale for the request) _____ (*Required)

Q41: Does the member have any contraindication(s) to metoprolol? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No (please provide clinical rationale for the request) _____ (*Required)

Q42: Has the member had an inadequate response or intolerance to at least 2-month trial of Elavil (amitriptyline) or Effexor (venlafaxine)? (Check only one that apply)

Yes (please specify drug name, corresponding intolerance experienced and the start and end date(s) of therapy (month/year)) _____ (*Required)

No

Q43: Does the member have any contraindication(s) to Elavil (amitriptyline)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____ (*Required)

No _____ (*Required)

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Q44: Does the member have any contraindication(s) to Effexor (venlafaxine)? (Check only one that apply)

Yes (please specify corresponding contraindication(s)) _____
(*Required)

No _____ (*Required)

Q45: Will the request be used in combination with another oral, calcitonin gene-related peptide (CGRP) inhibitor for the preventive treatment of migraines? (Check only one that apply)

Yes (please provide clinical rationale for the request) _____
(*Required)

No

Q46: Is the member at least 18 years old? (Check only one that apply)

Yes

No (please specify member's age and provide clinical rationale for the request)
_____ (*Required)

Q47: Is the medication prescribed by or in consultation with a neurologist, headache specialist, or pain specialist? (Check only one that apply)

Yes (please specify prescriber specialty) _____ (*Required)

No (please provide clinical rationale for the request) _____
(*Required)

Attestation: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan, Insurer, Medical Group or its designated representatives may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.

Signature of Prescriber or Authorized Representative:

Date:

Print Authorized Representative Name: