











Prior Authorization Request Administrative Information

Member Information						
Last name	First name		МІ			
Member ID	Date of birth					
Sex assigned at birth Female Male "X" or Intersex						
Current gender Female Male Transgender male Transgender female Other						
Place of residence						
Race/ethnicity Preferred spoken la	anguage	Preferred written lang	guage			
MassHealth does not exclude people or treat them differently because of race, color, national origin, age, disability, religion, creed, sexual orientation, or sex (including gender identity and gender stereotyping).						
Plan Contact Information						
Please indicate the member's MassHealth Plan and fax or submit this completed and signed form according to the Plan's contact information below.						
MassHealth Fee-For-Service (FFS) Plan, Primary Care Clinician (PCC) Plan, Primary Care Accountable Care Organization (PCACO) Plan, Children's Medical Security Plan, and Health Safety Net Plan						
☐ MassHealth Drug Utilization Review Prog	yram					
Pharmacy: Fax: (877) 208-7428 - Tel: (800)	745-7318					
MassHealth Managed Care Organization	(MCO) and Ac	countable Care Partnersh	ip Plans (ACPP)			
☐ Fallon Health						
Online Prior Authorization: go.covermymed	s.com/OptumRx					
Online Prior Authorization: providerportal.su	urescripts.net/Pr	oviderPortal/optum				
Pharmacy: Fax: (844) 403-1029 - Tel: (844) 720-0033						
☐ Health New England						
Online Prior Authorization: go.covermymeds.com/OptumRx						
Pharmacy: Fax: (800) 550-9246 - Tel: (800) 918-7545						
☐ Mass General Brigham Health Plan						
Online Prior Authorization (Non-Specialty Drugs): go.covermymeds.com/OptumRx						
Online Prior Authorization (Specialty/Medical Drugs): provider.massgeneralbrighamhealthplan.org						
Pharmacy: Fax: (844) 403-1029 - Tel: (800) 711-4555						
☐ Tufts Health Plan						
Online Prior Authorization: point32health.promptpa.com						
Pharmacy: Fax: (617) 673-0939 - Tel: (888) 257-1985						
☐ WellSense Health Plan						
Online Prior Authorization: wellsense.org/providers/ma/pharmacy/prior-authorizations						
Pharmacy: Fax: (833) 951-1680 - Tel: (877) 417-1822						

Antiemetics Prior Authorization Request

MassHealth reviews requests for prior authorization (PA) on the basis of medical necessity only. If MassHealth approves the request, payment is still subject to all general conditions of MassHealth, including current member eligibility, other insurance, and program restrictions. MassHealth will notify the requesting provider and member of its decision. Keep a copy of this form for your records. If faxing this form, please use black ink.

Additional information about these agents, including PA requirements and preferred products, can be found within the MassHealth Drug List at www.mass.gov/druglist.

Medication information	
Medication requested	
 ☐ Akynzeo (fosnetupitant/palonosetron injection) > 2 units/28 days ☐ Akynzeo (netupitant/palonosetron capsule) > 2 units/28 days ☐ Anzemet (dolasetron) ☐ aprepitant 40 mg, 125 mg capsule > 2 units/28 	 ☐ Emend (aprepitant 125 mg powder for oral suspension) > 6 units/28 days ☐ fosaprepitant injection > 2 vials/28 day ☐ granisetron tablet > 2 tablets/28 days ☐ ondansetron solution ☐ palonosetron 0.25 mg/2 mL injection > 2
days aprepitant 80 mg > 4 capsules/28 days aprepitant trifold pack > 2 packs/28 days Bonjesta (doxylamine/pyridoxine extended-release) Cinvanti (aprepitant injectable emulsion) doxylamine/pyridoxine delayed-release	units/28 days palonosetron 0.25 mg/5 mL injection > 2 units/28 days Sancuso (granisetron transdermal system) Sustol (granisetron extended-release injection) > 2 units/28 days
Dose, frequency and duration of requested medication	on Landson
Indication (Check all that apply or include ICD-10 code,Chemotherapy-induced nausea and vomiting (CINV)Hyperemesis gravidarum	if applicable.) Postoperative nausea and vomiting (PONV) Radiation-induced nausea and vomiting (RINV) Other
Section I. Please complete for Cinvanti requests	S.
Has the member had a trial of oral aprepitant or fosapreportion. Yes. Please list the dates/duration of trial and outcor	•
Dates/duration of use Did the member experience any of the following? Briefly describe details of adverse reaction, inadequa	
☐ No. (Please explain why.)	
Section II. Please complete for Akynzeo, aprepit palonosetron, and Sustol requests ex Please describe the medical necessity for exceeding the	9 ,

PA-44 (Rev. 05/24) over

	ion III. Please complete for ondansetron solution requests. ses the member have a medical condition in which they are unable to swallow tablets/capsules?
	Yes. (Please list reason.)
	 ☐ No. (Please provide clinical rationale why conventional dosage forms cannot be used.)
Sect	ion IV. Please complete for Sancuso requests.
Ha	s the member had a trial of ondansetron ODT?
	Yes. Please list the dates/duration of trial and outcomes below.
	Dates/duration of use
	Did the member experience any of the following? \Box Adverse reaction \Box Inadequate response \Box Other
	Briefly describe details of adverse reaction, inadequate response, contraindication, or other.
	☐ No. (Please explain why.)
	□ No. (Flease explain wity.)
Sect	ion V. Please complete for Bonjesta and doxylamine/pyridoxine delayed-release requests
1.	Has the member had a trial of pyridoxine?
	Yes. Please list the dates/duration of trial and outcomes below.
	Dates/duration of use
	Did the member experience any of the following? Adverse reaction Inadequate response Other
	Briefly describe details of adverse reaction, inadequate response, contraindication, or other.
	Energy describe details of daverse reaction, inadequate responses, contrainal eatien, or ether.
	☐ No. (Please explain why.)
2.	Has the member had a trial of doxylamine?
	Yes. Please list the dates/duration of trial and outcomes below.
	Dates/duration of use
	Did the member experience any of the following? Adverse reaction Inadequate response Otherse
	Briefly describe details of adverse reaction, inadequate response, contraindication, or other.
0	□ No. (Please explain why.) □
3.	For Bonjesta requests, has the member had a trial of doxylamine/pyridoxine delayed-release?
	Yes. Please list the dates/duration of trial and outcomes below.
	Dates/duration of use
	Did the member experience any of the following? \Box Adverse reaction \Box Inadequate response \Box Oth
	Briefly describe details of adverse reaction, inadequate response, contraindication, or other.
	No. (Places symbols why)
	☐ No. (Please explain why.) □

Sec	tion	VI. Please complete for Anzemet requests.
1.	Has	the member had a trial of granisetron tablet?
		Yes. Please list the dates/duration of trial and outcomes below.
		Dates/duration of use
		Did the member experience any of the following? Adverse reaction Inadequate response Other
		Briefly describe details of adverse reaction, inadequate response, contraindication, or other.
		No. (Please explain why.)
2.	⊢ Has	the member had a trial of ondansetron tablet or ondansetron ODT?
		Yes. Please list the dates/duration of trial and outcomes below.
		Dates/duration of use
		Did the member experience any of the following? Adverse reaction Inadequate response Other
		Briefly describe details of adverse reaction, inadequate response, contraindication, or other.
	_	
		No. (Please explain why.)
		on in, or physical or mental harm to the member? Yes No es, briefly describe details of contraindication, adverse reaction, or harm.
	clinica	alternative drug required under the step therapy protocol expected to be ineffective based on the known all characteristics of the member and the known characteristics of the alternative drug regimen? Yes No
	If y	es, briefly describe details of known clinical characteristics of member and alternative drug regimen.
	altern drug v	ne member previously tried the alternative drug required under the step therapy protocol, or another ative drug in the same pharmacologic class or with the same mechanism of action, and such alternative was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event? Yes \sum No es, please provide details for the previous trial.
	Dri	g name Dates/duration of use
		the member experience any of the following? Adverse reaction Inadequate response
		efly describe details of adverse reaction or inadequate response.
		, , , , , , , , , , , , , , , , , , , ,
4.	Is the	member stable on the requested prescription drug prescribed by the health care provider, and switching
		will likely cause an adverse reaction in or physical or mental harm to the member?
	•	·
	∐ Ye ∏ No	s. Please provide details.
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Prior Authorization Request Prescriber and Provider Information

Prescriber Information					
Last name*	First name*	MI			
NPI*	Individual MH Provider	ID			
DEA No.	Office Contact Name				
Address	City	State Zip			
Email address					
Telephone No.*	Fax No.*				
* Required					
Please also complete for professionally adm	inistered medications	, if applicable.			
Start date	End date				
Servicing prescriber/facility name		☐ Same as prescribing provider			
Servicing provider/facility address					
Servicing provider NPI/tax ID No.					
Name of billing provider					
Billing provider NPI No.					
Is this a request for recertification? Yes No					
CPT code No. of visits	J code	No. of units			
Prescribing provider's attestation, signature	e, and date				
I certify under the pains and penalties of perjury that I am the prescribing provider identified in the Prescriber information section of this form. Any attached statement on my letterhead has been reviewed and signed by me. I certify that the medical necessity information (per 130 CMR 450.204) on this form is true, accurate, and complete, to the best of my knowledge. I understand that I may be subject to civil penalties or criminal prosecution for any falsification, omission, or concealment of any material fact contained herein.					
Prescribing provider's signature					
Printed name of prescribing provider		Date			
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(The form can either be signed by hand and then scanned, or it can be signed electronically using DocuSign or Adobe Sign. For electronic signatures, the signer can upload a picture of their wet signature. The typed text of a signature is not an acceptable form of an electronic signature.)