











Prior Authorization Request Administrative Information

Member Information			
Last name	First name		МІ
Member ID	Date of birth		
	X" or Intersex		
Current gender Female Male Transge	ender male 🔲 Tra	nsgender female Othe	-
Place of residence Home Nursing facility	Other		
Race/ethnicity Preferred spoken la	anguage	Preferred written lang	uage
MassHealth does not exclude people or treat the disability, religion, creed, sexual orientation, or s			
Plan Contact Information			
Please indicate the member's MassHealth Plan according to the Plan's contact information belo		his completed and signed	form
MassHealth Fee-For-Service (FFS) Plan, Pr Care Organization (PCACO) Plan, Child			
☐ MassHealth Drug Utilization Review Prog	gram		
Pharmacy: Fax: (877) 208-7428 - Tel: (800) 745-7318		
MassHealth Managed Care Organization	n (MCO) and Acco	untable Care Partnershi	p Plans (ACPP)
☐ Fallon Health			
Online Prior Authorization: go.covermymed	ds.com/OptumRx		
Online Prior Authorization: providerportal.s	urescripts.net/Provi	derPortal/optum	
Pharmacy: Fax: (844) 403-1029 - Tel: (844) 720-0033		
☐ Health New England			
Online Prior Authorization: go.covermymed	ds.com/OptumRx		
Pharmacy: Fax: (800) 550-9246 - Tel: (800) 918-7545		
Online Prior Authorization (Non-Specialty D	rugs): go.covermyr	neds.com/OptumRx	
Online Prior Authorization (Specialty/Medica	al Drugs): provider.	massgeneralbrighamhealt	hplan.org
Pharmacy: Fax: (844) 403-1029 - Tel: (800)	711-4555		
☐ Tufts Health Plan			
Online Prior Authorization: point32health.pr	romptpa.com		
Pharmacy: Fax: (617) 673-0939 - Tel: (888	3) 257-1985		
□ WellSense Health Plan			
Online Prior Authorization: wellsense.org/p	roviders/ma/pharma	acy/prior-authorizations	
Pharmacy: Fax: (833) 951-1680 - Tel: (877) 417-1822		

Opioids/Acetaminophen Analgesic 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 opioid and acetaminophen analgesic agents, including PA requirements and preferred products, can be found within the MassHealth Drug List at www.mass.gov/druglist.

Medication i	nformation			
Drug name		Dose and frequency	D	ouration of therapy
Has the pres potential risk Has the men	factors for abuse/mis	sachusetts Prescription Av suse in their assessment of d/or given a prescription fo	f this member? 🗌 Ye	
specialty of t Is this memb If yes, Massl	he collaborating phys er a referral candidat	e for care coordination? Coordination services to the] Yes 🗌 No	
☐ Yes. Druç	nber tried a morphine	or oxycodone extende extended-release product	` '	
☐ No. If mo	rphine and fentanyl tr	ansdermal are contraindic	ated in this member,	please describe.
☐ Yes. I	member tried a morph Dose and frequency morphine is contrain	or methadone (Doloph nine extended-release pro Dates of u	duct?	Outcome
	member tried a fentar Dose and frequency	nyl transdermal product? Dates of u	use	Outcome

PA-29 (Rev. 04/24) over

	No. If fentanyl transdermal is c	ontraindicated in this men	nber, please describ	е.		
3.	3. If the answer to questions 1 and 2 is no, please provide clinical rationale for the use of methadone instead					
	other long-acting opioids.					
4. 5.			c interval? ☐ Yes ☐	¬Nο		
	That the member had a bassime E	oo showing a normal Q1				
Sec	-	requests for fentanyl tr a), oxymorphone imme	_	, ,,,		
1.	` ,	ed on a long-acting opioid	regimen?			
	☐ Yes. Drug	Dose and frequency		Dates of use		
2.		g agents? 🗌 Yes. Please	describe below.			
	hydromorphone IR Dose and	d frequency	Dates of use	Outcome		
	morphine IR Dose and	d frequency	Dates of use	Outcome		
	oxycodone IR Dose and No. If hydromorphone, morphin	d frequency	Dates of use	Outcome		
	☐ No. II Hydromorphone, morphil	le, and oxycodone are con	ntramulcated in this	member, piease describe.		
3.	If the request is for fentanyl bucca	I tablet, has the member t	ried fentanyl transm	ucosal system (Actiq)?		
	☐ Yes. Dose and frequency	Dates of use		Outcome		
☐ No. If fentanyl transmucosal system (Actiq) is contraindicated in this member, please desc						
				>		
Sec	capsule, hydromorph		ablet, oxycodone	ER), hydrocodone ER ER capsule (Xtampza),		
1.	Has the member tried the following	-	· •			
	fentanyl transdermal Dose and	d frequency	Dates of use	Outcome		
	morphine ER Dose and	d frequency	Dates of use	Outcome		
	oxycodone ER Dose and No. If fentanyl transdermal, mo	d frequency orphine ER, and oxycodon	Dates of use e ER are contraindic	Outcome cated in this member, please		
	describe.					
2.	For levorphanol tablet requests, pl	ease provide clinical ratio	nale for the use of le	evorphanol instead of other		
	long-acting opioids.					
	ction V. Please complete for m		equests.*			
1.	Has the member tried morphine ex ☐ Yes. Dose and frequency	Dates of use		Outcome		

over

	☐ No. If morphine extended-release tablets are contraindicated in this member or there is medical
	necessity for the requested formulation, please describe.
2.	Please provide medical necessity for once daily dosing.
	tion VI. Please complete for meperidine (Demerol) requests. ease attach documentation describing medical necessity due to allergy to morphine.
Ple	ction VII. Please complete for requests for benzhydrocodone/acetaminophen, dihydrocodeine/acetaminophen/caffeine, hydrocodone/acetaminophen 300mg, hydrocodone 5 mg, 10 mg/ibuprofen, and oxycodone/acetaminophen 300mg.* ease attach documentation of prior combination analgesic trials including hydrocodone/acetaminophen, ycodone/acetaminophen, codeine/acetaminophen, and hydrocodone/ibuprofen.
	tion VIII. Please complete for buprenorphine buccal film (Belbuca) requests.* as the member tried a morphine extended-release product?
	Yes. Dose and frequency Dates of use Outcome
	ease provide medical necessity for use of requested formulation instead of other strengths.
Ple or	ction X. Please complete for butorphanol nasal spray requests. ease attach documentation describing an adverse reaction or contraindication to all other short-acting opioids medical necessity for nasal spray formulation in addition to an adverse reaction or contraindication to or propring and oxycodone IR solutions.
Se	ction XI. Please complete for tramadol ER (Conzip) requests.
1.	Please provide medical necessity for use of an extended-release formulation.
2.	Please attach documentation describing an inadequate response or adverse reaction to tramadol IR.
PΙ	ction XII. Please complete for Seglentis and tramadol/acetaminophen (Ultracet) requests. ease provide medical necessity for use of the combination product instead of the separately available gredients.

	ction XIII. Please complete for tramadol 100 mg requests.
1.	Please provide medical necessity for use of the requested strength.
2.	Please attach documentation describing an inadequate response or adverse reaction to tramadol 50 mg at the requested dose.
	ction XIV. Please complete for requests for codeine and tramadol products for members < 12 years of age. ease provide clinical rationale for use of a codeine and tramadol-containing product in a member < 12 years of
ag	e.
ls	ction XV. Please complete for tramadol solution requests.* there a medical necessity for use of an oral solution formulation? Yes. Please explain. No. Please attach medical records documenting inadequate response or adverse reaction to a tramadol mediate-release tablet formulation that is available without PA.
	Ition XVI. Please complete for oliceridine (Olinvyk) MB requests.* Is the total course of therapy limited to 48 hours? Yes No Has the member tried the following agents? Yes. Please describe below.
	fentanyl injection Dose and frequency Dates of use Outcome
	hydromorphone injection Dose and frequency Dates of use Outcome
	morphine injection Dose and frequency Dates of use Outcome No. If fentanyl injection, hydromorphone injection, and morphine injection are contraindicated in this
	member, please describe.
3.	Please indicate billing preference. Prescriber in-office Hospital outpatient If applicable, please also complete section for professionally administered medications at end of form. MB This drug is available through the health care professional who administers the drug or in an outpatient of inpatient hospital setting. MassHealth does not pay for this drug to be dispensed through the retail pharmacular listed, prior authorization does not apply through the hospital outpatient and inpatient settings. Please refer to 130 CMR 433.408 for prior authorization requirements for other health care professionals. Notwithstanding the above, this drug may be an exception to the unified pharmacy policy; please refer to respective MassHealth Accountable Care Partnership Plans (ACPPs) and Managed Care Organizations (MCOs) for prior authorization status and criteria, if applicable.

Fo	ction XVIII. Please complete for requests above established dose limits. or all opioids, please provide medical records documenting treatment plan including clinical rationale for high use and titration of medication up to current dose. In addition, please provide a signed and dated patient-			
prescriber agreement and a consult from a pain specialist recommending the requested dose for this ma current pain consult is not available, please provide the anticipated date of upcoming pain consult. If the plans to initiate a taper of the requested medication within the next 90 days, please provide medical recommendation.				
us	cumenting treatment plan. For acetaminophen and aspirin products, please provide a clinical rationale for the e above 4 grams per day. For ibuprofen products, please provide a clinical rationale for the use above 3.2 ams per day.			
PΙ	ease provide medical records documenting treatment plan including clinical rationale for use of high dose			
an	ort-acting opioids without a long-acting opioid agent. In addition, please provide clinical rationale for high dose distration of medication up to current dose, a signed and dated patient-prescriber agreement, and a consult of a pain specialist recommending the requested dose for this member.			
	III a pain specialist recommending the requested dose for this member.			
_				
Ca	etion XX. Please complete for requests above established quantity limits. an the requested dose be obtained by using products within established quantity limits (i.e., for oxycodone ER mg, 2 tablets twice daily could be consolidated to one oxycodone ER 40 mg tablet twice daily)?			
	Yes No. If dose consolidation is not an option, please explain why.			
	tion XXI. Please complete for concurrent therapy with opioid dependence agents. Are you the prescriber of both buprenorphine/naloxone or buprenorphine and the opioid? Yes No			
2.				
3.	Please document the medical necessity for concurrent buprenorphine/naloxone or buprenorphine and opioid therapy. Please submit medical records supporting the medical necessity, including the specific pain that the current opioid is being used to treat.			
4.	Please document the complete treatment plan, including expected duration of therapy for this member in regard to acute pain management with concurrent buprenorphine/naloxone or buprenorphine and opioid therapy.			

*Attach a letter with additional information regarding medication trials as applicable. If MassHealth pharmacy claims history of required trials is not available, medical records documenting such trials may be required.

Section XXII. Concomitant Opioid and Benzodiazepine Polypharmacy. Please complete information for medications requested and clinical rationale for polypharmacy with opioids and benzodiazepines [≥ 15 days supply for one or more opioid(s) who are newly starting opioid therapy and one or more benzodiazepine(s), excluding clobazam, nasal and rectal diazepam, nasal midazolam, and injectable formulations for ≥ 15 days supply within a 45-day period].

Please document the indication or ICD-10 code(s), if applicable, for the agents requested.

1. Opioid				
Name/dose/frequency		Indica	ation	
Name/dose/frequency		Indica	ation	
Name/dose/frequency 2. Benzodiazepine		Indica	ation	
Name/dose/frequency		Indica	ation	
Name/dose/frequency		Indica	ation	
Name/dose/frequency Please document clinical rationale for concomitant use of opioids and benzodiazepines for this member.				
Please describe the ongoing treatment plan for continued use.				
Has the member had trials v		nerapies?	Outron	
Yes. Drug name	Dates		Outcome	
Drug name	Dates		Outcome	
Drug name	Dates		Outcome	
Other	Dates		Outcome	
☐ No. Please documen	nt clinical rationale for t	he use of opioids instea	d of non-opioid alternatives.	
		f hannadiananina ay ani	-:-IO	
Has consideration been given for possible taper of benzodiazepine or opioid?				
Yes. Please describe	e plan for taper and pla	an to reevaluate in the fu	iture.	
 No. Please describe why taper is not possible at this time and plan to reevaluate in the future. 				

H	las the member been offered and/or given a prescription for naloxone treatment?
	☐ Yes ☐ No. Please provide details.
,	Attach a letter with additional information regarding medication trials as applicable.
S e 1.	Is the alternative drug required under the step therapy protocol contraindicated, or will likely cause an adverse reaction in, or physical or mental harm to the member? If yes, briefly describe details of contraindication, adverse reaction, or harm.
2.	Is the alternative drug required under the step therapy protocol expected to be ineffective based on the known clinical characteristics of the member and the known characteristics of the alternative drug regimen? Yes No If yes, briefly describe details of known clinical characteristics of member and alternative drug regimen.
3.	Has the member previously tried the alternative drug required under the step therapy protocol, or another alternative drug in the same pharmacologic class or with the same mechanism of action, and such alternative drug was discontinued due to lack of efficacy or effectiveness, diminished effect, or an adverse event? Yes No If yes, please provide details for the previous trial.
	Drug name Dates/duration of use Did the member experience any of the following? Adverse reaction Inadequate response Briefly 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 drugs will likely cause an adverse reaction in or physical or mental harm to the member?
	☐ Yes. Please provide details. ☐ No

Please continue to next page and complete Prescriber and Provider Information section.

Prior Authorization Request Prescriber and Provider Information

Prescriber Information			
Last name*	First name*		МІ
NPI*	Individual MH Provide	er ID	
DEA No.	Office Contact Name		
Address	City	State	Zip
Email address			
Telephone No.*	Fax No.*		
* Required			
Please also complete for professionally	administered medication	ns, if applicab	le.
Start date	End date		
Servicing prescriber/facility name		☐ Same as	s 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, signal certify under the pains and penalties of perinformation section of this form. Any attached I certify that the medical necessity information complete, to the best of my knowledge. I under prosecution for any falsification, omission, or	jury that I am the prescribing point of the statement on my letterhead on (per 130 CMR 450.204) on derstand that I may be subject	has been review this form is true to civil penaltie	wed and signed by me e, accurate, and es or criminal
Prescribing provider's signature			
Printed name of prescribing provider		Date	
(The form can either be signed by hand and	then scanned, or it can be sig	ned electronica	Illy using DocuSign or

(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.)