## MASSACHUSETTS STANDARD FORM FOR CHEMOTHERAPY AND SUPPORTIVE CARE PRIOR AUTHORIZATION REQUESTS\*

\*Providers may use the health plan's portal in place of this form.

Request Date:			Treatment Start Date:				☐ Standard ☐ Expedite					
I.												
	Health Plan Name:											
Hea	Ith Plan P	hone: 800-708-441	4 Fax: 617	7-673-0988	Online Prior	Authorization	: <u>https://poir</u>	nt32health.pro	omptpa.com			
Mei	nber Info	rmation										
	First: Last: MI:											
DOB:				Gender: [	Gender: M F Unknown Other:							
				Weight:								
			ICD-10:					ge (0–4 or recurrent):				
			Line of Busines					mber ID:				
*ECOG Score:					*Information in attached office note Yes							
*Tur	mor Histol	ogy:										
¥ A II												
^Alle	ergies:											
*(0	 morbiditie	c·										
	morbiatic	J.										
IJΔ	nti-cance	r Treatment Reque	est New:	Retrospec	tive: T	Re-Authorizati	on: $\square$					
#	Billing Code/ J CODE	Administrative Code	Drug Name	•	Dose	Frequency and Schedule	Cycles or Refills	Billing Method (B = Buy and Bill or P = Pharmacy)	FDA Approved for the Diagnosis?	For single use vials, is provider willing to dose round?		
1								ВШР	□Y □N	☐ Y ☐ N ☐ Unknown		
2								В 🗆 Р	□Y □N	Y N Unknown		
3								□В□Р	□Y□N	☐Y ☐N ☐ Unknown		
4								ВПР	□Y □N	☐Y ☐N ☐ Unknown		

#	Billing Code/ J CODE	Administrative Code	Drug Name	Route	Dose	Frequency and Schedule	Condition (ex: Nausea)	Billing Method (B = Buy and Bill or P = Pharmacy)		
1								□В□Р		
2								□В□Р		
3								□В□Р		
4								□В□Р		
If bor	_	thening agents or l	bone antiresorptive agents are reque s	sted, select vant Breast						
If ESAs requested, select indication:  CKD Chemotherapy Induced Anemia (CIA) MDS Anemia of Chronic Disease (ACD)										
IV. Pı	rovider a	nd Place of Treatr	ment Information							
Orde	ring Provi	ider:				Specialty	/:			
NPI#	:		TIN #:			DEA #:				
Phon				Fax:						
Treating Provider: (if different)  Specialty:										
NPI #:					TIN #:					
Phon				Fax:						
		nent: (if different)		1						
NPI #:					TIN #:					
	Phone: Fax:									
		atment Center:	ated with the requested regimen(s)?	□Vos □	No □Unl	(nown				
	of Treatm		area with the requested regiments):	L res L	110 [ 011	KHOWH				
		s has the patient p	reviously tried?							
***************************************		- The tire patient p	reviously tried.							
Has t	he patier	it been screened fo	or tumor mutations/biomarkers/gene	etic testing?	☐ Yes ☐	No Unkn	own			
If so,	If so, what tumor mutations/biomarkers/genetic testing result has the patient been tested for?									
			st, is this provider the only available t ?	reating/ser	vicing provide	er within a reas	sonable distar	nce that can provide this		
Has t	he memb	per been receiving	cancer treatments from the requesti	ng treating	provider? 🗌	Yes 🗌 No	Unknown	1		
Is treating provider in-network?  Yes  Unknown										
Site of Service: Outpatient Hospital Home Infusion Other										
Attachments: Labs Imaging Chemo Orders Pathology Progress Notes										
Authorized Representative:										
Phon	e:			Fax:						

V. Exceptions to Step Therapy Please complete the applicable section(s).
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?   Yes No
If yes, briefly describe details of contraindication, adverse reaction, or harm:
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 regiment?
If yes, briefly describe details of known clinical characteristics of member and alternative drug regimen:
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? $\square$ Yes $\square$ 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:
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:
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? $\square$ Yes $\square$ No
If yes, briefly provide details of the adverse reaction or physical or mental harm:

Providers should consult the health plan's coverage policies, member benefits, and medical necessity guidelines to complete this form. Providers must attach any additional data required relevant to medical necessity criteria, including PROGRESS NOTES, CHEMO ORDERS, LABS, PATHOLOGY, AND IMAGING RESULTS WITH REQUEST.