## **HEALTHY U** MEDICAID

## PRIOR AUTHORIZATION REQUEST FORM

## **PARKINSON'S AGENTS**

Apomorphine hydrochloride injection, Duopa™, Kynmobi™, Neupro®, Nourianz™, Ongentys®, Rytary®, Tasmar®, tolcapone, Zelapar®

For authorization, please answer each question and fax this form PLUS chart notes back to the Healthy U Prior Authorization Department at 385-425-4052.

Failure to submit clinical documentation to support this request will result in a dismissal of the request.

If you have prior authorization questions, please call for assistance: 385-425-5094 Disclaimer: Prior Authorization request forms are subject to change in accordance with Federal and State notice requirements. Date: Member Name: ID#: DOB: Gender: Physician: Office Phone: Office Fax: Office Contact: Height/Weight: Member must try formulary preferred drugs before a request for a non-preferred drug may be considered. If treatment with preferred products has not been successful, you must submit which preferred products have been tried, dates of treatment, and reason for failure. Reasons for failure must meet the Health Plan medical necessity criteria. **Product being requested:**  $\square$  apomorphine hydrochloride injection,  $\square$  Duopa<sup>TM</sup> (levodopa/carbidopa enteral suspension),  $\square$ Kynmobi™, ☐ Neupro® (rotigotine patch), ☐ Nourianz™ (istradefylline), ☐ Ongentys® (opicapone), ☐ Rytary® (carbidopa/levodopa extended release), □ tolcapone, □ Zelapar® (selegiline hydrochloride ODT) Dosing/Frequency:

If the request is for reauthorization, proceed to reauthorization section.						
Questions		Yes	No	Comments/Notes		
1. Does the member have a diagnosis of Parkinson's disease?				Please provide documentation		
2. Is the prescriber a neurologist?						
Has the member had an inadequate response to oral levodopa/carbidopa therapy?				Please provide documentation		
APOMORPHINE HYDROCHLORIDE INJECTION						
1. Is the request for apomorphine hydr	ochloride injection?					
2. Will the member be concurrently tak	king levodopa/carbidopa					
with apomorphine hydrochloride inju	ection therapy?					
3. Is the member experiencing "off" ep	· · · · · · · · · · · · · · · · · · ·			Please provide documentation		
wearing off" and unpredictable "on/	off" episodes) associated					
with advanced Parkinson's disease?						
4. Has the member had a trial and failu	re or			Please provide documentation		
contraindication/intolerance to a pre	eferred dopamine agonist					
(pramipexole, ropinirole), COMT inhi	bitor (entacapone), or					
MAO-B inhibitor (selegiline)?						
<ol><li>Will the member be taking a 5HT3 ar apomorphine hydrochloride injection</li></ol>						

	DUOPA™		
1.	Is the request for Duopa <sup>TM?</sup>		
2.	Is the member responsive to levodopa with defined "on" periods?		Please provide documentation
3.	Is the member experiencing ≥3 hours of "off" episodes despite		Please provide documentation
	maximally tolerated levodopa/carbidopa and one other class of		
	anti-Parkinson's disease therapy (dopamine agonist,		
	pramipexole or ropinirole), COMT inhibitor (entacapone), or		
	MAO-B inhibitor (selegiline)?		
4.	Has the member undergone or has a planned placement of a PEG-J tube?		
	KYNMOBI™		
1.	Is the request for Kynmobi™?		
2.	Will the member be concurrently taking levodopa/carbidopa with Kynmobi™ therapy?		
3.	Is the member experiencing "off" episodes ("end-of-dose		Please provide documentation
	wearing off" and unpredictable "on/off" episodes) associated		·
	with advanced Parkinson's disease?		
4.	Has the member had a trial and failure or		Please provide documentation
	contraindication/intolerance to a preferred dopamine agonist		
	(pramipexole, ropinirole), COMT inhibitor (entacapone), or		
	MAO-B inhibitor (selegiline)?		
5.	Will the member be taking a 5HT3 antagonist concurrently with Kynmobi™?		
	NEUPRO®		
1.	Is the request for Neupro®?		
2.	Is the member unable to take medications by mouth or is oral		Please provide documentation
	therapy clinically inappropriate?		•
3.	Has the member had a trial and failure or		Please provide documentation
	contraindication/intolerance to at least two of the following,		
	one of which must be an extended release product: ropinirole,		
	pramipexole, bromocriptine?		
	NOURIANZ™		
	Is the request for Nourianz™?		
2.	Will the member be concurrently taking levodopa/carbidopa		
	with Nourianz <sup>™</sup> therapy?	 	
3.	Is the member experiencing ≥2 hours of "off" episodes		Please provide documentation
	associated with advanced Parkinson's disease despite		
	maximally tolerated levodopa/carbidopa and two other classes		
	of anti-Parkinson's disease therapy (dopamine agonist, pramipexole or ropinirole), COMT inhibitor (entacapone), or		
	MAO-B inhibitor (selegiline)?		
	ONGENTYS®		
1	Is the request for Ongentys®?		
	Will the member be concurrently taking levodopa/carbidopa		
۷.	with Ongentys® therapy?		
3.	Is the member experiencing ≥2 hours of "off" episodes		Please provide documentation
	associated with advanced Parkinson's disease despite	_	
	maximally tolerated levodopa/carbidopa and two other classes		
	of anti-Parkinson's disease therapy (dopamine agonist,		
	pramipexole or ropinirole), COMT inhibitor (entacapone), or		
	MAO-B inhibitor (selegiline)?		

RYTARY®							
	Is the request for Rytary®?						
2.	Has the member had at least a 3-month trial and failure or			Please provide documentation			
	contraindication to generic extended-release						
	carbidopa/levodopa?						
	TOLCAPONE						
1.	Is the request for tolcapone generic tablets?						
2.	Has the member had a 3-month trial and failure or			Please provide documentation			
	contraindication/intolerance to entacapone or			-			
	levodopa/carbidopa/entacapone?						
3.	Will the member be concurrently taking levodopa/carbidopa						
	with tolcapone therapy?						
	ZELAPAR®	,					
1.	Is the request for Zelapar®?						
	Has the member exhibited deterioration in the quality of their			Please provide documentation			
	response to levodopa/carbidopa?		_	•			
3.	Has the member had a trial and failure or contraindication/			Please provide documentation			
	intolerance to conventional selegiline tablets?			<b>,</b>			
4.	Will the member be concurrently taking levodopa/carbidopa	П					
	with Zelapar® therapy?						
REAUTHORIZATION							
1.	Is the requesting for reauthorization of therapy?						
	Has the therapy shown to be effective with a positive clinical		П	Please provide documentation			
	response?			r rease provide decame maner.			
w	hat medications and/or treatment modalities have been tried in	the pas	st for this	condition? Please document			
	me of treatment, reason for failure, treatment dates, etc.	•					
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Additional information:							
Physician Signature:							

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Policy PHARM-HU-089 Origination Date: 01/01/2022 Reviewed/Revised Date: 03/15/2023 Next Review Date: 03/15/2024 Current Effective Date: 04/01/2023

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