# HEALTHY U CHIP

## PRIOR AUTHORIZATION REQUEST FORM MYASTHENIA GRAVIS

Rystiggo<sup>®</sup>, Soliris<sup>®</sup>, Ultomiris<sup>®</sup>, Vyvgart<sup>®</sup>, Vyvgart<sup>®</sup> Hytrulo

For authorization, please answer each question and fax this form PLUS chart notes back to the Healthy U CHIP Prior Authorization Department at 801-213-1547.

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

If you have medical pharmacy prior authorization questions, please call for assistance: 833-434-4300

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:	HCPCS Code:	

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.

#### Preferred/Non-preferred

- 1. 1<sup>st</sup> line preferred agents:
  - A. Rystiggo<sup>®</sup> (rozanolixizumab-noli) subcutaneous infusion, Vyvgart<sup>®</sup> (efgartigimod alfa-fcab) intravenous infusion
- 2. 2<sup>nd</sup> line non-preferred agents; after trial and failure of the preferred first-line agents:
  - A. Ultomiris® (ravulizumab) intravenous infusion
- 3. Excluded/Not covered unless failure or contraindication to all other agents:
  - A. Soliris<sup>®</sup> (eculizumab) intravenous infusion; Vyvgart<sup>®</sup> Hytrulo (efgartigimod alfa/hyaluronidase) subcutaneous infusion

Product being requested: \_\_\_\_\_

Dosing/Frequency:\_\_\_\_\_

If the request is for reauthorization, proceed to reauthorization section						
Questions		Yes	No	Comments/Notes		
MYASTHENIA GRAVIS (gMG)						
1.						
	neurologist or other specialist in the treatment of gMG?					
2.	Does the member have a diagnosis of gMG?					
3.	Does the member have a positive serologic test for anti-			Please provide documentation		
	acetylcholine receptor (anti-AchR) antibodies?					
4.	If the request is for Rystiggo <sup>®</sup> , does the member have a positive			Please provide documentation		
	serologic test for anti-acetylcholine receptor (anti-AchR)					
	antibodies OR anti-muscle-specific kinase (anti-MuSK)					
	antibodies?					
5.	Has the member been diagnosed with class II to IV gMG			Please provide documentation		
	according to the Myasthenia Gravis Foundation of America?					

6.	Has the member tried and failed pyridostigmine AND at least			Please provide documentation	
	two immunosuppressive therapies (e.g. rituximab,				
	methotrexate, mycophenolate mofetil, azathioprine,				
-	cyclosporine) for a total duration of at least 12 months?				
7.	Has the member tried and failed intravenous immunoglobulin (IVIG)?			Please provide documentation	
8.	Will the requested therapy be used in combination with IVIG or				
0.	other biologic agents for gMG treatment?				
9.	If the request is for Rystiggo <sup>®</sup> , is the member's Myasthenia			Please provide documentation	
	Gravis Activities of Daily Living (MG-ADL) score $\geq$ 3?				
10.	If the request is for Vyvgart <sup>®</sup> , is the member's MG-ADL score ≥ 5?			Please provide documentation	
11.	If the request is for Soliris <sup>®</sup> or Ultomiris <sup>®</sup> , is the member's MG-ADL score $\geq 6$ ?			Please provide documentation	
12.	If the request is for Soliris <sup>®</sup> or Ultomiris <sup>®</sup> , is the prescribing				
	physician enrolled in Soliris <sup>®</sup> or Ultomiris <sup>®</sup> REMS program?				
REAUTHORIZATION					
1.	Is the request for reauthorization of therapy?				
2.	If the request is for reauthorization of Vyvgart <sup>®</sup> or Rystiggo <sup>®</sup> , has			Please provide documentation	
	the member had a positive clinical response to treatment shown				
	by a $\geq$ 2 points reduction in MG-ADL score?				
3.	If the request is for reauthorization of Soliris <sup>®</sup> or Ultomiris <sup>®</sup> , has			Please provide documentation	
	the member had a positive clinical response to treatment shown by $a \ge 2$ points reduction in MC ADL score or $a \ge 2$ points				
	by $a \ge 2$ points reduction in MG-ADL score or $a \ge 3$ points				
\ <b>\</b> /b	reduction in quantitative myasthenia gravis (QMG) score? The medications and/or treatment modalities have been tried in the second states in the second second second sec	o nast	for this	condition? Plaze document	
	ne of treatment, reason for failure, treatment dates, etc.	ie past		condition: Flease document	
IIai	ne of treatment, reason for failure, treatment dates, etc.				
Additional information:					
Physician's Signature:					

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Policy: PHARM-CHIP-M046 Origination Date: 07/01/2024 Reviewed/Revised Date: Next Review Date: Current Effective Date: 07/01/2024

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