

PHARMACY PRIOR AUTHORIZATION REQUEST FORM **SOLIRIS®**, **ULTOMIRIS®**

For authorization, please answer each question and fax this form PLUS chart notes back to the U of U Health Plans 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: Individual Exchange: 833-981-0214, Commercial Groups: 833-981-0213, MHC: 844-262-1500 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. **Product being requested:** □ Soliris® (eculizumab), □ Ultomiris® (ravilizumab) Dosing/Frequency: If the request is for reauthorization, proceed to reauthorization section Questions Yes No **Comments/Notes** PAROXYSMAL NOCTURNAL HEMOGLOBINURIA (PNH) 1. Is the member 18 years of age or older? 2. Is the diagnosis of Paroxysmal Nocturnal Hemoglobinuria (PNH) Please provide documentation П confirmed via flow cytometry? 3. Is the member transfusion dependent requiring at least four Please provide documentation transfusions in the past 12 months? 4. Does the member have a history of a major thrombotic event? Please provide documentation 5. Does the member have high lactate dehydrogenase (LDH) Please provide documentation activity with serum levels ≥1.5 times the upper limit of normal and have clinical symptoms? 6. Has the member had Neisseria meningitidis vaccination at least Please provide documentation 2 weeks prior to start date? 7. Is the prescribing physician enrolled in the Soliris® or Ultomiris® П П Risk Evaluation and Mitigation Strategies (REMS) program? 8. If the request is for Soliris®, has the member tried and failed Please provide documentation Ultomiris®, unless contraindicated? **ATYPICAL HEMOLYTIC UREMIC SYNDROME (aHUS)** 1. Does the member have a diagnosis of Atypical Hemolytic Uremic Syndrome (aHUS)?

2.	Has Shiga toxin-related hemolytic uremic syndrome been ruled out?			Please provide documentation		
3.	Does the member have a normal ADAMTS-13 level?			Please provide documentation		
4.	Has the member had the Neisseria meningitidis vaccination?			Please provide documentation		
5.	Is the prescribing physician enrolled in Soliris® or Ultomiris® REMS program?			-		
6.	If the request is for Soliris®, has the member tried and failed Ultomiris®, unless contraindicated?			Please provide documentation		
MYASTHENIA GRAVIS (gMG)						
1.	Does the member have a diagnosis of Myasthenia Gravis (gMG)?					
2.	Is the member 18 years of age or older?					
3.	Does the member have a positive serologic test for anti- acetylcholine receptor (AchR) antibodies?			Please provide documentation		
4.	Has the member been diagnosed with class II to IV gMG			Please provide documentation		
	according to the Myasthenia Gravis Foundation of America?					
	Is the member's Myasthenia Gravis Activities of Daily Living (MG-ADL) score > 6?			Please provide documentation		
6.	Has the member tried and failed at least two			Please provide documentation		
	immunosuppressive therapies (e.g. methotrexate,					
	corticosteroids, azathioprine, or cyclosporine) for a total duration of at least one year?					
7.	Has the member tried and failed at least one			Please provide documentation		
	immunosuppressive therapy and required chronic			P		
	plasmapheresis or IVIG for a total duration of at least one year?					
8.	Has the member had Neisseria meningitidis vaccination at least			Please provide documentation		
	2 weeks prior to start date?					
	Is the prescribing physician enrolled in Soliris® or Ultomiris® REMS program?					
10.	If the request is for Soliris®, has the member tried and failed					
	Ultomiris®, unless contraindicated? NEUROMYELITIS OPTICA SPECTRUM DIS	CORDE	D /NIN4C) (20)		
1	Is the prescribing provider a neurologist who specializes in			וטפע		
1.	treating NMOSD?					
2.	Does the member have a confirmed diagnosis of NMOSD			Please provide documentation		
	including both:			•		
	 Anti-aquaphorin-4 (AQP4) positive 					
	At least one of the core clinical characteristics					
3.	Has the member had at least one relapse requiring rescue			Please provide documentation		
	therapy in the last 12 months or two or more relapses requiring rescue therapy in the last 24 months?					
4.	Is there documentation of an Expanded Disability Status Score		П	Please provide documentation		
	(EDSS) of ≤8?			,		
5.	Has the member had an adequate trial and failure of Enspryng®,			Please provide documentation		
	Ruxience® AND Uplizna™?					
6.	Is the prescribing physician enrolled in Soliris® REMS program?					
REAUTHORIZATION TO THE PROPERTY OF THE PROPERT						
1.	Is the request for reauthorization of therapy?			Diagona anno del del constanti		
2.	Reauthorization of PNH treatment : Has a clinically significant response been demonstrated (e.g. decrease in LDH from			Please provide documentation		
	baseline, improvement in hemoglobin, or decrease in red blood					
	cell transfusion frequency)?					

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3.	Reauthorization of aHUS treatment: Has a clinically significant			Please provide documentation		
	response been demonstrated (e.g. decrease in LDH,					
	improvement in SCr/eGFR, increase in platelet count, or					
	decrease in plasmapheresis frequency from baseline)?					
4.	Reauthorization of gMG treatment: Has a clinically significant			Please provide documentation		
	response been demonstrated (e.g. MG-ADL score reduction of 2					
	points or more, QMG score reduction of 3 points or more)?					
5.	Reauthorization of NMOSD treatment: Has a clinically			Please provide documentation		
	significant response been demonstrated (e.g. decrease in					
	relapse rate, improvement or stabilization of symptoms					
	associated with relapse, improvement in EDSS score)?					
Wh	at medications and/or treatment modalities have been tried in the	ne past	for this	condition? Please document		
name of treatment, reason for failure, treatment dates, etc.						
Add	litional information:					
Dhy	rsician's Signature:					
гну	Siciali S Signature.					

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Policy: PHARM-M013

Origination Date: 12/29/2017 Reviewed/Revised Date: 2/17/2023 Next Review Date: 2/17/2024 Current Effective Date: 3/1/2023

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