

If this is an URGENT request, please call (800) 882-4462 (800.88.CIGNA)

## Bkemv (eculizumab-aeeb) Epysqli (eculizumab-aagh) Soliris (eculizumab)

PHYSICIAN INFORMATION			PATIENT INFORMATION				
* Physician Name:			*Due to privacy regulations we will not be able to respond via fax with the outcome of our review unless all asterisked (*) items on				
Specialty:	* DEA, NPI or TIN:			this form are completed. *			
Office Contact Person:			* Patient Name:				
Office Phone:			* Cigna ID:		* Date of Birth:		
Office Fax:			* Patient Street Address:				
Office Street Address:			City:	State:		Zip:	
City:	State:	Zip:	Patient Phone:				
Urgency: ☐ Standard	☐ Urge		ox, I attest to the fact that applying the standard review time frame may the customer's life, health, or ability to regain maximum function)				
Medication Requested:	Bkemv	☐ Epysql	qli 🔲 Soliris				
Dose:		Frequency of therap	py: Duration of therapy:				
J-Code:		ICD10:					
Will the patient be taking the requested medication in combination with Empaveli (pegcetacoplan subcutaneous injection) for greater than 4 weeks? ☐ Yes ☐ No							
Will the patient be taking the requested medication concomitantly with another complement Inhibitor (except Voydeya [danicopan tablets])? Please Note: Examples of complement inhibitors are Fabhalta (iptacopan capsules), PiaSky (crovalimab-akkz intravenous infusion or subcutaneous injection), and Ultomiris (ravulizumab-cwvz intravenous infusion).							
Will the patient be taking the requested medication concomitantly with a rituximab product, a Neonatal Fc Receptor Blocker, Zilbrysq (zilucoplan subcutaneous injection), Enspryng (satralizumab-mwge subcutaneous injection), or Uplizna (inebilizumabcdon intravenous infusion)? Please Note: Examples of Neonatal Fc receptor blockers are Imaavy (nipocalimab-aahu intravenous infusion), Rystiggo (rozanolixizumab-noli subcutaneous infusion), Vyvgart (efgartigimod alfa-fcab intravenous infusion), and Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc subcutaneous injection).							
Is this a new start or continuation of therapy**? If your patient has already begun treatment with samples, please choose "new start of therapy".   ☐ new start of therapy ☐ continuation of therapy:							
Where will this medication be obtained?  Accredo Specialty Pharmacy**  Hospital Outpatient Retail pharmacy Other (please specify):			☐ Home Health / Home Infusion vendor ☐ Physician's office stock (billing on a medical claim form)  **Cigna's nationally preferred specialty pharmacy				
**Medication orders can be placed with Accredo via E-prescribe - Accredo (1620 Century Center Pkwy, Memphis, TN 38134-8822   NCPDP 4436920), Fax 888.302.1028, or Verbal 866.759.1557							
Facility and/or doctor di Facility Name: Address (City, State, Zip Co		d administering m State:	nedication:	Tax ID#:			
Where will this drug be ☐ Patient's Home ☐ Hospital Outpatient	administered	?		sician's Office r (please spe			
NOTE: Per some Cigna plans, infusion of medication MUST occur in the least intensive, medically appropriate setting.							

Is this patient a candidate for re-direction to an alternate setting (such as alternate infusion site, physician's office, ho assistance of a Specialty Care Options Case Manager?	
Is the requested medication for a chronic or long-term condition for which the prescription medication may be necess the patient?	ary for the life of ☐ Yes ☐ No
What is your patient's diagnosis?  atypical hemolytic uremic syndrome generalized myasthenia gravis paroxysmal nocturnal hemoglobinuria neuromyelitis optica spectrum disorder All other indications	
Clinical Information	
If aHUS:	
Does the patient have Shiga toxin Escherichia coli related hemolytic uremic syndrome?	Yes 🗌 No 🗌
Is the requested medication being prescribed by or in consultation with a nephrologist?	Yes 🗌 No 🗌
If MG:	
Is the requested medication prescribed by or in consultation with a neurologist?	Yes 🗌 No 🗌
Is the patient currently receiving the requested medication?	Yes 🗌 No 🗌
(if not currently receiving) Is documentation being provided that the patient has confirmed anti-acetylcholine receptor generalized myasthenia gravis? - Please note: Documentation may include, but is not limited to, chart notes, laborate records, and/or other information. Medical documentation specific to your response to this question must be attached your request could be denied.	ory tests, claims
(if 18 years or older and not currently receiving) Does the patient have Myasthenia Gravis Foundation of America clast IV?	ssification of II to Yes  □ No □
(if 18 years or older and not currently receiving) Does the patient have a Myasthenia Gravis Activities of Daily Living of greater than or equal to 6?	(MG-ADL) score Yes
(if not currently receiving) Has the patient previously received pyridostigmine OR is the patient currently receiving pyridostigmine OR is the patient currently receiving pyridostigmine.	
(if no) Has the patient had inadequate efficacy, a contraindication, or significant intolerance to pyridostigmin	
(if not currently receiving) Has the patient previously received or is the patient currently receiving two different immun therapies for greater than or equal to 1 year? - Please Note: Examples of immunosuppressant therapies tried include azathioprine, cyclosporine, mycophenolate mofetil, methotrexate, tacrolimus, cyclophosphamide.	
(if no) Has the patient had inadequate efficacy, a contraindication, or significant intolerance to two different immunosuppressant therapies? - Please Note: Examples of immunosuppressant therapies tried include cortazathioprine, cyclosporine, mycophenolate mofetil, methotrexate, tacrolimus, cyclophosphamide.	ticosteroid, Yes
(if not currently receiving) Does the patient have evidence of unresolved symptoms of generalized myasthenia gravis Evidence of unresolved symptoms of generalized myasthenia gravis includes difficulty swallowing, difficulty breathing disability resulting in the discontinuation of physical activity (for example, double vision, talking, impairment of mobility	g, and a functional
(if currently receiving) According to the prescriber, is the patient continuing to derive benefit from eculizumab? Please of benefit include reductions in exacerbations of myasthenia gravis; improvements in speech, swallowing, mobility, are function.	
If NMOSD:	
Is this medication prescribed by, or in consultation with, a neurologist?	Yes 🗌 No 🗌
Is the patient currently receiving the requested medication?	Yes 🗌 No 🗌

(if, NMOSD, initial therapy) Is documentation being provided that the patient's diagnosis anti-aquaporin-4 antibody -poneuromyelitis optical spectrum disorder was confirmed by a blood serum test? - Please note: Documentation may inclimited to, chart notes, laboratory tests, claims records, and/or other information. Medical documentation specific to you this question must be attached to this case or your request could be denied.	ude, but is not				
(if NMOSD, currently receiving) Was the diagnosis of anti-aquaporin-4 antibody-positive neuromyelitis optica spectrum confirmed by a blood serum test?	n disorder Yes				
(if currently receiving) Has the patient had a clinical benefit from the use of eculizumab, according to the prescriber? Please Note: Examples of clinical benefit include reductions in relapse rate, reduction in symptoms (for example, pain, fatigue, motor function), or a slowing progression in symptoms.  Yes  No					
If PNH:					
Is this medication prescribed by or in consultation with a hematologist?	Yes 🗌 No 🗌				
Is the patient currently receiving the requested medication?	Yes 🗌 No 🗌				
(if not currently receiving) Is documentation being provided that the paroxysmal nocturnal hemoglobinuria diagnosis was confirmed by peripheral blood flow cytometry results showing the absence or deficiency of glycosylphosphatidylinositol (GPI)-anchored proteins on at least two cell lineages? - Please note: Documentation may include, but is not limited to, chart notes, laboratory tests, claims records, and/or other information. Medical documentation specific to your response to this question must be attached to this case or your request could be denied.					
(if currently receiving) According to the prescriber, is the patient continuing to derive benefit from eculizumab? Please of benefit include stabilization of hemoglobin levels, decreased transfusion requirements or transfusion independence hemolysis, improvement in Functional Assessment of Chronic Illness Therapy (FACIT)-Fatigue score.					
Additional pertinent information:					
Attestation: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan or insurer its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.					
Prescriber Signature: Date:					
Save Time! Submit Online at: <a href="https://www.covermymeds.com/main/prior-authorization-forms/cigna/">www.covermymeds.com/main/prior-authorization-forms/cigna/</a> or via SureScrip	ots in your EHR.				

Our standard response time for prescription drug coverage requests is 5 business days. If your request is urgent, it is important that you call us to expedite the request. View our Prescription Drug List and Coverage Policies online at cigna.com.

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