

Vyvgart and Vyvgart Hytrulo (efgartigimod alfa-fcab; efgartigimod alfa and hyaluronidase-qvfc) PAM-052

Iowa Medicaid Program:	Prior Authorization	Effective Date:	07/01/2022
Revision Number:	2	Last Rev Date:	01/19/2024
Reviewed By:	Medicaid Medical Director	Next Rev Date:	01/17/2025
Approved By:	Medicaid Clinical Advisory Committee	Approved Date:	01/20/2023

Overview

Medication:	efgartigimod alfa-fcab¹	efgartigimod alfa and hyaluronidase-qvfc ²		
Brand Name:	Vyvgart [®]	Vyvgart Hytrulo®		
Pharmacologic Category:	Neonatal Fc receptor blocker	Neonatal Fc receptor blocker + endoglycosidase		
FDA-Approved Indication(s):	Treatment of generalized myasthenia gravis (gMG) in adult patients who are antiacetylcholine receptor (AChR) antibody positive			
How Supplied:	Single-dose vial400 mg/20 mL	Single-dose vial 1,008 mg efgartigimod alfa and 11,200 units hyaluronidase per 5.6 mL		
Dosage and Administration:	 Intravenous (IV) infusion I0 mg/kg once weekly for 4 weeks* * Weight ≥ 120 kg: 1,200 mg per dose 	Subcutaneous (SC) injection by a healthcare professional only 1,008 mg/11,200 units SC injection once weekly for 4 weeks		
 Administer subsequent treatment cycles based on clinical evaluation. The safety of initiating subsequent cycles sooner than 50 days from the previous treatment cycle has not been established. 		es sooner than 50 days from the start of the		
Benefit Category:	Medical			

Descriptive Narrative

Neuromuscular junction (NMJ) disorders comprise several dysfunctions that ultimately lead to muscle weakness. Some of these diseases, such as congenital myasthenic syndromes, are genetic. Other NMJ disorders are acquired autoimmune forms, such as myasthenia gravis, the most prevalent neuromuscular junction disorder. Myasthenia gravis (MG) is characterized by muscle weakness and fatigue. The weakness is due to an antibody-mediated immunologic attack directed at proteins in the postsynaptic membrane of the neuromuscular junction.

Myasthenia gravis can affect all ages, but it is considered "a disease of young women and old men." The most common onset age is between 20 and 39 years in women and between 50 and 70 years in men.³ Autoimmune myasthenia gravis has a reported worldwide prevalence of 40

to 180 per million people, and an annual incidence of 4-12 per million people. In the United States, there are an estimated 60,000 patients diagnosed with myasthenia gravis.

Autoimmune myasthenia gravis is characterized by the presence of antibodies against several components of the neuromuscular junctions. The most common antibody found in autoimmune myasthenia gravis is directed against post-synaptic acetylcholine receptors (AChRs). Anti-AChR antibodies are present in approximately 80 percent of all autoimmune MG patients. Less frequent autoantibodies found in autoimmune MG include the anti-muscle-specific kinase (MuSK) antibody (4 percent of the cases) and the anti-lipoprotein receptor-related protein-4 (LRP4) antibody (2 percent of the cases) directed against LPR4. All these autoantibodies belong to the immunoglobulin G (IgG) class.⁴

Guidelines

The International Consensus Guidance for Management of Myasthenia Gravis was last updated in 2020. Vyvgart® and Vyvgart Hytrulo® are not yet included in the consensus guidance.

Goals of Therapy

The goals of therapy in myasthenia gravis (MG) are to render patients minimally symptomatic or better while minimizing side effects from medications. There are four primary therapies currently used to treat MG. The time of onset of clinical effect of each of these therapies for varies considerably, which plays a large role, in addition to the pace and severity of the disease, in choosing the appropriate therapy for a given patient. ⁶

Commonly used therapies for mya (estimated times are rough guidelines by	Time to onset of effect	Time to maximal effect	
Symptomatic therapy — increases the amount of acetylcholine available at the neuromuscular junction	pyridostigmine	10-15 minutes	2 hours
Chronic immunotherapies	prednisone	2 to 3 weeks	5 to 6 months
– target underlying immune	azathioprine	~12 months	I to 2 years
dysregulation	mycophenolate mofetil	6 to 12 months	I to 2 years
	cyclosporine	~6 months	~7 months
	tacrolimus	~6 months	~12 months
	efgartigimod alfa	I to 2 weeks	~4 weeks
	ravulizumab	I to 2 weeks	~4 to 10 weeks
Rapid immunotherapies	plasmapheresis	I to 7 days	I to 3 weeks
– manage myasthenia crisis	intravenous immune globulin (IVIG)	I to 2 weeks	I to 3 weeks
Surgery — therapeutic option in select patients	thymectomy	I to I0 years	I to 10 years

Outcomes Measurements⁸

The Myas	thenia Gravis Foundation of America Classification (MGFA)		
MGFA cla	MGFA classification separates patients in groups based on disease severity and the localization of the symptoms.		
Class I	Any ocular muscle weakness; may have weakness of eye closure. All other muscle strength is normal.		
Class II	Mild weakness affecting muscles other than ocular, <u>+</u> ocular muscle weakness of any severity.		
	IIa. Predominantly affecting limb, axial muscles, or both. May also have lesser involvement of		
	oropharyngeal muscles.		
	Ilb. Predominantly affecting oropharyngeal, respiratory muscles, or both. May also have lesser or		
	equal involvement of limb, axial muscles, or both.		
Class III	Moderate weakness affecting muscles other than ocular, \pm ocular muscle weakness of any severity.		
	IIIa. Predominantly affecting limb, axial muscles, or both. May also have lesser involvement of		
	oropharyngeal muscles.		
	IIIb. Predominantly affecting oropharyngeal, respiratory muscles, or both. May also have lesser or		
	equal involvement of limb, axial muscles, or both.		
Class IV	Severe weakness affecting muscles other than ocular, \pm ocular muscle weakness of any severity.		
	IVa. Predominantly affecting limb, axial muscles, or both. May also have lesser involvement of		
	oropharyngeal muscles.		
	IVb. Predominantly affecting oropharyngeal, respiratory muscles, or both. May also have lesser or		
	equal involvement of limb, axial muscles, or both.		
Class V	Defined as intubation, with or without mechanical ventilation, except when employed during routine		
	postoperative management. The use of a feeding tube without intubation places the patient in class IVb.		

The Myasthenia Gravis Activities of Daily Living (MG-ADL)

The MG-ADL assesses the impact of generalized myasthenia gravis (gMG) on daily functions by assigning a score to 8 signs or symptoms that are commonly affected in gMG, then totaling the individual scores for a composite MG-ADL score. Each item is measured on a 4-point scale, where a score of 0 represents normal function and a score of 3 represents the loss of ability to perform that function.

Grade	0	l 2		3
I. Talking	Normal	Intermittent slurring or	Constant slurring or nasal,	Difficult to
		nasal speech	but can be understood	understand speech
2. Chewing	Normal	Fatigue with solid food	Fatigue with soft food	Gastric tube
3. Swallowing	Normal	Rare episode of choking	Frequent choking necessitating changes in diet	Gastric tube
4. Breathing	Normal	Shortness of breath with exertion	Shortness of breath at rest	Ventilator dependence
5. Impairment of ability to brush teeth or comb hair	None	Extra effort, but no rest periods needed	Rest periods needed	Cannot do one of these functions
6. Impairment of ability to arise from a chair	None	Mild, sometimes uses arms	Moderate, always uses arms	Severe, requires assistance
7. Double vision	None	Occurs, but not daily	Daily, but not constant	Constant
8. Eyelid droop	None	Occurs, but not daily	Daily, but not constant	Constant

Criteria

Prior authorization is required.

Vyvgart® or Vyvgart Hytrulo® are considered medically necessary when <u>ALL</u> of the following are met:

- Documented diagnosis of acetylcholine receptor antibody-positive (AChR-Ab+) generalized myasthenia gravis (gMG); <u>AND</u>
- 2. Member has Myasthenia Gravis Foundation of America (MGFA) Clinical Classification of Class II to IV; **AND**
- 3. Member has a Myasthenia Gravis Activities of Daily Living (MG-ADL) score of at least 5 or higher, with at least 50 percent of the baseline MG-ADL score due to non-ocular symptoms; **AND**
- 4. Member is 18 years of age or older; **AND**
- 5. Member is on a stable dose (for the duration specified, if indicated) of at least ONE (or documentation of inadequate response, intolerance, or labeled contraindication to ALL) of the following standard-of-care treatments for gMG:
 - a. Acetylcholine inhibitors (e.g., pyridostigmine); and/or
 - b. Steroids (at least 3 months of treatment); and/or
 - c. Non-steroidal immunosuppressive therapy, e.g., azathioprine, cyclophosphamide, cyclosporine, mycophenolate mofetil, or tacrolimus (at least 6 months of treatment); **AND**
- 6. Vyvgart[®] or Vyvgart Hytrulo[®] are not prescribed concurrently with eculizumab (Soliris[®]), rituximab, ravulizumab (Ultomiris[®]), or maintenance immunoglobulin treatment; **AND**
- 7. Prescribed by, or in consultation with, an immunologist, neurologist, or rheumatologist; **AND**
- 8. Request meets one of the following (a or b):
 - a. The regimen prescribed is within the FDA-approved labeling:
 - i. Vyvgart®: Dose does not exceed 10 mg/kg (1,200 mg per infusion for members weighing 120 kg or more) once weekly for 4 weeks; or
 - ii. Vyvgart Hytrulo[®]: Dose does not exceed 1,008 mg/11,200 units subcutaneously once weekly for 4 weeks; **OR**
 - b. Regimen is supported by clinical practice guidelines. Supporting clinical documentation must be provided with any request for which the regimen or dosage prescribed does not align with FDA-approved labeling.

Vyvgart® or Vyvgart Hytrulo® are considered medically necessary for continuation of therapy when **ALL** of the following are met:

- I. Member is currently receiving medication through the Iowa Medicaid benefit or has previously met initial approval criteria; **AND**
- Documentation of positive clinical response to therapy, as demonstrated by at least a 2-point reduction in the total Myasthenia Gravis Activities of Daily Living (MG-ADL) score from pre-treatment baseline; <u>AND</u>
- 3. Vyvgart[®] or Vyvgart Hytrulo[®] are not prescribed concurrently with eculizumab (Soliris[®]), rituximab, ravulizumab (Ultomiris[®]), or maintenance immunoglobulin treatment; **AND**
- 4. Prescribed by, or in consultation with, an immunologist, neurologist, or rheumatologist; **AND**
- 5. Request meets one of the following (a or b):
 - a. The regimen prescribed is within the FDA-approved labeling:
 - i. Vyvgart[®]: Dose does not exceed 10 mg/kg (1,200 mg per infusion for members weighing 120 kg or more) once weekly for 4 weeks **AND** at least 50 days have passed since the start of the previous cycle; or
 - ii. Vyvgart Hytrulo[®]: Dose does not exceed 1,008 mg/11,200 units subcutaneously once weekly for 4 weeks **AND** at least 50 days have passed since the start of the previous cycle; **OR**
 - b. Regimen is supported by clinical practice guidelines. Supporting clinical documentation must be provided with any request for which the regimen or dosage prescribed does not align with FDA-approved labeling.

Approval Duration and Quantity Limits

	Initial Authorization	Subsequent Authorization(s)
Approval Duration	6 months	12 months
Quantity Limits - Vyvgart® -	Dose not to exceed 10 mg/kg (1,200 mg per infusion for members 120 kg or more): administered as an IV infusion once weekly for 4 weeks; AND (for subsequent cycles) at least 50 days have passed since the start of the previous cycle	
Quantity Limits - Vyvgart Hytrulo® -	Dose not to exceed 1,008 mg/11,200 units: administered subcutaneously once weekly for 4 weeks; AND (for subsequent cycles) at least 50 days have passed since the start of the previous cycle	

Coding and Product Information

The following list(s) of codes and product information are provided for reference purposes only and may not be all inclusive. The inclusion of a code does not imply any right to reimbursement or guarantee claim payment, nor does the exclusion of a code imply that its association to the HCPCS code is inappropriate.

HCPCS	Description
J9332	Injection, efgartigimod alfa-fcab, 2 mg [Vyvgart®]
J9334	Injection, efgartigimod alfa, 2 mg and hyaluronidase-qvfc [Vyvgart Hytrulo®] – code effective 1/1/2024

ICD-10	Description
G70.00	Myasthenia gravis without (acute) exacerbation
G70.01	Myasthenia gravis with (acute) exacerbation

NDC	Labeler	Dosage	Pkg Size	Pkg Qty	Units/Pkg
73475-3041-05 [Vyvgart [®]]	argenx US, Inc.	2 mg	I	EA	200
73475-3102-03 [Vyvgart Hytrulo®]	argenx US, Inc.	2 mg		EA	504

Compliance

- I. Should conflict exist between this policy and applicable statute, the applicable statute shall supersede.
- 2. Federal and State law, as well as contract language, including definitions and specific contract provisions or exclusions, take precedence over medical policy and must be considered first in determining eligibility for coverage.
- 3. Medical technology is constantly evolving, and Iowa Medicaid reserves the right to review and update medical policy on an annual or as-needed basis.

Medical necessity guidelines have been developed for determining coverage for member benefits and are published to provide a better understanding of the basis upon which coverage decisions are made. Medical necessity guidelines are developed for selected physician-administered medications found to be safe and proven to be effective in a limited, defined population or clinical circumstances. They include concise clinical coverage criteria based on current literature review, consultation with practicing physicians in the service area who are medical experts in the particular field, FDA and other government agency policies, and standards adopted by national accreditation organizations. Criteria are revised and updated annually, or more frequently if new evidence becomes available that suggests needed revisions.

References

Vyvgart prescribing information (04/2022). argenx US, Inc.: Boston, MA. Available online at www.vyvgarthcp.com. Accessed November 16, 2023.

² Vyvgart Hytrulo prescribing information (06/2023). argenx US, Inc.: Boston, MA. Available online at www.vyvgarthcp.com. Accessed July 31, 2023.

³ Bubuioc AM, Kudebayeva A, Turuspekova S, Lisnic V, Leone MA. The epidemiology of myasthenia gravis. J Med Life. 2021 Jan-Mar;14(1):7-16. PMID: 33767779.

⁴ Gilhus NE, Verschuuren JJ. Myasthenia gravis: subgroup classification and therapeutic strategies. Lancet Neurol. 2015 Oct;14(10):1023-36. PMID: 26376969.

Development of utilization management criteria may also involve research into other state Medicaid programs, other payer policies, consultation with experts and review by the Medicaid Clinical Advisory Committee (CAC). These sources may not be referenced individually unless they are specifically published and are otherwise applicable to the criteria at issue.

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Criteria Chan	ge History	
Change Date	Changed By	Description of Change Version
[mm/dd/yyyy]	CAC	
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Change Date	Changed By	Description of Change Version
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Change Date	Changed By	Description of Change Version
01/19/2024	CAC	Annual review. New SC formulation Vyvgart Hytrulo®, updated policy to
		include this formulation. Updated criteria with dosing information.
		Updated criteria, quantity limits, approval duration, and coding information to include Vyvgart Hytrulo [®] .
Signature		0.0000000000000000000000000000000000000
William (Bill) Jag	iello, DO	1000000
Change Date	Changed By	Description of Change Version
01/20/2023	CAC	Criteria implementation.
Signature		0.0000000000000000000000000000000000000
William (Bill) Jag	iello, DO	1000000gg

CAC = Medicaid Clinical Advisory Committee

⁵ Narayanaswami P, Sanders DB, et al. International Consensus Guidance for Management of Myasthenia Gravis: 2020 Update. Neurology. 2021 Jan 19;96(3):114-122. Epub 2020 Nov 3. PMID: 33144515.

⁶ Bird SJ. Overview of the treatment of myasthenia gravis. Goddeau RP, ed. UpToDate. Waltham, MA: UpToDate Inc. www.uptodate.com. Accessed December 9, 2023.

⁷ Bird SJ. Chronic immunotherapy for myasthenia gravis. Goddeau RP, ed. UpToDate. Waltham, MA: UpToDate Inc. <u>www.uptodate.com</u>. Accessed December 9, 2023.

⁸ Barnett C, Herbelin L, Dimachkie MM, Barohn RJ. Measuring Clinical Treatment Response in Myasthenia Gravis. Neurol Clin. 2018 May;36(2):339-353. PMID 29655453.