

PHARMACY / MEDICAL POLICY - 5.01.564

Pharmacotherapy of Miscellaneous Autoimmune Diseases

BCBSA Ref. Policy: 5.01.39

Effective Date: July 1, 2025*

Last Revised: Feb. 11, 2025
Replaces: Extracted from

5.01.550

*This policy has been updated. Click here to view the current

policy.

RELATED MEDICAL POLICIES:

5.01.550 Pharmacotherapy of Arthropathies

5.01.556 Rituximab: Non-oncologic and Miscellaneous Uses5.01.563 Pharmacotherapy of Inflammatory Bowel Disorder

5.01.575 C5 Complement Inhibitors

11.01.523 Site of Service: Infusion Drugs and Biologic Agents

The Site of Service Medical Necessity criteria within this policy DOES NOT apply to Alaska fully-insured members; refer to the infusion drug Medical Necessity criteria only.

Site of Service and the infusion drug Medical Necessity criteria apply to all other plan members.

Please contact Customer Service for more information.

Select a hyperlink below to be directed to that section.

POLICY CRITERIA | DOCUMENTATION REQUIREMENTS | CODING RELATED INFORMATION | EVIDENCE REVIEW | REFERENCES | HISTORY

Clicking this icon returns you to the hyperlinks menu above.

Introduction

The term "autoimmune disorders" refers to a number of conditions where a person's immune system is activated against a part of their body. Many of these diseases are grouped together based on what part of the body is affected. The cells involved are usually lymph cells, and disease develops consistent with long standing inflammation. Common autoimmune disorders include certain types of arthritis, some skin diseases, inflammatory bowel diseases and others. This policy discusses treatment for the following autoimmune diseases: hidradenitis suppurativa, systemic lupus erythematosus (lupus), pyoderma gangrenosum, Behcet's disease, giant cell arteritis, uveitis, neuromyelitis optica spectrum disorder, periodic fever syndromes, Still's disease,

recurrent pericarditis, deficiency of interleukin-1 receptor antagonist, primary immunoglobulin A nephropathy (IgAN), and other conditions. The policy describes which drugs need to be preapproved before they are covered by the plan.

Note: The Introduction section is for your general knowledge and is not to be taken as policy coverage criteria. The rest of the policy uses specific words and concepts familiar to medical professionals. It is intended for providers. A provider can be a person, such as a doctor, nurse, psychologist, or dentist. A provider also can be a place where medical care is given, like a hospital, clinic, or lab. This policy informs providers about when a service may be covered.

Policy Coverage Criteria

Site of Service Medical Necessity criteria does NOT apply to Alaska fully-insured members; refer to the infusion drug Medical Necessity criteria only. Please contact Customer Service for more information.

We will review specific intravenous (IV) and injectable drugs for medical necessity for all ages.

For individuals aged 13 and older, we also will review the site of service for medical necessity. Site of service is defined as the location where the drug is administered, such as a hospital-based outpatient setting, an infusion center, a physician's office, or at home.

Drugs subject to site of service review addressed in this policy are:

- Actemra (tocilizumab) IV
- Avsola (infliximab-axxq)
- Benlysta (belimumab)
- Inflectra (infliximab-dyyb)
- Infliximab (Janssen unbranded)
- Remicade (infliximab)
- Renflexis (infliximab-abda)
- Tofidence (tocilizumab-bavi) IV

• Uplizna (inebilizumab-cdon)

Note: Medications listed in this policy may also be subjected to quantity limits per the FDA labeled dosing.

Click on the links below to be directed to the related medical necessity criteria:

Behcet's Disease

Chronic Inflammatory Demyelinating Polyneuropathy (CIDP)

Cytokine Release Syndrome

Giant Cell Arteritis

Hidradenitis Suppurativa (HS)

Pyoderma Gangrenosum

Site of Service

Systemic Lupus Erythematosus (SLE) & Lupus Nephritis

Uveitis

Neuromyelitis Optica Spectrum Disorder (NMOSD)

Deficiency of Interleukin-1 Receptor Antagonist (DIRA)

Recurrent Pericarditis

Periodic Fever Syndromes & Still's Disease

Graft Versus Host Disease

Myasthenia Gravis

Primary Immunoglobulin A Nephropathy (IgAN)

Sarcoidosis

Site of Service	Medical Necessity
Administration	
Medically necessary sites of service • Physician's office • Infusion center • Home infusion	 IV infusion therapy of various medical or biologic agents will be covered in the most appropriate, safe, and cost-effective site: These are the preferred medically necessary sites of service for specified drugs.
Hospital-based outpatient setting Outpatient hospital IV infusion department	IV infusion therapy of various medical or biologic agents will be covered in the most appropriate, safe, and cost-effective site.
Hospital-based outpatient clinical level of care	 This site is considered medically necessary for the first 90 days for the following: The initial course of infusion of a pharmacologic or biologic agent OR Re-initiation of an agent after 6 months or longer following discontinuation of therapy* Note: *This does not include when standard dosing between infusions is 6 months or longer
	This site is considered medically necessary when there is no outpatient infusion center within 50 miles of the individual's home and there is no contracted home infusion agency that will travel to their home, or a hospital is the only place that offers infusions of this drug. This site is considered medically necessary only when the individual has a clinical condition which puts him or her at increased risk of complications for infusions, including any ONE of the following: • Known cardiac condition (e.g., symptomatic cardiac arrhythmia) or pulmonary condition (e.g., significant respiratory disease, serious obstructive airway disease, %FVC ≤ 40%) that may increase the risk of an adverse reaction



Site of Service	Medical Necessity	
Administration		
	 Unstable renal function which decreases the ability to respond to fluids Difficult or unstable vascular access Acute mental status changes or cognitive conditions that impact the safety of infusion therapy A known history of severe adverse drug reactions and/or anaphylaxis to prior treatment with a related or similar drug 	
	This site is considered medically necessary when the individual has cytokine release syndrome (CRS) and all the following are met: CRS is grade 3 or 4 as evidenced by ALL the following:	
	 Temperature ≥ 38 °C Hypotension that requires one or more vasopressors Hypoxia that requires oxygen through a high-flow nasal cannula, face mask, non-rebreather mask, or Venturi mask OR positive pressure (continuous positive airway pressure [CPAP], bilevel positive airway pressure [BiPAP], intubation, or mechanical ventilation) 	
	AND	
	The individual will be admitted into an inpatient setting as soon as possible	
Hospital-based outpatient	These sites are considered not medically necessary for infusion	
setting	and injectable therapy services of various medical and biologic	
Outpatient hospital IV	agents when the site-of-service criteria in this policy are not	
infusion department	met.	
Hospital-based outpatient		
clinical level of care		

Agent	Medical Necessity		
Hidradenitis Suppurativa (HS)			
First-line TNF-α Antagonists			
Adalimumab-adaz	Adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm		
(Hyrimoz unbranded) SC	(Cyltezo unbranded), adalimumab-ryvk (Simlandi unbranded),		



Agent

- Adalimumab-adbm (Cyltezo unbranded) SC
- Adalimumab-ryvk
 (Simlandi unbranded) SC
- Cyltezo (adalimumabadbm) SC
- Simlandi (adalimumabryvk) SC

Managed under pharmacy benefit

Medical Necessity

Cyltezo (adalimumab-adbm), and Simlandi (adalimumab-ryvk) may be considered medically necessary for the treatment of hidradenitis suppurativa when:

• The individual is aged 12 years or older

AND

 Has tried at least one other therapy (e.g., intralesional or oral corticosteroids, systemic antibiotics)

AND

The medication is prescribed by or in consultation with a dermatologist

Note: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine whether this policy criteria applies.

First-line IL-17 Antagonists

Cosentyx (secukinumab) SC

Managed under pharmacy benefit

Cosentyx (secukinumab) may be considered medically necessary for the treatment of hidradenitis suppurativa when:

• The individual is aged 18 years or older

AND

 Has tried at least one other therapy (e.g., intralesional or oral corticosteroids, systemic antibiotics)

AND

 The medication is prescribed by or in consultation with a dermatologist

Second-line TNF-α Antagonists

- Abrilada (adalimumabafzb) SC
- Adalimumab-aacf (Idacio unbranded) SC
- Adalimumab-aaty (Yuflyma unbranded) SC
- Adalimumab-fkjp (Hulio unbranded) SC

Abrilada (adalimumab-afzb), adalimumab-aacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab – fkjp (Hulio unbranded), Hadlima (adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab), Hyrimoz (adalimumab-adaz), Idacio (adalimumab-aacf), Yuflyma (adalimumab-aaty) and Yusimry (adalimumab-aqvh) may be considered medically necessary for the treatment of hidradenitis suppurativa when:



Agent	Medical Necessity
 Agent Amjevita (adalimumabatto) SC Hadlima (adalimumabbwwd) SC Hulio (adalimumab-fkjp) SC Humira (adalimumab) SC Hyrimoz (adalimumabadaz) SC Idacio (adalimumabadaz) SC Yuflyma (adalimumabadaty) SC Yusimry (adalimumabady) SC Yusimry (adalimumabady) SC Managed under pharmacy benefit 	 The individual is aged 12 years or older AND Has tried at least one other therapy (e.g., intralesional or oral corticosteroids, systemic antibiotics) AND Has had an inadequate response or intolerance to ALL the following agents: Cyltezo (adalimumab-adbm) OR adalimumab-adbm (Cyltezo unbranded) Adalimumab-adaz (Hyrimoz unbranded) Simlandi (adalimumab-ryvk) OR adalimumab-ryvk (Simlandi unbranded) AND The medication is prescribed by or in consultation with a dermatologist Note: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine
Systemic Lunus Erythemat	osus (SLE) & Lunus Nonhritis
	cosus (SLE) & Lupus Nephritis
Anti-CD20Rituxan (rituximab)Ruxience (rituximab-pvvr)Truxima (rituximab-abbs)	See policy 5.01.556 Rituximab: Non-oncologic and Miscellaneous Uses
BLyS Inhibitors	
Benlysta (belimumab) IV Managed under medical benefit	Benlysta (belimumab) IV is subject to review for site of service administration. Benlysta (belimumab) IV may be considered medically necessary for the treatment of active, autoantibody positive
Benlysta (belimumab) SC	SLE when the following conditions are met:The individual is aged 5 years or olderAND



Agent	Medical Necessity
Managed under pharmacy	Has a diagnosis of SLE confirmed using either the American
and medical benefit	College of Rheumatology (ACR or EULAR/ACR) or Systemic
	Lupus International Collaborating Clinics (SLICC) criteria
	AND
	Benlysta (belimumab) IV is being used as add-on-therapy
	following standard induction therapy with mycophenolate,
	cyclophosphamide, azathioprine, or immunosuppressant, plus a
	corticosteroid
	AND
	 Benlysta (belimumab) IV is not used concurrently with Saphnelo
	(anifrolumab-fnia) for the treatment of SLE
	(allifoldinab-filla) for the treatment of SLE
	Benlysta (belimumab) SC may be considered medically
	necessary for the treatment of active, autoantibody positive
	SLE when the following conditions are met:
	The individual is aged 5 years or older
	AND
	 Has a diagnosis of SLE confirmed using either the American
	College of Rheumatology (ACR or EULAR/ACR) or Systemic
	Lupus International Collaborating Clinics (SLICC) criteria
	AND
	Benlysta (belimumab) SC is being used as add-on-therapy
	following standard induction therapy with mycophenolate,
	cyclophosphamide, azathioprine, or immunosuppressant, plus a
	corticosteroid
	AND
	Benlysta (belimumab) SC is not used concurrently with
	Saphnelo (anifrolumab-fnia) for the treatment of SLE
	Benlysta (belimumab) IV may be considered medically
	necessary for the treatment of pediatric and adult individuals
	with active lupus nephritis who are receiving standard therapy
	when the following conditions are met:
	The individual is aged 5 years or older
	AND
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Agent	Medical Necessity
	Has a diagnosis of SLE confirmed using either the American
	College of Rheumatology (ACR or EULAR/ACR) or Systemic
	Lupus International Collaborating Clinics (SLICC) criteria
	AND
	 Is receiving standard therapy with mycophenolate,
	cyclophosphamide, azathioprine, or immunosuppressant, plus a corticosteroid
	AND
	 Has class III (focal proliferative), class IV (diffuse proliferative), and/or class V (membranous) lupus nephritis
	AND
	 No previous use of dialysis in the past 12 months
	AND
	 Benlysta (belimumab) is not used concurrently with Lupkynis
	(voclosporin) for the treatment of active lupus nephritis
	AND
	 Benlysta (belimumab) is prescribed by or in consultation with a
	nephrologist or rheumatologist
	nephrologist of meaniatologist
	Benlysta (belimumab) SC may be considered medically
	necessary for the treatment of adult individuals with active
	lupus nephritis who are receiving standard therapy when the
	following conditions are met:
	 The individual is aged 18 years or older
	AND
	 Has a diagnosis of SLE confirmed using either the American
	College of Rheumatology (ACR or EULAR/ACR) or Systemic
	Lupus International Collaborating Clinics (SLICC) criteria
	AND
	 Is receiving standard therapy with mycophenolate,
	cyclophosphamide, azathioprine, or immunosuppressant, plus a corticosteroid
	AND
	 Has class III (focal proliferative), class IV (diffuse proliferative), and/or class V (membranous) lupus nephritis
	and, or state 1 (membraness) rapus reprints



AND

Agent	Medical Necessity
	 No previous use of dialysis in the past 12 months AND Benlysta (belimumab) is not used concurrently with Lupkynis (voclosporin) for the treatment of active lupus nephritis AND Benlysta (belimumab) is prescribed by or in consultation with a nephrologist or rheumatologist
Calcineurin Inhibitors	
Calcineurin Inhibitor • Lupkynis (voclosporin) oral Managed under pharmacy benefit	Lupkynis (voclosporin) may be considered medically necessary for the treatment of adult individuals with active lupus nephritis when the following conditions are met: • The individual is aged 18 years or older AND • Has a diagnosis of SLE confirmed using either the American College of Rheumatology (ACR or EULAR/ACR) or Systemic Lupus International Collaborating Clinics (SLICC) criteria AND • Lupkynis (voclosporin) will be used in combination with mycophenolate, cyclophosphamide, azathioprine, or an immunosuppressant AND a corticosteroid AND • Has class III (focal proliferative), class IV (diffuse proliferative), and/or class V (membranous) lupus nephritis AND • No previous use of dialysis in the past 12 months AND • Lupkynis (voclosporin) is not used concurrently with Benlysta (belimumab) for the treatment of active lupus nephritis AND • The dose prescribed is ≤ 47.4 mg per day (taken as three 7.9 mg capsules twice daily)
	Lupkynis (voclosporin) is prescribed by or in consultation with a nephrologist or rheumatologist

Type I Interferon (IFN) Receptor Antagonist

Agent Type I IFN Receptor Antagonist • Saphnelo (anifrolumabfinia) IV Managed under medical benefit

Medical Necessity

Saphnelo (anifrolumab-fnia) may be considered medically necessary for the treatment of adult individuals with moderate to severe systemic lupus erythematosus (SLE) when the following conditions are met:

• The individual is aged 18 years or older

AND

 Has a diagnosis of SLE confirmed using either the American College of Rheumatology (ACR or EULAR/ACR) or Systemic Lupus International Collaborating Clinics (SLICC) criteria

AND

 Saphnelo (anifrolumab-fnia) is being used as add-on therapy following standard induction therapy with mycophenolate, azathioprine, or immunosuppressant, plus a corticosteroid

AND

 Does not have severe (IV cyclophosphamide and/or high dose IV pulse corticosteroid is not used) active central nervous system lupus

AND

Does not have severe (IV cyclophosphamide and/or high dose
 IV pulse corticosteroid is not used) active lupus nephritis

AND

 Saphnelo (anifrolumab-fnia) is not used concurrently with Benlysta (belimumab) for the treatment of SLE

Pyoderma Gangrenosum

First-line Agents

TNF-α Antagonists

- Adalimumab-adaz (Hyrimoz unbranded) SC
- Adalimumab-adbm (Cyltezo unbranded) SC
- Adalimumab-ryvk
 (Simlandi unbranded) SC
- Cyltezo (adalimumabadbm) SC
- Simlandi (adalimumabryvk) SC
- Enbrel (etanercept) SC

Adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-ryvk (Simlandi unbranded), Cyltezo (adalimumab-adbm), Simlandi (adalimumab-ryvk), and Enbrel (etanercept) may be considered medically necessary for the treatment of pyoderma gangrenosum when:

• The individual is aged 18 years or older

AND

 Has not responded to one standard non-biologic therapy (e.g., oral corticosteroids, systemic cyclosporine, topical tacrolimus, etc.)

AND



Agent	Medical Necessity		
Managed under pharmacy benefit	The medication is prescribed by or in consultation with a dermatologist		
	Note: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine whether this policy criteria applies.		
TNF-α Antagonists	Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded),		
Inflectra (infliximab-	and Remicade (infliximab) are subject to review for site of		
dyyb) IV	service administration.		
 Infliximab (Janssen – unbranded) IV 			
Remicade (infliximab) IV	Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded),		
	and Remicade (infliximab) may be considered medically		
Managed under medical	necessary for the treatment of pyoderma gangrenosum when:		
benefit	 The individual is aged 18 years or older AND		
	 Has not responded to one standard non-biologic therapy (e.g., oral corticosteroids, systemic cyclosporine, topical tacrolimus, etc.) 		
	AND		
	The medication is prescribed by or in consultation with a dermatologist		
Second-line Agents			
TNF- α Antagonists	Avsola (infliximab-axxq) and Renflexis (infliximab-abda) are		
Avsola (infliximab-axxq)	subject to review for site of service administration.		
IVRenflexis (infliximab-	Aveala (inflivingly and Dauflavia (inflivingly and Dauflavia		
abda) IV	Avsola (infliximab-axxq) and Renflexis (infliximab-abda) may be considered medically necessary for the treatment of		
	pyoderma gangrenosum when:		
Managed under medical	The individual is aged 18 years or older		
benefit	AND		

Agent	Medical Necessity
	 Has not responded to one standard non-biologic therapy (e.g., oral corticosteroids, systemic cyclosporine, topical tacrolimus, etc.) AND Has had an inadequate response or intolerance to Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded), or Remicade (infliximab) AND The medication is prescribed by or in consultation with a dermatologist
 TNF-α Antagonists Abrilada (adalimumabafzb) SC Adalimumab-aacf (Idacio unbranded) SC Adalimumab-aaty (Yuflyma unbranded) SC Adalimumab-fkjp (Hulio unbranded) SC Amjevita (adalimumabatto) SC Hadlima (adalimumabbwwd) SC Hulio (adalimumab-fkjp) SC Humira (adalimumab) SC Hyrimoz (adalimumabadaz) SC Idacio (adalimumabaacf) SC Yuflyma (adalimumabaaty) SC Yuflyma (adalimumabaaty) SC Yusimry (adalimumabaaqvh) SC 	Abrilada (adalimumab-afzb), adalimumab-aacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-fkjp (Hulio unbranded), Amjevita (adalimumab-atto), Hadlima (adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab), Hyrimoz (adalimumab-adaz), Idacio (adalimumab-aacf), Yuflyma (adalimumab-aaty), and Yusimry (adalimumab-aqvh) considered medically necessary for the treatment of pyoderma gangrenosum when: • The individual is aged 18 years or older AND • Has not responded to one standard non-biologic therapy (e.g., oral corticosteroids, systemic cyclosporine, topical tacrolimus, etc.) AND • Has had an inadequate response or intolerance to ALL the following agents: • Cyltezo (adalimumab-adbm) OR adalimumab-adbm (Cyltezo unbranded) • Adalimumab-adaz (Hyrimoz unbranded) • Simlandi (adalimumab-ryvk) OR adalimumab-ryvk (Simlandi unbranded)
Managed under pharmacy benefit	 The medication is prescribed by or in consultation with a dermatologist

Agent	Medical Necessity	
	Note:	This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine whether this policy criteria applies.

Uveitis

First-line Agents

TNF-α Antagonists

- Adalimumab-adaz (Hyrimoz unbranded) SC
- Adalimumab-adbm (Cyltezo unbranded) SC
- Adalimumab-ryvk (Simlandi unbranded) SC
- Cyltezo (adalimumabadbm) SC
- Simlandi (adalimumabryvk) SC

Managed under pharmacy benefit

Adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-ryvk (Simlandi unbranded), Cyltezo (adalimumab-adbm), and Simlandi (adalimumab-ryvk) may be considered medically necessary for the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis when:

The individual is aged 2 years or older

AND

- Has tried one of the following therapies:
 - o Periocular, intraocular, or systemic corticosteroids
 - Immunosuppressives

AND

Note:

The medication is prescribed by or in consultation with an ophthalmologist

This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy **5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies.** Please check the member Plan booklet or member ID card to determine whether this policy criteria applies.

Second-line Agents

TNF-α Antagonists

- Abrilada (adalimumabafzb) SC
- Adalimumab-aacf (Idacio unbranded) SC

Abrilada (adalimumab-afzb), adalimumab-aacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-fkjp (Hulio unbranded), Amjevita (adalimumab-atto), Hadlima (adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab), Hyrimoz (adalimumab-adaz), Idacio



Agent

- Adalimumab-aaty (Yuflyma unbranded) SC
- Adalimumab-fkjp (Hulio unbranded) SC
- Amjevita (adalimumabatto) SC
- Hadlima (adalimumabbwwd) SC
- Hulio (adalimumab-fkjp)
 SC
- Humira (adalimumab) SC
- Hyrimoz (adalimumabadaz) SC
- Idacio (adalimumab-aacf)
 SC
- Yuflyma (adalimumabaaty) SC
- Yusimry (adalimumabaqvh) SC

Managed under pharmacy benefit

Medical Necessity

(adalimumab-aacf), Yuflyma (adalimumab-aaty), and Yusimry (adalimumab-aqvh) may be considered medically necessary for the treatment of non-infectious intermediate uveitis, posterior uveitis, or panuveitis when:

• The individual is aged 2 years or older

AND

- Has tried one of the following therapies:
 - o Periocular, intraocular, or systemic corticosteroids
 - Immunosuppressives

AND

- Has had an inadequate response or intolerance to ALL the following agents:
 - Cyltezo (adalimumab-adbm) OR adalimumab-adbm (Cyltezo unbranded)
 - Adalimumab-adaz (Hyrimoz unbranded)
 - Simlandi (adalimumab-ryvk) OR adalimumab-ryvk (Simlandi unbranded)

AND

The medication is prescribed by or in consultation with an ophthalmologist

Note: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine whether this policy criteria applies.

Giant Cell Arteritis

IL-6 Antagonist

- Actemra (tocilizumab) SC,
 IV
- Tyenne (tocilizumabaazg) SC, IV
- Tofidence (tocilizumabbavi) IV

Actemra (tocilizumab) IV and Tofidence (tocilizumab-bavi) IV are subject to review for site of service administration.

Actemra (tocilizumab), Tyenne (tocilizumab-aazg), and Tofidence (tocilizumab-bavi) IV may be considered medically necessary for the treatment of giant cell arteritis when:

• The individual is aged 18 years or older

AND



Agent	Medical Necessity	
Managed under pharmacy and medical benefit	 Has tried one systemic corticosteroid AND The medication is prescribed by or in consultation with a rheumatologist Emyelinating Polyneuropathy (CIDP) 	
Vyvgart Hytrulo	Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc)	
(efgartigimod alfa and	may be considered medically necessary for the treatment of	
hyaluronidase-qvfc)	chronic inflammatory demyelinating polyneuropathy (CIDP)	
	when all the following criteria are met:	
Managed under medical benefit	 The individual is aged 18 years or older AND 	
	 Has been diagnosed with CIDP based on all the following: Individual has experienced progressive or relapsing motor and/or sensory symptoms of more than one limb AND hyporeflexia or areflexia in affected limbs is present for at least 2 months AND 	
	 Has electrophysiologic findings that meets 3 of the following 4 criteria per the American Academy of Neurology indicating demyelinating neuropathy Partial conduction block of ≥ 1 motor nerve Reduced conduction velocity of ≥ 2 motor nerves Prolonged distal latency of ≥ 2 motor nerves Prolonged F-wave latencies of ≥ 2 motor nerves or the absence of F waves 	
	 Other causes of demyelinating neuropathy have been excluded such as Borrelia burgdorferi infection (Lyme disease), diphtheria, drug or toxin exposure, hereditary demyelinating neuropathy, prominent sphincter disturbance, multifocal motor neuropathy (MMN), and IgM monoclonal gammopathy 	
	AND	
	 If available, results of other testing to support the diagnosis should be provided such as any of the following: Cerebrospinal fluid (CSF) examination demonstrating elevated CSF protein with leukocyte count <10/mm³ 	



Agent	Medical Necessity
	 MRI showing gadolinium enhancement and/or hypertrophy of the cauda equina, lumbosacral or cervical nerve roots, or the brachial or lumbosacral plexuses Nerve biopsy showing unequivocal evidence of demyelination and/or remyelination by electron microscopy or teased fiber analysis AND Has tried and had an inadequate response or intolerance to intravenous or subcutaneous immune globulin (e.g., Gammagard Liquid or Gammaked) AND Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) is
	prescribed by or in consultation with a neurologist
 Cytokine Release Syndrom IL-6 Antagonist Actemra (tocilizumab) IV Tofidence (tocilizumab-bavi) IV Tyenne (tocilizumab-aazg) IV Managed under medical benefit 	Actemra (tocilizumab) IV, Tofidence (tocilizumab-bavi) IV, and Tyenne (tocilizumab-aazg) IV may be considered medically necessary for adults and pediatric individuals when the following criteria are met: • The individual is aged 2 years or older AND • Has a documented treatment-induced grade 3, or 4 cytokine release syndrome (CRS) as evidenced by ALL of the following: • Temperature ≥ 38 °C • Hypotension that requires one or more vasopressors • Hypoxia requiring oxygen through a high-flow nasal cannula, face mask, non-rebreather mask, or Venturi mask OR requiring positive pressure (continuous positive airway pressure [CPAP], bilevel positive airway pressure [BiPAP],
Interleukin-1 Receptor Antagonist • Kineret (anakinra) SC Managed under pharmacy and medical benefit	intubation, or mechanical ventilation) Kineret (anakinra) may be considered medically necessary when the individual has a documented treatment-induced grade 3 or 4 cytokine release syndrome (CRS) as evidenced by all of the following: Temperature ≥ 38 °C Hypotension that requires one or more vasopressors

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Agent	Medical Necessity
	 Hypoxia requiring oxygen through a high-flow nasal cannula, face mask, non-rebreather mask, or Venturi mask OR requiring positive pressure (continuous positive airway pressure [CPAP], bilevel positive airway pressure [BiPAP], intubation, or mechanical ventilation)
Behcet's Disease	
Phosphodiesterase 4	Otezla (apremilast) may be considered medically necessary for
(PDE4) inhibitor	the treatment of oral ulcers associated with Behcet's Disease
Otezla (apremilast) Oral	when:
	The individual is aged 18 years or older
Managed under pharmacy	AND
benefit	Has tried one other systemic therapy (e.g., colchicine,
	corticosteroids, azathioprine)
	AND
	The medication is prescribed by or in consultation with a
	rheumatologist or dermatologist
Neuromyelitis Optica Spe	ctrum Disorder (NMOSD)
CD19-directed cytolytic	Uplizna (inebilizumab-cdon) is subject to review for site of
antibody	service administration.
Uplizna (inebilizumab-	service administration.
•	Service administration. Uplizna (inebilizumab-cdon) may be considered medically
Uplizna (inebilizumab- cdon) IV	
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically
Uplizna (inebilizumab- cdon) IV	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are anti-
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met:
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met: • The individual is aged 18 years or older
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met: • The individual is aged 18 years or older AND
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met: • The individual is aged 18 years or older AND • Has a documented diagnosis of NMOSD confirmed by:
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met: • The individual is aged 18 years or older AND • Has a documented diagnosis of NMOSD confirmed by: • At least one of the following core clinical characteristics:
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met: • The individual is aged 18 years or older AND • Has a documented diagnosis of NMOSD confirmed by: • At least one of the following core clinical characteristics: • Optic neuritis
Uplizna (inebilizumab- cdon) IV Managed under medical	Uplizna (inebilizumab-cdon) may be considered medically necessary for the treatment of neuromyelitis optica spectrum disorder (NMOSD) in adult individuals who are antiaquaporin-4 (AQP4) antibody positive when the following are met: • The individual is aged 18 years or older AND • Has a documented diagnosis of NMOSD confirmed by: • At least one of the following core clinical characteristics: • Optic neuritis • Acute myelitis



Acute brainstem syndrome

Agent	Medical Necessity
	 Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions Symptomatic cerebral syndrome with NMOSD-typical brain lesions
	AND
	 Positive test for AQP4-IgG antibodies
	AND Evaluaion of alternative diagnoses (e.g. multiple salaresis)
	 Exclusion of alternative diagnoses (e.g., multiple sclerosis) AND
	 History of at least 1 relapse in last 12 months or 2 relapses in
	the last 24 months
	AND
	• Expanded Disability Status Scale (EDSS) score ≤ 7.5
Interleukin-6 (IL-6)	Enspryng (satralizumab-mwge) may be considered medically
receptor antagonist	necessary for the treatment of neuromyelitis optica spectrum
Enspryng (satralizumab-	disorder (NMOSD) in adult individuals who are anti-
mwge) SC	aquaporin-4 (AQP4) antibody positive when the following are
Managed under pharmacy	met:
and medical benefit	 The individual is aged 18 years or older AND
	 Has a documented diagnosis of NMOSD confirmed by:
	 At least one of the following core clinical characteristics:
	Optic neuritis
	 Acute myelitis
	 Area postrema syndrome: Episode of otherwise
	unexplained hiccups or nausea and vomiting
	Acute brainstem syndrome
	 Symptomatic narcolepsy or acute diencephalic clinical syndrome with NMOSD-typical diencephalic MRI lesions
	 Symptomatic cerebral syndrome with NMOSD-typical brain lesions
	AND
	 Positive test for AQP4-IgG antibodies
	AND
	 Exclusion of alternative diagnoses (e.g., multiple sclerosis)



Agent	Medical Necessity
	AND
	History of at least 1 relapse in last 12 months or 2 relapses in
	the last 24 months
	AND
	• Expanded Disability Status Scale (EDSS) score ≤ 6.5
Deficiency of Interleukin-	l Receptor Antagonist (DIRA)
Interleukin-1 Blocker	Arcalyst (rilonacept) may be considered medically necessary
Arcalyst (rilonacept) SC	for the treatment of deficiency of interleukin-1 receptor
	antagonist (DIRA) when the following criteria are met:
Managed under pharmacy	Genetic testing has confirmed a mutation in the IL1RN gene
and medical benefit	AND
	 Individual weight is ≥ 10 kg
	AND
	Arcalyst (rilonacept) is prescribed by or in consultation with a
	rheumatologist, geneticist, or dermatologist
Interleukin-1 Receptor	Kineret (anakinra) may be considered medically necessary for
Antagonist	the treatment of deficiency of interleukin-1 receptor
Kineret (anakinra) SC	antagonist (DIRA) when the following criteria are met:
	Genetic testing has confirmed a mutation in the IL1RN gene
Managed under pharmacy	AND
and medical benefit	Kineret (anakinra) is prescribed by or in consultation with a
	rheumatologist, geneticist, or dermatologist
Recurrent Pericarditis	
Interleukin-1 Blocker	Arcalyst (rilonacept) may be considered medically necessary
Arcalyst (rilonacept) SC	for the treatment of recurrent pericarditis (RP) and reduction
	in risk of recurrence when the following criteria are met:
Managed under pharmacy	The individual is aged 12 years or older
and medical benefit	AND
	Has a documented prior episode of acute pericarditis
	AND
	 Has typical pleuritic chest pain plus ≥ 1 of the following:
	o Fever
	o Pericardial rub
	o ECG changes
	 New or worsening pericardial effusion

Agent	Medical Necessity
	 ○ Elevation of markers of inflammation (elevation in white blood cell count, erythrocyte sedimentation rate, or C-reactive protein) OR ○ There is evidence of pericardial inflammation on cardiovascular magnetic resonance (CMR) or computed tomography (CT) after a ≥ 4-week symptom-free interval AND Has received prior treatment for RP with an NSAID or corticosteroid unless contraindicated AND
	 Arcalyst (rilonacept) is prescribed by or in consultation with a cardiologist
Periodic Fever Syndromes	& Still's Disease
Interleukin-1 Blocker	Arcalyst (rilonacept) may be considered medically necessary
Arcalyst (rilonacept) SC	for the treatment of:
Managed under pharmacy and medical benefit	 Cryopyrin-associated periodic syndromes (CAPS), in adults and children aged 12 years and older, including: Familial cold auto-inflammatory syndrome (FCAS) Muckle-Wells syndrome (MWS) AND Arcalyst (rilonacept) is prescribed by or in consultation with a
	rheumatologist, geneticist, or dermatologist
Interleukin-1β blocker	Ilaris (canakinumab) may be considered medically necessary
 Ilaris (canakinumab) SC Managed under pharmacy and medical benefit 	 for the treatment of: Periodic Fever Syndromes: Cryopyrin-associated periodic syndromes (CAPS), in adults and children aged 4 years and older, including: Familial cold auto-inflammatory syndrome (FCAS) Muckle-Wells syndrome (MWS) Tumor necrosis factor receptor associated periodic syndrome (TRAPS) in adult and pediatric individuals aged 2
	years and older o Hyperimmunoglobulin D syndrome (HIDS)/mevalonate kinase deficiency (MKD) in adult and pediatric individuals aged 2 years and older



Agent	Medical Necessity
	 Familial Mediterranean fever (FMF) in adult and pediatric individuals aged 2 years and older OR Active Still's disease, including adult-onset Still's disease
	(AOSD) and systemic juvenile idiopathic arthritis (SJIA) in individuals aged 2 years and older AND
	Ilaris (canakinumab) is prescribed by or in consultation with a rheumatologist, geneticist, or dermatologist
Interleukin-1 Receptor	Kineret (anakinra) may be considered medically necessary for
Antagonist	the treatment of cryopyrin-associated periodic syndromes
Kineret (anakinra) SC	(CAPS) when the following criteria are met:
	The individual has been diagnosed with neonatal-onset
Managed under pharmacy	multisystem inflammatory disease (NOMID)
and medical benefit	AND
	Kineret (anakinra) is prescribed by or in consultation with a
	rheumatologist, geneticist, or dermatologist
Graft Versus Host Disease	
Niktimvo (axatilimab-csfr)	Niktimvo (axatilimab-csfr) may be considered medically
IV	necessary for the treatment of chronic graft versus host
	disease after failure of at least two prior lines of systemic
Managed under medical	therapy when the following conditions are met:
benefit	The individual weighs at least 40 kg
	AND
	Has tried and failed at least two systemic treatments such as
	cyclosporine, ibrutinib, mycophenolate mofetil, ruxolitinib,
	sirolimus, or tacrolimus
	AND
	Medication is being prescribed by or in consultation with an
	oncologist, hematologist, or a physician affiliated with a
	transplant center
	AND
	Dose is limited to 35 mg every 2 weeks
Orencia (abatacept)	Orencia (abatacept) may be considered medically necessary for
	the prevention of acute graft versus host disease when the
Managed under pharmacy	following conditions are met:
and medical benefit	The individual is aged 2 years or older

Agent	Medical Necessity
	 Will also receive standard therapy with a calcineurin inhibitor (cyclosporine or tacrolimus) AND Will also receive standard therapy with methotrexate AND Will undergo hematopoietic stem cell transplantation from a matched unrelated donor OR a 1-allele-mismatched unrelated donor AND The medication is being prescribed by or in consultation with an oncologist, hematologist, or a physician affiliated with a transplant center
Rezurock (belumosudil) Managed under pharmacy	Rezurock (belumosudil) may be considered medically necessary for the treatment of chronic graft versus host disease when the following conditions are met:
benefit	 The individual is aged 12 years or older AND Has tried and failed at least two systemic treatments such as cyclosporine, ibrutinib, mycophenolate mofetil, ruxolitinib, sirolimus, or tacrolimus AND The medication is being prescribed by or in consultation with an oncologist, hematologist, or a physician affiliated with a transplant center AND The dose is limited to 200 mg daily
Ryoncil (remestemcel-L-rknd)	Ryoncil (remestemcel-L-rknd) may be considered medically necessary for the treatment of pediatric individuals with steroid-refractory acute graft versus host disease (aGvHD)
Managed under medical benefit	 when all the following criteria are met: The individual is aged 2 months to 17 years AND Has been diagnosed with one of the following: Grade C or D aGvHD involving the skin, liver, or gastrointestinal tract Grade B aGvHD involving the liver or gastrointestinal tract



Agent	Medical Necessity
rigent	AND
	 Has tried and had an inadequate response or intolerance to
	systemic corticosteroid therapy
	AND
	Has tried and had an inadequate response or intolerance to
	one other therapy (e.g., ruxolitinib or mycophenolate mofetil)
	AND
	The medication is being prescribed by or in consultation with
	an oncologist, hematologist, or a physician affiliated with a
	transplant center
Myasthenia Gravis	
Rystiggo	Rystiggo (rozanolixizumab-noli) may be considered medically
(rozanolixizumab-noli)	necessary for the treatment of myasthenia gravis when the
	following criteria are met:
Managed under medical	The individual is aged 18 years or older
benefit	AND
	Has a diagnosis of myasthenia gravis with a serological test for
	anti-acetylcholine receptor (AChR) or anti-muscle-specific
	tyrosine kinase (MuSK) antibodies
	AND
	Is currently using the acetylcholinesterase inhibitor
	pyridostigmine, has tried and failed pyridostigmine or has
	contraindications to use of pyridostigmine
	AND
	Is currently using two or more immunosuppressive therapies
	(ISTs) (e.g., glucocorticoids, azathioprine, mycophenolate
	mofetil, cyclosporine) or has tried and failed two ISTs or has
	contraindications that prevent use of two ISTs
	AND
	For the treatment of AChR antibody positive myasthenia gravis
	the individual has tried and failed ≥ 1 of the following:
	Soliris (eculizumab)
	Ultomiris (ravulizumab-cwvz)
	Vyvgart (efgartigimod alfa-fcab), Vyvgart Uytrula (efgartigimad alfa and byaluranidasa gyfs)
	Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc)
	AND



Agent	Medical Necessity
	Medication is not being used concurrently with Vyvgart
	(efgartigimod alfa-fcab), Vyvgart Hytrulo (efgartigimod alfa and
	hyaluronidase-qvfc), Soliris (eculizumab), Ultomiris
	(ravulizumab-cwvz), or Zilbrysq (zilucoplan)
Vyvgart (efgartigimod	Vyvgart (efgartigimod alfa-fcab) may be considered medically
alfa-fcab)	necessary for the treatment of myasthenia gravis when the
	following criteria are met:
Managed under medical	The individual is aged 18 years or older
benefit	AND
	Has a diagnosis of myasthenia gravis with a serological test for
	anti-acetylcholine receptor (AChR) antibodies
	AND
	Is currently using two or more immunosuppressive therapies
	(ISTs) (e.g., glucocorticoids, azathioprine, mycophenolate
	mofetil, cyclosporine) or has tried and failed two ISTs or has
	contraindications that prevent use of two ISTs
	AND
	Medication is not being used concurrently with Vyvgart Hytrulo
	(efgartigimod alfa and hyaluronidase-qvfc), Rystiggo
	(rozanolixizumab-noli), Soliris (eculizumab), Ultomiris
	(ravulizumab-cwvz), or Zilbrysq (zilucoplan)
Vyvgart Hytrulo	Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc)
(efgartigimod alfa and	may be considered medically necessary for the treatment of
hyaluronidase-qvfc)	myasthenia gravis when the following criteria are met:
	The individual is aged 18 years or older
Managed under medical	AND
benefit	Has a diagnosis of myasthenia gravis with a serological test for
	anti-acetylcholine receptor (AChR) antibodies
	AND
	Is currently using two or more immunosuppressive therapies
	(ISTs) (e.g., glucocorticoids, azathioprine, mycophenolate
	mofetil, cyclosporine) or has tried and failed two ISTs or has
	contraindications that prevent use of two ISTs
	AND
	Medication is not being used concurrently with Vyvgart
	(efgartigimod alfa-fcab), Rystiggo (rozanolixizumab-noli), Soliris

Agent	Medical Necessity
Agent	(eculizumab), Ultomiris (ravulizumab-cwvz), or Zilbrysq
	(zilucoplan)
Primary Immunoglobulin A	
Fabhalta (iptacopan)	Fabhalta (iptacopan) may be considered medically necessary to
Managad under pharmage	reduce proteinuria in adults with primary immunoglobulin A
Managed under pharmacy benefit	nephropathy (IgAN) at risk of rapid disease progression when the following criteria are met:
bellefit	 The individual is aged 18 years or older
	AND
	 Has a documented diagnosis of biopsy-proven primary immunoglobulin A nephropathy (IgAN) AND
	 Has a documented urine protein-to-creatinine ratio (UPCR) ≥ 1.5 g/g
	AND
	 Has tried and failed an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB)
	AND
	 Has tried and failed Filspari (sparsentan) or Tarpeyo (budesonide)
	AND
	Fabhalta (iptacopan) is prescribed by or in consultation with a nephrologist
	AND
	The dose prescribed is limited to 400 mg per day
Filspari (sparsentan)	Filspari (sparsentan) may be considered medically necessary to
	slow kidney function decline in adults with primary
Managed under pharmacy	immunoglobulin A nephropathy (IgAN) at risk for disease
benefit	progression when the following criteria are met:
	The individual is aged 18 years or older
	AND
	Has a documented diagnosis of biopsy-proven primary
	immunoglobulin A nephropathy (lgAN)
	AND
	 Has a documented urine protein-to-creatinine ratio (UPCR) ≥
	1.5 g/g
	AND



Agent	Medical Necessity
	 Has tried and failed an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) AND Filspari (sparsentan) is not used concurrently with other ACE inhibitors, ARB, endothelin receptor antagonists (ERAs), and aliskiren
	 AND Filspari (sparsentan) is prescribed by or in consultation with a nephrologist AND The dose prescribed is limited to 400 mg per day
Tarpeyo (budesonide)	 Tarpeyo (budesonide) may be considered medically necessary to reduce the loss of kidney function with primary immunoglobulin A nephropathy (IgAN) at risk of disease progression when the following criteria are met: The individual is aged 18 years or older AND Has a documented diagnosis of biopsy-proven primary immunoglobulin A nephropathy (IgAN) AND Has a documented urine protein-to-creatinine ratio (UPCR) ≥ 0.8 g/g OR proteinuria ≥ 1 g/day AND Is used in combination with an angiotensin-converting enzyme (ACE) inhibitor or angiotensin receptor blocker (ARB) AND Tarpeyo (budesonide) is prescribed by or in consultation with a nephrologist AND The dose prescribed is limited to 16 mg daily AND The total duration of therapy is limited to 9 months
Sarcoidosis	
First-line Agents	
TNF-α Antagonists • Adalimumab-adaz (Hyrimoz unbranded) SC	Adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-ryvk (Simlandi unbranded),



Agent

- Adalimumab-adbm (Cyltezo unbranded) SC
- Adalimumab-ryvk
 (Simlandi unbranded) SC
- Cyltezo (adalimumabadbm) SC
- Simlandi (adalimumabryvk) SC

Managed under pharmacy benefit

Medical Necessity

Cyltezo (adalimumab-adbm), and Simlandi (adalimumab-ryvk) may be considered medically necessary for the treatment of sarcoidosis when:

• The individual is aged 18 years or older

AND

 Has tried and had an inadequate response or intolerance to one corticosteroid

AND

 Has tried and had an inadequate response or intolerance to one immunosuppressive medication (e.g., methotrexate, leflunomide, azathioprine, mycophenolate, cyclosporine, chlorambucil, cyclophosphamide, thalidomide, or chloroquine)

AND

 The medication is prescribed by or in consultation with a pulmonologist, ophthalmologist, or dermatologist

Note: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please

check the member Plan booklet or member ID card to determine

whether this policy criteria applies.

TNF-α Antagonists

- Inflectra (infliximabdyyb) IV
- Infliximab (Janssen unbranded) IV
- Remicade (infliximab) IV

Managed under medical benefit

Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded), and Remicade (infliximab) are subject to review for site of service administration.

Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded), and Remicade (infliximab) may be considered medically necessary for the treatment of sarcoidosis when:

• The individual is aged 18 years or older

AND

 Has tried and had an inadequate response or intolerance to one corticosteroid

AND

 Has tried and had an inadequate response or intolerance to one immunosuppressive medication (e.g., methotrexate,



Agent	Medical Necessity
Second-line Agents TNF-α Antagonists	leflunomide, azathioprine, mycophenolate, cyclosporine, chlorambucil, cyclophosphamide, thalidomide, or chloroquine) AND The medication is prescribed by or in consultation with a pulmonologist, ophthalmologist, or dermatologist Abrilada (adalimumab-afzb), adalimumab-aacf (Idacio
 Abrilada (adalimumabafzb) SC Adalimumabaacf (Idacio unbranded) SC Adalimumabaaty (Yuflyma unbranded) SC Adalimumabafkjp (Hulio unbranded) SC Amjevita (adalimumabatto) SC 	unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-fkjp (Hulio unbranded), Amjevita (adalimumab-atto), Hadlima (adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab), Hyrimoz (adalimumab-adaz), Idacio (adalimumab-aacf), Yuflyma (adalimumab-aaty), and Yusimry (adalimumab-aqvh) may be considered medically necessary for the treatment of sarcoidosis when: • The individual is aged 18 years or older AND
 Hadlima (adalimumab-bwwd) SC Hulio (adalimumab-fkjp) SC Humira (adalimumab) SC Hyrimoz (adalimumab-adaz) SC Idacio (adalimumab-aacf) SC 	 Has tried and had an inadequate response or intolerance to one corticosteroid AND Has tried and had an inadequate response or intolerance to one immunosuppressive medication (e.g., methotrexate, leflunomide, azathioprine, mycophenolate, cyclosporine, chlorambucil, cyclophosphamide, thalidomide, or chloroquine) AND
 Yuflyma (adalimumabaaty) SC Yusimry (adalimumabaqvh) SC Managed under pharmacy benefit 	 Has had an inadequate response or intolerance to ALL the following agents: Cyltezo (adalimumab-adbm) OR adalimumab-adbm (Cyltezo unbranded) Adalimumab-adaz (Hyrimoz unbranded) Simlandi (adalimumab-ryvk) OR adalimumab-ryvk (Simlandi unbranded)
	 unbranded) AND Medication is prescribed by or in consultation with a pulmonologist, ophthalmologist, or dermatologist



Agent	Medical Necessity
	Note: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine whether this policy criteria applies.
TNF- α Antagonists	Avsola (infliximab-axxq) and Renflexis (infliximab-abda) are
Avsola (infliximab-axxq) IV	subject to review for site of service administration.
Renflexis (infliximab- abda) IV	Avsola (infliximab-axxq) and Renflexis (infliximab-abda) may be considered medically necessary for the treatment of
Managed under medical benefit	sarcoidosis when:The individual is aged 18 years or olderAND
	Has tried and had an inadequate response or intolerance to one corticosteroid
	AND
	 Has tried and had an inadequate response or intolerance to one immunosuppressive medication (e.g., methotrexate, leflunomide, azathioprine, mycophenolate, cyclosporine, chlorambucil, cyclophosphamide, thalidomide, or chloroquine)
	AND
	Has had an inadequate response or intolerance to Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded), or Remicade (infliximab)
	 AND The medication is prescribed by or in consultation with a pulmonologist, ophthalmologist, or dermatologist

Agent	Investigational
As listed	The medications listed in this policy are subject to the product's US Food and Drug Administration (FDA) dosage and administration prescribing information.



Agent	Investigational
	All other uses of the above-named agents when used in
	combination with each other or for conditions not outlined in
	this policy or policies 5.01.550, 5.01.563, and 5.01.629 are
	considered investigational.

Length of Approval			
Approval	Criteria		
Initial authorization	Non-formulary exception reviews and all other reviews for all drugs listed in the policy may be approved up to 12 months.		
Re-authorization criteria	Non-formulary exception reviews and all other reviews for all drugs listed in the policy may be approved up to 12 months as long as the drug-specific coverage criteria are met, and chart notes demonstrate that the individual continues to show a positive clinical response to therapy.		

Documentation Requirements

The individual's medical records submitted for review for all conditions should document that medical necessity criteria are met. The record should include the following:

 Office visit notes that contain the diagnosis, relevant history, physical evaluation, and medication history

Coding

Code	Description			
HCPCS				
J0129	Injection, abatacept (Orencia), 10 mg (code may be used for Medicare when drug administered under the direct supervision of a physician, not for use when drug is self-administered)			
J0135	Injection, adalimumab (Humira), 20mg (code termed 01/01/25)			
J0139	Injection, adalimumab, 1 mg (new code effective 01/01/25)			
J0490	Injection, belimumab (Benlysta), 10 mg			
J0491	Injection, anifrolumab-fnia (Saphnelo), 1 mg			



Code	Description			
J0638	Injection, canakinumab, (llaris)1 mg			
J1438	Injection, etanercept (Enbrel), 25mg (code may be used for Medicare when drug administered under the direct supervision of a physician, not for use when drug is self-administered)			
J1745	Injection, infliximab, excludes biosimilar (Remicade or Janssen unbranded), 10mg			
J1823	Injection, inebilizumab-cdon, (Uplizna) 1 mg			
J2793	Injection, rilonacept, (Arcalyst) 1 mg			
J3262	Injection, tocilizumab, (Actemra) 1 mg			
J3590	Unclassified biologics (Use to report Abrilada, Amjevita, Bimzelx, Cyltezo, Enspryng, Hadlima, Hyrimoz, Hulio, Kineret, Rystiggo, Ryoncil, Sandoz, Simlandi, Yuflyma, Yusimry)			
J9332	Injection, efgartigimod alfa-fcab,(Vyvgart) 2 mg			
J9333	Injection, rozanolixizumab-noli (Rystiggo), 1 mg			
J9334	Injection, efgartigimod alfa, 2 mg and hyaluronidase-qvfc			
Q5103	Injection, infliximab-dyyb, biosimilar, (Inflectra), 10 mg			
Q5104	Injection, infliximab-abda, biosimilar, (Renflexis), 10 mg			
Q5121	Injection, infliximab-axxq, biosimilar, (Avsola), 10 mg			
Q5133	Injection, tocilizumab-bavi (Tofidence), biosimilar, 1 mg (new code effective 04/01/24)			
Q5135	Injection, tocilizumab-aazg (tyenne), biosimilar, 1 mg (new code effective 10/01/24)			
Q5140	Injection, adalimumab-fkjp, biosimilar, 1 mg (new code effective 01/01/25)			
Q5141	Injection, adalimumab-aaty, biosimilar, 1 mg (new code effective 01/01/25)			
Q5142	Injection, adalimumab-ryvk biosimilar, 1 mg (new code effective 01/01/25)			
Q5143	Injection, adalimumab-adbm, biosimilar, 1 mg (new code effective 01/01/25)			
Q5144	Injection, adalimumab-aacf (idacio), biosimilar, 1 mg (new code effective 01/01/25)			
Q5145	Injection, adalimumab-afzb (abrilada), biosimilar, 1 mg (new code effective 01/01/25)			

Note: CPT codes, descriptions and materials are copyrighted by the American Medical Association (AMA). HCPCS codes, descriptions and materials are copyrighted by Centers for Medicare Services (CMS).

Consideration of Age

Age limits specified in this policy are determined according to US Food and Drug Administration (FDA)-approved indications, where applicable.

For site of service for medical necessity the age described in this policy is 13 years of age or older. Site of service is defined as the location where the drug is administered, such as a hospital-based outpatient setting, an infusion center, a physician's office, or at home. The age criterion for site of service for medical necessity is based on the following: Pediatric individuals are not small adults. Pediatric individuals differ physiologically, developmentally, cognitively, and emotionally from adult individuals, and vary by age groups from infancy to teen. Children often require smaller doses than adults, lower infusion rates, appropriately sized equipment, the right venipuncture site determined by therapy and age, and behavioral management during administration of care. Specialty infusion training is therefore necessary for pediatric IV insertions and therapy. Due to pediatrics unique physiology and psychology, site of service review is limited to individuals above the age of 13.

Benefit Application

Pharmacy Benefit

Cosentyx (secukinumab), Filspari (sparsentan), Lupkynis (voclosporin), Otezla (apremilast), Rezurock (belumosudil), and Tarpeyo (budesonide) are managed through the pharmacy benefit.

Medical Benefit

Avsola (infliximab-axxq), Inflectra (infliximab-dyyb), Infliximab (Janssen – unbranded), Remicade (infliximab), Renflexis (infliximab-abda), Ryoncil (remestemcel-L-rknd), Rystiggo (rozanolixizumab-noli), Saphnelo (anifrolumab-fnia), Tofidence (tocilizumab-bavi), Uplizna (inebilizumab-cdon), Vyvgart (efgartigimod alfa-fcab), and Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) are managed through the medical benefit.

Medical / Pharmacy Benefit

Abrilada (adalimumab-afzb), Actemra (tocilizumab), adalimumab-aacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-ryvk (Simlandi), Amjevita (adalimumab-atto), Benlysta (belimumab), Cyltezo (adalimumab-adbm), Enbrel (etanercept), Enspryng (satralizumab-mwge), Hadlima (adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab), Hyrimoz (adalimumab-adaz), Ilaris (canakinumab), Kineret (anakinra), Orencia (abatacept), Simlandi (adalimumab-ryvk), Tyenne (tocilizumab-aazg), Yuflyma (adalimumab-aaty), and Yusimry (adalimumab-aqvh) are managed through both the pharmacy and medical benefit.

Table 1. Criteria for the International Bone Marrow Transplant Registry (IBMTR) Severity Index for Acute Graft versus Host Disease

Index*	Skin involvement		Liver involvement		Gastrointestinal involvement	
						Volume of
						diarrhea
						(ml/d)
Α	1	< 25%	0	< 34	0	< 500
В	2	25 – 50%	1 – 2	34 – 102	1 – 2	550 – 1500
С	3	> 50%	3	103 – 255	3	> 1500
D	4	Bullae	4	> 255	4	Severe pain and ileus

^{*}Assign Index based on maximum involvement in an individual organ system.

Evidence Review

Miscellaneous Autoimmune Diseases

TNF inhibitors, rituximab and various other agents have been used off-label to treat a variety of autoimmune diseases. Most of this use represents significant unmet medical needs for chronic diseases with few treatment options.



Hidradenitis Suppurativa

Hidradenitis Suppurativa (HS) is an inflammatory skin disease affecting an estimated 1 to 4% of the world population. The main features of HS include painful and chronically recurring, deep-seated follicular nodules, papules, pustules, and abscesses, scarring, sinus tracts, and recurrent discharge. The area's most commonly affected are the under the arms, groin, buttocks, and under the breasts. The disease is variable and recurrent. It may occur as solitary or multiple lesions in one area, or in many areas. In more severe cases, there may be large areas of skin affected by recurrent, draining lesions.

The FDA approved Humira (adalimumab) to treat individuals with HS.

Two randomized, double-blind, placebo-controlled studies (Studies HS-I and II) evaluated the safety and efficacy of Humira in a total of 633 adult subjects with moderate to severe hidradenitis suppurativa (HS) with Hurley Stage II or III disease and with at least 3 abscesses or inflammatory nodules. In both studies, subjects received placebo or Humira at an initial dose of 160 mg at Week 0, 80 mg at Week 2, and 40 mg every week starting at Week 4 and continued through Week 11. Subjects used topical antiseptic wash daily. Concomitant oral antibiotic use was allowed in Study HS-II.

Both studies evaluated Hidradenitis Suppurativa Clinical Response (HiSCR) at Week 12. HiSCR was defined as at least a 50% reduction in total abscess and inflammatory nodule count with no increase in abscess count and no increase in draining fistula count relative to baseline (see Table below). Reduction in HS-related skin pain was assessed using a Numeric Rating Scale in individuals who entered the study with an initial baseline score of 3 or greater on a 11-point scale.

In both studies, a higher proportion of Humira than placebo-treated subjects achieved HiSCR (see **Table 1** below).

Table 1. Efficacy Results at 12 Weeks in Subjects with Moderate to Severe Hidradenitis Suppurativa

HS Study I		HS Study II*	
Placebo			Humira 40 mg
	Weekly		Weekly



	HS Study I		HS Study II*	
Hidradenitis Suppurativa Clinical Response (HiSCR)	N=154, 40 (26%)	N=153, 64 (42%)	N=163, 45 (28%)	N=163, 96 (59%)

^{*19.3%} of subjects in Study HS-II continued baseline oral antibiotic during the study.

In both studies, from Week 12 to Week 35 (Period B), subjects who had received Humira were re-randomized to 1 of 3 treatment groups (Humira 40 mg every week, Humira 40 mg every other week, or placebo). Subjects who had been randomized to placebo were assigned to receive Humira 40 mg every week (Study HS-I) or placebo (Study HS-II).

During Period B, flare of HS, defined as ≥25% increase from baseline in abscesses and inflammatory nodule counts and with a minimum of 2 additional lesions, was documented in 22 (22%) of the 100 subjects who were withdrawn from Humira treatment following the primary efficacy time point in two studies.

Cosentyx (secukinumab)

Two randomized, double-blind, placebo-controlled 52-week Phase 3 trials (i.e., HS Trial 1 [NCT03713619] and HS Trial 2 [NCT03713632]) assessed the efficacy and safety of Cosentyx in the treatment of adult individuals with moderate to severe hidradenitis suppurativa (HS). In both trials, subjects were randomized to placebo or Cosentyx 300 mg by subcutaneous injection at Weeks 0, 1, 2, 3 and 4, followed by 300 mg every 2 weeks or every 4 weeks. At Week 16, subjects who were randomized to placebo were reassigned to receive Cosentyx 300 mg at Weeks 16, 17, 18, 19, and 20 followed by either Cosentyx 300 mg every 2 weeks (Q2W) or Cosentyx 300 mg every 4 weeks (Q4W). In HS Trial 1 and HS Trial 2, a statistically significantly higher proportion of subjects treated with Cosentyx 300 mg every 2 weeks (after the first four weeks) achieved a HiSCR50 response at Week 16 compared to individuals treated with placebo. In both HS trials, a higher proportion of subjects treated with Cosentyx 300 mg every 4 weeks (after the first four weeks) achieved HiSCR50 at Week 16 compared to subjects treated with placebo, where statistical significance was reached in HS Trial 2. In both trials, the onset of action of Cosentyx occurred as early as Week 2 and the efficacy progressively increased up to Week 16.



Lupus – Systemic Lupus Erythematosus (SLE)

Systemic lupus erythematosus (SLE) is a chronic, complicated, progressive autoimmune disease impacting multiple organ systems. It is a condition characterized by auto-reactive B-cells. Autoantibody production from such abnormal B lymphocyte function leads to chronic inflammation and cellular, tissue and organ damage. Diverse in presentation, individuals with SLE experience mild to life-threatening manifestations and unpredictable clinical course of exacerbations and remissions. As symptoms are non-specific, the identification of SLE is oftentimes delayed. It has been reported that individuals visit a mean of three different physicians and an average of 4 years after the onset of symptoms before a correct diagnosis is reached.

The mucocutaneous (rash), articular (arthritis), serosal (pleuritis, pericarditis), renal (proteinuria) and neurologic (seizures, psychosis) clinical features, as well as hematologic and immunologic laboratory findings, incorporated in the American College of Rheumatology SLE diagnosis classification criteria reflects the heterogeneity of the disease. Most commonly involved organs include the skin, musculoskeletal, renal, nervous, cardiovascular and pulmonary systems. Over 75% of SLE individuals have debilitating, generally non-fatal mucocutaneous (rash) and musculoskeletal involvement (arthritis). A smaller SLE population (50%-66%) is afflicted with renal disorders and is associated with poorer outcome and mortality. About 2/3 of SLE individuals also present with varying severity of neuropsychiatric manifestations ranging from mood disorders, anxiety, psychosis to seizures. Other less common but serious manifestations include serositis (16 to 64%), neurological disorders (9 to 36%), and immune-mediated cytopenia's (4 to 43%). Depression is common among people with chronic autoimmune disease. Overall, SLE individuals have a 2-5 times greater mortality rate.

As endogenous female sex hormone is identified to have a role in SLE development, SLE is found primarily in women (90% of SLE population are female, 6-10 female:1 male), typically 15-44 years of age. In the US, more than 300,000 people have SLE and an annual incident rate of 15,000. 4 million people are impacted worldwide.

While SLE individuals have at least twice the mortality risk relative to the general population, survival rate at 15 years improved dramatically from 50% in the 1950s to currently greater than 80%. Most common causes of death are cardiovascular disease, infections, renal disease and complications due to SLE disease activity.

In addition to gender, ethnicity has an influence on the development of SLE. Mestizo, indigenous Americans, Blacks and Asians have more severe SLE disease and poorer clinical progression. Blacks are three times more likely than Caucasians to have SLE. Asian and African American SLE individuals develop renal disease more frequently than those of European descent (60-70%, 50%, 20-30%, respectively).



SLE is characterized by auto-reactive B-cells. Autoantibody production from such abnormal B lymphocyte function leads to chronic inflammation. Autoantibody complex, cytokines and complement activation represent mediators of tissue damage in SLE individuals. Anti-nuclear antibody (ANA) is found present in more than 90% of individuals. Those positive are more likely to have active lupus associated with B-cell dysfunction. Anti-dsDNA, a type of ANA, is one of the diagnosis criteria established by the American College of Rheumatology and is monitored as gauge of SLE disease response to treatment. Consistent with existing pathophysiology, inhibition of BlyS, an endogenous protein responsible for B-cell homeostasis, decreases autoreactive B-cell activity and serological changes. Transgenic animals overexpressing BlyS have lupus-like syndrome, increased immunoglobulins and immune complex depositions. BlyS is also found elevated in human autoimmune diseases such as rheumatoid arthritis, multiple sclerosis and Sjogren's.

Most individuals present with generalized symptoms of fatigue, fever, anorexia, weight loss, photosensitivity, malar rash, oral ulcers, arthralgia, and hair loss. Incompletely controlled SLE can progress to end-stage organ involvement; SLE activity of 60% of SLE individuals is found to worsen within 2-7 years of diagnosis. Irreversible cellular and tissue damages can accumulate to result in life-threatening renal, cardiac, pulmonary, CNS and hematological system toxicities. The subsequent development of pleuritis, pericarditis, stroke, seizure, nephritis, vasculitis, anemia, thrombocytopenia and other blood dyscrasias present significant mortality and morbidity risks.

Aside from these autoimmune mediated disease manifestations, SLE individual are in high risk for infections of the respiratory and urinary systems, cardiovascular diseases, hematological and solid tumors, maternal and fetal morbidity and mortality (spontaneous abortions, pre-eclampsia, intrauterine growth impairment, premature birth). Most common causes of death are infections, renal disease, cardiovascular disease and complications due to SLE disease activity.

The current SLE standard of care is similar across the world. Treatment of mild-to-moderate symptoms involves the use of non-steroidal anti-inflammatory drugs (NSAIDs), antimalarial drugs such as hydroxychloroquine and corticosteroids such as prednisone and its equivalent. For life-threatening manifestations such as the renal, CNS, cardiovascular and pulmonary systems, aggressive single or combination of treatments with high dose corticosteroids and immunosuppressive agents such as cyclophosphamide, azathioprine, methotrexate and mycophenolate is used. Corticosteroids, hydroxychloroquine and aspirin have FDA approved SLE indications.

Particularly for individuals with active and life-threatening disease activity, SLE remains an unmet medical disease. The very treatments used to alleviate lupus symptoms have poor tolerability and short- and long-term morbidity risks. Ones used for mild/mod SLE flares involves nonspecific immune system suppression. Aggressive treatments such as cyclophosphamide is



associated with gonadal toxicity, whereas high dose corticosteroids (>7.5 mg/day, cumulative doses >365g) can lead to cataracts, osteoporosis, metabolic disorders, increased infections, edema, weight gain and hyperlipidemia. This is especially concerning as SLE individuals tend to be young women of childbearing age, have lower immune system and greater cardiovascular risks due to the nature of the underlying autoimmune disease. Currently there is no approved SLE treatment shown to prolong survival or reverse the course of the disease.

Benlysta (belimumab)

Benlysta (belimumab) is an FDA-approved 147kDa, recombinant fully human $IgG1\lambda$ monoclonal antibody. It targets a novel pathway to potentially treat SLE by binding to soluble, endogenous human B-lymphocyte stimulator BlyS (also known as B-cell activating factor or BAFF, TALL-1, THANK, TNFSF13B, zTNF4). The binding inhibits BlyS biological activity of B-cell selection, survival, differentiation and eventual antibody formation of native, activated plasmacytoid and plasma cells.

The efficacy of belimumab was studied in two Phase III trials. SLE Responder Index (SRI) response at 52 weeks, the primary endpoint, was met for belimumab 10 mg/kg treatment arm in both BLISS 52 [1.83 OR (1.30-2.59), p=0.0006] and BLISS76 [1.52 OR (1.07-2.15), p=0.0207]. Overall, secondary endpoints of reduction in severe flare, steroid use, autoantibodies, B-cell subsets, normalization of complement levels and improvement in quality of life were also achieved. 66% of the FDA Arthritis Advisory Committee (10 out of 15) felt the clinical data provided support of efficacy. Concerns were cited over the lack of study consistency within and between the phase 3 studies, lack of statistical significance for some populations and the exclusion of SLE individuals with severe renal or central nervous system diseases. The representative nature of the SLE individuals sampled was also questioned.

The two-Phase III studies were set-up nearly identically, though differences in baseline demographics, serological activity, geographical location and concurrent SLE medication use necessitate their separate analyses. Bliss 76 was conducted in North America and Europe, with 70% Caucasian and 14% African American. Relative to BLISS 52, BLISS 76 had a lower baseline SLE activity (less of SS score >=10, proteinuria>= 2g/24 hours, 1A or 2B BILAG, auto-antibodies, much less prescribed corticosteroid, while using greater NSAIDS and immunosuppressive agents). The data from BLISS 76 clinical trial was less convincing, with its narrower incremental benefit of belimumab over placebo in SRI response, steroid use and SLE flare reduction, lack of efficacy for African American groups, and later onset of significant SS score improvement (32 weeks versus 16 weeks in BLISS 52). With the exception of African American groups, the evidence from BLISS 52 clinical trial was stronger, more robust and consistent across different



ethnicities. A lower number of BLISS 52 participants receiving 10mg/kg belimumab required an increase of corticosteroids. Reduction in flares and prolongation to first flare were seen only in this ex-US-conducted study.

For both studies, disease manifestation resolution often seen in organ systems were those commonly involved at baseline: mucocutaneous (rash, oral ulcers, alopecia), immunologic (serological measures of disease activity, anti-dsDNA and complements) and musculoskeletal (arthritis). SLE activity reduction was also observed with the vascular (vasculitis) and central nervous system (lupus headache), both systems of which were less commonly involved at study initiation. However, resolution of similarly less frequently involved hematology abnormalities and fever was not observed in the belimumab group. The statistically significant difference in improvement from baseline as benchmarked by SRI response was driven largely by improvement of the mucocutaneous and musculoskeletal systems, and not organ systems more associated with poor SLE outcome and mortality (kidneys, central nervous system, blood vessels). Observations of these serious organ manifestations were too uncommon to assess treatment effects.

Subgroup analyses revealed a lack demonstrated efficacy in African American subjects in both Phase III studies, which contradicted the positive treatment response previously observed in LBS02 Phase II trial. Similarly, Native Americans were found more associated with favorable disease activity reduction in BLISS 52 but not its counterpart trial. There was some geographical dependence, as participants from US and Canada had smaller treatment effect compared to some other regions. Since belimumab is to be administered chronically, durability and onset of response are of concern. Of note, differences in efficacy endpoint at the conclusion of BLISS 76 were no longer statistically significant between treatment arms [PLO 32%, 10mg/kg 39%, 1.3 (0.9, 1.9), p=0.13], which was a drop from PLO 34%, 10mg/kg 43% 1.5 (1.07, 2.15), p=0.0207 in the preceding 24 weeks. Dose-response was not consistent; throughout the studies, 1mg/kg was noticed at times to be more, or just as effective as the more potent proposed formulation. Individuals with severe renal or central nervous system (CNS) diseases were not evaluated and therefore efficacy is not known. A disclaimer to this effect was included in the final approved product label.

As safety data were pooled from the three intravenous belimumab clinical studies (LBS02, BLISS 52 and BLISS76) in an attempt to generate a sufficiently large sample of rare events, the ability to detect safety trend concerning specific ethnicity and geological populations was lost. Overall, headache, upper respiratory tract infection and arthralgia were some of the common adverse events experienced by belimumab participants. Pyrexia was the most reported serious adverse event. The investigational drug was found to be associated with greater risk of infection, mortality and psychiatric events ranging from depression, suicidal ideation to suicide. Notably,



no such neuropsychiatric adverse events were seen in those receiving only SLE standard therapy. Malignancy and hypersensitivity rates were comparable to the placebo group. While belimumab has safety signals, its safety profile is favorable and relatively minor compared to the side effects experienced by those on current SLE standard-of-care. 14 of the 15 Advisory Committee members agreed that the clinical data provided adequate safety evidence.

In Trial 4 the safety and efficacy of Benlysta IV was evaluated in an international, randomized, double-blind, placebo-controlled, 52-week, pharmacokinetics (PK), efficacy and safety study conducted in 93 pediatric individuals with a clinical diagnosis of SLE according to the American College of Rheumatology classification criteria. Individuals had active SLE disease, defined as a SELENA-SLEDAI score ≥6 and positive autoantibodies at screening as defined in the adult trials. Individuals were on a stable SLE treatment regimen (standard of care) and had similar inclusion and exclusion criteria as in the adult studies. The median age was 15 years (range: 6 to 17). The majority (95%) of individuals were female. More than 50% of individuals had 3 or more active organ systems involved at baseline. The most common active organ systems at baseline based on SELENA-SLEDAI were mucocutaneous (91%), immunologic (74%), and musculoskeletal (73%). Overall, 19% of pediatric individuals had some degree of renal activity and less than 7% had activity in the cardio-respiratory, hematologic, CNS or vascular systems. Randomization into age-related treatment cohorts was stratified by screening SELENA-SLEDAI scores (6 to 12 vs >13) and age (5 to 11 years vs 12 to 17 years).

The primary efficacy endpoint was the SLE Responder Index (SRI-4) at Week 52. There was a numerically higher proportion of pediatric individuals achieving a response in SRI-4 and its components in pediatric individuals receiving Benlysta IV plus standard therapy compared with placebo plus standard therapy.

At baseline, 95% of pediatric individuals were receiving prednisone. Among those pediatric individuals, 20% of pediatric individuals receiving Benlysta IV plus standard therapy reduced their average prednisone dose by at least 25% per day during Weeks 44 through 52 compared with 21% of pediatric individuals on placebo plus standard therapy.

In Trial 4, the probability of experiencing a severe SLE flare, as measured by the modified SELENA-SLEDAI Flare Index, excluding severe flares triggered only by an increase of the SELENA-SLEDAI score to >12, was calculated. The proportion of pediatric individuals reporting at least one severe flare during the study was numerically lower in pediatric individuals receiving Benlysta IV plus standard therapy (23%) compared with those receiving placebo plus standard therapy (43%). Pediatric individuals receiving Benlysta IV 10 mg/kg plus standard therapy had a 62% lower risk of experiencing a severe flare during the 52 weeks of observation, relative to the placebo plus standard therapy group. Of the pediatric individuals experiencing a severe flare, the median time to the first severe flare was 160 days in pediatric individuals receiving Benlysta



IV plus standard therapy compared with 82 days in pediatric individuals receiving placebo plus standard therapy.

Saphnelo (anifrolumab-fnia)

Saphnelo (anifrolumab-fnia) is a human IgG1 κ monoclonal antibody that binds to subunit 1 of the type I interferon receptor (IFNAR) with high specificity and affinity. This binding inhibits type I IFN signaling, thereby blocking the biologic activity of type I IFNs. Anifrolumab also induces the internalization of IFNAR1, thereby reducing the levels of cell surface IFNAR1 available for receptor assembly. Blockade of receptor mediated type I IFN signaling inhibits IFN responsive gene expression as well as downstream inflammatory and immunological processes. Inhibition of type I IFN blocks plasma cell differentiation and normalizes peripheral T-cell subsets. Type I IFNs play a role in the pathogenesis of SLE. Approximately 60-80% of adult individuals with active SLE express elevated levels of type I IFN inducible genes.

Anifrolumab has been studied in two Phase 3 trials for SLE and a Phase IIIb trial extension of a Phase II trial. Anifrolumab is also under study for lupus nephritis (Phase II) and in a subcutaneous format (Phase II).

The TULIP-1 and TULIP-2 trials were 52-week, multicenter, double-blind, randomized, placebo-controlled, Phase III studies. Both trials included individuals 18-70 years of age who met ACR criteria for SLE and who had moderate to severe active disease. This was defined as a Systemic Lupus Erythematosus Dis-ease Activity Index-2000 (SLEDAI-2K) score ≥ 6 excluding points related to fever, lupus-related headache (HA), or organic brain syndrome and a clinical SLEDAI-2K score without laboratory results of ≥ 4 . Additionally, severe disease activity in ≥ 1 organ or moderate in ≥ 2 organs as defined by the BILAG-2004 index (organ domain scores ≥ 1 A item or ≥ 2 B items) and physician's global assessment (PGA) ≥ 1 on a four-point scale visual analogue scale (VAS) scale were required. Individuals were also stable on ≥ 1 SLE treatment. Individuals with severe lupus nephritis or neuropsychiatric lupus were excluded.

The TULIP-1 trial randomized 457 individuals to anifrolumab 300 mg IV every 4 weeks, anifrolumab 150 mg IV every 4 weeks, or placebo.1 All comparisons were conducted between anifrolumab 300 mg and placebo only. The primary efficacy measure was SRI-4 at 52 weeks while the key secondary endpoints were reduction in steroid dose \leq 7.5 mg from week 40-52 if the baseline dose of steroid was \geq 10 mg, \geq 50% re-duction in Cutaneous Lupus Erythematosus Disease Area and Severity Index (CLASI) score at week 12 in individuals with moderate to severe cutaneous activity (CLASI \geq 10 at baseline), annualized flare rate at Week 52, SRI-4 at Week 24, and SRI-4 at Week 52 in individuals with high IFN gene signature (IFNGS) status. BICLA response



at Week 52 was assessed as an "other" secondary endpoint. The primary outcome of SRI-4 response was defined as ≥4 point reduction in SLEDAI-2K from baseline, no new disease activity in any organ (defined as ≥1 new BILAG A item or ≥2 BILAG B items), no worsening in PGA score (defined as ≥0.3 points increase from baseline), and no study treatment discontinuation or use of restricted medications beyond protocol-allowed thresholds. Anifrolumab did not meet the primary outcome of SRI-4 at 52 weeks (36% anifrolumab vs 40% placebo, p=0.412); therefore, all secondary endpoints were considered nominal. Key secondary outcomes of reduction steroid dose, annualized flare rate, SRI-4 response at 24 weeks, and SRI-4 response in individuals with high IFNGS did not reach significance. However, more individuals in the anifrolumab group achieved ≥50% reduction in CLASI score from baseline at week 12 than placebo (42% vs 25%, nominal p=0.005). Of note, the original study protocol considered individuals with new NSAIDs or an NSAID dose change as nonresponders. The authors stated these original rules were inconsistent with the intention of the protocol and were inappropriate. The sponsor and a group of SLE experts revised the study rules and instituted a post-hoc amendment which considered individuals non-responders only if changes in NSAID use occurred during the last 2 weeks of the study. However, no significant difference in the primary outcome of SRI-4 at 52 weeks was identified between groups despite the amendment (47% anifrolumab vs 43% placebo, p=0.455).

The TULIP-2 trial randomized 365 individuals to anifrolumab 300 mg IV every 4 weeks or placebo. The primary efficacy measure was changed during the study from SRI-4 to the difference in BICLA response between groups at week 52. This occurred before unblinding of the data and was done in response to the results of the TULIP-1 trial. BICLA response was defined as all of the following: 1) reduction of all severe or moderately severe (BILAG A or B) disease activity at baseline to lower levels and no worsening in other organ systems (worsening defined as ≥1 new BILAG A item or ≥2 BILAG B items); 2) no worsening in disease activity per SLEDAI-2K score and PGA score (defined as no increase of ≥0.3 from baseline); 3) no discontinuation of trial intervention; and 4) no use of restricted medications beyond protocol-allowed thresholds. Key secondary endpoints included BICLA response at Week 52 in individuals with high IFNGS at baseline, reduction in steroid dose to ≤7.5 mg/day from week 40-52 if baseline dose was ≥10 mg/d; ≥50% reduction in CLASI at week 12 in individuals with moderate to severe cutaneous activity defined as CLASI ≥10, ≥50% reduction in swollen or tender joints at week 52 in individuals with ≥6 swollen and ≥6 tender joints at baseline, and annualized flare rate at Week 52. NSAID rules consistent with the post-hoc amendment from the TULIP-1 trial were used in the TULIP-2 trial. Anifrolumab significantly increased the primary outcome of the BICLA response at 52 weeks compared to placebo (47.8% vs 31.5%, p=0.001). Additionally, anifrolumab significantly improved the key secondary outcomes of BICLA at 52 weeks in individuals with high IFNGS, reduced steroid dose, reduction in CLASI activity, and annualized fare rate compared to placebo. There was no difference between groups in reduction in swollen and tender joints



(p=0.55). SRI-4 results were not considered key and were not multiplicity adjusted. The difference be-tween groups in SRI-4 at 52 weeks was 18.2% (95% confidence interval [CI] 8.1-28.3), favoring anifrolumab.

Pyoderma Gangrenosum

Pyoderma gangrenosum is an inflammatory disease with dermatologic manifestations including painful ulcerations with erythematous borders. It is presumed to be autoimmune in origin, though the mechanism is not well understood. Lesions usually develop at sites of minor skin injury, usually on the lower extremities. These lesions can grow in size and become necrotic. Underlying fasciitis may occasionally develop from them. Some individuals develop pustular, bullous or vegetative lesions. Other common sites are colostomies and paraneoplastic lesions in individuals with hematologic malignancies. Progress of the lesions is highly variable, and individual response to treatment is heterogeneous. Obesity, diabetes or edema may be contributing factors.

Due to the infrequent occurrence and heterogeneity of pyoderma gangrenosum, the treatment approach is empiric and individual specific. First-line options include topical tacrolimus, nicotine, and 5-ASA, systemic corticosteroids and immunosuppressant agents such as azathioprine, cyclosporine, methotrexate and mycophenolate. When these approaches fail, biologic therapy is usually tried. Successful treatment with TNF inhibitors (etanercept, adalimumab, infliximab) has been reported. Response to ustekinumab and various investigational interleukin inhibitors has also been reported. Surgical management is another option.

Wegener's Granulomatosis and Microscopic Polyangiitis

Wegener's granulomatosis (WG) is an autoimmune vasculitis that may affect various internal organs and can be potentially life-threatening. Symptoms vary and can mimic a variety of other diseases, making it difficult to diagnose. These include rhinitis, glomerulonephritis, pulmonary nodules and hemorrhage, neuropathies, gastrointestinal symptoms and various other inflammatory manifestations. The disease can occur at any age, usually in adults.

WG can be recognized by the distinctive triad of granulomatous inflammation, necrosis, and vasculitis of the respiratory tract. Vasculitis in other regions is also common. It can follow a varied clinical course that is strongly influenced by treatment. Untreated, generalized WG is usually lethal. Historically, treatment with immunosuppressants has been used. Glucocorticoids and cyclophosphamide have been a standard therapy, but this is limited by cyclophosphamide

toxicity. If remission is achieved, less toxic agents such as azathioprine may be employed for maintenance.

The FDA has approved rituximab in combination with glucocorticoids, to treat individuals with WG and microscopic polyangiitis (MPA). Both of these diseases affect people of all ages and ethnicities, and both genders. The causes of these disorders are unknown, and both are considered orphan diseases because they each affect less than 200,000 people in the United States.

Giant Cell Arteritis

Giant cell arteritis (GCA) is an inflammation of the lining of the arteries. It affects the arteries in the head, especially those in the temples. Temporal arteritis is another name for this disease. GCA frequently causes headaches, scalp tenderness, jaw pain, and vision problems.

The safety of subcutaneous Actemra (tocilizumab) has been studied in one Phase III study (WA28119) with 251 GCA individuals. The total individual years duration in the Actemra GCA all exposure population was 138.5 individual years during the 12-month double blind, placebocontrolled phase of the study. The overall safety profile observed in the Actemra treatment groups was generally consistent with the known safety profile of Actemra. There was an overall higher incidence of infections in GCA individuals relative to RA individuals. The rate of infection/serious infection events was 200.2/9.7 events per 100 individual years in the Actemra weekly group and 160.2/4.4 events per 100 individual years in the Actemra every other week group as compared to 156.0/4.2 events per 100 individual years in the placebo + 26-week prednisone taper and 210.2/12.5 events per 100 individual years in the placebo + 52-week taper groups.

Neuromyelitis Optica Spectrum Disorders

Neuromyelitis optica spectrum disorders (NMOSD), previously known as Devic disease or neuromyelitis optica (NMO) are CNS inflammatory disorders characterized by severe, immunemediated demyelination and axonal damage predominantly targeting optic nerves and spinal cord. Differential diagnosis is from RRMS. Presentation is generally bilateral and monophasic and may be difficult to distinguish from MS due to variability in presentation and clinical course, but once diagnosed, a different treatment strategy is indicated. Hallmark features include acute attacks of bilateral or rapidly sequential optic neuritis (leading to severe visual loss) or transverse myelitis (often causing limb weakness, sensory loss, and bladder dysfunction) with a typically



relapsing course. Attacks most often occur over days, with variable degrees of recovery over weeks to months. Other suggestive symptoms include episodes of intractable nausea, vomiting, hiccups, excessive daytime somnolence or narcolepsy, reversible posterior leukoencephalopathy syndrome, neuroendocrine disorders, and (in children) seizures. While no clinical features are disease-specific, some are highly characteristic. Optic neuritis presents with varying degrees of vision loss and is almost always associated with eye pain that worsens with movement of the eye.

Reported prevalence of NMOSD ranges from 0.5 to 10 per 100,000. The reported incidence of NMOSD in women is 5-10 times higher than in men. Median age of onset is 32 to 40, it sometimes occurs in children or older adults. It may be overrepresented in some non-European populations, including Africans, East Asians, and Latin Americans, MS is less prevalent. Reported prevalence is higher among black compared with white individuals, but the evidence for this is relatively weak. In Japan, optic-spinal multiple sclerosis (OSMS), represents approximately 15 to 40 percent of MS. Whether NMOSD and Asian OSMS are the same remains uncertain. NMOSD is usually sporadic, though a few familial cases have been reported.

NMOSD has a relapsing course in most cases. In some individuals, optic neuritis and transverse myelitis occur concurrently; in others, clinical episodes are separated by a variable time delay. Relapse occurs within the first year following an initial event in 60 percent of individuals and within three years in 90 percent. As a rule, severe residual deficits follow initial and subsequent attacks, leading to rapid development of disability due to blindness and paraplegia within five years.

MS is mostly cell-mediated, while NMOSD is thought to be primarily mediated by the humoral immune system. Damage is to both gray and white matter of the optic nerves and associated spinal segments. A disease-specific serum NMO-immunoglobulin G (IgG) antibody selectively binds aquaporin-4 (AQP4), previously known as NMO IgG. Presence of aquaporin-4 (AQP4)-immunoglobulin G (IgG) antibodies is required for definitive diagnosis. Serum anti-AQP4 titers correlate with clinical disease activity, drop after immunotherapy, and remain low during remissions. Titers at the nadir of attacks correlate with spinal cord damage. AQP4 is a water channel protein. AQP4-IgG antibodies that bind to astrocyte AQP4 water channels, leading to astrocyte dysfunction and the clinical manifestations of nausea and vomiting. A potential subset of individuals have anti-myelin oligodendrocyte glycoprotein (MOG).

NMOSD is frequently associated with systemic autoimmune disorders, including hypothyroidism, pernicious anemia, ulcerative colitis, myasthenia gravis, and idiopathic thrombocytopenic purpura; systemic lupus erythematosus, antiphospholipid syndrome, and Sjögren syndrome, and sometimes with neoplasms.



Myasthenia Gravis

Myasthenia gravis (MG) is a chronic autoimmune disease mainly characterized by fatigue and muscle weakness in ocular, limb, and respiratory muscles. Many individuals also experience bulbar weakness, which refers to an impairment of the lower cranial nerves. This results in difficulty talking, chewing, swallowing, and holding up the head. The degree of muscle weakness can fluctuate and vary in severity from person to person; however, it will generally improve with rest and worsen with physical activity. Other precipitating factors include pregnancy, infection, surgery, and stress. The cause of MG is unknown, but it is usually diagnosed in young women (20 to 30 years of age) or men ≥50 years of age. The life expectancy for MG individuals is near normal. The mortality rate is now about 3%, mainly due to the risk of myasthenic crisis, a potentially life-threatening complication in which muscle weakness causes respiratory failure. The muscle weakness presenting in MG is due to an antibody-mediated immunologic attack directed at proteins in the postsynaptic membrane of the neuromuscular junction. Myasthenia gravis has been associated with antibodies against 3 postsynaptic proteins: acetylcholine receptor (AChR), muscle-specific kinase (MuSK), and low-density lipoprotein receptor-related protein 4 (LRP4). AChR antibody-positive individuals represent the vast majority of gMG individuals.

Vyvgart (efgartigimod alfa-fcab)

Vyvgart is a first-in-class human immunoglobulin G1 (IgG1) antibody fragment that binds the neonatal Fc receptor (FcRn), keeping antibodies in circulation and preventing FcRn from recycling IgG back into the blood. This causes a reduction in overall levels of IgG, including the abnormal AChR antibodies that are present in most individuals with gMG. Vyvgart was evaluated in the Phase 3 ADAPT trial, a 26-week randomized, double-blind, placebo-controlled study that was conducted in North America, Europe, and Japan. Study participants were ≥18 years of age with class II to IV gMG. These individuals were eligible to participate in the study regardless of AChR antibody status if they had a Myasthenia Gravis Activities of Daily Living (MG-ADL) score of at least 5 (>50% non-ocular) and were on a stable dose of at least 1 treatment for gMG. The primary analysis of ADAPT was completed in a modified intention-to-treat population of all AChR antibody—positive individuals who had a valid baseline MG-ADL assessment and at least 1 post-baseline MG-ADL assessment. Participants were randomly assigned (1:1) to Vyvgart (10 mg/kg) or matching placebo, administered as 4 infusions per cycle (1 infusion per week), repeated as needed depending on clinical response no sooner than 8 weeks after initiation of the previous cycle. The efficacy of Vyvgart was measured using the Myasthenia Gravis-Specific



Activities of Daily Living scale (MG-ADL) which assesses the impact of gMG on daily functions of 8 signs or symptoms that are typically affected in gMG. Each item is assessed on a 4-point scale where a score of 0 represents normal function and a score of 3 represents loss of ability to perform that function. A total score ranges from 0 to 24, with the higher scores indicating more impairment. In this study, an MGADL responder was defined as an individual with a 2-point or greater reduction in the total MG-ADL score compared to the treatment cycle baseline for at least 4 consecutive weeks, with the first reduction occurring no later than 1 week after the last infusion of the cycle. The primary efficacy endpoint was the comparison of the percentage of MG-ADL responders during the first treatment cycle between treatment groups in the AChR-Ab positive population. A statistically significant difference favoring Vyvgart was observed in the MG-ADL responder rate during the first treatment cycle [67.7% in the Vyvgart-treated group vs 29.7% in the placebo-treated group (p<0.0001)].

The safety analysis included all randomly assigned individuals who received at least 1 dose or partial dose of Vyvgart or placebo. In the ADAPT trial, 77% of individuals in the Vyvgart group and 84% of individuals in the placebo group had treatment-emergent adverse events; the most frequent of which were headache (Vyvgart [29%] versus and nasopharyngitis (Vyvgart [12%] versus placebo [18%]). In addition, 4 (5%) Vyvgart-treated individuals and 7 (8%) individuals in the placebo group had a serious adverse event; 3 individuals in each treatment group (4%) discontinued treatment during the study. There were no deaths.

Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc)

Vyvgart Hytrulo is a first-in-class neonatal F c receptor blocker, which is administered as subcutaneous (SC) injection that is approved for gMG by the FDA. It contains efgartigimod alfa, a human immunoglobulin G1 (IgG1) antibody fragment that binds the neonatal Fc receptor (FcRn), keeping antibodies in circulation and preventing FcRn from recycling IgG back into the blood. It also contains recombinant human hyaluronidase PH20, which is Halozyme Therapeutics' Enhanced drug delivery technology that facilitates the SC delivery. The safety and efficacy of Vyvgart Hytrulo was evaluated in a phase 3, randomized, multicenter, open-label, parallel group bridging study to the phase 3 ADAPT study. Individuals were randomized 1:1 to receive Vyvgart Hytrulo or Vyvgart once a week for four weeks. The primary efficacy endpoint was to compare the mean IgG reduction between two groups. At the end of the treatment period, the mean total IgG reduction was 66.6% in the Vyvgart group compared to 62.2% in the Vyvgart group, with p-value < 0.0001. Similar responses were found in the Myasthenia Gravis Activities of Daily Living (MG-ADL) and Quantitative Myasthenia Gravis (QMG). Also, the safety profile of Vyvgart was similar to Vyvgart Hytrulo other than injection site reactions, which were higher in the Vyvgart Hytrulo group.



Rystiggo (rozanolixizumab-noli)

Rystiggo, administered as a subcutaneous (SC) infusion, is a humanized immunoglobulin G4 monoclonal antibody that binds to neonatal Fc receptor (FcRn), which reduces the levels of circulating IgG. It is FDA-approved for the treatment of generalized myasthenia gravis (gMG) in adult individuals who are anti-AChR or anti-MuSK Ab+. The efficacy of Rystiggo for the treatment of gMG in adults who are anti-AChR Ab+ or anti-MuSK Ab+ was established in the Phase 3 MycarinG trial (Study 1; NCT03971422), a multicenter, randomized, double-blind, placebo-controlled study. The study included a 4-week screening period and a 6-week treatment period, followed by 8 weeks of observation. During the treatment period, Rystiggo or placebo were administered as an SC infusion once a week for 6 weeks. In the MycarinG study, 200 individuals were randomly assigned (1:1:1) to receive SC infusions of Rystiggo 7 mg/kg, Rystiggo 10 mg/kg, or placebo once a week for 6 weeks. Treatment with Rystiggo resulted in a greater reduction in the Myasthenia Gravis Activities of Daily Living (MG-ADL) total score at Day 43 than placebo (−3.4 versus −0.8 points). The most common adverse reactions reported in ≥10% of individuals receiving Rystiggo were headache, infections, diarrhea, pyrexia, hypersensitivity reactions, and nausea.

Graft versus Host Disease

Graft-versus-host disease (GVHD) is a potentially fatal complication following allogeneic hematopoietic stem cell transplantation (HSCT) and occurs when immune cells transplanted from a non-identical donor (graft) recognize the transplant recipient (host) as foreign. This initiates an immune reaction, causing damage across different organs and tissues. Acute graft-versus-host disease (aGVHD) classically presents within 100 days of HSCT (usually 2 to 3 weeks post-transplant) and primarily affects the skin, liver, and gastrointestinal (GI) tract. This marker of 100 days is not absolute; some individuals may experience persistent, recurrent, or late-onset aGVHD > 100 days after HSCT. Individuals can experience clinical manifestations of aGVHD such as rash, persistent nausea and vomiting, abdominal cramping, and diarrhea. It is estimated that there are approximately 10,000 allogeneic HSCTs performed in the United States every year. Despite the use of current prophylactic regimens, aGVHD occurs in 20% to 80% of HSCT individuals. Even in fully human leukocyte antigen (HLA)–matched (preferred donor source) allogeneic HSCT, the incidence of aGVHD is estimated at about 30% to 50%. The overall survival rate of individuals has improved over the past 2 decades with new advances in technology and antiinfectives. The overall 5-year survival rate in aGVHD individuals is now estimated to be up to



72%. Individuals with aGVHD usually die due to infection or severe GI complications, which are usually resistant to steroid therapy.

Orencia (abatacept)

Orencia is an immunomodulator that inhibits T-cell activation by binding to CD80 and CD86 on antigen-presenting cells; therefore, it can block the signaling processes that would otherwise induce T cells to attack the host. Orencia was studied in acute Graft Versus Host Disease (GVHD) in 2 phase 2 studies: GVHD-1 and GVHD-2. GVHD-1 was a Phase 2, multicenter, 2-cohort clinical trial of 186 individuals ≥6 years of age who underwent HSCT from a matched unrelated donor and received Orencia (or placebo) on Days −1, 5, 14, and 28 in combination with a calcineurin inhibitor (e.g. cyclosporine or tacrolimus) on Day −2 through at least Day 100 and methotrexate on Days 1, 3, 6, and 11. Grade III-IV aGVHD free survival rate was 87% in the Orencia arm and 75% in the placebo arm. The rate of grade II-IV aGVHD free survival was 50% in the Orencia arm and 32% in the placebo arm. Overall survival rate was 97% in the Orencia arm versus 84% in the placebo arm.

GVHD-2, the second study supporting Orencia's approval in aGVHD, used real-world data from the Center for International Blood and Marrow Transplant Research (CIBMTR). This observational study included individuals ≥6 years of age who underwent HSCT from a 1 allele–mismatched unrelated donor between 2011 and 2018 and analyzed the outcomes of individuals who had received Orencia in combination with CNI and methotrexate (n = 54) versus individuals who received CNI and methotrexate alone (n = 162) for the prophylaxis of aGVHD. Forty-two individuals from the GVHD-1 study were included in the Orencia group in the GVHD-2 study. Efficacy was established based on overall survival at Day 180 post-transplant; the overall survival rate at Day 180 in the Orencia group was 98% (95% confidence interval [CI]: 78%, 100%) versus 75% (95% CI: 67%, 82%) in the comparator group (P = 0.0028). Efficacy for Orencia was established based on overall survival and moderate GFS (grade II–IV) results. Orencia did not significantly improve severe GFS (grade III–IV) in the GVHD-1 trial. However, overall survival rates were similar between the GVHD-1 trial and the real-world data analysis from CIBMTR.

n the GVHD-1 study, serious adverse reactions reported up to Day 225 post-transplant included fever (20%), pneumonia (8%), acute kidney injury (7%), diarrhea (6%), hypoxia (5%), and nausea (5%). Common adverse reactions included anemia, hypertension, cytomegalovirus (CMV) reactivation/infection, fever, pneumonia, nosebleed, decrease in CD4 lymphocytes, hypermagnesemia, and acute kidney injury. Individuals receiving Orencia should be monitored for Epstein-Barr virus reactivation before starting treatment and for 6 months post-transplant and CMV infection/reinfection for 6 months post-transplant.



Rezurock (belumosudil)

Rezurock is a rho-associated, coiled-coil kinase 2 (ROCK2) inhibitor. ROCK2 is a signaling pathway that modulates inflammatory response and fibrotic processes. By inhibiting ROCK2, Rezurock is thought to restore immune homeostasis and reduce fibrosis in affected organs. Rezurock was approved based on the results of the Phase 2 randomized, multicenter ROCKstar clinical trial, which enrolled individuals ≥ 12 years of age with chronic graft versus host disease who had received 2–5 previous lines of systemic therapy (including Imbruvica and Jakafi). The primary endpoint of overall response rate was met by 75% of individuals receiving Rezurock 200 mg once daily and was consistent across all organ systems; 69% (n = 45) of individuals displayed a partial response and 6% (n = 4) displayed a complete response. Overall, Rezurock was well-tolerated with adverse effects similar to corticosteroids and other immunosuppressants.

Ryoncil (remestemcel-L-rknd)

The approval of Ryoncil was supported by data from the Phase 3, single-arm MSB-GVHD001 trial, which included pediatric individuals with aGVHD who failed to respond to systemic corticosteroid therapy. MSB-GVHD001 was a single-arm, prospective study that enrolled 55 pediatric individuals 2 months to 17 years of age with grade B-D (as defined by the International Bone Marrow Transplant Registry system) SR-aGVHD. All participants received IV infusions of Ryoncil at 2 × 106 MSCs/kg (actual body weight at screening) twice per week for 4 consecutive weeks. Individuals could continue receiving stable doses of steroid therapy (until eligible for steroid tapering, as determined by the treating physician) and continue their established prophylactic aGVHD regimen. No other medications for the treatment of SR-aGVHD could be introduced to individuals during the initial 28 days of Ryoncil administration unless disease progression occurred. The overall response rate at Day 28 was 70%, including 30% of individuals who achieved complete response. Overall response rates (ORRs) at Day 28 and overall survival (OS) rates at Day 100 were similar across age groups (0–7 years, 8–12 years, 13–17 years), severity of disease (grade C, grade D), risk of disease (standard vs. high), and organs involved at baseline (skin, lower GI, multiple organs). Thirty individuals received >8 initial infusions: of those, 25 individuals received continued therapy for partial response/mixed response, and five individuals received additional therapy for aGVHD flares. Among 21 individuals who achieved partial response at Day 28 and who received continued therapy after Day 28, 16 (76.2%) achieved an overall response (OR) by Day 56 and 19 (90.5%) achieved an OR by Day 100. Survival rates at Day 100 were comparable between individuals who received the initial treatment (4–8 infusions) and those who continued treatment (9–12 infusions) (76.2% vs. 72.0%).



The most common adverse reactions (incidence ≥20%) were infectious disorders, pyrexia, hemorrhage, edema, abdominal pain, and hypertension. Serious adverse reactions occurred in 35 individuals (65%), with eight individuals (15%) discontinuing treatment due to adverse reactions or death.

Primary Immunoglobulin A nephropathy (IgAN)

Immunoglobulin A nephropathy (IgAN) is an autoimmune kidney disease where immunoglobulin A deposits in the glomerular mesangium of the kidneys and attacks the glomeruli. This diminishes the kidney's capacity to filter, resulting in the leakage of blood and protein into the urine. Over many years, the damage may progress slowly, leading to scarring of the nephrons. Eventually IgA nephropathy can lead to end-stage renal disease (ESRD). Individuals can experience clinical manifestations of IgAN such as hematuria with or without proteinuria, acute kidney injury, and rapidly progressive glomerulonephritis. There are approximately 150,000 people affected with IgAN in the United States. The management of primary IgAN includes supportive care such as lifestyle modifications, reducing blood pressure to an optimal level, reducing proteinuria to an optimal level through renin-angiotensin system inhibition, and immunosuppressive therapy.

Filspari (sparsentan)

Filspari is a dual-acting angiotensin II type 1 (AT₁R) and endothelin type A (ET_AR) receptor antagonist that selectively blocks the action of two vasoconstrictor and mitogenic agents to reduce proteinuria in adults with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression. Endothelin-1 and angiotensin II are believed to participate in the pathogenesis of immunoglobulin A nephropathy (IgAN) via the ET_AR and AT₁R pathway. The approval of Filspari for IgAN has been granted under the accelerated approval pathway due to observed reduction in proteinuria.

PROTECT study was randomized, Double-blind, parallel-group, multicenter, active-control study to determine the efficacy and safety of sparsentan compared to irbesartan in the treatment of IgAN. This study included 404 individuals \geq 18 years of age with persistent proteinuria (total urine protein \geq 1.0 g/ day despite being on maximized stable dose of RAS inhibitor treatment (\geq 50% of maximum labeled dose). These individuals were randomized 1:1 to receive Filspari 400 mg once daily following 200 mg once daily for 14 days or irbesartan 300 mg once daily dose following 150 mg once daily for 14 days. The trial protocol allowed for the initiation of rescue immunosuppressive treatment at the investigator's discretion. However, the usage of SGLT2



inhibitors was prohibited during the trial. The primary endpoint of the study was the change, relative to baseline, in urine protein/creatinine ratio (UPCR) at week 36. Following a 36-week treatment period, individuals in the sparsentan group exhibited a mean reduction in proteinuria of 49.8% from baseline, while individuals in the irbesartan treatment group demonstrated a mean reduction in proteinuria of 15.1% from baseline. The secondary endpoint was overall change in eGFR from baseline, change in eGFR over 104-week period and change in eGFR over a 52-week period.

Sparsentan was overall well tolerated. Most common adverse events were peripheral edema, dizziness, hypotension, anemia, and hyperkalemia. An increase in ALT/AST level of at three times the upper limit of normal was observed in 2.5% of individuals in the clinical trial, and evidence of fetal harm was detected in animal reproduction studies. There are two specific reasons have resulted in Filspari being available only through the Filspari REMS (Risk Evaluation and Mitigation Strategy) program.

Tarpeyo (budesonide)

Tarpeyo was approved based on the results from the first part of the Phase 3 NeflgArd study (NCT03643965), a randomized, double-blind trial in adult patients with biopsy-verified IgAN, reduced kidney function (estimated glomerular filtration rate [eGFR] \geq 35 mL/min/1.73 m2), and proteinuria (\geq 1 g/day or urine protein to creatinine ratio [UPCR] \geq 0.8) who were receiving a stable dose of a maximally tolerated renin-angiotensin system (RAS) inhibitor therapy, either angiotensin-covering enzyme (ACE) inhibitors or angiotensin receptor blockers (ARBs). In Part A of the study, individuals were randomized to receive Tarpeyo 16 mg once daily (n = 97) or placebo (n = 102) for 9 months, followed by a 2-week taper of either Tarpeyo 8 mg once daily or placebo. The primary endpoint of the study was percentage reduction in UPCR from baseline. At 9 months, a 34% reduction in UPCR was observed in individuals receiving Tarpeyo versus a 5% reduction in the placebo group (31% [95% confidence interval, 16% to 42%]; P = 0.0001). Adverse effects were mild or moderate in severity in Part A of the NeflgArd study. Common adverse reactions (>5%) included hypertension (16%), peripheral edema (14%), muscle spasms (13%), acne (11%), dermatitis (7%), weight increase (7%), dyspnea (6%), and face edema (6%).

2019 Update

Reviewed prescribing information and conducted literature search for all drugs listed in policy. Updated criteria for Benlysta (belimumab) IV for use in individuals aged 5 years and older.



2020 Update

Reviewed prescribing information for all drugs listed in policy and conducted a literature search on the management of hidradenitis suppurativa, pyoderma gangrenosum, and systemic lupus erythematosus. No new evidence found that would change this policy. Added links to the ACR, EULAR/ACR, and SLICC criteria.

2021 Update

Reviewed prescribing information for all drugs listed in policy and conducted a literature search on the management of pyoderma gangrenosum, giant cell arteritis, and neuromyelitis optica spectrum disorder. No new evidence found that would change this policy. Added Arcalyst (rilonacept) to policy for the FDA-approved indications which is treatment of cryopyrin-associated periodic syndromes (CAPS), maintenance of remission of deficiency of interleukin-1 receptor antagonist (DIRA), and treatment of recurrent pericarditis (RP). Updated llaris (canakinumab) criteria adding requirement the drug is prescribed by or in consultation with a rheumatologist, geneticist, or dermatologist which brings drug criteria in alignment with Kineret (anakinra) and Arcalyst (rilonacept) for the management of CAPS. Updated the investigational table adding restrictions on combination therapy and for drug quantities that exceed the FDA labeled dosing for condition.

2022 Update

Reviewed prescribing information and conducted literature search for all drugs listed in policy. No new evidence found that would change this policy. Added criteria for Vyvgart for the treatment of generalized myasthenia gravis (gMG) in adult individuals who are antiacetylcholine receptor (AChR) antibody positive. Added criteria for Orencia for the prophylaxis of acute graft versus host disease (aGVHD).

2023 Update

Reviewed prescribing information and conducted literature search for all drugs listed in policy. No new evidence found that would change this policy. Added criteria for Filspari for the



treatment of proteinuria in adults with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression. Added criteria for Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qyfc) for the treatment of generalized myasthenia gravis (gMG) in adult individuals who are anti-acetylcholine receptor (AChR) antibody positive. Added coverage for the biosimilars Hyrimoz LCF (adalimumab-adaz) SC, Abrilada (adalimumab-afzb) SC, Hulio ((adalimumab-fkjp) SC, Yusimry (adalimumab-aqvh) SC, Hadlima (adalimumab-bwwd) SC, and Yuflyma (adalimumab-aaty) SC for the treatment of HS, PG, and uveitis as non-preferred products and with the identical coverage criteria as Amjevita (adalimumab-atto) [NDCs starting with 72511]. Added coverage for Cyltezo LCF (adalimumab-adbm), Hyrimoz HCF (adalimumabadaz) and Adalimumab-adaz HCF (Sandoz - unbranded) SC for the treatment of HS, PG, and uveitis as preferred products and with the identical coverage criteria as Amjevita (adalimumabatto) [NDCs starting with 55513]. Moved Avsola to 1st line (preferred) with the effective date of 01/01/2024. Added Avsola to the list of preferred infliximab products to be tried and failed prior to non-preferred infliximab products with the effective date of 01/01/2024. Moved Inflectra to 2nd line (non-preferred) infliximab products with the effective date of 01/01/2024. Removed Inflectra from the list of preferred infliximab products to be tried and failed prior to trying nonpreferred infliximab products with the effective date of 01/01/2024. Added Humira biosimilars Adalimumab-fkjp (Biocon-unbranded) and Idacio (adalimumab-aacf) as non-preferred products with similar criteria as Amjevita (adalimumab-atto) [NDCs starting with 72511]. Updated criteria for Actemra for the treatment of CRS to require documentation confirming the diagnosis. Added criteria for Rystiggo (rozanolixizumab-noli) for the treatment of gMG. Updated Amjevita [NDCs starting with 55513] to a non-preferred product effective January 1, 2024. Added Hyrimoz (Cordavis) [NDCs starting with 83457] and adalimumab-aacf (Idacio) as a non-preferred product effective January 1, 2024. Added adalimumab-adbm (Cyltezo unbranded) as a preferred product effective January 1, 2024. Updated Hyrimoz LCF (Sandoz) from a non-preferred to a preferred product effective January 1, 2024.

2024 Update

Reviewed prescribing information and conducted literature search for all drugs listed in policy. Added coverage criteria for Cosentyx (secukinumab) for the treatment of adults with moderate to severe hidradenitis suppurativa. Updated Vyvgart (efgartigimod alfa-fcab) criteria to require that medication is not being used concurrently with Vyvgart Hytrulo, Rystiggo, Soliris, Ultomiris, or Zilbrysq. Updated Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) criteria to require that medication is not being used concurrently with Vyvgart, Rystiggo, Soliris, Ultomiris, or Zilbrysq. Added coverage criteria for Tarpeyo (budesonide) for the treatment of adults with primary immunoglobulin A nephropathy (IgAN). Added coverage criteria for Rezurock



(belumosudil) for the treatment of chronic graft versus host disease. Updated coverage criteria for Cosentyx (secukinumab) and removed adalimumab step therapy requirement for the treatment of adults with moderate to severe hidradenitis suppurativa. Added Humira (adalimumab) (Cordavis) [NDCs starting with 83457] as a non-preferred product. Added adalimumab-aaty (Yuflyma unbranded) as a non-preferred product. Added Simlandi (adalimumab-ryvk) and adalimumab-ryvk (Simlandi unbranded) as preferred products. Updated Lupkynis (voclosporin) coverage criteria to clarify that the requirement is for Lupkynis (voclosporin) to be used in combination with mycophenolate, cyclophosphamide, azathioprine, or an immunosuppressant and a corticosteroid. Updated Benlysta (belimumab) SC for systemic lupus erythematosus (SLE) coverage criteria to include coverage of pediatric individuals 5 years and older. Updated non-preferred adalimumab coverage criteria to require trial and treatment failure with all preferred adalimumab products. Updated Rystiggo (rozanolixizumab-noli) coverage criteria to require that the medication not being used concurrently with Vyvgart (efgartigimod alfa-fcab), Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-gvfc), Soliris (eculizumab), Ultomiris (ravulizumab-cwvz), or Zilbrysq (zilucoplan). Added adalimumab and infliximab coverage criteria for the treatment of certain individuals with sarcoidosis. Minor correction to indicate that Actemra (tocilizumab) IV requires site of service review. Clarified the use of Lupkynis (voclosporin) without changes to policy statements. Added Tofidence (tocilizumab-bavi) and Tyenne (tocilizumab-bavi) coverage criteria for the treatment of certain individuals with cytokine release syndrome and giant cell arteritis. Updated Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) coverage criteria to include treatment of certain individuals with chronic inflammatory demyelinating polyneuropathy (CIDP). Added site of service review for Tofidence (tocilizumab-bavi) IV. The following changes are effective January 3, 2025. Changed Inflectra (infliximab-dyyb) to a first-line agent. Changed Avsola (infliximab-axxq) to a second-line agent. Updated coverage criteria for Avsola and Renflexis to require the individual to have an adequate trial and failure with Inflectra, Infliximab (Janssen – unbranded), or Remicade. Updated Rystiggo criteria to require for AChR antibody positive myasthenia gravis the individual has tried and failed Soliris, Ultomiris, Vyvgart, or Vyvgart Hytrulo. Updated Hyrimoz (Sandoz) (adalimumab-adaz) [NDCs starting with 61314] from a preferred product to a non-preferred product. Updated Humira (AbbVie) (adalimumab) [NDCs starting with 00074] to require that the individual has had an inadequate response or intolerance to a preferred product for new starts. Clarified that the medications listed in this policy are subject to the product's FDA dosage and administration prescribing information. Added the following to note to all criteria for adalimumab products: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please



check the member Plan booklet or member ID card to determine whether this policy criteria applies.

2025 Update

Reviewed prescribing information and conducted literature search for all drugs listed in policy. Policy updated to indicate that Site of Service Medical Necessity criteria does not apply to Alaska fully-insured members pursuant to Alaska HB 226 (accessed January 3, 2025). Clarified that non-formulary exception review authorizations for all drugs listed in this policy may be approved up to 12 months. Clarified that the Filspari (sparsentan) coverage criteria is for slowing kidney function decline in adults with primary immunoglobulin A nephropathy (IgAN) who are at risk for disease progression per the updated prescribing information language. Added coverage criteria for Niktimvo (axatilimab-csfr) and Fabhalta (iptacopan). Updated Tarpeyo (budesonide) urine protein-to-creatinine ratio from ≥ 1.5 g/g to ≥ 0.8 g/g or proteinuria ≥ 1 g/day. Updated Actemra (tocilizumab) IV, Tofidence (tocilizumab-bavi) IV, and Tyenne (tocilizumab-aazg) IV cytokine release syndrome (CRS) coverage criteria to indicate that coverage is medically necessary if CRS is treatment-induced (not limited to CAR-T products) and grade 3-4. Updated Kineret (anakinra) coverage criteria to include treatment of certain individuals with cytokine release syndrome. Clarified the use of Lupkynis (voclosporin) without changes to policy statements. Added an exception to the site-of-service requirements for certain individuals receiving treatment for cytokine release syndrome (CRS). Updated Abrilada (adalimumab-afzb), adalimumab-aacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-fkjp (Hulio unbranded), Amjevita (adalimumab-atto), Hadlima (adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab) (Cordavis) [NDCs starting with 83457], Hyrimoz (adalimumab-adaz), Idacio (adalimumab-aacf), Yuflyma (adalimumab-aaty), Yusimry (adalimumab-aqvh), adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-ryvk (Simlandi unbranded), Cyltezo (adalimumab-adbm), Simlandi (adalimumab-ryvk), Enbrel (etanercept), Inflectra (infliximab-dyyb), infliximab (Janssen – unbranded), Remicade (infliximab), Avsola (infliximab-axxq), Renflexis (infliximab-abda), and Humira (adalimumab) (AbbVie) [NDCs starting with 00074] to include an age requirement for pyoderma gangrenosum coverage criteria. Updated Abrilada (adalimumab-afzb), adalimumabaacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-fkjp (Hulio unbranded), Amjevita (adalimumab-atto), Hadlima (adalimumab-bwwd), Hulio (adalimumabfkjp), Humira (adalimumab) (Cordavis) [NDCs starting with 83457], Hyrimoz (adalimumab-adaz), Idacio (adalimumab-aacf), Yuflyma (adalimumab-aaty), Yusimry (adalimumab-agvh), adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumabryvk (Simlandi unbranded), Cyltezo (adalimumab-adbm), Simlandi (adalimumab-ryvk), Inflectra



(infliximab-dyyb), infliximab (Janssen – unbranded), Remicade (infliximab), Avsola (infliximab-axxq), Renflexis (infliximab-abda), and Humira (adalimumab) (AbbVie) [NDCs starting with 00074] to include an age requirement for sarcoidosis coverage criteria. Updated Humira (adalimumab) (AbbVie) [NDCs starting with 00074] from a preferred to a non-preferred adalimumab product. Added coverage criteria for Ryoncil (remestemcel-L-rknd).

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History

Date	Comments
07/01/16	New policy approved June 14, 2016, add to Prescription Drug section. Policy information on drug treatment for miscellaneous autoimmune diseases extracted from 5.01.550. Medical necessity review criteria for site of service IV therapy added.
10/01/16	Interim Update, approved September 13, 2016: inclusion of a new indication for Humira; changing criteria for Benlysta (defining "adequate" trial of previous therapies).
11/01/16	Interim review, approved October 11, 2016. Clarified age criteria language indicating that site of service review is applicable to only those age 13 and older; drug criteria review applies to all ages. Coding update, added HCPCS Q5102.
07/01/17	Annual review, approved June 13, 2017. Added coverage criteria for Actemra in the setting of giant cell arteritis, added HCPCS code J3262. Formatting update; added hyperlinks to Medical Necessity criteria sections.
08/15/17	Interim Review, approved August 15, 2017. Added Benlysta SC.
09/01/17	Interim review, approved August 15, 2017. Added Infliximab-abda (Renflexis) to coverage criteria and coding section. Clarified pyoderma gangrenosum first-line/second-line treatment.
11/01/17	Interim Review, approved October 3, 2017. Clarified site of service exception criterion related to access: There is no outpatient infusion center within 50 miles of the individual's home and there is no contracted home infusion agency that will travel to their home, or a hospital is the only place that offers infusions of this drug. Removed HCPCS codes J3490 and J3590.
02/14/18	Interim Review, approved February 13, 2018. Update hospital-based outpatient coverage from 30 days to 90 days.
04/01/18	Coding update: added new HCPCS codes Q5103 and Q5104 (effective 4/1/18), noted that Q5102 terminated 4/1/18.
07/01/18	Annual Review, approved June 22, 2018. Dosage and quantity limit prescribing table was removed. Two related medical policies were added in related medical policy section.
11/01/18	Minor update, the Site of Service criteria was updated for clarity.

Date	Comments
12/01/18	Interim Review, approved November 21, 2018. Updated pediatric indications for Humira: uveitis and hidradenitis.
01/01/19	Coding update, added new HCPCS code Q5109 (new code effective 1/1/19).
04/01/19	Coding update: removed HCPCS code Q5102 as it terminated 4/1/18.
08/01/19	Annual Review, approved July 25, 2019. Updated criteria for Benlysta (belimumab) IV. Removed HCPCS code J9310.
09/01/19	Interim Review, approved August 22, 2019. Added criteria for Otezla (apremilast) for Bechet's Disease.
01/01/20	Interim Review, approved December 17, 2019, effective for dates of service on or after April 3, 2020, following provider notification. Added Ruxience (rituximab-pvvr) with Rituxan.
10/01/20	Annual Review, approved September 8, 2020. Added coverage criteria for Uplizna (inebilizumab-cdon) for the treatment of NMOSD. Added coverage criteria for Enspryng (satralizumab-mwge) for the treatment of NMOSD. Added Avsola (infliximab-axxq) as a second-line agent for the treatment pyoderma gangrenosum along with site-of-service requirement. Added HCPCS codes Q5121 and J3590 Effective for dates of service on or after January 1, 2021, after provider notification:
	Added Ilaris (canakinumab) to policy with coverage criteria for periodic fever syndromes and Still's disease. Added HCPCS code J0638.
01/01/21	Interim Review, approved December 17, 2020. Added coverage criteria for Actemra (tocilizumab) for the treatment of cytokine release syndrome. Added HCPCS code J1823.
02/01/21	Interim Review, approved January 12, 2021. Added coverage criteria for Benlysta (belimumab) for the treatment of lupus nephritis. Removed HCPCS J0717 and Q5109.
06/01/21	Interim Review, approved May 11, 2021. Added Kineret (anakinra) for the treatment of cryopyrin-associated periodic syndromes and the deficiency of interleukin-1 receptor antagonist. Added Lupkynis (voclosporin) for the treatment of lupus nephritis. Updated Benlysta (belimumab) criteria for the treatment of lupus nephritis removing prior use of Benlysta in the prior 12 months and adding restriction on combination therapy with Lupkynis.
09/01/21	Annual Review, approved August 10, 2021. Updated Ilaris (canakinumab) criteria adding requirement the drug is prescribed by or in consultation with a rheumatologist, geneticist, or dermatologist. Updated the investigational table adding restrictions on combination therapy and for drug quantities that exceed the FDA labeled dosing for condition. Added Arcalyst (rilonacept) for the treatment of DIRA, CAPS, and RP. Coverage criteria for Arcalyst (rilonacept) (HCPCS code J2793) becomes effective for dates of service on or after December 2, 2021, following 90-day provider notification.
11/01/21	Interim Review, approved October 12, 2021. Added coverage criteria for Saphnelo (anifrolumab-fnia) for the treatment of adult individuals with SLE. Updated Benlysta



Date	Comments
	(belimumab) criteria regarding concurrent use with Saphnelo (anifrolumab-fnia) for the treatment of SLE. Added site of service review for Uplizna (inebilizumab-cdon) for dates of service on or after February 4, 2022.
01/01/22	Interim Review, approved December 14, 2021. Updated Humira criteria for the treatment of hidradenitis suppurativa to include individual has tried at least one other therapy and prescriber specialty. Updated Humira criteria for the treatment of uveitis to include individual has tried at least one other therapy and prescriber specialty. For pyoderma gangrenosum added prescriber specialty to Humira, Enbrel, Remicade, Inflectra, Renflexis, and Avsola. Updated Actemra criteria for the treatment of giant cell arteritis to include individual has tried at least one other therapy and prescriber specialty. Updated Otezla criteria for the treatment of Behcet's Disease to include individual has tried at least one other therapy and prescriber specialty. Added HCPCS code C9086.
04/01/22	Annual Review, approved March 8, 2022. Added criteria for Vyvgart for the treatment of generalized myasthenia gravis in adult individuals who are AChR antibody positive. Added criteria for Orencia for the prophylaxis of acute graft versus host disease. Added HCPCS code J0129. Added term date to HCPC code C9086. Added code J0491.
06/01/22	Interim Review, approved May 10, 2022. Added Infliximab (Janssen – unbranded) to policy with identical site-of-service requirements and coverage criteria as brand Remicade (infliximab) for the treatment of pyoderma gangrenosum. Moved Inflectra (infliximab-dyyb) to a first-line TNF-α antagonists for the treatment of pyoderma gangrenosum. Updated coverage criteria for Renflexis (infliximab-abda) and Avsola (infliximab-axxq) for the treatment of pyoderma gangrenosum to require the individual has had an inadequate response or intolerance to Infliximab (Janssen – unbranded), Inflectra (infliximab-dyyb), or Remicade (infliximab).
07/01/22	Coding update. Added HCPCS code J9332.
10/01/22	Interim Review, approved September 13, 2022. Updated Benlysta IV and Benlysta SC criteria for the treatment of SLE to require the drug is being used as add-on-therapy following standard induction. Updated Benlysta IV criteria for the treatment of active lupus nephritis from 18 years of age or older to 5 years of age or older. Changed the wording from "patient" to "individual" throughout the policy for standardization.
02/01/23	Interim Review, approved January 10, 2023. Added coverage for the biosimilar Amjevita (adalimumab-atto) for the treatment of hidradenitis suppurativa, pyoderma gangrenosum, and uveitis with the identical coverage criteria as Humira (adalimumab). Added HCPC code J0135. Added Amjevita to HCPC code J3590.
04/01/23	Interim Review, approved March 14, 2023. Added criteria for Filspari (sparsentan) for the treatment of proteinuria in adults with primary immunoglobulin A nephropathy (IgAN) at risk of rapid disease progression.
08/01/23	Annual Review, approved at MPC, July 11, 2023. Reviewed prescribing information for all drugs in the policy. Added criteria for Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) for the treatment of generalized myasthenia gravis (gMG) in adult



Date	Comments
	individuals who are anti-acetylcholine receptor (AChR) antibody positive. Added coverage for the biosimilars Hyrimoz LCF (adalimumab-adaz) SC, Abrilada (adalimumab-afzb) SC, Hulio ((adalimumab-fkjp) SC, Yusimry (adalimumab-aqvh) SC, Hadlima (adalimumab-bwwd) SC, and Yuflyma (adalimumab-aaty) SC for the treatment of HS, PG, and uveitis as non-preferred products and with the identical coverage criteria as Amjevita (adalimumab-atto) [NDCs starting with 72511]. Added coverage for Cyltezo LCF (adalimumab-adbm), Hyrimoz HCF (adalimumab-adaz) and Adalimumab-adaz HCF (Sandoz – unbranded) SC for the treatment of HS, PG, and uveitis as preferred products and with the identical coverage criteria as Amjevita (adalimumab-atto) [NDCs starting with 55513]. Added Cyltezo, Hyrimoz HCF, Adalimumab-adaz HCF (Sandoz – unbranded), Abrilada, Hadlima, Hulio, Hyrimoz LCF, Yuflyma and Yusimry to code J3590.
09/01/23	Interim Review, approved August 8, 2023. The following policy changes are effective September 1, 2023: added Humira biosimilars Adalimumab-fkjp (Biocon-unbranded) and Idacio (adalimumab-aacf) as non-preferred products with similar criteria as Amjevita (adalimumab-atto) [NDCs starting with 72511]. The following policy changes are effective January 1, 2024 following 90-day provider notification due to changes in the preferred medical benefit drugs: moved Avsola to 1st line (preferred); added Avsola to the list of preferred infliximab products to be tried and failed prior to non-preferred infliximab products; removed Inflectra from the list of preferred infliximab products to be tried and failed prior to trying non-preferred infliximab products.
11/01/23	Interim Review, approved October 10, 2023. Updated criteria for Actemra for the treatment of CRS to require documentation confirming the diagnosis.
12/01/23	Interim Review, approved November 14, 2023. Added criteria for Rystiggo (rozanolixizumab-noli) for the treatment of gMG. Added drug name Rystiggo to HCPCS code J3590.
01/01/24	Interim Review, approved December 12, 2023. Updated Amjevita [NDCs starting with 55513] to a non-preferred product. Added Hyrimoz (Cordavis) [NDCs starting with 83457] and adalimumab-aacf (Idacio) as a non-preferred product. Added adalimumab-adbm (Cyltezo unbranded) as a preferred product. Updated Hyrimoz LCF (Sandoz) from a non-preferred to a preferred product. Added new HCPCS codes J9333 and J9334.
02/01/24	Annual Review, approved January 9, 2024. Added coverage criteria for Cosentyx (secukinumab) for the treatment of adults with moderate to severe hidradenitis suppurativa. Updated Vyvgart (efgartigimod alfa-fcab) criteria to require that medication is not being used concurrently with Vyvgart Hytrulo, Rystiggo, Soliris, Ultomiris, or Zilbrysq. Updated Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) criteria to require that medication is not being used concurrently with Vyvgart, Rystiggo, Soliris, Ultomiris, or Zilbrysq. Added coverage criteria for Tarpeyo (budesonide) for the treatment of adults with primary immunoglobulin A nephropathy



Date	Comments
	(IgAN). Added coverage criteria for Rezurock (belumosudil) for the treatment of chronic graft versus host disease.
03/01/24	Interim Review approved February 13, 2024. Updated coverage criteria for Cosentyx (secukinumab) and removed adalimumab step therapy requirement for the treatment of adults with moderate to severe hidradenitis suppurativa.
05/01/24	Interim Review, approved April 9, 2024. Added Humira (adalimumab) (Cordavis) [NDCs starting with 83457] as a non-preferred product.
07/01/24	Interim Review, approved June 11, 2024. Added adalimumab-aaty (Yuflyma unbranded) as a non-preferred product. Added Simlandi (adalimumab-ryvk) and adalimumab-ryvk (Simlandi unbranded) as preferred products. Updated Lupkynis (voclosporin) coverage criteria to clarify that the requirement is for Lupkynis (voclosporin) to be used in combination with mycophenolate, cyclophosphamide, azathioprine, or an immunosuppressant and a corticosteroid. Updated Benlysta (belimumab) SC for systemic lupus erythematosus (SLE) coverage criteria to include coverage of pediatric individuals 5 years and older. Updated non-preferred adalimumab coverage criteria to require trial and treatment failure with all preferred adalimumab products. Updated Rystiggo (rozanolixizumab-noli) coverage criteria to require that the medication not being used concurrently with Vyvgart (efgartigimod alfa-fcab), Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc), Soliris (eculizumab), Ultomiris (ravulizumab-cwvz), or Zilbrysq (zilucoplan). Added drug name Simlandi to HCPCS code J3590.
09/01/24	Interim Review, approved August 13, 2024. Added adalimumab and infliximab coverage criteria for the treatment of certain individuals with sarcoidosis. Minor correction to indicate that Actemra (tocilizumab) IV requires site of service review. Clarified the use of Lupkynis (voclosporin) without changes to policy statements. Added Tofidence (tocilizumab-bavi) and Tyenne (tocilizumab-bavi) coverage criteria for the treatment of certain individuals with cytokine release syndrome and giant cell arteritis. Updated Vyvgart Hytrulo (efgartigimod alfa and hyaluronidase-qvfc) coverage criteria to include treatment of certain individuals with chronic inflammatory demyelinating polyneuropathy (CIDP). The following policy change is effective December 5, 2024, following 90-day provider notification. Added site of service review for Tofidence (tocilizumab-bavi) IV. Added HCPCS code Q5133 for Tofidence.
10/01/24	Interim Review, approved September 10, 2024. The following policy changes are effective January 3, 2025, following a 90-day provider notification. Changed Inflectra (infliximab-dyyb) to a first-line agent. Changed Avsola (infliximab-axxq) to a second-line agent. Updated coverage criteria for Avsola and Renflexis to require the individual to have an adequate trial and failure with Inflectra, Infliximab (Janssen – unbranded), or Remicade. Updated Rystiggo criteria to require for AChR antibody positive myasthenia gravis the individual has tried and failed Soliris, Ultomiris, Vyvgart, or Vyvgart Hytrulo. Updated Hyrimoz (Sandoz) (adalimumab-adaz) [NDCs starting with 61314] from a preferred product to a non-preferred product. Updated Humira (AbbVie) (adalimumab) [NDCs starting with 00074] to require that the individual has had an

Date	Comments
	inadequate response or intolerance to a preferred product for new starts. Coding update. Added new HCPCS code Q5135 effective 10/1/2024.
11/14/24	Minor update made to EDSS link.
01/01/25	Interim Review, December 10, 2024. Clarified that the medications listed in this policy are subject to the product's FDA dosage and administration prescribing information. Added the following to note to all criteria for adalimumab products: This medical necessity criteria does not apply to one Open formulary (Formulary ID: 6062; Rx Plan F1) and one Incentive formulary (Formulary ID: 6064; Rx Plan G3). The criteria for members with these custom Open and Incentive formulary plans can be found in policy 5.01.647 Medical Necessity Criteria for Custom Incentive and Open Formularies. Please check the member Plan booklet or member ID card to determine whether this policy criteria applies. Added drug name Bimzelx to unclassified HCPCS code, J3590. Added new HCPCS codes J0139, Q5140, Q5141, Q5142, Q5143, Q5144, Q5145.
02/01/25	Annual Review, approved January 14, 2025. Policy updated to indicate that Site of Service Medical Necessity criteria does not apply to Alaska fully-insured members; only Medical Necessity criteria for the infusion drug applies pursuant to Alaska HB 226 (link added). Clarified that non-formulary exception review authorizations for all drugs listed in this policy may be approved up to 12 months. Clarified that the Filspari (sparsentan) coverage criteria is for slowing kidney function decline in adults with primary immunoglobulin A nephropathy (IgAN) who are at risk for disease progression per the updated prescribing information language. Added coverage criteria for Niktimvo (axatilimab-csfr) and Fabhalta (iptacopan). Updated Tarpeyo (budesonide) urine protein-to-creatinine ratio from ≥ 1.5 g/g to ≥ 0.8 g/g or proteinuria ≥ 1g/day. Updated Actemra (tocilizumab) IV, Tofidence (tocilizumab-bavi) IV, and Tyenne (tocilizumab-aazg) IV cytokine release syndrome (CRS) coverage criteria to indicate that coverage is medically necessary if CRS is treatment-induced (not limited to CAR-T products) and grade 3-4. Updated Kineret (anakinra) coverage criteria to include treatment of certain individuals with cytokine release syndrome. Clarified the use of Lupkynis (voclosporin) without changes to policy statements. Updated Abrilada (adalimumab-afzb), adalimumab-aacf (Idacio unbranded), adalimumab-aaty (Yuflyma unbranded), adalimumab-bwwd), Hulio (adalimumab-fkjp), Humira (adalimumab-atto), Hadlima (adalimumab-bwwd), Hulio (adalimumab-adbm), Gotlezo unbranded), adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-adaz (Hyrimoz unbranded), Amjevita (adalimumab-aqvh), adalimumab-adaz (Hyrimoz unbranded), Cyltezo (adalimumab-adbm), Simlandi (adalimumab-ryvk), Enbrel (etanercept), Inflectra (infliximab-dyyb), infliximab dbase abda), and Humira (adalimumab) (AbbVie) (NDCs starting with 00074) to include an age requirement for pyoderma gangrenosum coverage criteria. Updated Abrilada (adalimumab-afzb), ad



Date	Comments
	(Cordavis) [NDCs starting with 83457], Hyrimoz (adalimumab-adaz), Idacio (adalimumab-aacf), Yuflyma (adalimumab-aaty), Yusimry (adalimumab-aqvh), adalimumab-adaz (Hyrimoz unbranded), adalimumab-adbm (Cyltezo unbranded), adalimumab-ryvk (Simlandi unbranded), Cyltezo (adalimumab-adbm), Simlandi (adalimumab-ryvk), Inflectra (infliximab-dyyb), infliximab (Janssen – unbranded), Remicade (infliximab), Avsola (infliximab-axxq), Renflexis (infliximab-abda), and Humira (adalimumab) (AbbVie) [NDCs starting with 00074] to include an age requirement for sarcoidosis coverage criteria. Added an exception to the site-of-service requirements for certain individuals receiving treatment for cytokine release syndrome (CRS).
03/01/25	Interim Review, approved February 11, 2025. Added coverage criteria for Ryoncil (remestemcel-L-rknd). The following policy changes are effective July 1, 2025, following a 90-day provider notification. Updated Humira (adalimumab) (AbbVie) [NDCs starting with 00074] from a preferred to a non-preferred adalimumab product.

Disclaimer: This medical policy is a guide in evaluating the medical necessity of a particular service or treatment. The Company adopts policies after careful review of published peer-reviewed scientific literature, national guidelines and local standards of practice. Since medical technology is constantly changing, the Company reserves the right to review and update policies as appropriate. Member contracts differ in their benefits. Always consult the member benefit booklet or contact a member service representative to determine coverage for a specific medical service or supply. CPT codes, descriptions and materials are copyrighted by the American Medical Association (AMA). ©2025 Premera All Rights Reserved.

Scope: Medical policies are systematically developed guidelines that serve as a resource for Company staff when determining coverage for specific medical procedures, drugs or devices. Coverage for medical services is subject to the limits and conditions of the member benefit plan. Members and their providers should consult the member benefit booklet or contact a customer service representative to determine whether there are any benefit limitations applicable to this service or supply. This medical policy does not apply to Medicare Advantage.

