



Original Effective Date: 10/27/2021
Current Effective Date: 12/28/2025
Last P&T Approval/Version: 10/29/2025
Next Review Due By: 10/2026
Policy Number: C21970-A

Saphnelo (anifrolumab-fnia)

PRODUCTS AFFECTED

Saphnelo (anifrolumab-fnia)

COVERAGE POLICY

Coverage for services, procedures, medical devices and drugs are dependent upon benefit eligibility as outlined in the member's specific benefit plan. This Coverage Guideline must be read in its entirety to determine coverage eligibility, if any. This Coverage Guideline provides information related to coverage determinations only and does not imply that a service or treatment is clinically appropriate or inappropriate. The provider and the member are responsible for all decisions regarding the appropriateness of care. Providers should provide Molina Healthcare complete medical rationale when requesting any exceptions to these guidelines.

Documentation Requirements:

Molina Healthcare reserves the right to require that additional documentation be made available as part of its coverage determination; quality improvement; and fraud; waste and abuse prevention processes. Documentation required may include, but is not limited to, patient records, test results and credentials of the provider ordering or performing a drug or service. Molina Healthcare may deny reimbursement or take additional appropriate action if the documentation provided does not support the initial determination that the drugs or services were medically necessary, not investigational or experimental, and otherwise within the scope of benefits afforded to the member, and/or the documentation demonstrates a pattern of billing or other practice that is inappropriate or excessive.

DIAGNOSIS:

Moderate to severe systemic lupus erythematosus (SLE)

REQUIRED MEDICAL INFORMATION:

This clinical policy is consistent with standards of medical practice current at the time that this clinical policy was approved. If a drug within this policy receives an updated FDA label within the last 180 days, medical necessity for the member will be reviewed using the updated FDA label information along with state and federal requirements, benefit being administered and formulary preferencing. Coverage will be determined on a case-by case basis until the criteria can be updated through Molina Healthcare, Inc. clinical governance. Additional information may be required on a case-by-case basis to allow for adequate review. When the requested drug product for coverage is dosed by weight, body surface area or other member specific measurement, this data element is required as part of the medical necessity review. The Pharmacy and Therapeutics Committee has determined that the drug benefit shall be a mandatory generic and that generic drugs will be dispensed whenever available.

A. SYSTEMIC LUPUS ERYTHEMATOSUS (SLE):

1. Documented diagnosis of systemic lupus erythematosus
AND

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2. Saphnelo (anifrolumab) will be used concurrently with standard therapy (i.e., corticosteroids, hydroxychloroquine, and/or immunosuppressive agents [azathioprine, cyclophosphamide, mycophenolate mofetil, methotrexate])
AND
3. Prescriber attests to, or clinical reviewer has found, no evidence of ANY of the following:
(a) Active central nervous system (CNS) lupus [such as seizures, psychosis, organic brain syndrome, cerebrovascular accident, cerebritis]
AND
(b) Active lupus nephritis
AND
4. Prescriber attests to (or the clinical reviewer has found that) the member not having any FDA labeled contraindications that haven't been addressed by the prescriber within the documentation submitted for review [Contraindications to Saphnelo (anifrolumab-fnia) include: patients with a history of anaphylaxis with anifrolumab-fnia, avoid initiating treatment during an active infection, avoid use of live or live-attenuated vaccines.]
AND
5. Documentation of prescriber baseline disease activity evaluation and goals for treatment to be used to evaluate efficacy of therapy at renewal (e.g., corticosteroid and immunosuppressant dose, frequency and severity of flares, flares which require systemic corticosteroids, etc.)
[DOCUMENTATION REQUIRED]
AND
6. IF THIS IS A NON-FORMULARY/NON-PREFERRED PRODUCT: Documentation of trial/failure of or serious side effects to a majority (not more than 3) of the preferred formulary alternatives for the given diagnosis. Documentation of medication(s) tried, dates of trial(s) and reason for treatment failure(s) is required.
MOLINA REVIEWER NOTE: For Illinois Marketplace, please see Appendix.

CONTINUATION OF THERAPY:

A. SYSTEMIC LUPUS ERYTHEMATOSUS (SLE):

1. Adherence to therapy at least 85% of the time as verified by the prescriber or member medication fill history OR adherence less than 85% of the time due to the need for surgery or treatment of an infection, causing temporary discontinuation
AND
2. Prescriber attests to or clinical reviewer has found no evidence of intolerable adverse effects or drug toxicity
AND
3. Documentation of positive clinical response as demonstrated by low disease activity and/or improvements in the condition's signs and symptoms (e.g., a dose reduction of corticosteroid or immunosuppressant therapy, zero flares or a decrease in amount of significant clinical flares which required a systemic corticosteroid boost during treatment, etc.) [DOCUMENTATION REQUIRED]
AND
4. Member continues standard therapy (prednisone or equivalent, hydroxychloroquine, azathioprine, mycophenolate mofetil, or methotrexate)

DURATION OF APPROVAL:

Initial authorization: 6 months, Continuation of Therapy: 12 months

MOLINA REVIEWER NOTE: For Texas Marketplace, please see Appendix.

PRESCRIBER REQUIREMENTS:

Prescribed by or in consultation with a board-certified rheumatologist or physician who specializes in the treatment of systemic autoimmune diseases [If prescribed in consultation, consultation notes must be submitted with initial request and reauthorization requests]

AGE RESTRICTIONS:

18 years of age and older

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QUANTITY:

300 mg every 4 weeks

PLACE OF ADMINISTRATION:

The recommendation is that infused medications in this policy will be for pharmacy or medical benefit coverage administered in a place of service that is a non-inpatient hospital facility-based location as per the Molina Health Care Site of Care program.

Note: Site of Care Utilization Management Policy applies for Saphnelo (anifrolumab). For information on site of care, see [Specialty Medication Administration Site of Care Coverage Criteria \(molinamarketplace.com\)](https://molinamarketplace.com)

DRUG INFORMATION

ROUTE OF ADMINISTRATION:

Intravenous

DRUG CLASS:

Type I Interferon (IFN) Receptor Antagonists

FDA-APPROVED USES:

Indicated for the treatment of adult patients with moderate to severe systemic lupus erythematosus (SLE), who are receiving standard therapy.

Limitations of Use: The efficacy of Saphnelo has not been evaluated in patients with severe active lupus nephritis or severe active central nervous system lupus. Use of Saphnelo is not recommended in these situations.

COMPENDIAL APPROVED OFF-LABELED USES:

None

APPENDIX

Reserved for State specific information. Information includes, but is not limited to, State contract language, Medicaid criteria and other mandated criteria.

State Specific Information

State Marketplace

Illinois (Source: Illinois General Assembly)

"(215 ILCS 134/45.1) Sec. 45.1. Medical exceptions procedures required. (c) An off-formulary exception request shall not be denied if: (1) the formulary prescription drug is contraindicated; (2) the patient has tried the formulary prescription drug while under the patient's current or previous health insurance or health benefit plan and the prescribing provider submits evidence of failure or intolerance; or (3) the patient is stable on a prescription drug selected by his or her health care provider for the medical condition under consideration while on a current or previous health insurance or health benefit plan. (d) Upon the granting of an exception request, the insurer, health plan, utilization review organization, or other entity shall authorize the coverage for the drug prescribed by the enrollee's treating health care provider, to the extent the prescribed drug is a covered drug under the policy or contract up to the quantity covered. (e) Any approval of a medical exception request made pursuant to this Section shall be honored for 12 months following the date of the approval or until renewal of the plan."

Texas (Source: [Texas Statutes, Insurance Code](#))

"Sec. 1369.654. PROHIBITION ON MULTIPLE PRIOR AUTHORIZATIONS.

(a) A health benefit plan issuer that provides prescription drug benefits *may not require an enrollee to receive more than one prior authorization annually* of the prescription drug benefit for a *prescription drug prescribed to treat an autoimmune disease, hemophilia, or Von Willebrand disease.*

(b) This section does not apply to:

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- (1) opioids, benzodiazepines, barbiturates, or carisoprodol;
- (2) prescription drugs that have a typical treatment period of less than 12 months;
- (3) drugs that:
 - (A) have a boxed warning assigned by the United States Food and Drug Administration for use; and
 - (B) must have specific provider assessment; or
- (4) the use of a drug approved for use by the United States Food and Drug Administration in a manner other than the approved use.”

BACKGROUND AND OTHER CONSIDERATIONS

BACKGROUND:

Systemic lupus erythematosus (SLE) is a chronic, occasionally life-threatening, multisystem immune-mediated disorder. Patients may present with a wide array of symptoms, signs, and laboratory findings and have a variable prognosis that depends upon the disease severity and type of organ involvement. Due to the variable disease course, effective management of SLE requires regular clinical and laboratory monitoring to assess disease activity; guide therapy to alleviate symptoms and reduce progressive organ damage; prevent and treat relapses; assess side effects related to drug therapy; encourage adherence with medications; and coordinate care with the patient's other providers.

The diagnosis of systemic lupus erythematosus (SLE) is based upon the judgment of an experienced clinician who recognizes characteristic constellations of symptoms and signs in the setting of supportive serologic studies after excluding alternative diagnoses. This is often challenging due to the great variability in the expression of SLE. Although the classification criteria were designed for research purposes, many clinicians refer to aspects of these criteria when making the diagnosis of SLE.

1997 ACR criteria – The classification criteria that were developed by the American Rheumatism Association (ARA, now the ACR) for the classification of SLE were established by cluster analyses, primarily in academic centers and in White American patients. The patient is classified with SLE using the ACR criteria if four or more of the manifestations are present, either serially or simultaneously, during any interval of observations. A positive lupus erythematosus cell test, used in older criteria, was replaced by the presence of antiphospholipid antibodies. When tested against other rheumatic diseases, these criteria have a sensitivity and specificity of approximately 96 percent.

2012 SLICC criteria – In 2012, the SLICC proposed classification criteria that were developed to address inherent weaknesses of the 1997 ACR classification criteria. Classification as having SLE by the SLICC criteria requires either that a patient satisfy at least 4 of 17 criteria, including at least 1 of the 11 clinical criteria and one of the six immunologic criteria, or that the patient has biopsy-proven nephritis compatible with SLE in the presence of ANA or anti-double-stranded DNA (anti-dsDNA) antibodies. Despite the improved sensitivity compared with the ACR criteria, the SLICC criteria might delay the diagnosis of SLE in a significant number of patients, and some patients might not be classified at all. These situations were demonstrated in a study in which patients with SLE from two large cohorts were grouped according to whether the SLICC criteria were met before, at the same time as, or after the ACR criteria, and the groups were then compared [57]. Among the patients diagnosed later with the SLICC criteria, in the majority of cases, the delay was due to the fact that the combination of malar rash and photosensitivity both fall within the acute cutaneous SLE category and thus only count as one criterion.

2019 EULAR/ACR criteria – The European League Against Rheumatism (EULAR)/American College of Rheumatology (ACR) classification criteria for SLE were developed to improve detection of early- or new-onset SLE as well as improve the sensitivity and specificity compared with previous criteria. The classification for SLE requires the presence of a positive antinuclear antibodies (ANA) as an entry criterion. Additive criteria consist of seven clinical (ie, constitutional, hematologic, neuropsychiatric, mucocutaneous, serosal, musculoskeletal) and three immunologic (ie, antiphospholipid antibodies, complement proteins, SLE-specific antibodies) categories, each of which are weighted from 2 to 10. Patients are classified as having SLE with a score of 10 or more points. In the validation cohort, which included patients with early disease, the EULAR/ACR criteria had a sensitivity of 96.1 percent and a specificity of 93.4 percent, compared with the 96.7 percent sensitivity and 83.7 percent specificity of the Systemic Lupus International Collaborating Clinics (SLICC) criteria and the 82.8 percent sensitivity and 93.4 percent specificity of the ACR criteria.

The goals of therapy for patients with systemic lupus erythematosus (SLE) are to ensure long-term survival, achieve the lowest possible disease activity, prevent organ damage, minimize drug toxicity, improve quality of

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life, and educate patients about their role in disease management. Treatment of SLE is individualized based upon patient preferences, clinical manifestations, disease activity and severity, and comorbidities. Patients require monitoring at regular intervals by a rheumatologist or other provider to optimize both nonpharmacologic and pharmacologic therapies and achieve treatment goals. Patients often have multiorgan system involvement and may require multidisciplinary care.

Saphnelo (anifrolumab-fnia)

Saphnelo is the first type I interferon (IFN) receptor agonist to be FDA-approved and the first drug for SLE since 2011, when GlaxoSmithKline's Benlysta (belimumab) was approved.

While Benlysta inhibits B-cell stimulating factor, Saphnelo binds to subunit 1 of the type I IFN receptor, blocking the activity of type I IFNs involved in regulating the inflammatory pathways implicated in SLE. Generally, IFNs signal for the immune system to be activated when there is an infection; however, in SLE, these signals are occurring unnecessarily, which leads to inflammation throughout the body. Saphnelo is thought to dampen these signals and decrease inflammation and systemic manifestations of SLE. Most adults (up to 80%) and children (90%) with SLE have increased type I IFN signaling, which is associated with increased disease activity and severity.

Saphnelo's efficacy and safety data were evaluated in 3 trials: MUSE (Trial 1; NCT01438489), TULIP-1 (Trial 2; NCT02446912), and TULIP-2 (Trial 3; NCT02446899). All 3 studies were randomized, double-blind, placebo-controlled trials in patients ≥ 18 years of age diagnosed with SLE according to the American College of Rheumatology (ACR) classification criteria and who were receiving standard therapy (at least one of the following: oral corticosteroids (OCSs), antimalarials, and immunosuppressants [methotrexate, azathioprine, or mycophenolate mofetil]).

MUSE (Trial 1; NCT01438489)-

Methods: Patients (n = 305) were randomized to receive intravenous anifrolumab (300 mg or 1,000 mg) or placebo, in addition to standard therapy, every 4 weeks for 48 weeks. Randomization was stratified by SLE Disease Activity Index 2000 score (<10 or ≥ 10), oral corticosteroid dosage (<10 or ≥ 10 mg/day), and type I IFN gene signature test status (high or low) based on a 4-gene expression assay. The primary end point was the percentage of patients achieving an SLE Responder Index (SRI[4]) response at week 24 with sustained reduction of oral corticosteroids (<10 mg/day and less than or equal to the dose at week 1 from week 12 through 24). Other end points (including SRI[4], British Isles Lupus Assessment Group [BILAG]-based Composite Lupus Assessment [BICLA], modified SRI[6], and major clinical response) were assessed at week 52. The primary end point was analyzed in the modified intent-to-treat (ITT) population and type I IFN-high subpopulation. The study result was considered positive if the primary end point was met in either of the 2 study populations. The Type I error rate was controlled at 0.10 (2-sided), within each of the 2 study populations for the primary end point analysis.

Results: The primary end point was met by more patients treated with anifrolumab (34.3% of 99 for 300 mg and 28.8% of 104 for 1,000 mg) than placebo (17.6% of 102) (P = 0.014 for 300 mg and P = 0.063 for 1,000 mg, versus placebo), with greater effect size in patients with a high IFN signature at baseline (13.2% in placebo-treated patients versus 36.0% [P = 0.004] and 28.2% [P = 0.029]) in patients treated with anifrolumab 300 mg and 1,000 mg, respectively. At week 52, patients treated with anifrolumab achieved greater responses in SRI(4) (40.2% versus 62.6% [P < 0.001] and 53.8% [P = 0.043] with placebo, anifrolumab 300 mg, and anifrolumab 1,000 mg, respectively), BICLA (25.7% versus 53.5% [P < 0.001] and 41.2% [P = 0.018], respectively), modified SRI(6) (28.4% versus 49.5% [P = 0.002] and 44.7% [P = 0.015], respectively), major clinical response (BILAG 2004 C or better in all organ domains from week 24 through week 52) (6.9% versus 19.2% [P = 0.012] and 17.3% [P = 0.025], respectively), and several other global and organ-specific end points. Herpes zoster was more frequent in the anifrolumab-treated patients (2.0% with placebo treatment versus 5.1% and 9.5% with anifrolumab 300 mg and 1,000 mg, respectively), as were cases reported as influenza (2.0% versus 6.1% and 7.6%, respectively), in the anifrolumab treatment groups. Incidence of serious adverse events was similar between groups (18.8% versus 16.2% and 17.1%, respectively).

TULIP-1 (Trial 2; NCT02446912), TULIP-2 (Trial 3; NCT02446899)

Methods: TULIP-1 and TULIP-2 were randomized, placebo-controlled, 52-week trials of intravenous anifrolumab (300 mg every 4 weeks for 48 weeks). For patients receiving baseline glucocorticoid ≥ 10 mg/day, attempted taper to ≤ 7.5 mg/day prednisone or equivalent from Weeks 8-40 was required and defined as sustained reduction when maintained through Week 52. Flares

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were defined as ≥ 1 new BILAG-2004 A or ≥ 2 new BILAG-2004 B scores versus the previous visit. Flare assessments were compared for patients receiving anifrolumab versus placebo.

Results: Compared with placebo (n = 366), anifrolumab (n = 360) was associated with lower annualized flare rates (rate ratio 0.75, 95% confidence interval [CI] 0.60-0.95), prolonged time to first flare (hazard ratio 0.70, 95% CI 0.55-0.89), and fewer patients with ≥ 1 flare (difference -9.3%, 95% CI -16.3 to -2.3), as well as flares in organ domains commonly active at baseline (musculoskeletal, mucocutaneous). Fewer BILAG-based Composite Lupus Assessment responders had ≥ 1 flare with anifrolumab (21.1%, 36/171) versus placebo (30.4%, 34/112). Of patients who achieved sustained glucocorticoid reductions from ≥ 10 mg/day at baseline, more remained flare free with anifrolumab (40.0%, 76/190) versus placebo (17.3%, 32/185). TULIP-1 (Trial 2; NCT02446912) did not meet its primary endpoint on the SRI-4 composite measure.

Prior to the unblinding of the TULIP-1 data, the primary endpoint of TULIP-2 was changed from SRI-4 to BICLA response. Combined data from all 3 trials demonstrated benefits in overall disease activity, skin and joints manifestations, and the ability to taper down steroid doses; however, these endpoints were not formally assessed for statistical significance

Safety

Combined data from all 3 clinical trials found adverse reactions were reported in 87% of patients receiving Saphnelo versus 79% of patients receiving placebo. Common adverse reactions included upper respiratory infection (43%), bronchitis (11%), infusion-related reactions (9.4%), herpes zoster (6.1%), and cough (5%).

Due to Saphnelo's mechanism of action in inhibiting the IFN pathway, infections such as herpes zoster are expected to some degree. Overall, 28 patients treated with Saphnelo developed herpes zoster, with 2 patients requiring hospitalization. Shingrix (zoster vaccine recombinant, adjuvanted) was approved in late July 2021 for the prevention of shingles in adults ≥ 18 years of age who are or will be at increased risk of shingles due to immunodeficiency or immunosuppression caused by a disease or therapy; however, the CDC's Advisory Committee on Immunization Practices (ACIP) has not yet endorsed this recommendation. Therefore, Shingrix could be considered in eligible patients prior to initiating Saphnelo to mitigate the risk of herpes zoster.

CONTRAINDICATIONS/EXCLUSIONS/DISCONTINUATION:

All other uses of Saphnelo (anifrolumab-fnia) are considered experimental/investigational and therefore, will follow Molina's Off- Label policy. Contraindications to Saphnelo (anifrolumab-fnia) include: patients with a history of anaphylaxis with anifrolumab-fnia, avoid initiating treatment during an active infection, avoid use of live or live-attenuated vaccines.

Exclusions/Discontinuation:

Do not use concurrently with another biologic [i.e., Actemra, Benlysta, Cimzia, Enbrel, Humira, Kineret, Orencia, Remicade, Rituxan, Simponi, Stelara].

OTHER SPECIAL CONSIDERATIONS:

The recommended dosage of Saphnelo is 300 mg, administered as an intravenous infusion over a 30- minute period, every 4 weeks. If a planned infusion is missed, administer Saphnelo as soon as possible. Maintain a minimum interval of 14 days between infusions.

CODING/BILLING INFORMATION

CODING DISCLAIMER. Codes listed in this policy are for reference purposes only and may not be all-inclusive or applicable for every state or line of business. Deleted codes and codes which are not effective at the time the service is rendered may not be eligible for reimbursement. Listing of a service or device code in this policy does not guarantee coverage. Coverage is determined by the benefit document. Molina adheres to Current Procedural Terminology (CPT®), a registered trademark of the American Medical Association (AMA). All CPT codes and descriptions are copyrighted by the AMA; this information is included for informational purposes only. Providers and facilities are expected to utilize industry-standard coding practices for all submissions. Molina has the right to reject/deny the claim and recover claim payment(s) if it is determined it is not billed appropriately or not a covered benefit. Molina reserves the right to revise this policy as needed.

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HCPCS CODE	DESCRIPTION
J0491	Injection, anifrolumab-fnia, 1 mg

AVAILABLE DOSAGE FORMS:

Saphnelo SOLN 300MG/2ML single-dose vial

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SUMMARY OF REVIEW/REVISIONS	DATE
REVISION- Notable revisions: Required Medical Information Continuation of Therapy Appendix Contraindications/Exclusions/Discontinuation References	Q4 2025

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REVISION- Notable revisions: Coding/Billing Information Template Update Required Medical Information	Q4 2024
REVISION- Notable revisions: Required Medical Information Continuation of Therapy Contraindications/Exclusions/Discontinuation Available Dosage Forms References	Q4 2023
REVISION- Notable revisions: Required Medical Information Continuation of Therapy Place of Administration Contraindications/Exclusions/Discontinuation Coding/Billing Information	Q4 2022
Q2 2022 Established tracking in new format	Historical changes on file