# DRUG DETERMINATION POLICY

Title: DDP-19 Benlysta

Effective Date: 08/31/2021



Physicians Health Plan PHP Insurance Company PHP Service Company

### Important Information - Please Read Before Using This Policy

The following policy applies to health benefit plans administered by PHP and may not be covered by all PHP plans. Please refer to the member's benefit document for specific coverage information. If there is a difference between this general information and the member's benefit document, the member's benefit document will be used to determine coverage. For example, a member's benefit document may contain a specific exclusion related to a topic addressed in a coverage policy.

Benefit determinations for individual requests require consideration of:

- 1. The terms of the applicable benefit document in effect on the date of service.
- 2. Any applicable laws and regulations.
- 3. Any relevant collateral source materials including coverage policies.
- 4. The specific facts of the particular situation.

Contact PHP Customer Service to discuss plan benefits more specifically.

### 1.0 Policy:

This policy describes the determination process for coverage of specific drugs.

This policy does not guarantee or approve benefits. Coverage depends on the specific benefit plan. Drug Determination Policies are not recommendations for treatment and should not be used as treatment guidelines.

#### 2.0 Background or Purpose:

Benlysta is a specialty drug indicated for Systemic Lupus Erythematosus (SLE) and Lupus Nephritis and is useful for specific organ system symptoms. These criteria were developed and implemented to ensure appropriate use for the specific symptoms detailed below.

#### 3.0 Clinical Determination Guidelines:

Document the following with chart notes

- I. Systemic Lupus Erythematosus and Lupus Nephritis [must meet all listed below]:
  - A. Age: at least five years.
  - B. Diagnosis and severity [must meet both listed below]:
    - 1. Active moderate to severe systemic lupus erythematosus refractory or intolerant to other immunosuppressive drugs.
    - Autoantibody positive: ANA at or above1:80 and/or anti-dsDNA at or above 30 units per mL.
  - C. Chronic other therapies: contraindication, inadequate response after four months or significant adverse effects to triple therapy listed under moderate or severe disease (see Appendix I). [must meet one listed below]:
    - 1. Moderate disease: prednisolone (at least 7.5mg per day); hydroxychloroquine; and azathioprine, methotrexate, mycophenolate mofetil OR cyclosporine.

2. Severe disease: prednisolone (at least 7.5mg per day); hydroxychloroquine; and mycophenolate OR cyclosporine.

## D. Dosage regimen:

- 1. Adults: Benlysta intravenous (belimumab IV): 10mg per Kg every two weeks times three, then every four weeks; or Benlysta subcutaneous (belimumab SQ): 200mg every week.
- 2. Pediatrics: Benlysta intravenous 10mg per Kg every two weeks, then 10mg per Kg every four weeks.
- II. Lupus Nephritis [must meet all listed below]:
  - A. Age: at least 18 years.
  - B. Diagnosis and severity [must meet both listed below]:
    - 1. Urinalysis: proteinuria, microscopic hematuria, kidney impairment and/or hypertension.
    - 2. Biopsy results: focal (class III), diffuse (class IV) or membranous nephropathy (class V LN).
  - C. Other therapies in combination with systemic glucocorticoids: contraindication, inadequate response after four months or significant adverse reactions to each listed below:
    - 1. Disease modifying anti-rheumatic drugs: mycophenolate or cyclophosphamide intravenous
    - 2. Calcineurin inhibitor triple therapy: tacrolimus added to one above.
  - D. Benlysta intravenous (belimumab IV):
    - 1. Adults: Benlysta intravenous (belimumab IV): 10mg per Kg every two weeks times three, then every four weeks; or Benlysta subcutaneous (belimumab SQ): 200mg every week.

#### III. Approval.

- A. Initial: six months.
- B. Re-approval: one year; decrease signs and symptoms and/or laboratory values of disease.
- C. Administration: medication falls under site of care policy (see DDP-08).

#### IV. Exclusions:

- A. Concurrent Disease: central nervous system lupus.
- B. Concurrent Medications: other biologics or intravenous cyclophosphamide.

#### 4.0 Coding:

AFFECTED CODES							
HCPCS Code	Brand Name	Generic Name	Billing Units (1U)	Prior Approval			
J0490	Benlysta	belimumab	10mg	Y			

## 5.0 References, Citations & Resources:

- 1. Lexi comp Online®, Lexi-Drugs®, Hudson, Ohio: Lexi-Comp, Inc.; Benlysta, accessed July 2021.
- 2. The British Society for Rheumatology guideline for the management of SLE in adults: Executive Summary. Rheumatology 2018;57.
- 3. DDP-08 Site of Care for Administration of Parenteral Specialty Medications.

## 6.0 Appendices:

See page 4.

## 7.0 Revision History:

Original Effective Date: 12/1/2011

Next Review Date: 07/28/2022

<b>Revision Date</b>	Reason for Revision
7/19	Annual review; replaced abbreviations
6/20	Annual review; removed mild disease other therapies replaced abbreviations, approved by P&T Committee 8/26/20.
6/21	Annual review, reformatted, added indication Lupus nephritis, revised SLE age

Appendix I: SLE Treatment Strategies for Mild, Moderate and Sever Non-renal Lupus<sup>2</sup>

ltem	Mild activity/flare BILAG C scores or single B score; SLEDAI <6	Moderate activity/flare BILAG 2 or more systems with B scores, SLEDAI 6-12	Severe activity/flare (non-renal) BILAG 1 or more A scores; SLEDAI >12	
Typical manifest- ations attributed to lupus	Fatigue, malar rash, diffuse alopecia, mouth ulcers, arth- ralgia, myalgia, platelets 50-149 × 10 <sup>9</sup> /l	Fever, lupus-related rash up to 2/9 body surface area, cutaneous vasculitis, alopecia with scalp inflammation, arthritis, pleurisy, pericarditis, hepatitis, platelets 25–49 × 10 <sup>9</sup> /l	Rash involving >2/9 body surface area, myositis, severe pleurisy and/or pericarditis with effusion, ascites, enteritis, myelopathy, psychosis, acute confusion, optic neuritis, platelets <25 × 10 <sup>9</sup> /l	
Initial typical drugs and target doses if no contra- indications	CSs <sup>a</sup> : topical preferred or oral prednisolone ≤20 mg daily for 1-2 weeks or l.m. or IA methyl-prednisolone 80-120 mg and HCQ ≤6.5 mg/kg/day and/or MTX 7.5-15 mg/week and/or NSAIDs (for days to few weeks only)	$\begin{array}{lll} Prednisolone^a &\leqslant 0.5 \text{ mg/day} \\ \text{or} & \text{i.v.} & \text{methyl-} & \text{prednisolone} \\ &\leqslant 250 \text{ mg} \times 13 \\ \text{or} & \text{i.m.} & \text{methyl-prednisolone} \\ &80120 \text{ mg} \\ \text{and AZA } 1.52.0 \text{ mg/kg/day} \\ \text{or MTX } (1025 \text{ mg/week}) \\ \text{or MMF } (23 \text{ g/day}) \text{ or} \\ &\text{ciclosporin} &\leqslant 2.0 \text{ mg/kg/day} \\ \text{and HCQ } &\leqslant 6.5 \text{ mg/kg/day} \end{array}$	Prednisolone <sup>a</sup> ≤ 0.5 mg/day and/or i.v. methyl-prednisolone 500 mg × 1-3 or prednisolone ≤ 0.75-1 mg/kg/day and AZA 2-3 mg/kg/day or MMF 2-3 g/day or CYC i.v. or ciclosporin ≤ 2.5 mg/kg/day and HCQ ≤ 6.5 mg/kg/day	
Aiming for typical maintenance drugs/doses providing no contra- indications	Prednisolone <sup>a</sup> ≤ 7.5 mg/day and HCQ 200 mg/day and/or MTX 10 mg/week	Prednisolone <sup>a</sup> ≤7.5 mg/day and AZA 50-100 mg/day or MTX 10 mg/week or MMF 1 g/day or ciclosporin 50-100 mg/day and HCQ 200 mg/day;	Prednisolone <sup>a</sup> ≤7.5 mg/day and MMF 1.0-1.5 g/day or AZA 50-100 mg/day or ciclosporin 50-100 mg/day and HCQ 200 mg/day;	
	Aim to reduce and stop drugs except HCQ eventually when in stable remission	Aim to reduce and stop drugs except HCQ eventually when in stable remission	Aim to reduce and stop drugs except HCQ eventually when in stable remission	

<sup>&</sup>lt;sup>a</sup>The lowest effective dose of prednisolone or other CSs should be used at all times.

# Appendix II: Monitoring & Patient Safety

Drug	Adverse Reactions	Monitoring	REMS
Benlysta belimumab	<ul> <li>Gastrointestinal: nausea (15%), diarrhea (12%)</li> <li>Miscellaneous: infusion related reaction (17%), hypersensitivity (13%)</li> <li>Pregnancy: IgG molecules cross placenta with increased amount through pregnancy (use contraception during and 4 mos. post use)</li> </ul>	<ul> <li>Central Nervous System:         worsening depression, mood         changes, suicidal thought</li> <li>Hypersensitivity, infusion reactions</li> <li>Infections</li> </ul>	None needed