## **HEALTHY U CHIP**

## PRIOR AUTHORIZATION REQUEST FORM **BENLYSTA**®

For authorization, please answer each question and fax this form PLUS chart notes back to the Healthy U CHIP Prior Authorization Department.

- For **Medical Pharmacy** please fax requests to: 801-213-1547
- For **Retail Pharmacy** please fax requests to: 385-425-4052

Failure to submit clinical documentation to support this request will result in a dismissal of the request.

| If you have prior authorization questions, please call for Pharmacy Customer Service for assistance at 385-425-5094   |   |        |                 |                              |  |  |  |  |
|---|---|--------|-----------------|------------------------------|--|--|--|--|
| Disclaimer: Prior authorization request forms are subject to change in accordance with Federal and State notice requirements.   |   |        |                 |                              |  |  |  |  |
|   |   |        |                 |                              |  |  |  |  |
| ate: Member Name:   |   |        | ID#:            |                              |  |  |  |  |
| DB: Gender:   |   |        | Physician:      |                              |  |  |  |  |
| Office Phone: Office Fax:   |   |        | Office Contact: |                              |  |  |  |  |
| Height/Weight:  |   |        | HCPCS Code:     |                              |  |  |  |  |
| Member must try formulary preferred drugs before a request for a non-preferred drug may be considered. If treatment with preferred products has not been successful, you must submit which preferred products have been tried, dates of treatment, and reason for failure. Reasons for failure must meet the Health Plan medical necessity criteria.  Product being requested:   Benlysta® (belimumab)  Dosing/Frequency: |   |        |                 |                              |  |  |  |  |
| If the request is for reauthorization, proceed to reauthorization section.  |   |        |                 |                              |  |  |  |  |
| Questions   |   | Yes    | No              | Comments/Notes               |  |  |  |  |
|   | SYSTEMIC LUPUS ERYTHEN                                | IATOSL | JS              |                              |  |  |  |  |
| <ol> <li>Does the member have a confirmed<br/>moderate to severe systemic lupus</li> </ol>  | _   |        |                 | Please provide documentation |  |  |  |  |
| 2. Is the request made by, or in consultrheumatologist?   | tation with, a  |        |                 |                              |  |  |  |  |
| <ol> <li>Does the member have a Safety of E<br/>Assessment-Systemic Lupus Erythen<br/>Index (SELENA-SLEDAI) score of ≥ 6?</li> </ol>  | natosus Disease Activity                              |        |                 | Please provide documentation |  |  |  |  |
| <ol> <li>Does the member have active musc<br/>disease that is unresponsive to stan<br/>glucocorticoids and/or other immur</li> </ol>  | dard therapy with                                     |        |                 | Please provide documentation |  |  |  |  |
| <ol> <li>Is there documentation of corticoste<br/>(prednisone equivalent dose ≥10mg<br/>both hydroxychloroquine AND at lea<br/>(e.g., azathioprine, methotrexate, m</li> </ol>  | /day) OR trial and failure of ast 1 immunosuppressant |        |                 | Please provide documentation |  |  |  |  |
| 6. Has the member been at least 80% months with their baseline therapy immunosuppressants)?   | -   |        |                 | Please provide documentation |  |  |  |  |
| 7. Will Benlysta® be used concurrently  | with baseline therapy?                                |        |                 | Please provide documentation |  |  |  |  |

| 8.   | Does the member have documentation of active central   |         |    | Please provide documentation   |
|--|--|---------|----|--|
|  | nervous system lupus (e.g. generalized seizures, psychosis,  |         |    |  |
|  | stroke, peripheral neuropathies)?  |         |    |  |
| 9.   | Has the member received any other biologics,   |         |    | Please provide documentation   |
|  | immunoglobulins, IV cyclophosphamide, or prednisone >100mg   |         |    |  |
|  | daily within the last 6 months?  |         |    |  |
|  | LUPUS NEPHRITIS  | 1       |    |  |
|  | Does the member have a confirmed diagnosis of lupus nephritis?   |         |    | Please provide documentation   |
| 2.   | Is the request made by, or in consultation with, a nephrologist or rheumatologist?   |         |    |  |
| 3.   | Did the member have a kidney biopsy showing a histological diagnosis of lupus nephritis Class III, IV or V?  |         |    | Please provide documentation   |
| 4.   | Does documentation show a recent eGFR ≥ 30 mL/min/1.73m <sup>2</sup> ?   |         |    | Please provide documentation   |
| 5.   | Has the member had dialysis in the past 12 months?   |         |    |  |
| 6.   | Is the member currently receiving standard immunosuppressive therapy for systemic lupus erythematosus?   |         |    | Please provide documentation   |
| 7.   | Will Benlysta® be used concurrently with baseline therapy?   |         |    | Please provide documentation   |
| 8.   | Does the member have active central nervous system lupus   |         |    | Please provide documentation   |
|  | (e.g. generalized seizures, psychosis, stroke, peripheral neuropathies)?   |         |    | ·  |
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|  | immunoglobulins, IV cyclophosphamide, or prednisone >100mg   |         |    |  |
|  | daily within the last 6 months?  |         |    |  |
|  |  |         |    |  |
|  | REAUTHORIZATION  | N       |    | <u> </u>   |
|  | REAUTHORIZATION SYSTEMIC LUPUS ERYTHEN   |         | JS |  |
| 1.   | SYSTEMIC LUPUS ERYTHEN   |         |    |  |
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| Additional information: |  |  |
|-------------------------|--|--|
|                         |  |  |
|                         |  |  |
|                         |  |  |
| Physician Signature:    |  |  |
|                         |  |  |

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Policy: PHARM-CHIP-081 Origination Date: 07/01/2024 Reviewed/Revised Date: 09/18/2024 Next Review Date: 09/18/2025 Current Effective Date: 10/01/2024

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