## **HEALTHY U CHIP**

## PRIOR AUTHORIZATION REQUEST FORM **SYNAGIS**®

For authorization, please answer each question and fax this form PLUS chart notes back to the Healthy U CHIP Prior Authorization Department at 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 assistance: 385-425-5094						
Disclaimer: Prior Authorization request forms are subject to change in accordance with Federal and State notice requirements.						
Date: Member Name:			ID#:			
DOB:	Gender:		Physician:			
Office Phone:	Office Fax:		Office Contact:			
Gestational Age at Birth (give weeks & days):		Member Weight:				
Product being requested: ☐ Synagis® (palivizumab)  Dosing/Frequency:  Please note:  Requests may be approved for up to a maximum of 4 to 5 doses at a dosing interval of not less than 28 days between injections.  Requests will only be authorized for treatment during the State Health Department's declared official Synagis® season.  Approved requests will be authorized to start on the first date of the official Synagis® season.  If the member has tested positive for RSV, further requests for Synagis® will not be approved.						
Questio	ns	Yes	No	Comments/Notes		
<ol> <li>Was the member's age ≤ 12 month season? If no, skip to question #7.</li> </ol>						
2. If the member is < 6 months of age available for administration?						
<ul> <li>Was the member born before 29 w</li> <li>Note: Synagis prophylaxis is no well infants &gt; 29 weeks gestati</li> </ul>	t recommended for otherwise					
<ol> <li>Was the member diagnosed with of prematurity, defined as gestational supplemental oxygen for at least the</li> </ol>	l age < 32 weeks AND required					
<ul> <li>5. Is the member diagnosed with hen congenital heart disease with one congenital heart disease, receiving congestive heart failure?; OR</li> <li>Member will require cardiac sure.</li> <li>Member has moderate to seven Note: Synagis® prophylaxis is not rechemodynamically insignificant heart septal defect, small ventricular septal uncomplicated aortic stenosis, mild aductus arteriosus.</li> <li>6. Does the member have anatomic properties.</li> </ul>	of the following: g medication to control  rgical procedures?; OR re pulmonary hypertension? commended for infants with disease, such as secundum atrial al defect, pulmonic stenosis, coarctation of the aorta, or patent					
neuromuscular disorders that impa secretions from the upper airway?	irs the ability to clear					

7.	Will the member be profoundly immunocompromised during the					
	respiratory syncytial virus (RSV) season?					
8.	Was the member's age ≥ 12 months and <24 months at the start					
	of the RSV season?					
9.	Is the member <20 months and expected to receive a heart					
	transplant during the current RSV season?					
10.	Is the member <20 months and expected to be profoundly					
	immunocompromised during the current RSV season?					
11.	Was the member born at less than 32 weeks 0 days gestation and					
	required at least 28 days of oxygen after birth and continues to					
	require medical intervention with supplemental oxygen, chronic					
	corticosteroids, or diuretic therapy in the 6 months prior to the					
	start of the current RSV season?					
	Note: Synagis prophylaxis is not recommended for otherwise well					
	infants with chronic lung disease of prematurity who are 12 to 24					
	months old.					
12.	Has the member had a respiratory syncytial virus-related					
	hospitalization during this RSV season?					
13.	Has the member received Beyfortus during this RSV season?					
1/1	Was Synagis® given while the member was in the hospital (e.g.,	П	П			
14.	NBICU, NICU)?					
	· · · · · · ·					
	If yes, please list dates given:					
Phy	sician's Signature:	ı	I			
** Failure to submit clinical documentation to support this request will result in						

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

- 1. Synagis® (palivizumab) therapy is authorized according to current guidelines for treatment of RSV as published by the American Academy of Pediatrics Committee on Infectious Diseases; American Academy of Pediatrics Bronchiolitis Guidelines Committee. The current guidelines may be found online at <a href="http://pediatrics.aappublications.org/content/pediatrics/134/2/415.full.pdf">http://pediatrics.aappublications.org/content/pediatrics/134/2/415.full.pdf</a>.
- 2. Synagis® injections may be authorized during the RSV season, as defined by the local State Department of Health.
- 3. Up to 4-5 monthly doses may be authorized. Infants born during the RSV season, and who are approved for Synagis® therapy, may receive monthly doses until end date determined by the local State Health Department.
- 4. Synagis® therapy will not be approved with active RSV infection.
- 5. Synagis® prophylaxis will be discontinued if the member is hospitalized for RSV infection while being treated with monthly prophylaxis.
- 6. Synagis® therapy will be provided by the preferred pharmacy vendor.
- 7. Synagis® season information is available on the CDC website: <a href="https://www.cdc.gov/surveillance/nrevss/rsv/state.html">https://www.cdc.gov/surveillance/nrevss/rsv/state.html</a>

Policy: PHARM-CHIP-073 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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