## SENTARA COMMUNITY PLAN (MEDICAID)

## MEDICAL PRIOR AUTHORIZATION/STEP-EDIT REQUEST\*

<u>Directions:</u> The prescribing physician must sign and clearly print name (preprinted stamps not valid) on this request. All other information may be filled in by office staff; <u>fax to 1-844-305-2331</u>. No additional phone calls will be necessary if all information (including phone and fax #s) on this form is correct. <u>If information provided is not complete, correct, or legible, authorization can be delayed</u>.

<u>Drug Requested</u>: Zynteglo<sup>®</sup> (betibeglogene autotemcel) (J3590/C9399) (Medical)

MEMBER & PRESCRIBER IN	FORMATION: Authorization may be delayed if incomplete.
Member Name:	
Member Sentara #:	Date of Birth:
Prescriber Name:	
Prescriber Signature:	Date:
Office Contact Name:	
Phone Number:	
DEA OR NPI #:	
DRUG INFORMATION: Authori	
Drug Form/Strength:	
	Length of Therapy:
Diagnosis:	ICD Code, if applicable:
Weight:	Date:

## **Dosing Limits**

- A. Quantity Limit (max daily dose) [NDC Unit]:
  - Zynteglo up to 4 infusion bags, 20 mL/infusion bag, overwrap, and metal cassette: 73554-3111-xx
  - A single dose of Zynteglo containing a minimum of  $5.0 \times 10^6$  CD34+ cells/kg of body weight, in one or more infusion bags
- B. Max Units (per dose and over time) [HCPCS Unit]:
  - A single dose of Zynteglo containing a minimum of  $5.0 \times 10^6$  CD34+ cells/kg of body weight, in one or more infusion bags

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**CLINICAL CRITERIA:** Check below all that apply. All criteria must be met for approval. To support each line checked, all documentation, including lab results, diagnostics, and/or chart notes, must be provided or request may be denied.

<u>Authorization Criteria</u>: Coverage will be provided for one treatment course (1 dose of Zynteglo) and may not be renewed.

	mber has a documented diagnosis of beta thalassemia (excludes alpha-thalassemia and hemoglobin thalassemia variants) as outlined by at least <b>ONE</b> of the following:
	Member diagnosis is confirmed by HBB sequence gene analysis showing biallelic pathogenic variants
	Member has severe microcytic hypochromic anemia, anisopoikilocytosis with nucleated red blood cells on peripheral blood smear, and hemoglobin analysis that reveals decreased amounts or complete absence of hemoglobin A and increased amounts of hemoglobin F
	Member has transfusion-dependent disease defined as a history of transfusions of at least 100 mL/kg/year of packed red blood cells (pRBCs) or with 8 or more transfusions of pRBCs per year in the 2 years preceding therapy (Note: Detailed medical records of transfusion program/schedule recording dates of administration and volume administered are required. Last two (2) years of records MUST be provided)
	$\underline{\mathbf{L}}$ of the following have been assessed, and confirmation is noted that the member does $\underline{\mathbf{NOT}}$ have any he following:
	Severely elevated iron in the heart (i.e., patients with cardiac T2* less than 10 msec by magnetic resonance imaging [MRI])
	Advanced liver disease
	MRI of the liver with results demonstrating liver iron content $\geq$ 15 mg/g (unless biopsy confirms absence of advanced disease)
Member is of sufficient weight to at least accept the minimum number of cells required to initiate the manufacturing process	
viru	mber has been screened for hepatitis B virus (HBV), hepatitis C virus (HCV), human T-lymphotrophic is 1 & 2 (HTLV-1/HTLV-2), and human immunodeficiency virus (HIV) in accordance with clinical delines prior to collection of cells (leukapheresis)
to n	mber has <u>NOT</u> used prophylactic HIV anti-retroviral medication or hydroxyurea within 30 days prior nobilization (or for the expected duration for elimination of those medications) and until all cycles of eresis are completed (Note: if a patient requires anti-retrovirals for HIV prophylaxis, confirm a gative test for HIV before beginning mobilization)
	n chelation therapy has been discontinued for at least 7 days prior to initiating myeloablative ditioning therapy
	nales of reproductive potential have a negative pregnancy test prior to start of mobilization and re- firmed prior to conditioning procedures and again before administration of betibeglogene autotemcel
	quested medication will be used as single agent therapy (not applicable to lymphodepleting or bridging rapy while awaiting manufacture)

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**Zynteglo (Medical)(Medicaid)** (Continued from previous page)

	Member will receive periodic life-long monitoring for hematological malignancies
	Member is eligible to undergo hematopoietic stem cell transplant (HSCT) and has $\underline{NOT}$ had prior HSCT or other gene-therapy
	Member does NOT have availability of a willing 10/10 HLA-matched sibling donor
Med	lication being provided by a Specialty Pharmacy - PropriumRx
	Location/site of drug administration:
	NPI or DEA # of administering location:
	OR
	Specialty Pharmacy - PropriumRx
review	gent reviews: Practitioner should call Sentara Pre-Authorization Department if they believe a standard would subject the member to adverse health consequences. Sentara's definition of urgent is a lack of ent that could seriously jeopardize the life or health of the member or the member's ability to regain num function.
**	Use of samples to initiate therapy does not meet step edit/preauthorization criteria.**
*Pre	vious therapies will be verified through pharmacy paid claims or submitted chart notes.*