

State of Oklahoma SoonerCare

Adcetris® (Brentuximab Vedotin) Prior Authorization Form

Member Name:	Date of Birth:	Member ID#:
	Drug Information	
Physician billing (HCPCS code:	Start Date (or da	te of next dose):
Dose:	Regimen:	
	Billing Provider Informatio	n
Provider NPI:		ne:
Provider Phone:		
Prescriber NPI:		
		Specialty:
	Criteria	
Page 1 of 2—Please complete and re		all pages will result in processing delays.
For Initial Authorization:1. Please indicate the requested inf		
	used as a single-agent? Yes	
	used as a primary treatment? Yes used in relapsed/refractory diseas	
		psphamide, doxorubicin, and prednisone
(CHP)? Yes No	docum combination with cyclophic	ophamiae, develablem, and preameene
2. Please indicate the diagnosis and	d information:	
	phoma (ALCL), Primary Cutane	
	Itifocal lesions or regional nodes?	Yes No
☐ Anaplastic Large Cell Lym		
A. Is the diagnosis previou	one or more lines of therapy? Yes_	No
☐ Adult Classical Hodgkin Ly		110
	ntreated Stage III or IV? Yes	No
 B. Will brentuximab vedotii 		orubicin, vinblastine, and dacarbazine?
YesNo	gous stem cell transplant (SCT) ca	andidate with failure of 2 or more
multi-agent chemothera	py regimens? Yes No	andidate with failable of 2 of more
 D. Has member failed auto 	ologous SCT? Yes No	
		ation with nivolumab, bendamustine, or
multi-agent chemothera	py?	
YesNo	scalidation after autologous SCT w	ith a high risk of relapse or progression?
Yes No	solidation after autologous SCT w	itir a riigir risk or relapse or progression:
☐ Pediatric Classical Hodgki A. Is cHL previously untrea	in Lymphona (cHL), (age 2-21 ye ated? Yes No	ears)
		V per Ann Arbor Staging System?
C. Will brentuximab vedoti	n be used in combination with dox amide (AVE-PC)? Yes No	orubicin, vincristine, etoposide, predni-
	Page 1 of 2	

Please complete and return all pages. Failure to complete all pages will result in processing delays.

PLEASE PROVIDE THE INFORMATION REQUESTED AND RETURN TO:

University of Oklahoma College of Pharmacy Pharmacy Management Consultants Product Based Prior Authorization Unit

> Fax: 1-800-224-4014 Phone: 1-800-522-0114 Option 4

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3/1/2023 Pharm-123



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 Member Name:
 ______ Date of Birth:
 ______ Member ID#:

Criteria Criteria	
Page 2 of 2—Please complete and return <u>all pages</u> . Failure to complete all pages will result in processing delays.	
 Please indicate the diagnosis and information, continued: 	
☐ Primary Cutaneous Lymphomas – Mycosis Fungoides (MF)/Sézary Syndrome (SS)	
= 1 milary cutamodus = ymphomus myssols 1 angeluss (m.)/20=ary cymuloms (20)	
□ Diffuse Large B-Cell Lymphoma (DLBCL) or High Grade Lymphoma	
A. Is disease CD30+? Yes No	
B. Is member a non-autologous stem cell transplant (SCT) candidate? Yes No	
C. Has member transformed to DLBCL from follicular lymphoma or marginal zone lymphoma and	
received 2 or more lines of therapy for indolent or transformed disease? Yes No	
□ Peripheral T-Cell Lymphoma (PTCL)	
A. Previously untreated CD30+ disease? Yes No	
B. Has member received one or more lines of therapy? Yes No	
□ Adult T-Cell Leukemia/Lymphoma	
A. Is disease CD30+? Yes No	
B. Is member a nonresponder to first-line therapy with chronic/smoldering subtype? Yes No	
C. Will brentuximab vedotin be used for first-line therapy for acute or lymphoma subtype?	
YesNo	
D. Will brentuximab vedotin be used for continued treatment in responders to first-line therapy for acute	
or lymphoma subtype? Yes No E. Has member received one or more lines of therapy? Yes No	
□ T-Cell Lymphoma, Extranodal NK/T-Cell Lymphoma, Nasal Type	
A. Is disease CD30+? Yes No	
B. Is disease relapsed/refractory following additional therapy with an alternate combination	
chemotherapy regimen not previously used? Yes No	
☐ If answer is none of the above, please indicate diagnosis:	
Additional Information:	
For Continued Authorization:	
Date of last dose:	
Does member have any evidence of progressive disease while on brentuximab vedotin? Yes No	
3. Has the member experienced any adverse drug reactions related to brentuximab vedotin therapy?	
Yes No	
If yes, please specify adverse reactions:	
Additional Information:	
Page 2 of 2	
Please complete and return <u>all</u> pages. Failure to complete all pages will result in processing delays.	
Please do not send in chart notes. Specific information will be requested if necessary.	
Prescriber Signature: Date:	
I certify that the indicated treatment is medically necessary and all information is true and correct to the best of m	
knowledge.	
-	

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