ADAKVEO <sup>®</sup> (crizanlizumab) Medication Precertification Request Page 1 of 2 (All fields must be completed and legible for precertification review.)						Aetna Precertification NotificationPhone: <u>1-866-752-7021</u> (TTY: <u>711</u> )FAX: <u>1-888-267-3277</u> For Medicare Advantage Part B:Please Use Medicare Request Form		
Please indicate:	Start of tre				,			
		-	of last treatment/	' /				
					:	Fax:		
A. PATIENT INFOR								
First Name:			Last Name:			DOB:		
Address:				City:		State:	ZIP:	
Home Phone:		Work Phone:		Cell Phone:		Email:	211.	
	-		ent Height: inches	or <u>cms</u>	Allergies:			
B. INSURANCE IN								
Aetna Member ID #: Group #: Insured:			Does patient have other coverage?					
Medicare: 🗌 Yes	□ No If ves	provide ID #:	Med	licaid: □Yes [	□ No If yes, pro	ovide ID #:		
C. PRESCRIBER I								
First Name:			Last Name:		(Check	One): 🗌 M.D.	🗌 D.O. 🗌 N.P. 🗌 P.A.	
Address:				City:		State:	ZIP:	
Phone:	Fax	(:	St Lic #:	NPI #:	DEA #:		UPIN:	
Provider Email:			Office Contact Name:			Phone:		
	ne) <sup>,</sup> Hem	atologist 🔲 Other						
	-							
Place of Administr			SINMATION	Dispensing P	Provider/Pharma	cv: Patient Se	lected choice	
Self-administere			Physician's Office					
_		Specialty Pharmacy						
Center Nar	me:				,			
Home Infusion C	Center	Phone:						
	me:							
Address:				TIN:		PIN:		
E. PRODUCT INFORMATION								
•	•	anlizumab) Dose: _		Frequency				
			ary ICD code and specify	-				
-			Secondary ICD Code			r ICD Code:		
		-	mation must be complete	d in its <u>entirety</u> fo	or all precertificat	on requests.		
For All Requests (cl			an ital a attice a					
☐ Yes ☐ No Is this infusion request in an outpatient hospital setting? ☐ Yes ☐ No Has the patient experienced an adverse event with the requested product that has not responded to conventional								
interventions (e.g., acetaminophen, steroids, diphenhydramine, fluids, other pre-medications or slowing of infusion rate) or a severe adverse event (anaphylaxis, anaphylactoid reactions, myocardial infarction, thromboembolism, or seizures) during or immediately after an infusion?								
	Yes No Does the patient have severe venous access issues that require the use of special interventions only available in the outpatient hospital setting?							
Yes No Does the patient have significant behavioral issues and/or physical or cognitive impairment that would impact the safety of the infusion therapy AND the patient does not have access to a caregiver?								
Please provide a description of the behavioral issue or impairment:   Yes No Is the patient medically unstable which may include respiratory, cardiovascular, or renal conditions that may limit the								
patient's ability to tolerate a large volume or load or predispose the patient to a severe adverse event that cannot be managed in an alternate setting without appropriate medical personnel and equipment?								
Please provide a description of the condition:  Cardiopulmonary:								
				· · -			· · · · · · · · · · · · · · · · · · ·	
☐ Other:								



## ADAKVEO<sup>®</sup> (crizanlizumab) Medication Precertification Request Page 2 of 2

Aetna Precertification NotificationPhone:1-866-752-7021 (TTY: 711)FAX:1-888-267-3277For Medicare Advantage Part B:Please Use Medicare Request Form

(All fields must be completed and legible for precertification review.)

Patient First Name	Patient Last Name	Patient Phone	Patient DOB					
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.								
☐ Yes ☐ No Does the patient have a documented diagnosis of sickle cell disease?								
☐ Yes ☐ No Will the requested drug be used to reduce the frequency of vaso-occlusive crises (VOCs)?								
Yes No Is the requested drug being prescribed by or in consultation with a hematologist or specialist in sickle cell disease?								
For Initial Requests (clinical documentation required):								
Please indicate the sickle cell genotype:								
🗌 Homozygous hemoglobin S (HbSS) or 🔲 Sickle beta0-thalassemia (HbSbeta0)								
Yes No Has the patient experienced, at any time in the past, an inadequate response or intolerance to a trial of hydroxyurea?								
$\longrightarrow$ Yes $\square$ No Does the patient have a contraindication to hydroxyurea?								
$\Box \rightarrow \Box$ Yes $\Box$ No Will the patient be using the requested drug with concurrent hydroxyurea therapy?								
$\rightarrow$ Please specify: $\Box$ Patient had an inadequate response $\Box$ Patient had an intolerance								
Sickle hemoglobin C (HbSC)								
☐ Sickle beta+-thalassemia (HbSbeta+)								
Other:								
Yes No Has the patient experienced a vaso-occlusive crisis (VOC) in the past 12 months?								
For Continuation Requests (clinical documentation required):								
☐ Yes ☐ No Has the patient experie	enced a reduction in the frequency of vas	o-occlusive crises (VOCs), or has the pat	ient maintained a reduction in the					
	nce initiating therapy with the requested d							
H. ACKNOWLEDGEMENT								
Request Completed By (Signature	Required):		Date: /					
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent								

insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.