

Cinryze® (C1 esterase inhibitor, human) Medication Precertification Request

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

Aetna Precertification Notification

Phone: 1-866-752-7021 **FAX:** 1-888-267-3277

For Medicare Advantage Part B: Please Use Medicare Request Form

Please indicate:	☐ Start of treatment: S☐ Continuation of ther	_		/	/					·
Precertification R	equested By:							Fax	C :	
A. PATIENT INFOR									-	
First Name:				Last	Name:					
Address:				City:				State:		ZIP:
Home Phone:		Work	Phone:			Cell	Phone:	I		
DOB:	Allergies:					Ema	ail:			
	lbs or	kas	Height:		inches or		cms			
B. INSURANCE INF										
	# :		Does patient have	other	coverage?	Yes	□No			
Group #:			If yes, provide ID#: Carrier Name:							
			Insured:							
Medicare: Tes	☐ No If yes, provide ID) #:		Medi	icaid: 🗌 Yes 🏻	□No	If yes, pro	vide ID #:		
C. PRESCRIBER IN	IFORMATION									
First Name:			Last Name:			(Check One	e): 🔲 M.[D. 🔲 🛭	D.O. 🗌 N.P. 🔲 P.A.
Address:					City:			State:		ZIP:
Phone:	Fax:		St Lic #:		NPI #:	I	DEA #:		UP	IN:
Provider Email:			Office Contact Nan	ne:				Pho	ne:	
Specialty (Check o	nne): Allergist 🗌	Immunolo	gist					•		
Center Nar Home Infusion C Agency Na	on Center Phone: me: enter Phone: me: de(s) (CPT):				Physician's Specialty Ph Name: Address: Phone: TIN:	narmacy	/ □ Oth			
	nryze (C1 esterase inhibi	tor, huma	n) Dose:			Frequ	ency:			
F. DIAGNOSIS INFO	ORMATION - Please indica	ate primary l	CD Code and specify	any	other where applic	able.				
Primary ICD Code:		Second	lary ICD Code:			Ot	her ICD Co	ode:		
G. CLINICAL INFO	RMATION – Required clinic	al information	on must be completed	l in its	entirety for all pre	certifica	ation reques	sts.		
Yes No Is tr	severe adver immediately: Yes	tpatient hos int experien (e.g., aceta se event (ar after an infu tent have se spital settinient have signy AND the de a descrip medically utility to toleran alternate de a descrip	pital setting? ced an adverse event minophen, steroids, c naphylaxis, anaphylac sion? vere venous access i g? gnificant behavioral is a patient does not hav tion of the behavioral nstable which may in te a large volume or setting without appro tion of the condition:	dipher ctoid r ssues sues re acco issue clude load co priate	ahydramine, fluids, eactions, myocard to that require the unand/or physical or ess to a caregiver or impairment:	other p lial infar se of sp cognitiv ? ovascula nember el and ed	re-medicati ction, thron pecial interv re impairme ar, or renal to a severe quipment?	entions or slovenbolistic entions on entions on entithat work conditions adverse e	wing of sm, or selly availuld imp	infusion rate) or a seizures) during or lable in the pact the safety of the ay limit the nat cannot be
☐ Yes ☐ No Will atta	ne requested medication be the requested medication b cks? number of hereditary angion	pe used in c	ombination with any c	ther r	medication used fo			f hereditar	y angic	pedema (HAE)



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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 Is the requested medication prescribed by or in consultation with a prescriber who specializes in management of hereditary angioedema (HAE)?										
Which of the following applies to the patient?										
Hereditary angioedema (HAE) with C1 inhibitor deficiency or dysfunction confirmed by laboratory testing Please indicate which of the following conditions the patient has/had at the time of diagnosis:										
☐ A C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by the laboratory performing the test										
☐ A normal C1-INH antigenic level and a low C1-INH functional level (functional C1-INH less than 50% or C1-INH functional level below the lower limit of normal as defined by the laboratory performing the test)										
Other										
Hereditary angioedema (HAE) with normal C1 inhibitor confirmed by laboratory testing										
Please indicate which of the following conditions the patient has/had at the time of diagnosis:										
☐ F12, angiopoietin-1, plasminogen, kininogen-1 (KNG1), heparan sulfate-glucosamine 3-0 sulfotransferase 6 (HS3ST6) or myoferlin (MYOF) gene mutation as confirmed by genetic testing										
□ Both of the following: 1). Angioedema refractory to a trial of high-dose antihistamine therapy (i.e., cetirizine at 40 mg per day or the equivalent) for at least one month AND 2). Family history of angioedema										
☐ Other										
For Continuation of Therapy Requests (clinic	cal documentation required for all reque	<u>sts)</u> :								
☐ Yes ☐ No Has the patient experienced a significant reduction in frequency of acute attacks (e.g., >= 50%) since starting treatment?										
Yes No Has the patient reduced the use of medications to treat acute attacks since starting treatment with the requested medication?										
H. ACKNOWLEDGEMENT										
Request Completed By (Signature Requir	ed):		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.