

BlueShield. ICATIBANT Federal Employee Program. PRIOR APPROVAL REQUEST

Send completed form to:
Service Benefit Plan
Prior Approval
P.O. Box 52080 MC 139
Phoenix, AZ 85072-2080
Attn. Clinical Services

Attn. Clinical Services Fax: 1-877-378-4727

Additional information is required to process your claim for prescription drugs. Please complete the patient portion, and have the prescribing physician complete the physician portion and submit this completed form.

	atient Infor	mation	(required)			Provider In	nformation (re	equired)
Date:					Provider Name	:		
Patient Name:					Specialty:		NPI:	
Date of Birth:		Sex:	■Male	Female	Office Phone:		Office Fax:	
Street Address:		•			Office Street A	ddress:		
City:		State	:	Zip:	City:		State:	Zip:
Patient ID: R	1 1	ı	1 1		Physician Signa	ature:		
			P	HYSICIAN C	OMPLETES			
Standard				-		-	referred products. at no cost in the b	
NOTE: Form must be completed in its entirety for processing								
Please select m	edication:		□Fira	azyr (icatibant)		□Sajaz	ir (icatibant)	
**Check www.fepbl	ue.org/formulary	to confirm	which medic	eation is part of the	patient's benefit			
1. Has the patient been on this medication continuously for the last 6 months, excluding samples? Please select answer below:								
☐ YES – this is a PA renewal for CONTINUATION of therapy, please answer the questions on PAGE 3								
□ NO – this is INITIATION of therapy, please answer the questions below:								
2. Is this request for brand or generic? ☐ Brand ☐ Generic								
3. BRAND Firazyr Request (Standard Option) : Would you like to switch the patient to a preferred product, GENERIC Firazyr (icatibant) or Sajazir? □Yes, switch to generic Firazyr (icatibant) □Yes, switch to Sajazir □No, do not switch* *If NO, does the patient have an intolerance or contraindication to or have they had an inadequate treatment response to generic Firazyr (icatibant) or Sajazir? Please select answer below: □Yes, specify drug and result: □								
□No: Is there a clinical reason for not trying generic Firazyr (icatibant) or Sajazir? □Yes* □No *If YES, please specify:								
4. What is the pa	_							
•	Angioedema (F			_				
5. Is this medication being used to treat acute attacks or for the routine prevention of hereditary angioedema? <i>Select answer below:</i> Acute attacks OR Routine prevention							t answer below:	
6. Does the patie				onfirmed by labor	ratory testing? S	Select answer b	elow:	
□Yes: Please answer the following questions: a. Does the patient have a F12, angiopoietin-1, plasminogen, or kininogen-1 (KNG1) gene mutation as confirmed by								
•	genetic testing? \(\text{Yes} \) \(\text{No} \)							
 b. Does the patient have a documented family history of angioedema? □Yes* □No *If YES, was the angioedema refractory to a trial of high-dose antihistamine such as cetirizine for at least one month? □Yes □No 								
			estions:					
	■No: Please answer the following questions: a. Does the patient have a C1 inhibitor deficiency or dysfunction as confirmed by laboratory testing? ■Yes ■No							
b. Is the patient's C4 level below the lower limit of normal as defined by the laboratory performing the test? \(\sigma \text{Yes} \square \square \text{No}\)								
c. Does the patient have a normal C1-INH antigenic level as defined by the laboratory performing the test? ☐ Yes: Does the patient have a C1-INH functional level less than 50% or a C1-INH functional level below the lower limit of normal as defined by the laboratory performing the test? ☐ Yes ☐ No								
□No: Is the patient's C1 inhibitor (C1-INH) antigenic level below the lower limit of normal as defined by the laboratory performing the test? □Yes □No								
	DITACE	DDACI	CED TO D.	A CE 2 EOD AD	DITIONIAT IN	TITTA TION O	TIECTIONS	

PLEASE PROCEED TO <u>PAGE 2</u> FOR ADDITIONAL INITIATION QUESTIONS

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PAGE 2 - PHYSICIAN COMPLETES					
F	Patient Name:	DOB:	Patient ID: R		
7.	Kalbitor, Ruconest)? □Yes* □	No	or treating acute attacks of hereditary	angioedema (e.g., Berina	ert,
8.	Is this medication being requested	as a change from BRAND Fira	azyr so the member can access their c	opay benefit? □Yes □	□No

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physician portion and submit this completed form.

Federal Employee Program.

OR

☐Acute attacks

Kalbitor, Ruconest)? □Yes*

*If YES, specify the medication: ___

□Routine prevention

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Patient In	nformation (required)		Provider Information (required)			
Date:			Provider Name:			
Patient Name:			Specialty:	NPI:		
Date of Birth:	Sex: Male	Female	Office Phone:	Office Fax:		
Street Address:			Office Street Address:			
City:	State:	Zip:	City:	State:	Zip:	
Patient ID: R			Physician Signature:	<u> </u>		
	P	HYSICIAN C	COMPLETES			
For Stan	dard Option patients GEN	NERIC Firazyr (id	catibant) and Sajazir are the p	preferred products		
		• .	azir will be eligible for 2 copay	-		
	CONTINUATIO	ON OF THI	ERAPY (PA RENE	WAL)		
	NOTE: Form m	ust be completed	d in its entirety for processing	<u>g</u>		
Please select medication	: □Fira	azyr (icatibant)	□Saja	zir (icatibant)		
*Check www.fepblue.org/form	ulary to confirm which medic	ation is part of the	patient's benefit			
□NO – this is INITIA' □YES – this is a PA re Is this request for brand BRAND Firazyr Requ (icatibant) or Sajazir? *If NO, does the patic Firazyr (icatibant) or □Yes, specify drug a □No: Is there a clinic	TION of therapy, please and the series of the continuation of generic? Brand est (Standard Option): Values, switch to generic lent have an intolerance of Sajazir? Please select and the continuation of t	answer the quest TON of therapy, Generic Would you like the Firazyr (icatibant contraindications wer below: eneric Firazyr (i	please answer the questions to switch the patient to a prefet DYes, switch to Sajazir at to or have they had an inade	below: erred product GEN No, do not sy equate treatment re	NERIC Firazyr witch*	
					_	
 What is the patient's dia ☐Hereditary Angioeder 	•					
☐Other diagnosis (pleas						
•		s or for the routi	ne prevention of hereditary a	ngioedema? Selec	t answer below:	

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7. Will this medication be used in combination with another agent for treating acute attacks of hereditary angioedema (e.g., Berinert,

6. Has the patient experienced a reduction in severity and/or duration of hereditary angioedema attacks? □Yes □No



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Message:

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Attached is a Prior Authorization request form.

For your convenience, there are 3 ways to complete a Prior Authorization request:

Electronically Online (ePA) Results in 2-3 minutes FASTEST AND EASIEST	Now you can get responses to drug Prior Authorization requests securely online. Online submissions may receive instant responses and do not require faxing or phone calls. Requests can be made 24 hours a day, 7 days a week. For more information on electronic prior authorization (ePA) and to register, go to Caremark.com/ePA.
Phone (4-5 minutes for response)	The FEP Clinical Call Center can be reached at (877)-727-3784 between the hours of 7AM-9PM Eastern Time. A live representative will assist with the Prior Authorization, asking for the same information contained on the attached form. Please review the form and have your answers ready for faster service. The process over the phone takes on average between 4 and 5 minutes.
Fax (3-5 days for response)	Fax the attached form to (877)-378-4727. Requests sent via fax will be processed and responded to within 5 business days. The form must be filled out completely, if there is any missing information the Prior Authorization request cannot be processed. Please only fax the completed form once as duplicate submissions may delay processing times.

faster... easier... better...

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