

PRIOR AUTHORIZATION REQUEST FORM CROHN'S DISEASE MEDICATIONS

Cimzia®, Hadlima™, Humira®, Rinvoq®, Skyrizi®, Stelara®

For authorization, please answer each question and fax this form PLUS chart notes back to Real Rx at 385-425-4052.

Failure to submit clinical documentation to support this request will result in a dismissal of the request.

If you have prior authorization questions, please call for assistance 385-425-5094.								
Disc	claimer: Prior authorization request fo	rms are subject to change in acc	ordance v	vith Feder	al and State notice requirements.			
Dat	e:	Member Name:		ID#:				
DOB:		Gender:		Physician:				
Office Phone:		Office Fax:		Office Contact:				
Height/Weight:			HCPCS Code:					
reason for failure. Reasons for failure must meet the Health Plan medical necessity criteria. Preferred/Non-preferred 1. Preferred Brands: A. Cimzia® (certolizumab), Hadlima™ (adalimumab-bwwd), Humira® (adalimumab), Rinvoq® (upadacitinib), Skyrizi® (risankizumab-rzaa), Stelara® (ustekinumab) Product being requested:								
Dosing/Frequency:								
If the request is for reauthorization, proceed to reauthorization section								
	Questions		Yes	No	Comments/Notes			
1.	Is the requested medication being provider's office and to be billed u ('buy-and-bill')?	•						
2.	Is this request for an expedited red By checking the "Yes" box to reque hours), you are certifying that app time frame (72 hours) may place to ability to regain maximum function	est an expedited review (24 lying the standard review he member's life, health, or						
3.	Is the request being made by or in gastroenterologist?							
4.	Does documentation include resul colonoscopy, MRI, CT scan?	ts from studies such as			Please provide documentation			

5.	Does the member have severe Crohn's Disease evidenced by at least one of the following: • A Crohn's Disease Activity Score (CDAI) >220 AND as shown on imaging • Active fistulizing disease			Please provide documentation				
6.	Does the member have moderate to severe Crohn's Disease evidenced by the following: • Persistent fistulizing disease or active ulcerative disease as shown on imaging and via CDAI > 150 despite an adequate trial with an immunomodulating medication such as methotrexate, azathioprine or 6-mercaptopurine, unless contraindicated to all.			Please provide documentation				
7.	Has the provider performed tuberculosis (TB) screening prior to therapy initiation?			Please provide documentation				
8.	If the request is for a Tumor Necrosis Factor Inhibitor, has the provider performed hepatitis B screening prior to therapy initiation?			Please provide documentation				
REAUTHORIZATION								
1.	Is the request for reauthorization of therapy?							
2.	Does documentation show a stabilization or decrease in the CDAI score of at least 70 points compared to baseline, endoscopic improvement in mucosa and/or no new fistulizing disease information?			Please provide documentation				
3.	Has the provider performed continued tuberculosis monitoring during therapy?			Please provide documentation				
4.	Has the provider performed continued Hepatitis B monitoring in HBV carriers?			Please provide documentation				
What medications and/or treatment modalities have been tried in the past for this condition? Please document name of treatment, reason for failure, treatment dates, etc.								
Additional information:								
Physician Signature:								

** Failure to submit clinical documentation to support this request will result in a dismissal of the request.**

Policy: PHARM-019

Origination Date: 03/14/2018 Reviewed/Revised Date: 09/13/2023 Next Review Date: 09/13/2024 Current Effective Date: 10/01/2023

Confidentiality Notice

This document and any accompanying document contain confidential information and is intended for the use of the individual or entity named on this transmission sheet. If you are not the intended recipient, you are hereby notified that any disclosure, copying, distribution or the taking of any action in reliance on the contents of this information is strictly prohibited and the document should be returned to this office immediately. If you have received this facsimile in error, please notify us by telephone immediately and destroy document received.