## HEALTHY U CHIP

### PRIOR AUTHORIZATION REQUEST FORM ULCERATIVE COLITIS

Avsola<sup>®</sup>, Entyvio<sup>®</sup>, Hadlima<sup>™</sup>, Inflectra<sup>®</sup>, infliximab, Remicade<sup>®</sup>, Renflexis<sup>®</sup>, Rinvoq<sup>®</sup>, Simlandi<sup>®</sup>, Simponi<sup>®</sup>, Skyrizi<sup>®</sup>, Tremfya<sup>®</sup>, Xeljanz<sup>®</sup>, Zeposia<sup>®</sup>, Yesintek<sup>™</sup>

For authorization, please answer each question and fax this form PLUS chart notes back to the Healthy U CHIP Prior Authorization Department.

- For Medical Pharmacy please fax requests to: 801-213-1547
- For **Retail Pharmacy** please fax requests to: 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 Pharmacy Customer Service for assistance at 385-425-5094

Disclaimer: Prior authorization request forms are subject to change in accordance with Federal and State notice requirements.

Date:	Member Name:	ID#:
DOB:	Gender:	Physician:
Office Phone:	Office Fax:	Office Contact:
Height/Weight:		HCPCS Code:

Member must try formulary preferred drugs before a request for a non-formulary drug may be considered. If treatment with preferred products has not been successful, you must submit which preferred products have been tried, dates of treatment, and reason for failure. Reasons for failure must meet the Health Plan medical necessity criteria.

#### Preferred/Non-Preferred/Non-Formulary:

- 1. 1<sup>st</sup> Line Preferred Agents:
  - A. Hadlima<sup>™</sup> (adalimumab-bwwd), Simlandi<sup>®</sup> (adalimumab-ryvk)
  - B. Infliximab products: Avsola<sup>®</sup> (infliximab-axxq), Inflectra<sup>®</sup> (infliximab-dyyb), infliximab, Remicade (infliximab), Renflexis<sup>®</sup> (infliximab-abda)
  - C. Wezlana<sup>™</sup> (ustekinumab-auub), Yesintek<sup>™</sup> (ustekinumab-kfce)
- 2. 2nd line preferred agents with single step; after trial and failure of an adalimumab product, an ustekinumab product and an infliximab product:
  - A. Enytvio<sup>®</sup> (vedolizumab) IV, Xeljanz<sup>®</sup>/XR (tofacitinib)
- 3. Non-Formulary Agents with a triple step; after trial and failure of an adalimumab product, an ustekinumab product, an infliximab product and 2 second line agents:
  - A. Rinvoq<sup>®</sup> (upadacitinib), Simponi<sup>®</sup> (golimumab)
- 4. Non-Formulary Agent after trial and failure of all the above:
  - A. Entyvio<sup>®</sup> (vedolizumab) subcutaneous injection, Skyrizi<sup>®</sup> (risankizumab-rzaa), Tremfya<sup>®</sup> (guselkumab), Zeposia<sup>®</sup> (ozanimod)

Product being requested: \_\_\_\_\_

Dosing/Frequency:

If the request is for reauthorization, proceed to reauthorization section					
Questions		No	Comments/Notes		
MODERATE TO SEVERE ULCERATIVE COLITIS					
1. Has the member been diagnosed with moderate to severe Ulcerative Colitis?			Please provide documentation		

2. Is the prescribing provider a gastroenterologist or in						
consultation with a gastroenterologist?						
3. Has the provider performed tuberculosis (TB) screening prior to therapy initiation?			Please provide documentation			
4. If the request is for Tumor Necrosis Factor Inhibitors (TNFIs),			Please provide documentation			
Rinvoq or Xeljanz/XR, has the provider performed hepatitis B screening prior to therapy initiation?						
5. Has the member had an adequate trial and failure of at least one			Please provide documentation			
of the following, or contraindication to all:						
<ul> <li>High dose oral 5-aminosalicyclic acid drug</li> </ul>						
<ul> <li>Topical 5-aminosalicylic acid drug</li> </ul>						
6. If the request is for Rinvoq or Xeljanz/XR <sup>®</sup> , does documentation			Please provide documentation			
show inadequate response or intolerance to at least one tumor						
necrosis factor (TNF) blocker such as an infliximab product,						
Cimzia, an adalimumab product and/or Simponi and does						
documentation show the member will not be receiving Rinvoq						
or Xeljanz/XR in combination with a potent immunosuppressant						
(e.g., azathioprine or cyclosporine )?						
SEVERE ULCERATIVE COL	LITIS					
1. Has the member been diagnosed with severe Ulcerative Colitis?			Please provide documentation			
• Has the patient had more than six stools per day with blood OR		_	•			
has systemic symptoms (fever, tachycardia, anemia or						
erythrocyte sedimentation rate > 30mm/h)?						
2. Is the prescribing provider a gastroenterologist or in			Please provide documentation			
consultation with a gastroenterologist?			· · · · · · · · · · · · · · · · · · ·			
3. Has the provider performed tuberculosis (TB) screening prior to			Please provide documentation			
therapy initiation?			· · · · · · · · · · · · · · · · · · ·			
4. Has the provider performed hepatitis B screening prior to			Please provide documentation			
therapy initiation?			•			
5. If the request is for Rinvoq or Xeljanz/XR <sup>®</sup> , does documentation			Please provide documentation			
show inadequate response or intolerance to at least one tumor						
necrosis factor (TNF) blocker such as an infliximab product,						
Cimzia, Humira and/or Simponi and does documentation show						
the member will not be receiving Rinvoq or Xeljanz/XR in						
combination with a potent immunosuppressant (e.g.,						
azathioprine or cyclosporine )?						
FULMINANT COLITIS						
1. Has the member been diagnosed with fulminant colitis?			Please provide documentation			
• Has the member had more than 10 bowel movements per day						
with continuous bleeding OR has colonic dilation, transfusion						
requirement, or toxicity?						
2. Is the prescribing provider a gastroenterologist or in			Please provide documentation			
consultation with a gastroenterologist?						
3. Has the provider performed tuberculosis (TB) screening prior to			Please provide documentation			
therapy initiation?						
4. Has the provider performed hepatitis B screening prior to			Please provide documentation			
therapy initiation?						
5. If the request is for Rinvoq or Xeljanz/XR <sup>®</sup> , does documentation			Please provide documentation			
show inadequate response or intolerance to at least one tumor						
necrosis factor (TNF) blocker such as an infliximab product,						
Cimzia, Humira and/or Simponi and does documentation show						
the member will not be receiving Rinvoq or Xeljanz/XR in	1	1				

combination with a potent immunosuppressant (e.g.,							
azathioprine or cyclosporine )?							
1. Is the request for reauthorization of therapy?							
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2. Does updated clinical documentation show a positive response			Please provide documentation				
to therapy, such as a decrease or stabilization in the Disease							
Activity Index (DAI) score?							
3. Has the provider performed continued tuberculosis monitoring during therapy?			Please provide documentation				
4. Has the provider performed continued Hepatitis B monitoring in			Please provide documentation				
HBV carriers?							
What medications and/or treatment modalities have been tried in the	ne past	for this	condition? Please document				
name of treatment, reason for failure, treatment dates, etc.							
Additional information:							
Dhucician's Signatura							
Physician's Signature:							

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

Policy: PHARM-CHIP-075 Origination Date: 07/01/2024 Reviewed/Revised Date: 04/09/2025 Next Review Date: 04/09/2026 Current Effective Date: 05/01/2025

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