

## NC Medicaid Pharmacy Prior Approval Request Immunomodulators: Xeljanz XR

## **Beneficiary Information**

1. Beneficiary Last Name:	2. First Name:		
	4. Beneficiary Date of Birth:		
Prescriber Information			
6. Prescribing Provider NPI #:			
			Ext
Drug Information	_		
8. Drug Name:	9. Strength:		10. Quantity Per 30 Days:
11. Length of Therapy (in days):			
Other			
Clinical Information			
Request for Ankylosing Spondyli	tis		
1. Does the beneficiary have a dia		dylitis? 🗆 Yes 🗆 No	
2. Is the beneficiary not on anoth	er injectable biologic immu	nomodulator? 🗆 Yes	□ No
3. Has the beneficiary individual risks and benefits been considered prior to initiating or continuing therapy in those at			
higher risk for malignancy and/or	major adverse cardiovascu	ılar events (MACE)? □	] Yes □ No
4. Is the beneficiary NOT consider	red to be at high risk for thr	rombosis? $\square$ Yes $\square$ N	0
5. Has the beneficiary been considered and screened for the presence of latent tuberculosis infection?   Yes  No			
6. Has the beneficiary been teste	d with Hep B SAG and Core	Ab? ☐ <b>Yes</b> ☐ <b>No</b>	
7. Will the beneficiary NOT receive live vaccines during therapy?   Yes   No			
8. Has the beneficiary tried at least one Tumor Necrosis Factor Blocker with inadequate response or unable to take			
these therapies due to intolerance or contraindications?   Yes  No			
9. Has the beneficiary had a trial and failure of Cosentyx, Enbrel or Humira or a clinical reason beneficiary cannot try			
Cosentyx, Enbrel or Humira? 🗆 Y	es ⊔ No		
Request for Psoriatic Arthritis			
1. Does the beneficiary have a do	cumented definitive diagno	osis of Psoriatic Arthri	tis? 🗆 <b>Yes</b> 🗆 <b>No</b>
2. Is the beneficiary 18 years of a	ge or older? 🗆 Yes 🗆 No		
3. Is the beneficiary not on anoth	er injectable biologic immu	nomodulator? 🗆 Yes	□ No
4. Has the beneficiary individual risks and benefits been considered prior to initiating or continuing therapy in those at			
higher risk for malignancy and/or	major adverse cardiovascu	ılar events (MACE)? □	] Yes □ No
5. Is the beneficiary NOT conside	red to be at high risk for thr	rombosis? $\square$ Yes $\square$ N	0
6. Has the beneficiary been consi	dered and screened for the	presence of latent tu	berculosis infection? $\square$ Yes $\square$ No
7. Has the beneficiary been teste	•		
8. Does the beneficiary have a do		onse, intolerance or o	contraindication to at least one
Tumor Necrosis Factor Blocker?			
		orei or Humira or a cli	nical reason beneficiary cannot try
Cosentyx, Enbrel or Humira? 🗆 Y	es ⊔ No		



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ignature of Prescriber: Date:
8. Has the beneficiary had a trial and failure of Humira or a clinical reason beneficiary cannot try Humira?   Yes  No
7. Will the beneficiary NOT receive live vaccines during therapy? ☐ <b>Yes</b> ☐ <b>No</b>
6. Has the beneficiary been tested with Hep B SAG and Core Ab?   Yes  No
5. Has the beneficiary been considered and screened for the presence of latent tuberculosis?   Yes  No
4. Is the beneficiary NOT considered to be at high risk for thrombosis? ☐ <b>Yes</b> ☐ <b>No</b>
higher risk for malignancy and/or major adverse cardiovascular events (MACE)?   Yes   No
3. Has the beneficiary individual risks and benefits been considered prior to initiating or continuing therapy in those at
2. Is the beneficiary not on another injectable biologic immunomodulator? ☐ <b>Yes</b> ☐ <b>No</b>
1. Does the beneficiary have a diagnosis of ulcerative colitis? ☐ Yes ☐ No
Request for Ulcerative Colitis (Adult)
Humira? ☐ Yes ☐ No
10. Has the beneficiary had a trial and failure of Enbrel or Humira or a clinical reason beneficiary cannot try Enbrel or
_ Yes □ No
9. Is the beneficiary unable to receive Tumor Necrosis Factor Blocker due to contraindications or intolerabilities?
Blocker? ☐ Yes ☐ No
8. Has the beneficiary experienced a therapeutic failure/inadequate response with at least one Tumor Necrosis Factor
7. Will the beneficiary NOT receive live vaccines during therapy?   Yes   No
6. Has the beneficiary been tested with Hep B SAG and Core Ab? $\square$ Yes $\square$ No
5. Has the beneficiary been considered and screened for the presence of latent tuberculosis?   Yes   No
4. Is the beneficiary NOT considered to be at high risk for thrombosis? ☐ <b>Yes</b> ☐ <b>No</b>
higher risk for malignancy and/or major adverse cardiovascular events (MACE)?   Yes   No
3. Has the beneficiary individual risks and benefits been considered prior to initiating or continuing therapy in those at
2. Is the beneficiary not on another injectable biologic immunomodulator? ☐ <b>Yes</b> ☐ <b>No</b>
1. Does the beneficiary have a diagnosis of Rheumatoid Arthritis? ☐ <b>Yes</b> ☐ <b>No</b>
Request for Rheumatoid Arthritis

(Prescriber Signature Mandatory)

I certify that the information provided is accurate and complete to the best of my knowledge, and I understand that any falsification, omission, or concealment of material fact may subject me to civil or criminal liability.

Pharmacy PA Call Center: 1-866-799-5318