Trillium Health Resources Pharmacy Prior Approval Request for



Immunomodulators: Cimzia

Member Information								
1. Member Last Name:	2. First Name:							
3. Member ID #:	4. Member Date of Birth:5. Member Gender:							
Prescriber Information								
6. Prescribing Provider NPI #:								
			Ext					
Drug Information								
8. Drug Name:	9. Strength: _	10. 0	Quantity Per 30 Days:					
11. Length of Therapy (in days):								
Other								
Clinical Information								
Request for Ankylosing Spond	ylitis							
1. Does the member have a dia	agnosis of Ankylosing Spondyli	tis? 🗆 Yes 🗆 No						
2. Is the member not on anoth	•							
3. Has the member been consi	•		losis infection? 🗆 Yes 🗆 No					
4. Has the member been teste	•							
			east two NSAIDS? Yes No					
	6. Is the member unable to receive treatment with NSAIDS due to contraindications? Yes No							
7. Does the member have clinical evidence of severe or rapidly progressing disease								
	8. Has the member had a trial and failure of Cosentyx, Enbrel or Humira or a clinical reason member cannot try Cosentyx, Enbrel or Humira? Yes No							
Request for Crohn's Disease (A	\dult)							
1. Does the member have a dia	ignosis of moderate to severe	Crohn's Disease? 🗆 Yes						
2. Is the member not on anoth	•							
3. Has the member been consi	•		losis infection? 🗆 Yes 🗆 No					
4. Has the member been tested with Hep B SAG and Core Ab? 🗆 Yes 🗆 No								
5. Has the member had a trial a	and failure of Humira or a clini	ical reason member canr	not try Humira? 🗆 Yes 🗆 No					
Request for Plaque Psoriasis (Adult)							
	cumented definitive diagnosis	of moderate-to-severe	Chronic Plaque Psoriasis? 🗆 Yes					
□ No								
2. Is the member 18 years of age or older? Yes No								
3. Is the member not on another injectable biologic immunomodulator? Yes No								
4. Has the member been considered and screened for the presence of latent tuberculosis infection (not required for Otezla)? Yes No								



- 5. Has the member been tested with Hep B SAG and Core Ab? \Box Yes \Box No
- 6. Does the member have a body surface area (BSA) involvement of at least 3%?

 Yes
 No

7. Does the member have involvement of the palms, soles, head and neck, or genitalia, causing disruption in normal daily activities and/or employment?
Yes
No

8. Has the member failed to respond to, or has been unable to tolerate phototherapy and **ONE** of the following medications or member has contraindications to these treatments: Soriatane (acitretin), Methotrexate, and/or Cyclosporine?
Yes No

9. Has the member had a trial and failure of Cosentyx, Enbrel or Humira or a clinical reason member cannot try Cosentyx, Enbrel or Humira?
Yes
No

10. Are the beneficiaries, Providers, and Pharmacies utilizing Siliq registered appropriately in the Siliq Risk Evaluation and Mitigation Strategy Program (REMS program)?
Yes
No

Request for Psoriatic Arthritis

1. Does the member have a documented definitive diagnosis of Psoriatic Arthritis? \Box Yes \Box No

- 2. Is the member 18 years of age or older (OR 2 years or older for Simponi Aria)?
 Yes
 No
- 3. Is the member not on another injectable biologic immunomodulator? \Box Yes \Box No

4. Has the member been considered and screened for the presence of latent tuberculosis infection (not required for Otezla?

Yes
No

5. Has the member been tested with Hep B SAG and Core Ab (not required for Otezla?
Yes
No

- 6. Does the member have a documented inadequate response or inability to take methotrexate?
- 7. Has the member had a trial and failure of Cosentyx, Enbrel or Humira or a clinical reason member cannot try Cosentyx, Enbrel or Humira?
 Yes
 No

Request for Rheumatoid Arthritis

1. Does the member have a diagnosis of Rheumatoid Arthritis? \Box Yes \Box No

- 2. Is the member not on another injectable biologic immunomodulator? \Box Yes \Box No
- 3. Has the member been considered and screened for the presence of latent tuberculosis? \Box Yes \Box No

4. Has the member been tested with Hep B SAG and Core Ab? \Box Yes \Box No

5. Has the member experienced a therapeutic failure/inadequate response with methotrexate or at least one disease modifying antirheumatic drug (e.g. leflunomide, hydroxychloroquine, minocycline, sulfasalazine)? □ Yes □ No
6. Is the member unable to receive methotrexate or disease modifying antirheumatic drug due to contraindications or intolerabilities? □ Yes □ No

7. Does the member have clinical evidence of severe or rapidly progressing disease?

Yes
No

8. Has the member had a trial and failure of Enbrel or Humira or a clinical reason member cannot try Enbrel or Humira?

Yes
No

Request for Non-Radiographic Axial Spondyloarthritis

1. Does the member have a diagnosis of Non-Radiographic Axial Spondyloarthritis? \Box Yes \Box No

2. Is the member not on another injectable biologic immunomodulator? \Box Yes \Box No

3. Has the member failed an adequate trial of a Non-Steroidal Anti-Inflammatory Drug (NSAID) unless

contraindicated?
Ves
No



4.	Has the membe	er been considered	l and screened fo	or the presence of	latent tuberculosis?	🗆 Yes 🗆 No
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- 5. Has the member been tested with Hep B SAG and Core Ab? \Box Yes \Box No
- 6. Has the member had a trial and failure of Cosentyx? \Box Yes \Box No

Date:_____

Signature of Prescriber: ______ (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.