Caterpillar Prescription Drug Benefit Phone: 877-228-7909 Fax: 800-424-7640

MEMBER'S LAST NAME:		MEMBER'S FIRST	NAME:	
Instructions: Please fill outhat is important for the revontained in this form is Pr	iew (e.g., chart notes o	or lab data, to support t		
				URGENT
MEMBER INFORMATION	١			
LAST NAME:		FIRST NAME:		
PHONE NUMBER:		DATE OF BIRT	H:	
STREET ADDRESS:				
CITY:		STATE:	ZIP CODE:	
PATIENT INSURANCE II	NUMBER:	1		
☐ MALE ☐ FEMALE H	HEIGHT (IN/CM):	WEIGHT (LB/KG)	: ALLERGIES: _	
IF YOU ARE NOT THE PADISCLOSURE AUTHORIZE FOLLOWING LINK: PRIM	ATION FORM WITH TETHERAPEUTICS.CO	THIS REQUEST WHIC DM/NOPP (IF APPLICABLE):	H CAN BE FOUND AT TH	
AUTHORIZED REPRESEI	NTATIVE'S PHONE N	UMBER:		
PRESCRIBER INFORMA	TION			
LAST NAME:		FIRST NAME:		
PRESCRIBER SPECIAL	гү:	EMAIL ADDRE	SS:	
NPI NUMBER:		DEA NUMBER	:	
PHONE NUMBER:		FAX NUMBER		
STREET ADDRESS:				
CITY:		STATE:	ZIP CODE:	
REQUESTER (if differen	t than prescriber):	OFFICE CONT	ACT PERSON:	
MEDICATION OR MEDIC	CAL DISPENSING INF	ORMATION		
MEDICATION NAME:				
DOSE/STRENGTH:	FREQUENCY:	LENGTH OF THERAPY/REI	QUANTITY:	
☐ NEW THERAPY	—	IF RENEWAL: DATE		
DURATION OF THERAP	Y (SPECIFIC DATES):			
Continued on next page				

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MEMBER'S LAST NAME:	MEMBER'S FIRST N	AME:		
1. HAS THE PATIENT TRIED ANY OTHER MEDICATIONS FOR THIS CONDITION? YES (if yes, complete below) NO				
MEDICATION/THERAPY (SPECIFY DRUG NAME AND DOSAGE):	DURATION OF THERAPY (SPECIFY DATES):	RESPONSE/REASON FOR FAILURE/ALLERGY:		
2. LIST DIAGNOSES:		ICD-10:		
 ☐ Rheumatoid arthritis(RA) ☐ Moderate to severe Atopic Der ☐ Psoriatic Arthritis (PsA) ☐ Ulcerative Colitis(UC) ☐ Crohn's Disease(CD) ☐ Ankylosing Spondylitis ☐ Non-radiographic Axial Spond ☐ Atopic Dermatitis ☐ Giant Cell Arteritis ☐ Other diagnosis: 	• •			
3. REQUIRED CLINICAL INFORMA	ATION: PLEASE PROVIDE ALL REL	EVANT CLINICAL INFORMATION		
TO SUPPORT A PRIOR AUTHORIZ	ZATION.			
Is patient going to be using drug	in combination with a clinical trial?	? ☐ Yes ☐ No		
Does patient have difficulty swallowing? □ Yes □ No Please submit documentation. Does patient have an enteral feeding tube? □ Yes □ No Please submit documentation. Is patient taking any other oral tablets or capsules(*sprinkle caps ok)? □ Yes □ No Please submit documentation.				
Is the patient currently being trea	ted with another biologic or immur	nomodulatory agent? □ Yes □		
If on another biologic therapy, will No	Il that biologic be stopped when st	arting the RinvoqER? □ Yes □		
	nadequate response to a three (3) of the contract of the contr			
Does patient have a absolute con ☐ No Please submit documentation	traindication to the biosimilar for F on.	lumira-adalimumab-aacf? □ Yes		
	nadequate response to a 4- month No Please submit documentati			
Does patient have a absolute con	traindication to the biosimilar for	Stelara- <u>Otulfi(ustekinumb-aauz</u>)?		



MEMBER'S LAST NAME: MEMBER'S FIRST NAME:	
For diagnosis of Rheumatoid Arthritis only:	
Is the prescriber a rheumatologist? □ Yes □ No	
Does the patient have a diagnosis of moderately to severely active rheumatoid arthritis? \hdots No	
Has the patient had a trial of methotrexate or another oral non-biologic disease modifying anti-rheumatic agent (DMARD) such as Imuran, Ridaura, Plaquenil, sulfasalazine or Arava? Please submit documentation with dates of service.	No
Does patient have chronic alcohol abuse/alcoholism, chronic liver disease such as chronic hepatity liver, nonalcoholic steatohepatitis/NASH, elevated liver enzymes) (Please provide documentation.)? □ Yes □ No Please submit documentation.	atitis,
For renewal only: Does the patient continue to have a positive clinical response and remission of disease maintain with continued use of the medication? Yes No Please submit chart documentation.	ned
Is the patient currently being treated with another biologic or immunomodulatory agent? $\ \ \Box$ Yes No	
Is the prescriber a rheumatologist? □ Yes □ No	
For diagnosis of <u>Atopic Dermatitis</u> only:	
Is the prescriber a dermatologist or allergist? □ Yes □ No	
Has the patient had the diagnosis of atopic dermatitis for at least 12 months? \Box Yes \Box No *Plosubmit documentation.	ease
Does the patient have atopic dermatitis on at least 10% or more of their body surface area? No *Please submit documentation.	es 🗆
Has the patient tried at least two different topical steroids? ☐ Yes ☐ No *Please submit documentation.	
If patient has not had at least 2 different topical steroids, has the patient tried at least one topical steroid AND one topical calcineurin inhibitor (tacrolimus or pimecrolimus)? — Yes — No *Please submit documentation.	
If patient has not had at least 2 different topical steroids, has the patient tried at least one topica steroid AND Eucrisa(crisaborole)? □ Yes □ No *Please submit documentation.	I
If patient has not had at least 2 different topical steroids, has the patient tried at least one topica steroid AND Zoryve(roflumilast)? Yes No *Please submit documentation.	I



MEMBER'S LAST NAME: MEMBER'S FIRST NAME:
If patient has not had at least 2 different topical steroids, has the patient tried at least one topical steroid AND Vtama(tapinarof)? □ Yes □ No *Please submit documentation.
Has patient tried and failed a 3-month trial of Dupixent(dupilumab)? ☐ Yes ☐ No *Please submit documentation.
Has patient tried and failed a 3-month trial of Adbry(tralokinumab-ldrm)? \Box Yes \Box No *Please submit documentation.
Has patient tried and failed a 3-month trial of Cibinqo(abrocitinib)? ☐ Yes ☐ No *Please submit documentation.
Will RinvoqER(upadacitinib) be used in combination with Cibinqo(abrocitinib), Olumiant(baracitinib), Opzelura(ruxolitinib), Dupixent(dupilumab), Adbry(tralokinumab), Xolair(omalizumab), Nucala(mepolizumab) or Fasenra(benralizumab? Yes No
For renewal only: Does the patient continue to have a positive clinical response and remission of disease maintained with continued use of the medication? Yes No Please submit chart documentation.
Is the patient currently being treated with another biologic or immunomodulatory agent? $\ \ \Box$ Yes $\ \ \Box$ No
Is RinvoqER(upadacitinib) being used in combination with Cibinqo(abrocitinib), Olumiant(baracitinib), Opzelura(ruxolitinib), Dupixent(dupilumab), Adbry(tralokinumab), Xolair(omalizumab), Nucala(mepolizumab) or Fasenra(benralizumab? □ Yes □ No
Is the prescriber a dermatologist or allergist? □ Yes □ No
For diagnosis of <u>Psoriatic Arthritis</u> only:
Is the prescriber a rheumatologist or dermatologist? □ Yes □ No
Does the patient have documented moderately to severely active disease? — Yes — No Please submit documentation
Has the patient had a trial and failed previous therapy with oral disease modifying anti-rheumatic agents (DMARDs, e.g., methotrexate, sulfasalazine (Azulfidine), leflunamide (Arava), or cyclosporine)? No Please submit documentation with dates of service.
For renewal only: Does the patient continue to have a positive clinical response and remission of disease maintained with continued use of the medication? No Please submit chart documentation.



MEMBER'S LAST NAME: MEMBER'S FIRST NAME:	
Is the patient currently being treated with another biologic or immunomodul No	latory agent? Yes
Is the prescriber a rheumatologist or dermatologist? ☐ Yes ☐ No	
For diagnosis of <u>Ulcerative Colitis and Crohn's Disease</u> Only: Is prescriber a gastroenterologist? □ Yes □ No Has patient tried and failed at least one of the following three therapies: cor azathioprine, and/or 6-mercaptopurine? □ Yes □ No	ticosteroids,
Has patient tried and failed at least three months of another intravenous, su therapy? □ Yes □ No Please submit documentation with dates of treatment	
For renewal only: Does the patient continue to have a positive clinical response and remission with continued use of the medication? Yes No Please submit chart do	
Is the patient currently being treated with another biologic or immunomodul No	latory agent? □ Yes □
Is the prescriber a rheumatologist or gastroenterologist? ☐ Yes ☐ No	
For diagnosis of Ankylosing Spondylitis only:	
Is the prescriber a rheumatologist? □ Yes □ No	
Does the patient have documented active disease? □ Yes □ No Please suit	bmit documentation
Has the patient had a trial and failed previous therapy with at least two (2) n inflammatory agents (NSAIDS), unless use is contraindicated? Please submit documentation with dates of service.	
For renewal only: Does the patient continue to have a positive clinical response and remission with continued use of the medication? Yes No Please submit chart do	
Is the patient currently being treated with another biologic or immunomodul No	latory agent? □ Yes □
Is the prescriber a rheumatologist? □ Yes □ No	
For diagnosis of Non-radiographic Axial Spondyloarthritis only: Is the prescriber a rheumatologist? Yes No	
Does the patient have objective signs of inflammation by presence of sacro results and/or elevated C-reactive protein level? Yes No Please submit report.	



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Has patient had an inadequate responsibility to the submit documentation.	onse to at least two different NSAIDs? □ Yes □ No Please
	contraindication or intolerance to a 3-month trial with at least TNF inhibitor or an IL-17 inhibitor? □ Yes □ No Please submit
	positive clinical response and remission of disease maintained n? Yes No Please submit chart documentation.
Is the patient currently being treated No	with another biologic or immunomodulatory agent? □ Yes □
Is the prescriber a rheumatologist?	□ Yes □ No
For Giant Cell Arteritis:	
Is prescriber a rheumatologist? □ Ye	es 🗆 No
Does patient have a diagnosis of Giadocumentation	ant Cell Arteritis? Yes No Please submit chart
Is patient currently on a tapering dos Will Rinvoq be used as monotherapy For renewal only:	/? □ Yes □ No
	positive clinical response and remission of disease maintained n? Yes No Please submit chart documentation.
Is the patient currently being treated No	with another biologic or immunomodulatory agent? □ Yes □
Is the prescriber a rheumatologist?	□ Yes □ No
Are there any other comments, diagrinformation the physician feels is im	noses, symptoms, medications tried or failed, and/or any other portant to this review?
Please note: Not all drugs/diagnosis arequired information is received.	re covered on all plans. This request may be denied unless all
understand that the Health Plan, insure	n provided is true and accurate to the best of my knowledge. I er, Medical Group or its designees may perform a routine audit and sary to verify the accuracy of the information reported on this form.

Prime THERAPEUTICS**

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MEMBER'S LAST NAME:	MEMBER'S FIRST NAME:		
Prescriber Signature or Electronic I.D. Verification	on: Date:	_	
	ompanying this transmission contain confidential health		
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MAIL REQUESTS TO: Prime Therapeutics Management Prior Authorization Program

Attn: CP-4201 P.O. Box 64811 St. Paul, MN 55164-0811 **Phone**: 877-228-7909

