

Remicade[®] (infliximab) Injectable **Medication Precertification Request**

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For New Jersey HMO D-SNP: FAX: 1-833-322-0034 PHONE: 1-844-362-0934

For other lines of business: Please use other form.

Note: Remicade is preferred for

	Start of treatment: Start date			.)	MAPD	plans varies base ion. See section	ed on
	uested By:		Phone:		Fax	:	
A. PATIENT INFORM	MATION						
First Name:			Last Name:				
Address:			City:		State:	ZIP:	
Home Phone:		Work Phone:		Cell Phone:			
DOB:	Allergies:			Email:			
-	3	gs Height: _	inches or				
B. INSURANCE INF		jo rioigiit.		0			
		Does patient have	other coverage?				
			(
nsured:		Insured:					
C. PRESCRIBER INI	FORMATION						
First Name:		Last Name:		(Check O	ne): 🗌 M.[D. 🗌 D.O. 🗌 N.P.	🗆 P.A.
Address:			City:		State:	ZIP:	
Phone:	Fax:	St Lic #:	NPI #:	DEA #:		UPIN:	
Provider Email:		Office Contact Name:	1	Phone:			
	OVIDER/ADMINISTRATION IN						
Outpatient Infusi Center Nan Agency Nan Administration co Address: City: Phone: TIN: NPI: PRODUCT INFOR Request is for: Rem DIAGNOSIS INFO Primary ICD Code: G. CLINICAL INFOR	d Physician's Office on Center Phone:	ZIP: edication being requested Freq mary ICD Code and specify Secondary ICD Code: prmation must be completed		Retail Pl An Ail Orce An Ail	er Fax: PIN: HC ode:	_ ZIP:	
Note: Remicade, Infle ulcerative colitis and I on indication. Yes No Has Yes No Has Please explain if ther diagnosis (select all t Yes No Wil Yes No Has bio	sts (clinical documentation record ctra, Entyvio, and Simponi Aria a Enbrel, Humira, Kevzara, Otezla, s the patient had prior therapy w s the patient had a trial and failu Enbrel (etanercept) ☐ Humira Skyrizi (risankizumab-rzaa) ☐ e are any other medical reason(hat apply) Enbrel (etanercept) ☐ Humira Skyrizi (risankizumab-rzaa) ☐ IRemicade (infliximab) be used s the patient been tested for TB logic therapy? eck all that apply): ☐ PPD test	re the preferred products for Rinvoq, Skyrizi, and Xeljanz/ ith Remicade (infliximab) wil re, intolerance, or contraindi (adalimumab)	Xeljanz XR are preferred f thin the last 365 days? cation to any of the follow (sarilumab)	ving? (select all apremilast)	that apply) Rinvoq (up when indica Rinvoq (up (e.g., adalin	ed products vary b adacitinib) ated for the patient adacitinib) numab, certolizum	ased 's

- Please enter results of the TB test:
 positive negative negative
 - If positive, Does the patient have latent or active TB? 🗌 latent 📋 active

If latent TB, Yes No Will TB treatment be started before initiation of therapy with Remicade (infliximab)?



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Patient First Nam	е	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INI	FORMATION (continued) – R	equired clinical information must	be completed in its <u>entirety</u> for all	precertification requests.
Ankylosing Spor Please select wh	ndylitis and Other Spondyloa	rthropathies ne patient: □ Ankylosing spondyl	itis 🔲 Other spondyloarthropath	
🗌 Yes 🗌 No	Is there evidence of inflamma	tory disease?		
	Please provide the names and		-steroidal anti-inflammatory drugs	s (NSAIDs)?
	NSAID #2:			
$ $ \rightarrow	Is the disease refractory to co Please indicate: Corticoste Please provide the name of d	rticosteroids or immunosuppress roids 🔲 immunosuppressive dri rug tried:	ugs	
Behcet's Uveitis				
	Is the disease refractory?			
🖵 Yes 🔲 No	bus/Pulmonary Sarcoidosis Has the patient remained sym Please provide the daily dose	ptomatic despite treatment with s of steroids: Dose:mg	teroids?	
	Please select: 🗌 azathioprine	ptomatic despite treatment with in cyclophosphamide	mmunosuppressants? hotrexate	ain:
		osis of fistulizing Crohn's disease	?	
│ └────────────────────────────────────	Please indicate how long the p Does the patient have a diagn	patient has been diagnosed with f osis of Crohn's disease?	istulizing Crohn's disease:	
$ $ \longrightarrow	Yes No Does the pati	f the patient's disease: ☐ mild [ent have a documented diagnosis ct all signs/symptoms that apply: al pain ☐ arthritis ☐ bleeding	」 moderate L_ severe s of active Crohn's disease?	□ intestinal obstruction
	☐ megacolo ☐ Yes ☐ No Have the Cro	on	dylitis weight loss None o I active despite treatment with 6-r	f the above
		k all medications that apply: 🗌 6	-mercaptopurine	e ylprednisolone 🗌 Other:
Hidradenitis Sup	opurativa		-	
	he stage of hidradenitis suppuration the patient completed a tr	ativa: Hurley stage I (mild dise Hurley stage III (severe ial of antibiotics?		derate disease)
$ \longrightarrow$	☐ Yes ☐ No Does the pati ☐ Yes ☐ No Was the treat	ent have a contraindication to ora ment with antibiotics ineffective?		
			trial: Less than 1 month 1 2 months 3 months (
Please indicate th	oint Inhibitor-Induced Toxicit lerapy used:	les		
🗌 PD-1		:		
PD-L1		olizumab 🔲 Other: Iumab 🔲 durvalumab 🔲 Othe		
Other Please explain	-			
Yes No		nibitor-induced toxicities persist de ab, ipilimumab, nivolumab, pembr		checkpoint inhibitors that target CTLA-4 or
	he toxicity, (check all that ap			
Ple Colitis Ple	ease select: 🔲 arrhythmias 🗌] impaired ventricular function	uced colitis. 🗌 mild 🔲 modera	
P	Yes \Box No Has the patient by \rightarrow Please indicate the plane in	een treated with corticosteroids? ne corticosteroid name:		
	Yes I No Did the patient sh	ow improvement after 48 hours o	t corticosteroids?	



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G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.					
Please indicate the toxicity, (check all that ap	ply):				
Elevated serum creatinine/acute renal failure					
Please indicate the severity of the disease					
Severe (creatinine greater than 3 tim	/				
Life-threatening (creatinine greater th	nan 6 times baseline; dialysis indicated)				
□ None of the above					
Yes No Has the patient been tre	eated with corticosteroids? me and length of therapy: Name:	Longth: 🗖 Loo	a than 1 weak \Box 1 weak or greater		
	remain greater than 2 to 3 times above bas				
☐ Inflammatory arthritis					
	efractory or severe disease? 🔲 refractory o	disease 🔲 severe disease			
	g to corticosteroids or anti-inflammatory ag		ts 🔲 corticosteroids		
Please indicate the severity of the disea					
	eated with corticosteroids for pneumonitis?				
Please indicate the co	provement after 48 hours of corticosteroids	2			
Juvenile Idiopathic Arthritis (Juvenile Rheum		•			
Please indicate the severity of the patient's disea					
☐ Yes ☐ No Does the patient have clinical		pathic arthritis (JRA)?			
Yes No Is there evidence that the dise					
Yes No Was treatment with Enbrel (eta					
	mented intolerance to Enbrel (etanercept)?				
Yes No Does the patient have a docur	mented contraindication to Enbrel (etanerce	ept)?			
Noninfectious Uveitis					
Yes No Was the treatment with cortico					
Please indicate the corticoster	oid name:				
│	nosuppressive drugs (e.g., azathioprine, cvo	closporine, or methotrexate) inef	fective?		
Yes No Does the patient have a docur	mented intolerance to corticosteroids or imr	nunosuppressive drugs?	70		
☐ Yes ☐ No Does the patient have a docur	e patient has intolerance to: Corticostero		gs		
			e druas		
Please indicate the drug(s) the patient has contraindication to: Corticosteroids immunosuppressive drugs					
Please indicate the severity of the patient's disea					
Yes No Is there evidence that the dise					
Yes No Is there clinical documentation					
Yes No Is the patient a candidate for systemic therapy or phototherapy?					
Please select: ☐ phototherapy ☐ systemic therapy ☐ phototherapy and systemic therapy Please provide the patient's Psoriasis Area and Severity Index (PASI) score:					
Please indicate the percentage of body surface area affected by plaque psoriasis:%					
☐ Yes ☐ No Does the plaque psoriasis involve sensitive areas? <i>If yes</i> , please select: ☐ hands ☐ feet ☐ face ☐ genitals					
🖵 Yes 🔲 No 🛛 Was the trial with systemic conventional DMARD(s) (e.g., methotrexate, acetretin, or cyclosporine) ineffective?					
\square Yes \square No Was the trial with systemic conventional DMARD(s) not tolerated?					
Yes No Are systemic conventional DMARDs contraindicated?					
Please select: acetretin cyclosporine methotrexate mycophenolate None of the above					
□ Yes □ No Was the trial with phototherapy ineffective?					
☐ Yes ☐ No Was the trial with phototherapy not tolerated? ☐ Yes ☐ No Is phototherapy contraindicated?					
Please check all that apply: Psoralens (methoxsalen, trioxsalen) with UVA light (PUVA)					
Please check all that apply: Provident (methoxsalen, thoxsalen) with OVA light (POVA)					
UVB (standard or narrow-band)					
□ None of the above					
	 rial: 🔲 Less than 1 month 🛛 1 month 🗌	2 months 3 months or area	ater		
g					



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G. CLINICAL INFORMATION (continued) – R	Required clinical information musi	be completed in its <u>entirety</u> for a	all precertification requests.	
Psoriatic Arthritis				
Yes No Is there evidence that the disc				
Yes ☐ No Does the patient have axial p ☐ Yes ☐ No Was the treat		al anti inflommatory druga (NSA)	IDa) inoffactiva?	
	de the names and length of treat			
NSAID #1:	to the number and length of treat			
NSAID #2:				
☐ Yes ☐ No Does the patient have non-a				
		entation, defined as severe disal	bility at onset with erosive disease involving	
multiple joints				
$ \longrightarrow \Box \operatorname{Yes} \sqcup $	No Was the treatment with meth		rated or contraindicated?	
		tment with methotrexate not tole select: not tolerated cont	traindicated	
		\square No Was treatment with and	other conventional DMARD ineffective?	
			ophosphamide 🔲 cyclosporine	
		· _ ·	oxychloroquine 🔲 leflunomide	
		🗌 sulfa	asalazine 🛛 Other, please explain:	
Pyoderma Gangrenosum				
Yes No Does the patient have a docu				
Reactive Arthritis (Reiter's syndrome) or Infla				
Please select which applies to the patient: re		e) [] Inflammatory bowel disea	ase arthritis (enteropathic arthritis)	
\square res \square No Was the treatment with metric		ated?		
	ient have a contraindication to m			
Yes No Was the treatment with sulfas				
└───> □ Yes □ No Was the trea	tment with sulfasalazine not toler	ated?		
	ient have a contraindication to su			
Yes No Was the treatment with non-s				
Yes No Was the treatment with non-steroidal anti-inflammatory drugs (NSAIDs) not tolerated? Yes No Does the patient have a contraindication to non-steroidal anti-inflammatory drugs (NSAIDs)?				
· · · · · · · · · · · · · · · · · · ·			ugs (INSAIDS)?	
Retinal Vasculitis				
Yes No Was treatment with a conven	tional DMARD ineffective?			
└───> □ Yes □ No Was treatme	nt with a conventional DMARD n	ot tolerated or contraindicated?	not tolerated contraindicated	
Rheumatoid Arthritis		_		
Please indicate the severity of the patient's rheu		derate 🔲 severe		
Yes No Is there evidence that the disc		with methotrovete?		
Yes ☐ No Will the patient be using Rem ☐ Yes ☐ Yes ☐ No Was treatme		with metholiexate?		
		xate not tolerated or contraindica	ated? 🔲 not tolerated 🛛 contraindicated	
			IARD (other than methotrexate) ineffective?	
			hloroquine 🔲 leflunomide 🗋 sulfasalazine	
Sarcoidosis				
☐ Yes ☐ No Is the disease refractory to co	rticosteroids?			



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□ Yes No Is there evider □ Yes No Is the patient r □ Yes No Yes No Please indication □ Yes No Was treatment □ Yes □ No Yes No	active fulminant ulcerative colitis? the patient's ulcerative colitis:	moderate severe costeroids (e.g., hydrocortisone, munosuppression with corticost Dose: IV Dose: athioprine, 6-mercaptopurine) ind int agent (e.g., azathioprine, 6-m	methylprednisolone, prednisone)? eroids (e.g., hydrocortisone,
☐ Yes ☐ No Was treatmen		alsalazide, mesalamine, sulfasa cid agents (e.g., balsalazide, me ntraindicated cal, Delzicol, Lialda, Pentasa, Re please explain:	salamine, sulfasalazine) owasa, Canasa (mesalamine)
Please indicate the length of time on Remicade (,		
□ Yes No Is there clinical documentation □ Yes No Is there clinical documentation □ Yes No Is there clinical documentation □ Yes No Does the patient have any risk □ Yes No Has the patient □ Yes No Has the patient received Reminer □ Yes No Does the patient received Reminer □ Yes No Does the patient received Reminer	used concomitantly with apremilast, tofacities supporting disease stability? supporting disease improvement? factors for TB? it had a TB test within the past year? apply): PPD test interferon-gamma he results of the TB test: positive in cade (infliximab) within the past 6 months? ent have a documented severe and/or pote	nib, or other biologic DMARDs (a assay (IGRA)	vent that occurred during or following
For Crohn's disease, Juvenile idiopathic arthr	itis, Plaque psoriasis, and Rheumatoid a	arthritis, Ulcerative colitis only	/:
Please indicate the severity of the disease at bas	senne (pretreatment with Remicade (inflixin	nap)): 📋 mild 📋 moderate 🗋	J severe
H. ACKNOWLEDGEMENT			
Request Completed By (Signature Require	d):		Date: / /
Any person who knowingly files a request for a insurance company by providing materially finsurance act, which is a crime and subjects s	false information or conceals material i	information for the purpose c	

The plan may request additional information or clarification, if needed, to evaluate requests.