

Remicade® (infliximab) Injectable Medication Precertification Request

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(All fields must be completed and legible for precertification review.)

For other lines of business:
Please use other form.
Note: Remicade is preferred

FAX:

Note: Remicade is preferred for MA plans. Preferred status for MAPD plans varies based on

For Medicare Advantage Part B:

1-844-268-7263

PHONE: 1-866-503-0857

	☐ Start of treatment: Star☐ Continuation of therapy			,			indication	on. See section	n G below.
Precertification Requ		7. Buto or luc	t troutmont		Phone:		Fax:		
A. PATIENT INFORM	IATION								
First Name:				Last	Name:				
Address:				City			State:	ZIP:	
Home Phone:		Work	Phone:			Cell Phone:	ı		
DOB:	Allergies:					Email:			
	lbs or	kgs	Height:		inches or		<u> </u>		
B. INSURANCE INFO		ngo	rioight.			OTTIC			
			Does patient have	other	coverage?	П No			
Group #:					C				
Insured:			Insured:						
C. PRESCRIBER INF	ORMATION								
First Name:			Last Name:			(Check O	ne): 🔲 M.D.	. 🔲 D.O. 🔲 N.I	P. 🗌 P.A.
Address:					City:		State:	ZIP:	
Phone:	Fax:		St Lic #:		NPI #:	DEA #:		UPIN:	
Office Contact Name:	II.		1		1	Phone:		1	
D. DISPENSING PRO	OVIDER/ADMINISTRATIO	N INFORMA	ATION						
Place of Administra	tion:			Disp	ensing Provider/P	harmacy:			
☐ Self-administered	d Physician's Office	:			Physician's Office	☐ Retail P	harmacy		
	on Center Phone:				Specialty Pharmacy	y 🔲 Mail Ord	ler		
	ne:				Other:				
Agency Nar	enter Phone:			Nam	ne:				
	ode(s) (CPT):				ress:				
Address:									
City:	State	: Z	IP:	-	ne:				
	Fax:								
	PIN: _						PIN:		
NPI:				NPI:					
	MATION – Please select								
	icade (infliximab) Dose:				-		НСР	CS Code:	
F. DIAGNOSIS INFO	RMATION - Please indica	te primary I	CD Code and specify	any c	other where applicabl	le.			
Primary ICD Code:		Second	ary ICD Code:			_ Other ICD C	ode:		
G. CLINICAL INFOR	MATION – Required clinic	al informatio	n must be completed	l in its	entirety for all prece	rtification reque	ests.		
For Initiation Reques	sts (clinical documentation	on required	for all requests):		-	-			
	, Remicade, and Simponi								
ulcerative colitis and Enbrel, Humira, Rinvoq, Skyrizi, and Xeljanz/Xeljanz XR are preferred for other indications. Preferred products vary based on indication.									
Yes No Has the patient had prior therapy with Remicade (infliximab) within the last 365 days?									
☐ Yes ☐ No Has the patient had a trial, intolerance, or contraindication to any of the following? (select all that apply) ☐ Enbrel (etanercept) ☐ Humira (adalimumab) ☐ Rinvoq (upadacitinib) ☐ Skyrizi (risankizumab-rzaa) ☐ Xeljanz/Xeljanz XR (tofacitinib)									
Please explain if there are any other medical reason(s) that the patient cannot use any of the following preferred products when indicated for the patient's									
diagnosis (select all that apply) □ Enbrel (etanercept) □ Humira (adalimumab) □ Rinvoq (upadacitinib) □ Skyrizi (risankizumab-rzaa) □ Xeljanz/Xeljanz XR (tofacitinib)									
□ E	Enbrel (etanercept) L H	umira (adalir	numab) L Rinvoq ((upada	acitinib) 🔲 Skyrizi (risankizumab-r	zaa) 📙 Xel	janz/Xeljanz XF	R (tofacitinib)
☐ Yes ☐ No Will Remicade (infliximab) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)? ☐ Yes ☐ No Has the patient been tested for TB with a PPD test, interferon-release assay (IGRAs) or chest x-ray within 6 months of initiation a biologic therapy? ☐ (check all that apply): ☐ PPD test ☐ interferon-gamma assay (IGRA) ☐ chest x-ray ☐ PPD test ☐ positive ☐ negative ☐ unknown									
•	ositive, Does the patient h I tent TB, ☐ Yes ☐ No		_			Remicade (infl	iximab\?		
II Ia	103 _ 100	ıb iicali	De started belor	- 11 III	addit of thorapy with	Translate (IIII			

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Note: Remicade is preferred for MA plans. Preferred status for MAPD plans varies based on indication. See section G.

Patient First N	lame	Patient Last Name	Patient Phone	Patient DOB		
G CLINICAL	INFORMATION (continued) - R	l equired clinical information must be co	ompleted in its entirety for all n	recertification requests		
			ompieted in its <u>entirety</u> for all p	recertification requests.		
Please select Yes N Yes N	Ankylosing Spondylitis and Other Spondyloarthropathies Please select which of the following applies to the patient: Ankylosing spondylitis Other spondyloarthropathy I sthere evidence that the disease is active? I yes No Is there evidence of inflammatory disease? I yes No Has the patient had an ineffective response to two or more non-steroidal anti-inflammatory drugs (NSAIDs)?					
Yes Lik	Please provide the names and NSAID #1:	tive response to two or more non-ster d length of treatment:		(NOAIDS)?		
Behcet's Dise						
Yes N	→ Please indicate: ☐ corticoste	rticosteroids or immunosuppressive d roids				
Behcet's Uve						
	lo Is the disease refractory?					
	neous/Pulmonary Sarcoidosis		4-0			
	Please provide the daily dose	-				
Yes N	lo Has the patient remained sym → Please select: ☐ azathioprine	ptomatic despite treatment with immu ☐ cyclophosphamide ☐ methotrexa	nosuppressants? te			
Crohn's Dise						
		osis of fistulizing Crohn's disease? patient has been diagnosed with fistuli	zing Crohn's disease:			
☐ Yes ☐ N	lo Does the patient have a diagn	osis of Crohn's disease?				
		the patient's disease: mild mo				
		ent have a documented diagnosis of a ct all signs/symptoms that apply:	ictive Cronn's disease?			
	abdomina abdomina	al pain arthritis bleeding				
		on perianal disease spondylitis				
	Yes No Have the Cro	hn's disease symptoms remained acti	ve despite treatment with 6-me	ercaptopurine, azathioprine,		
		k all medications that apply:	captopurine azathioprine			
	☐ corticoste			prednisolone 🗌 Other:		
Hidradenitis		ativa: Hurley stage I (mild disease)	□ Hurlov stage II (mode	orato diagona)		
	lo Has the patient completed a tr	☐ Hurley stage III (severe disea		erate disease)		
├────────────────────────────────────						
Yes No Was the treatment with antibiotics ineffective? Please indicate the duration of the medication trial: Less than 1 month 1 month 2 months 3 months (90 days) or greater						
	ckpoint Inhibitor-Induced Toxicit e therapy used:			days) of ground		
CTLA-4	e therapy useu.					
	ect drug: ipilimumab Other:					
Please select drug: 🗌 nivolumab 🗎 pembrolizumab 🔲 Other:						
☐ PD-L1 Please select drug: ☐ atezolizumab ☐ avelumab ☐ durvalumab ☐ Other:						
☐ Other Please explain:						
	lo Do the immune checkpoint inh	nibitor-induced toxicities persist despite ab, ipilimumab, nivolumab, pembrolizu		neckpoint inhibitors that target CTLA-4 or		
Please indicate the toxicity, (check all that apply):						
☐ Cardiac	Please select: arrhythmias	eckpoint inhibitor-induced cardiac toxi impaired ventricular function	ocarditis			
☐ Colitis		immune checkpoint inhibitor-induced				
Please indicate which of the following symptoms the patient exhibits: ☐ 7 or more stools per day over baseline ☐ ileus ☐ fever ☐ None ☐ Yes ☐ No Has the patient been treated with corticosteroids?						
	Please indicate the	· · · · · · · · · · · · · · · · · · ·				
	☐ Yes ☐ No Did the patient sh	ow improvement after 48 hours of cor	ticosteroids?			



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G. CLINICAL INFORMATION (continued) – R	equired clinical information must be comple	tod in its antiroty for all proceptif	ication requests		
Please indicate the toxicity, (check all that ap		led in its <u>entirety</u> for all precentil	ication requests.		
☐ Elevated serum creatinine/acute renal failure					
Please indicate the severity of the disease					
☐ Severe (creatinine greater than 3 times					
☐ Life-threatening (creatinine greater the	nan 6 times baseline; dialysis indicated)				
☐ None of the above					
Yes No Has the patient been tre					
Please indicate the na	me and length of therapy: Name: remain greater than 2 to 3 times above base	Length:Less	s than 1 week		
☐ Inflammatory arthritis	Terriain greater than 2 to 5 times above bas	cline after 1 week of treatment v	viii cornoosteroids:		
☐ Yes ☐ No Does the patient have r	efractory or severe disease? refractory of	lisease 🗌 severe disease			
	g to corticosteroids or anti-inflammatory age	ents? 🔲 anti-inflammatory agen	its Corticosteroids		
Pneumonitis	and Dunille Duna danata Danisana				
Please indicate the severity of the disea	se: mild moderate severe eated with corticosteroids for pneumonitis?				
Please indicate the col	rticosteroid name:				
☐ Yes ☐ No Did the patient show im	provement after 48 hours of corticosteroids	?			
Juvenile Idiopathic Arthritis (Juvenile Rheum					
Please indicate the severity of the patient's disease		andhin andhidir (IDA)2			
Yes No Does the patient have clinical Yes No Is there evidence that the dise		Dathic arthritis (JRA)?			
Yes No Was treatment with Enbrel (et					
Yes No Does the patient have a docur					
Yes No Does the patient have a docur		pt)?			
Noninfectious Uveitis					
Yes No Was the treatment with cortico					
Please indicate the corticoster	oid name:				
☐ Yes ☐ No Was the treatment with immur	nosuppressive drugs (e.g., azathioprine, cyc	closporine, or methotrexate) inef	fective?		
Please provide the name:		<u> </u>			
□ Ves □ No. Does the nationt have a document	mented intolerance to corticosteroids or imn	nunosunnressive drugs?			
Yes No Does the patient have a docur	e patient has intolerance to: Corticosteroids of infine	ds immunosuppressive dru	qs		
Yes No Does the patient have a docur	mented contraindication to corticosteroids of	r immunosuppressive drugs?			
	e patient has contraindication to: 🗌 corticos	steroids	e drugs		
Plaque Psoriasis Please indicate the severity of the patient's disea	ase: □ mild □ moderate □ severe				
Yes No Is there evidence that the dise					
Yes No Is there clinical documentation of chronic disease?					
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?					
☐ Yes ☐ 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)					
UVB with coal tar or dithranol					
│ UVB (standard or narrow-band) │ Home UVB					
☐ None of the above					
Please indicate the length of trial: Less than 1 month					
Ficase indicate the length of the	iai. 🔲 LESS IIIAII I IIIOIIIII 🔲 I IIIOIIIII 📙	izmontas 🗀 smontas or grea	alGi		

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G. CLINICAL INFORMATION (continued) – R	equired clinical information must be comp	leted in its <u>entirety</u> for all precer	tification requests.		
Psoriatic Arthritis					
☐ Yes ☐ No Is there evidence that the dise	ease is active?				
Yes No Does the patient have axial ps					
	ment with 2 or more non-steroidal anti-infl	ammatory drugs (NSAIDs) ineff	ective?		
	e the names and length of treatment:				
NSAID #2:					
Yes No Does the patient have non-ax	ial psoriatic arthritis?				
	ent have severe disease at presentation,	defined as severe disability at o	nset with erosive disease involving		
multiple joints		<i></i>			
→ ☐ Yes ☐ I	No Was the treatment with methotrexate i				
	→ Yes No Was treatment with	n methotrexate not tolerated or	contraindicated?		
	Please select:	not tolerated	led		
		Was treatment with another con			
		Please select: Cyclophospha	oquine		
			Other, please explain:		
Pyoderma Gangrenosum		☐ SullaSalaZille	U Other, please explain.		
Yes No Does the patient have a docur	mented diagnosis of refractory pyoderma	rangrenosum?			
Reactive Arthritis (Reiter's syndrome) or Infla	, , ,	, ,			
Please select which applies to the patient:			itis (enteronathic arthritis)		
Yes No Was the treatment with metho		laminatory bower disease artific	ius (enteropatric artificis)		
	ment with methotrexate not tolerated?				
	ent have a contraindication to methotrexa	te?			
Yes No Was the treatment with sulfast					
	ment with sulfasalazine not tolerated?				
	ent have a contraindication to sulfasalazir	e?			
Yes No Was the treatment with non-steroidal anti-inflammatory drugs (NSAIDs) ineffective?					
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)?					
			,		
Retinal Vasculitis					
☐ Yes ☐ No Was treatment with a convent					
☐ Yes ☐ No Was treatmen	nt with a conventional DMARD not tolerate	ed or contraindicated? 🗌 not to	lerated		
Rheumatoid Arthritis					
Please indicate the severity of the patient's rheumatoid arthritis: mild moderate severe					
Yes No Is there evidence that the disease is active?					
Yes No Will the patient be using Remicade (infliximab) in combination with methotrexate?					
Yes No Was treatment with methotrexate ineffective?					
Yes No Was treatment with methotrexate not tolerated or contraindicated? In not tolerated contraindicated					
Yes No Was treatment with another conventional DMARD (other than methotrexate) ineffective?					
	─────────────────────────────────────	hioprine 🔲 hydroxychloroquir	ne 🗌 leflunomide 🔲 sulfasalazine		
Sarcoidosis					
☐ Yes ☐ No Is the disease refractory to cor	ticosteroids?				

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G CLINICAL INFORMATION (continued) - R	equired clinical information must be comp	leted in its entirety for all pred	pertification requests		
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Ves					
Please select: ☐ not tolerated ☐ contraindicated → Please select: ☐ Colazal (balsalazide) ☐ Ariso, Asacal, Delzicol, Lialda, Pentasa, Rowasa, Canasa (mesalamine) ☐ Azulfidine (sulfasalazine) ☐ Other, please explain: → Please select the symptoms the patient exhibit: ☐ more than 10 stools per day ☐ continuous bleeding ☐ abdominal pain ☐ distension ☐ acute, severe toxic symptoms, including fever and anorexia					
For Continuation of Therapy (clinical docume Please indicate the length of time on Remicade					
Yes No Is this continuation request a result of the patient receiving samples of Remicade (infliximab)? Yes No Will Remicade (infliximab) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)? Is there clinical documentation supporting disease stability? Is there clinical documentation supporting disease improvement? Yes No Does the patient have any risk factors for TB? Yes No Has the patient had a TB test within the past year? Yes No Please enter the results of the TB test: positive negative unknown Has the patient received Remicade (infliximab) within the past 6 months? Yes No Does the patient have a documented severe and/or potentially life-threatening adverse event that occurred during or following the previous infusion? Yes No Could the adverse reaction be managed through pre-medication in the home or office setting?					
For Crohn's disease, Juvenile idiopathic arthritis, Plaque psoriasis, and Rheumatoid arthritis, Ulcerative colitis only: Please indicate the severity of the disease at baseline (pretreatment with Remicade (infliximab)): mid moderate severe					
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Require	ed):		Date:/ /		
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.					

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