

## Remicade® (infliximab) Injectable **Medication Precertification Request**

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

plans. Preferred status for MAPD plans varies based on indication. ☐ Start of treatment: Start date \_\_\_\_/ Please indicate: See section G below. Continuation of therapy: Date of last treatment \_\_\_\_/\_\_/ Phone: Precertification Requested By: Fax: A. PATIENT INFORMATION First Name: Last Name: City: Address: State: ZIP: Home Phone: Work Phone: Cell Phone: DOB: Allergies: Email: Current Weight: Height: \_\_\_\_\_ inches or \_\_ \_\_\_\_\_ lbs or \_\_\_\_\_ kgs B. INSURANCE INFORMATION Member ID #: Does patient have other coverage? ☐ Yes ☐ No If yes, provide ID#: \_\_\_\_\_ Carrier Name: \_\_\_\_ Group #: Insured: C. PRESCRIBER INFORMATION First Name: Last Name: (Check One): ☐ M.D. ☐ D.O. ☐ N.P. ☐ P.A. State: ZIP: Address: City: Phone: NPI#: Fax: St Lic #: DEA #: UPIN: Provider Email: Office Contact Name: Phone: D. DISPENSING PROVIDER/ADMINISTRATION INFORMATION Place of Administration: **Dispensing Provider/Pharmacy:** ☐ Self-administered ☐ Physician's Office ☐ Physician's Office ☐ Retail Pharmacy Outpatient Infusion Center Phone: ☐ Specialty Pharmacy ☐ Mail Order Center Name: \_\_\_\_ Other: Home Infusion Center Phone: Agency Name: \_\_\_ Address: Administration code(s) (CPT): City: \_\_\_\_\_ State: \_\_\_\_ ZIP: \_\_\_\_ Phone: \_\_\_\_\_ Fax: \_\_\_\_\_ **TIN:** \_\_\_\_\_ PIN: \_\_\_\_ TIN: \_\_\_\_\_ PIN: NPI: \_\_\_\_\_ **E. PRODUCT INFORMATION** – Please select the medication being requested Request is for: Remicade (infliximab) Dose: HCPCS Code: \_\_\_\_\_ Frequency: F. DIAGNOSIS INFORMATION – Please indicate primary ICD Code and specify any other where applicable. Primary ICD Code: Secondary ICD Code: Other ICD Code: G. CLINICAL INFORMATION - Required clinical information must be completed in its entirety for all precertification requests. For Initiation Requests (clinical documentation required for all requests): Note: Remicade, Inflectra, Entyvio, Simponi Aria and unbranded infliximab are the preferred products for MA plans. For MAPD plans, Remicade, Inflectra, Entyvio, and unbranded infliximab are preferred for ulcerative colitis and Enbrel, Humira, Kevzara, Otezla, Rinvoq, Skyrizi, Stelara, Tremfya 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 and failure, intolerance, or contraindication to any of the following? (select all that apply) ☐ Enbrel (etanercept) ☐ Humira (adalimumab) ☐ Kevzara (sarilumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib) ☐ Skyrizi (risankizumab-rzaa) ☐ Stelara (ustekinumab) ☐ Tremfya (guselkumab) ☐ 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) ☐ Kevzara (sarilumab) ☐ Otezla (apremilast) ☐ Rinvoq (upadacitinib) ☐ Skyrizi (risankizumab-rzaa) ☐ Stelara (ustekinumab) ☐ Tremfya (guselkumab) ☐ Xeljanz/Xeljanz XR (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 Please enter results of the TB test: positive negative unknown 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)?

Continued on next page

For Medicare Advantage Part B:

FAX: 1-844-268-7263

Please use other form.

For other lines of business:

Phone: 1-866-503-0857 (TTY: 711)

Note: Remicade is preferred for MA



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) – R	l equired clinical information must be comple	ted in its entirety for all pr	ecertification requests		
		rod iir ko <u>oriaroty</u> for ali pr	soortiiisation roquoeto.		
Ankylosing Spondylitis and Other Spondyloarthropathies  Please select which of the following applies to the patient: ☐ Ankylosing spondylitis ☐ Other spondyloarthropathy ☐ Yes ☐ No ☐ Is there evidence that the disease is active? ☐ Yes ☐ No ☐ Is there evidence of inflammatory disease? ☐ Yes ☐ No ☐ Has the patient had an ineffective response to two or more non-steroidal anti-inflammatory drugs (NSAIDs)?  Please provide the names and length of treatment:					
NSAID #1: NSAID #2:		_			
Behcet's Disease		_			
Please provide the name of d	rticosteroids or immunosuppressive drugs? roids				
Behcet's Uveitis					
Yes No Is the disease refractory?					
Chronic Cutaneous/Pulmonary Sarcoidosis					
Yes No Has the patient remained sym  Please provide the daily dose					
☐ Yes ☐ No ☐ Has the patient remained sym  Please select: ☐ azathioprine			ı:		
Crohn's Disease	and of first divine One built divine on O				
Yes No Does the patient have a diagn  Please indicate how long the p		Crohn's disease:			
☐ Yes ☐ No Does the patient have a diagn	osis of Crohn's disease?				
Please indicate the severity of					
	ent have a documented diagnosis of active ct all signs/symptoms that apply:	Cronn's disease?			
□ abdomina	al pain  arthritis  bleeding  diarrh	nea 🔲 internal fistulae 「	☐ intestinal obstruction		
=	on 🗌 perianal disease 🔲 spondylitis 🔲	=			
	hn's disease symptoms remained active de	spite treatment with 6-me	rcaptopurine, azathioprine,		
	oids? k all medications that apply:		orednisolone □ Other		
Hidradenitis Suppurativa	product demany.   production in the production i	yarooorasone 🖂 mearyip	Tearniscienc 🗆 euror.		
Please indicate the stage of hidradenitis suppura	☐ Hurley stage III (severe disease)	☐ Hurley stage II (mode	rate disease)		
Yes No Has the patient completed a tr	เลเ or antibiotics ? ent have a contraindication to oral antibiotic	rs?			
Yes No Was the treat					
	te the duration of the medication trial: $\Box$ Le	ess than 1 month 🔲 1 mo	onth		
Laurence Observation land land land land land land land lan		months 3 months (90	days) or greater		
Immune Checkpoint Inhibitor-Induced Toxicit Please indicate therapy used:	ies				
☐ CTLA-4					
Please select drug: ☐ ipilimumab ☐ Other:					
☐ PD-1 Please select drug: ☐ nivolumab ☐ pembr	olizumab 🔲 Other:	<u> </u>			
│					
Other Please explain:	iumab 🗀 durvaiumab 🗀 Omer				
☐ Yes ☐ No Do the immune checkpoint inh	nibitor-induced toxicities persist despite disc ab, ipilimumab, nivolumab, pembrolizumab)		eckpoint inhibitors that target CTLA-4 or		
Please indicate the toxicity, (check all that ap					
	eckpoint inhibitor-induced cardiac toxicities				
Please select: ☐ arrhythmias ☐ impaired ventricular function ☐ myocarditis ☐ pericarditis ☐ Colitis Please indicate the severity of the immune checkpoint inhibitor-induced colitis. ☐ mild ☐ moderate ☐ severe					
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?					
	een treated with corticosteroids'? ne corticosteroid name:				
	ow improvement after 48 hours of corticost	eroids?			



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FAX: <u>1-844-268-7263</u>

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See section G.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) - Re	equired clinical information must be comple	ited in its entirety for all precertif	ication requests			
		ted in its <u>entirety</u> for all precentin	ication requests.			
Please indicate the toxicity, (check all that apply):						
Elevated serum creatinine/acute renal failure  Please indicate the severity of the disease:						
☐ Severe (creatinine greater than 3 time						
☐ Life-threatening (creatinine greater that						
☐ None of the above						
☐ Yes ☐ No Has the patient been treation	ated with corticosteroids?	<u>_</u>	_			
Please indicate the nam	ne and length of therapy: Name: remain greater than 2 to 3 times above base	Length: Les	s than 1 week			
☐ Yes ☐ No Did the creatinine level re	emain greater than 2 to 3 times above base	eline after i week of treatment v	with corticosteroids?			
1 <del>-</del> _ , _	efractory or severe disease?   refractory of	disease □ severe disease				
	g to corticosteroids or anti-inflammatory age		its Corticosteroids			
☐ Pneumonitis						
Please indicate the severity of the diseas						
Please indicate the cort	ated with corticosteroids for pneumonitis?					
	provement after 48 hours of corticosteroids	?				
Juvenile Idiopathic Arthritis (Juvenile Rheuma	atoid Arthritis)					
Please indicate the severity of the patient's disea						
Yes No Does the patient have clinical of		pathic arthritis (JRA)?				
☐ Yes ☐ No Is there evidence that the disea	ase is active?					
Naminfortions Unaitio						
Noninfectious Uveitis	steroids ineffective?					
Please indicate the corticostero						
Yes No Was the treatment with immuno	osuppressive drugs (e.g., azathioprine, cyc	losporine, or methotrexate) inef	fective?			
Please provide the hame.		<del></del>				
☐ Yes ☐ No Does the patient have a docum	nented intolerance to corticosteroids or imm	nunosuppressive drugs?				
Please indicate the drug(s) the patient has intolerance to:   criticosteroids immunosuppressive drugs						
☐ Yes ☐ No Does the patient have a documented contraindication to corticosteroids or immunosuppressive drugs?  Please indicate the drug(s) the patient has contraindication to: ☐ corticosteroids ☐ immunosuppressive drugs						
Plaque Psoriasis	patient has contraindication to.   Gordoos	steroids 🖂 illillidilosuppressive	2 drugs			
Please indicate the severity of the patient's disea	se:  mild moderate severe					
☐ Yes ☐ No Is there evidence that the disea						
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 convenional binards contraindicated?  Please select: acetretin convenional binards contraindicated?  Please select: acetretin convenional binards contraindicated?						
☐ 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						
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FAX: 1-844-268-7263

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) - F	equired clinical information must	be completed in its entirety for all pro-	ecertification requests.		
Psoriatic Arthritis					
☐ Yes ☐ No Is there evidence that the disc	ease is active?				
Yes No Does the patient have axial p					
		al anti-inflammatory drugs (NSAIDs) i	neffective?		
The state of the s	de the names and length of treatr				
NSAID #2: _					
Yes No Does the patient have <b>non-a</b>		antation defined as assume dischiller.	at amount with amount of all amount investigation		
multiple joints		entation, defined as severe disability	at onset with erosive disease involving		
' '	No Was the treatment with meth	otrexate ineffective?			
7 2 1		tment with methotrexate not tolerated	d or contraindicated?		
		select:  not tolerated contraine			
	☐ Yes	☐ No Was treatment with another	conventional DMARD ineffective?		
		——→ Please select: ☐ cyclopho	, .		
			hloroquine 🗌 leflunomide		
		☐ sulfasala:	zine		
Pyoderma Gangrenosum					
Yes No Does the patient have a docu					
Reactive Arthritis (Reiter's syndrome) or Infla					
Please select which applies to the patient:		e)	irthritis (enteropathic arthritis)		
Yes ☐ No Was the treatment with method Yes ☐ Yes ☐ No Was the treatment with method Yes ☐ Yes ☐ No Was the treatment with method Yes ☐ Yes ☐ No Was the treatment with method Yes ☐		ratad?			
	ient have a contraindication to m				
Yes No Was the treatment with sulfas		ethotrexate:			
Yes No Was the trea		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 pat	ent have a contraindication to no	n-steroidal anti-inflammatory drugs (	NSAIDs)?		
· ·					
Retinal Vasculitis					
Yes No Was treatment with a conven		at talayatad ay asytysiy disatad2 🗆 y	et televeted		
· — —	ni with a conventional diviard n	ot tolerated or contraindicated? 🗌 no	of tolerated		
Rheumatoid Arthritis  Please indicate the severity of the nationt's rheumatoid arthritis	ımatoid arthritis: □ mild □ mo	derate $\square$ severe			
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? ☐ not tolerated ☐ contraindicated					
Yes No Was treatment with another conventional DMARD (other than methotrexate) ineffective?					
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Sarcoidosis

 $\square$  Yes  $\square$  No Is the disease refractory to corticosteroids?



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued)	<ul> <li>Required clinical information must</li> </ul>	be completed in its entirety for all	precertification requests.		
Ulcerative Colitis	with active fulminant ulcerative colitic	2			
Yes No Is the patient hospitalized  Please indicate the severi	with active ruiminant dicerative collis:	s? I mild □ moderate □ severe			
	vidence that the disease is active?				
☐ Yes ☐ No Is the pati	ent refractory to immunosuppression	with corticosteroids (e.g., hydroc	ortisone, methylprednisolone, prednisone)?		
☐ ☐ ☐ Yes			corticosteroids (e.g., hydrocortisone,		
	methylprednisolone, prednisone)?  Name and dose: Name: Dose: Dose:				
	Please indicate the route:				
	r reads indicate the reats.	, e.a			
Name a	nd dose: Name:	Dose:			
Please	indicate the route:  ☐ Oral ☐ IV				
□ Voc □ No. Wee tree	tment with immunosuppressant ager	et (o.g. ezethioprine 6 mercenten	vurina) inoffactiva?		
			prine, 6-mercaptopurine) not tolerated		
	or contraindicated?	эсарр, сосалт аделт (с.д., адаптер			
	Please select: not tolera				
Please s	select: 🗌 6-mercaptopurine 🔲 azat	hioprine			
☐ Yes ☐ No. Was trea	tment with 5-aminosalicylic acid age	nts (e.g. balsalazide mesalamine	e sulfasalazine) ineffective?		
	☐ No Was treatment with 5-amino				
	not tolerated or contraindica		·		
Pleases	Please select: not tolera		ntasa, Rowasa, Canasa (mesalamine)		
/ I loade o			masa, nowasa, Ganasa (mesalamine)		
	,				
Please select the sympto	ms the patient exhibit: more than				
		acute, severe toxic symptoms	s, including fever and anorexia		
For Continuation of Therapy (clinical doc		<u>ts):</u>			
Please indicate the length of time on Remica  ☐ Yes ☐ No Is this continuation reques		nnles of Reminede (inflivimen)?			
☐ Yes ☐ No Will Remicade (infliximab)			MARDs (e.g., adalimumah, certolizumah)?		
☐ Yes ☐ No Is there clinical document		dot, toldottillo, of other biologic bi	vii (125 (e.g., addiimamas, sertolizamas):		
Yes No 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?					
Check all that apply): ☐ PPD test ☐ interferon-gamma assay (IGRA) ☐ chest x-ray  Please enter the results of the TB test: ☐ positive ☐ negative ☐ unknown					
☐ Yes ☐ No Has the patient received F					
Yes No Does the	patient have a documented severe a		dverse event that occurred during or following		
	ous infusion?		in the beauty on efficiency		
☐ 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)):   mild moderate severe					
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Req	uired):		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.