

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

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

For Ohio MMP: 1-855-734-9389 FAX: PHONE: 1-855-364-0974

For other lines of business: Please use other form.

Note: Remicade is preferred for MA plans. Preferred status for

Please indicate:		tart date <u>/</u>			on review.)		MAPD	plans varies b tion. See section	ased on
Precertification	Requested By:				Phone:		Fax	::	
A. PATIENT INFO	ORMATION								
First Name:				Last Name:					
Address:				City:			State:	ZIP:	
Home Phone:		Work F	Phone:			Cell Phone:		I	
DOB:	Allergies:	1				Email:			
Current Weight:	lbs or	kgs	Height:	inc	hes or	cms			
B. INSURANCE									
Member ID #:			Does patient have	other coverage	? 🗌 Yes	🗌 No			
			If yes, provide ID#:		Ca	arrier Name:			
nsured:			Insured:						
C. PRESCRIBER									
First Name:			Last Name:			(Check Or	<i>e):</i> 🗌 M.[D. 🗌 D.O. 🗌 N	I.P. 🗌 P.A.
Address:				City:			State:	ZIP:	
Phone:	Fax:		St Lic #:	NPI #:		DEA #:		UPIN:	
Office Contact Na	me:					Phone:		·	
D. DISPENSING	PROVIDER/ADMINISTRAT	ION INFORMAT	ΓΙΟΝ						
Place of Admini	stration:			Dispensing F	Provider/Pl	harmacy:			
Self-administ	ered 🛛 🗌 Physician's Off	ice		Physician	i's Office	Retail Ph	armacy		
Outpatient Infusion Center Phone:			Specialty Pharmacy 🔲 Mail Order						
	Name:			Other:					
Home Infusio				Name:					
	Name: n code(s) (CPT):								
Address:								ZIP:	
	Sta	ite: ZIF	D:						
-	Fax								
	PIN					F	PIN:		
NPI:				NPI:					
E. PRODUCT INI	FORMATION – Please sele	ct the medication	being requested						
Request is for: F	Remicade (infliximab) Dos	e:	Fred	quency:			нс	PCS Code:	
F. DIAGNOSIS IN	FORMATION – Please ind	icate primary ICI	D Code and specify	any other wher	e applicable	9.			
Primary ICD Code	:	Secondar	ry ICD Code:			Other ICD C	ode:		
	ORMATION – Required cli								
For Initiation Rec Note: Avsola, Ent ulcerative colitis a	uests (clinical documenta yvio, Remicade, and Simpon and Enbrel, Humira, Otezla, F Has the patient had prior th	ation required fo i Aria are the pre- kinvoq, Skyrizi, ar	or all requests): ferred products for nd Xeljanz/Xeljanz X	MA plans. For N R are preferred	IAPD plans, for other inc	Avsola, Entyvi	o, and Rem		
	Has the patient had a trial, Enbrel (etanercept)	intolerance, or co Humira (adalimu	ontraindication to ar	ny of the followi	ng? (select a		Skyrizi (ris	sankizumab-rza	ıa)
Please explain if t diagnosis (select	there are any other medical all that apply)	reason(s) that th	•	2	0.1	·			
	☐ Enbrel (etanercept) ☐ ☐ Xeljanz/Xeljanz XR (tofa	· ·	umab) 🔲 Otezla (a	apremilast) 🗌	Rinvoq (up	adacitinib) 🗌	Skyrizi (ris	sankizumab-rza	.a)
☐ Yes ☐ No	Will Remicade (infliximab) Has the patient been tested biologic therapy? (check all that apply): PP Please enter results of the <i>If positive,</i> Does the patier	l for TB with a Pf ⊃D test □ inter TB test:	PD test, interferon-r feron-gamma assa ive negative [release assay (I y (IGRA)	GRAs) or cl	-	-		:umab)?



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) – R	Required clinical information must be co	mpleted in its entirety for all pr	ecertification requests			
Ankylosing Spondylitis and Other Spondyloa						
Please select which of the following applies to the		Other spondyloarthropathy				
Yes No Is there evidence that the dise	ease is active?					
Yes No Is there evidence of inflamma	5					
Yes No Has the patient had an ineffect		oidal anti-inflammatory drugs (۱	NSAIDs)?			
Please provide the names an						
NSAID #1:						
Behcet's Disease						
☐ Yes ☐ No Is the disease refractory to co	orticosteroids or immunosuppressive d	ugs?				
	eroids 🔲 immunosuppressive drugs	0				
	Irug tried:					
Behcet's Uveitis						
☐ Yes ☐ No Is the disease refractory?						
Chronic Cutaneous/Pulmonary Sarcoidosis		4-0				
Yes No Has the patient remained syn		15 ?				
	ngng					
Yes No Has the patient remained syn						
	e 🗌 cyclophosphamide 🔲 methotrexa	te 🗌 Other, please explain:				
Crohn's Disease	and a finitulizing Oraba's disease?					
☐ Yes ☐ No Does the patient have a diagr	patient has been diagnosed with fistuli	zing Crohn's disease				
☐ Yes ☐ No Does the patient have a diagr						
\rightarrow Please indicate the severity o						
	ient have a documented diagnosis of a	ctive Crohn's disease?				
	ct all signs/symptoms that apply: al pain arthritis bleeding	diarrhea 🔲 internal fistulae [intestinal obstruction			
	on perianal disease spondylitis					
-	hn's disease symptoms remained activ	-				
or corticoster		·				
	ck all medications that apply: 🗌 6-mer		radniaciana 🗖 Other:			
Hidradenitis Suppurativa	eroids- please identify: 🗌 prednisone					
Please indicate the stage of hidradenitis suppur	ativa: 🗌 Hurley stage I (mild disease)	Hurley stage II (model	rate disease)			
	Hurley stage III (severe disea	se) 🔲 Unknown				
Yes No Has the patient completed a t						
\square Yes \square No Was the treat	ient have a contraindication to oral ant tment with antibiotics ineffective?	DIOLICS				
/	ate the duration of the medication trial:	Less than 1 month 1 mo	onth			
		$2 \text{ months} \square 3 \text{ months} (90)$				
Immune Checkpoint Inhibitor-Induced Toxici	ties					
Please indicate therapy used:						
Please select drug: ipilimumab Other						
DP-1						
Please select drug: 🗌 nivolumab 📋 pembrolizumab 🔲 Other:						
PD-L1 Please select drug: atezolizumab avelumab durvalumab Other:						
Please explain:						
Yes No Do the immune checkpoint inhibitor-induced toxicities persist despite discontinuation of immune checkpoint inhibitors that target CTLA-4 or PD-1/PD-L1 (e.g., atezolizumab, ipilimumab, nivolumab, pembrolizumab)?						
Please indicate the toxicity, (check all that apply):						
	eckpoint inhibitor-induced cardiac toxic					
] impaired ventricular function myo					
Colitis Please indicate the severity of the immune checkpoint inhibitor-induced colitis. I mild moderate severe Please indicate which of the following symptoms the patient exhibits: 7 or more stools per day over baseline lieus fever None						
☐ Yes ☐ No Has the patient been treated with corticosteroids?						
Please indicate t	he corticosteroid name:					
☐ Yes ☐ No Did the patient sh	now improvement after 48 hours of cor	icosteroids?				



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G. CLINICAL INFORMATION (continued) – R	aguirad alinical information must be comple	tod in its optiraty for all proportif	ention requests	
Please indicate the toxicity, (check all that ap		ted in its <u>entirety</u> for all precertin	cation requests.	
Elevated serum creatinine/acute renal failure				
Please indicate the severity of the disease				
Severe (creatinine greater than 3 tim				
Life-threatening (creatinine greater the second	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:	Length: Diles	s than 1 week . 🗖 1 week or greater	
\square Yes \square No Did the creatinine level	remain greater than 2 to 3 times above bas	eline after 1 week of treatment	vith corticosteroids?	
Inflammatory arthritis	-			
	efractory or severe disease? refractory of the second sec			
	g to corticosteroids or anti-inflammatory age	ents? 📋 anti-inflammatory agen	ts Corticosteroids	
Please indicate the severity of the disea	se: mild moderate			
Yes No Has the patient been tre	eated with corticosteroids for pneumonitis?			
\Box	ticosteroid name:			
	provement after 48 hours of corticosteroids	<i>?</i>		
Juvenile Idiopathic Arthritis (Juvenile Rheuma Please indicate the severity of the patient's disea	atold Arthritis) ase:			
☐ Yes ☐ No Does the patient have clinical		pathic arthritis (JRA)?		
Yes No Is there evidence that the dise		ζ, γ		
Yes No Was treatment with Enbrel (et				
Yes No Does the patient have a docur		-4/2		
	nented contraindication to Enbrel (etanerce	pt)?		
Noninfectious Uveitis	osteroids ineffective?			
	oid name:			
│ │ Yes │ No Was the treatment with immur	nosunnressive drugs (e.g. azathionrine, cyc	losporine or methotrevate) inef	fective?	
Please indicate the drug(s) the	e patient has intolerance to: Corticosteroids or imn		as	
☐ Yes ☐ No Does the patient have a docur			90	
	e patient has contraindication to: 🗌 corticos	steroids 🔲 immunosuppressive	e drugs	
Plaque Psoriasis Please indicate the severity of the patient's disea	ase [.] I mild I moderate I severe			
\Box Yes \Box No Is there evidence that the dise	ase is active?			
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? If yes, 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				
\longrightarrow Yes \square 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)				
□ None of the above				
Please indicate the length of the	rial: 🗌 Less than 1 month 📋 1 month 📋	2 months 🔲 3 months or grea	ater	



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G. CLINICAL INFORMATION (continued) – F	Required clinical information must	be completed in its <u>entirety</u> for all pre	ecertification requests.	
Psoriatic Arthritis				
Yes No Is there evidence that the dis				
Yes No Does the patient have axial p			a affa a thu a D	
	de the names and length of treatr	al anti-inflammatory drugs (NSAIDs) i	nellective?	
NSAID #1:	ue the hames and length of treat			
NSAID #1:				
Yes No Does the patient have non-a				
\square Yes \square No Does the pat	ient have severe disease at pres	entation, defined as severe disability	at onset with erosive disease involving	
multiple joint	s?		-	
\square Yes \square	No Was the treatment with meth			
		tment with methotrexate not tolerated		
		select: I not tolerated I contrained		
		□ No Was treatment with another		
		Please select: Cyclophos	nloroquine	
			zine Other, please explain:	
Pyoderma Gangrenosum				
Yes No Does the patient have a docu	mented diagnosis of refractory p	/oderma gangrenosum?		
Reactive Arthritis (Reiter's syndrome) or Infla	ammatory Bowel Disease Arthr	itis (Enteropathic Arthritis)		
Please select which applies to the patient:	-	,	rthritis (enteropathic arthritis)	
Yes INO Was the treatment with method	otrexate ineffective?	,		
└───> □ Yes □ No Was the trea				
	tient have a contraindication to me	ethotrexate?		
Yes No Was the treatment with sulfas		1.10		
	tient have a contraindication to su			
□ 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 conven			_	
	nt with a conventional DMARD n	ot tolerated or contraindicated? 🗌 no	ot tolerated	
Rheumatoid Arthritis				
Please indicate the severity of the patient's rheu		derate 📋 severe		
Yes No Is there evidence that the disc Yes No Will the patient be using Rem		with mothetroyate?		
\rightarrow Yes \square No Was treatme		with methodexate?		
		kate not tolerated or contraindicated?	not tolerated Contraindicated	
	$ ightarrow$ \Box Yes \Box No $$ Was treatme	nt with another conventional DMARD	(other than methotrexate) ineffective? quine I leflunomide I sulfasalazine	
Sarcoidosis				
☐ Yes ☐ No Is the disease refractory to co	rticosteroids?			
			Continued on next page	



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
G. CLINICAL INFORMATION (continued) - Re	equired clinical information must be con	pleted in its <u>entirety</u> for all precer	ification requests.				
Ulcerative Colitis	active fulminant ulcerative colitis?						
	the patient's ulcerative colitis: mild	🗌 moderate 🔲 severe					
Yes No Is there eviden	nce that the disease is active?						
	refractory to immunosuppression with c						
$ \qquad \qquad$	No Does the patient require continuous	immunosuppression with corticos	steroids (e.g., hydrocortisone,				
	methylprednisolone, prednisone)?						
	Name and dose: Name: Dose: Please indicate the route: □ Oral □ IV						
Name and d	ose: Name: ate the route: Oral IV	Dose:					
Please indic	ate the route:						
□ Yes □ No Was treatmen	nt with immunosuppressant agent (e.g.,	azathioprine 6-mercantopurine)	neffective?				
	No Was treatment with immunosuppre						
	or contraindicated?						
	\rightarrow Please select: \Box not tolerated \Box						
Please selec	t: 🗌 6-mercaptopurine 🔲 azathioprine						
☐ Yes ☐ No Was treatmen	nt with 5-aminosalicylic acid agents (e.g	balsalazide, mesalamine, sulfas	alazine) ineffective?				
	No Was treatment with 5-aminosalicyli						
	not tolerated or contraindicated?						
	\rightarrow Please select: \Box not tolerated \Box	contraindicated	Rowasa, Canasa (mesalamine)				
Please select: Colazal (balsalazide) Ariso, Asacal, Delzicol, Lialda, Pentasa, Rowasa, Canasa (mesalamine) Azulfidine (sulfasalazine) Other, please explain:							
\rightarrow Please select the symptoms the symptoms the symptoms the symptoms the symptoms the symptometry of the s	he patient exhibit: 🔲 more than 10 stoc						
	distension distension	ute, severe toxic symptoms, inclue	ling fever and anorexia				
For Continuation of Therapy (clinical docume	ntation required for all requests):						
Please indicate the length of time on Remicade							
Yes No Is this continuation request a r							
 ☐ Yes ☐ No Will Remicade (infliximab) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)? ☐ Yes ☐ No Is there clinical documentation supporting disease stability? 							
	supporting disease stability?						
□ Yes □ No Does the patient have any risk factors for TB?							
└────────────────────────────────────							
→ (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 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 in	nfusion?						
\square Yes \square No	Could the adverse reaction be mana	ged 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 bas	seline (pretreatment with Remicade (inf	iximab)): 🗋 mild 📋 moderate	severe				
H. ACKNOWLEDGEMENT							
Request Completed By (Signature Require	ed):		Date: / /				
Any person who knowingly files a request for	authorization of coverage of a medica	I procedure or service with the i	ntent to injure, defraud or deceive any				
insurance company by providing materially	false information or conceals mater	al information for the purpose					
insurance act, which is a crime and subjects s	such person to criminal and civil pena	Ities.					

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