

# Remicade<sup>®</sup> (infliximab) Injectable Medication Precertification Request

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For Michigan MMP: FAX: 1-844-241-2495 **PHONE:** 1-855-676-5772

For other lines of business: Please use other form.

Note: Remicade is preferred for

Start of treatment: Star	t date / /		/iew.)	MAPD pla	ns varies based on See section G below.
	· · · · · · · · · · · · · · · · · · ·		e:	Fax:	
RMATION					
		Last Name:			
		City:		State:	ZIP:
	Work Phone:		Cell Phone:		
Allergies:			-		
0	kas Height	. inches (			
			<u> </u>		
	Does natient have	e other coverage?	Yes 🗆 No		
		-			
	Insured:				
NFORMATION					
	Last Name:		(Check O	ne): 🗌 M.D. [	] D.O. 🗌 N.P. 🗌 P.A.
		City:		State:	ZIP:
Fax:	St Lic #:	NPI #:	DEA #:	ι	JPIN:
	Office Contact Name:		Phone:		
ROVIDER/ADMINISTRATIO			i nono.		
ed Physician's Office sion Center Phone: Center Phone: ame: code(s) (CPT): State Fax: PIN: PRMATION – Please select micade (infliximab) Dose: ORMATION – Please indica	: ZIP: :he medication being requested the primary ICD Code and speci Secondary ICD Code: al information must be complete	Physician's Of     Specialty Phar     Other:	fice Retail Ph macy Mail Ord State State	er ZI =ax: ZI =NN: HCPC: ode:	P:
ectra, Entyvio, and Simponi d Enbrel, Humira, Kevzara, O as the patient had prior thera as the patient had a trial and Enbrel (etanercept) Skyrizi (risankizumab-rzaa ere are any other medical re- that apply) Enbrel (etanercept) Skyrizi (risankizumab-rzaa	Aria are the preferred products f tezla, Rinvoq, Skyrizi, and Xeljan apy with Remicade (infliximab) v d failure, intolerance, or contrain umira (adalimumab)	IZ/Xeljanz XR are prefer within the last 365 days idication to any of the for ra (sarilumab)	red for other indications? pollowing? (select all tages) g preferred products zla (apremilast)	hat apply) Rinvoq (upada when indicated Rinvoq (upada	ncitinib) I for the patient's
	□ Start of treatment: Star   □ Continuation of therapy   equested By:   NATION     □ Allergies:   □ Ibs or   FORMATION   ■ Fax:   ROVIDER/ADMINISTRATIO   ration:   ed   □ Physician's Office   sion Center   Phone:   □ Center   Phone:   □ Center   Phone:   □ Code(s) (CPT):   □ State   □ Pin:   PIN: <td>□ Start of treatment: Start date       / / / / / / / / / / / / / / / / / / /</td> <td>Start of treatment: Start date       / / _ / _ / _ / _ / _ / _ / / /</td> <td>□ continuation of therapy: Date of last treatment       /         quested By:Phone:Phone:</td> <td>Start of treatment: Start date       / / / / / / / / / / / / / / / / / / /</td>	□ Start of treatment: Start date       / / / / / / / / / / / / / / / / / / /	Start of treatment: Start date       / / _ / _ / _ / _ / _ / _ / / /	□ continuation of therapy: Date of last treatment       /         quested By:Phone:Phone:	Start of treatment: Start date       / / / / / / / / / / / / / / / / / / /

- Please enter results of the TB test: Dositive 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)?



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Patient First Name	e	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INF	<b>FORMATION</b> (continued) – R	Required clinical information must	be completed in its <u>entirety</u> for all pre	certification requests.
	ndylitis and Other Spondyloa			
Please select whi	ich of the following applies to the	he patient: 🗌 Ankylosing spondyl	itis 🔲 Other spondyloarthropathy	
🗌 Yes 🔲 No	Is there evidence that the dise	ease is active?		
	Is there evidence of inflamma			
			-steroidal anti-inflammatory drugs (N	SAIDs)?
	Please provide the names an NSAID #1			
	NSAID #2:			
Behcet's Disease				
🔲 Yes 🔲 No	Is the disease refractory to co	orticosteroids or immunosuppressi	ive drugs?	
$  \longrightarrow$		roids 🔲 immunosuppressive dr		
Bahaatia Uwaitia	Please provide the name of d	Irug tried:		
Behcet's Uveitis	le the diagona refrectory?			
	Is the disease refractory?			
	-	nptomatic despite treatment with s	toroide?	
	Please provide the daily dose			
	······			
		nptomatic despite treatment with in		
Crohn's Disease	-	e 📋 cyclophosphamide 📋 met	hotrexate D Other, please explain:	
		nosis of fistulizing Crohn's disease	2	
	Please indicate how long the	patient has been diagnosed with f	istulizing Crohn's disease:	
🗋 Yes 🔲 No	Does the patient have a diagn	nosis of Crohn's disease?		
$  \longrightarrow$		f the patient's disease: 🗌 mild 🛛		
		ient have a documented diagnosis ct all signs/symptoms that apply:	s of active Crohn's disease?	
			🗌 diarrhea 🔲 internal fistulae 🗌	l intestinal obstruction
			 dylitis	
			active despite treatment with 6-merc	
	or corticoster			
			-mercaptopurine 🗌 azathioprine	ednisolone 🗌 Other:
Hidradenitis Sup		erolus- please identity. 🗋 prednis		
Please indicate th	ne stage of hidradenitis suppur	ativa: 🔲 Hurley stage I (mild dise		ate disease)
		Hurley stage III (severe	disease) 🔲 Unknown	
	Has the patient completed a till $\Box$ Vos $\Box$ No. Doos the patient	rial of antibiotics? ient have a contraindication to ora	l antibiotics?	
		tment with antibiotics ineffective?		
			trial: 🗌 Less than 1 month 🛛 1 mor	nth
			🗌 2 months 🔲 3 months (90 d	
Immune Checkpo Please indicate the	oint Inhibitor-Induced Toxici	ties		
CTLA-4	erapy used.			
	drug: 🔲 ipilimumab 🛛 Other			
🗌 PD-1		_		
	drug: 🔲 nivolumab 🛛 pembr	rolizumab 🔲 Other:		
PD-L1	drug: 🗋 atezolizumah 🗌 ave	elumab 🔲 durvalumab 🔲 Othe	r	
☐ Other				
Please explain				
🗌 Yes 🗌 No		hibitor-induced toxicities persist de ab, ipilimumab, nivolumab, pembr		ckpoint inhibitors that target CTLA-4 or
	he toxicity, (check all that ap			
		eckpoint inhibitor-induced cardiac		
		] impaired ventricular function	」myocarditis            pericarditis uced colitis.           mild         moderate	
			s: $\Box$ 7 or more stools per day over b	
<b>Д</b>	Yes I No Has the patient b	een treated with corticosteroids?		
	Please indicate the	he corticosteroid name: now improvement after 48 hours o	f a sufficiente un inte 2	
		iow improvement after 48 hours o	i conticosteroids?	



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		ted in its <u>entirety</u> for all precertin	ication requests.
Please indicate the toxicity, (check all that ap			
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?		
$\square$ Ves $\square$ No. Did the creating level	me and length of therapy: Name: remain greater than 2 to 3 times above bas	Length: Les	s than 1 week $\square$ 1 week or greater
☐ Inflammatory arthritis			
	efractory or severe disease? 🔲 refractory o	disease 🔲 severe disease	
	g to corticosteroids or anti-inflammatory ag	ents? 🔲 anti-inflammatory ager	its Corticosteroids
Please indicate the severity of the disea	se:midmoderatesevere eated with corticosteroids for pneumonitis?		
Please indicate the cor	rticosteroid name:		
	provement after 48 hours of corticosteroids	?	
Juvenile Idiopathic Arthritis (Juvenile Rheuma			
Please indicate the severity of the patient's disea			
Yes No Does the patient have clinical Yes No Is there evidence that the dise	documentation of polyarticular juvenile idio	pathic arthritis (JRA)?	
☐ Yes ☐ No Was treatment with Enbrel (et			
	mented intolerance to Enbrel (etanercept)?		
	mented contraindication to Enbrel (etanerce	ept)?	
Noninfectious Uveitis			
Yes No Was the treatment with cortico			
Please indicate the corticoster	oid name:		
☐ Yes ☐ No Was the treatment with immur		closporine, or methotrexate) inef	ifective?
Please provide the name:			
Yes No Does the patient have a docur	mented intolerance to corticosteroids or imr	nunosuppressive drugs?	
$\rightarrow$ Please indicate the drug(s) the	e patient has intolerance to: 🗌 corticostero	ids 🔲 immunosuppressive dru	gs
Yes No Does the patient have a docur			
Please indicate the drug(s) the Plaque Psoriasis	e patient has contraindication to: 🗌 cortico	steroids 📋 immunosuppressive	e drugs
Please indicate the severity of the patient's disea	ase: 🗌 mild 🔲 moderate 🔲 severe		
Yes 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 invo			-
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?			
	cyclosporine methotrexate mycc	phenolate 🔲 None of the above	ve
Yes No Was the trial with phototherapy			
$\longrightarrow$ Yes $\square$ No Was the trial v			
Second Se			
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 D 3 months or greater	ater



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G. CLINICAL INFORMATION (continued) – R	equired clinical information mus	t be completed in its <u>entirety</u> for all pr	ecertification requests.
Psoriatic Arthritis			
Yes No Is there evidence that the dise			
Yes No Does the patient have <b>axial</b> p			
	le the names and length of treat	al anti-inflammatory drugs (NSAIDs)	ΙΝΕΠΕCLIVE ?
NSAID #1:	le the hames and length of treat		
NSAID #1:			
Yes No Does the patient have <b>non-a</b>	<b>tial</b> psoriatic arthritis?		
		entation, defined as severe disability	at onset with erosive disease involving
multiple joints			
$ \qquad \qquad$	No Was the treatment with meth	notrexate ineffective?	
		atment with methotrexate not tolerate	
		select: not tolerated contrain	
		$\longrightarrow$ Please select: $\Box$ cyclopho	
			chloroquine 🗌 leflunomide
			izine Other, please explain:
Pyoderma Gangrenosum		_	
Yes No Does the patient have a docu	mented diagnosis of refractory p	yoderma gangrenosum?	
Reactive Arthritis (Reiter's syndrome) or Infla			
Please select which applies to the patient:	active arthritis (Reiter's syndrom	ie) 🔲 inflammatory bowel disease a	arthritis (enteropathic arthritis)
Yes No Was the treatment with method		1.10	
$ \qquad \qquad$	ent have a contraindication to m		
Yes No Was the treatment with sulfas			
$\square$ Yes $\square$ No Was the treat		rated?	
	ent have a contraindication to su		
Yes DNo Was the treatment with non-st	eroidal anti-inflammatory drugs	(NSAIDs) ineffective?	
		ammatory drugs (NSAIDs) not tolerat	
		on-steroidal anti-inflammatory drugs (	(NSAIDs)?
Please provide the name:			
Yes No Was treatment with a convent	ional DMARD ineffective?		
		not tolerated or contraindicated? 🔲 n	ot tolerated  Contraindicated
Rheumatoid Arthritis			
Please indicate the severity of the patient's rheu	ımatoid arthritis: 🗌 mild 🛛 mo	derate 🔲 severe	
Yes No Is there evidence that the dise	ease is active?		
Yes No Will the patient be using Rem		with methotrexate?	
└───> ☐ Yes ☐ No Was treatment	nt with methotrexate ineffective?		
			?  not tolerated  contraindicated (other than methotrexate) ineffective?
			oquine 🗌 leflunomide 🗍 sulfasalazine
Sarcoidosis			
Yes No Is the disease refractory to co	rticosteroids?		
			Continued on next page



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G. CLINICAL INFORMATION (continued) – Re	equired clinical information must be comple	eted in its <u>entirety</u> for all precertif	ication requests.
Ulcerative Colitis	active fulminant ulcerative colitis? the patient's ulcerative colitis:	moderate	methylprednisolone, prednisone)?
□ Yes       No       Was treatment         □ Yes       □ Yes       N         □ Yes       □ Yes       N         □ Please select       □ Yes       □ Yes         □ Yes       □ No       Was treatment	<ul> <li>ht with immunosuppressant agent (e.g., aza or contraindicated?</li> <li>→ Please select: □ not tolerated □ contraindicated?</li> <li>→ Int of the end of th</li></ul>	ant agent (e.g., azathioprine, 6-m ntraindicated ] cyclosporine alsalazide, mesalamine, sulfasa cid agents (e.g., balsalazide, me ntraindicated	nercaptopurine) not tolerated lazine) ineffective? salamine, sulfasalazine)
Please select the symptoms the symptoms the symptome of Therapy (clinical docume)	ntation required for all requests):	please explain:	☐ abdominal pain
☐ 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 Reminities         ☐ Yes       No       Has the patient received Reminities         ☐ Yes       No       Does the patient received Reminities <td>esult of the patient receiving samples of Reused concomitantly with apremilast, tofacities supporting disease stability? supporting disease improvement? factors for TB? thad a TB test within the past year? tapply): PPD test interferon-gamment the results of the TB test: positive in cade (infliximab) within the past 6 months? ent have a documented severe and/or potent fusion? Could the adverse reaction be managed itis, Plaque psoriasis, and Rheumatoid a</td> <td>nib, or other biologic DMARDs ( a assay (IGRA)</td> <td>vent that occurred during or following nome or office setting?</td>	esult of the patient receiving samples of Reused concomitantly with apremilast, tofacities supporting disease stability? supporting disease improvement? factors for TB? thad a TB test within the past year? tapply): PPD test interferon-gamment the results of the TB test: positive in cade (infliximab) within the past 6 months? ent have a documented severe and/or potent fusion? Could the adverse reaction be managed itis, Plaque psoriasis, and Rheumatoid a	nib, or other biologic DMARDs ( a assay (IGRA)	vent that occurred during or following nome or office setting?
H. ACKNOWLEDGEMENT			
Request Completed By (Signature Require			Date: / /
Any person who knowingly files a request for insurance company by providing materially insurance act, which is a crime and subjects s	false information or conceals material	information for the purpose c	

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