

# Inflectra® (infliximab-dyyb) Injectable **Medication Precertification Request**

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

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

For other lines of business: Please use other form.

Note: Inflectra is preferred for MA plans. Preferred status for MAPD

	of treatment: Start date	/ / last treatment /	/		•	es based on indication. on G below.
Precertification Requested By			Phone:		Fax:	
A. PATIENT INFORMATION						
First Name:		Last Name:			DOB:	
Address:		City:			State:	ZIP:
Home Phone:	Work Phone:	Cell	Phone:		Email:	
Current Weight: lbs or	kgs Height:	inches or cms	Allergies:			
<b>B. INSURANCE INFORMATIO</b>	N					
Aetna Member ID #:		Does patient have othe	r coverage?	🗌 Yes 🗌 No		
Group #:			-	_ Carrier Name:		
Insured:		Insured:				
C. PRESCRIBER INFORMATION	ON					
First Name:		Last Name:	1	(Check		□ D.O. □ N.P. □ P.A.
Address:	1		City:		State:	ZIP:
Phone:	Fax:	St Lic #:	NPI #:	DEA #:		UPIN:
Provider Email:	Of	ffice Contact Name:		Phone:		
D. DISPENSING PROVIDER/A	DMINISTRATION INFOR	MATION				
Outpatient Infusion Center     Center Name:     Home Infusion Center     Agency Name:     Administration code(s) (CF Address:     City:     Phone:     TIN:     NPI:     E. PRODUCT INFORMATION Request is for: Inflectra (inflix)	Phone:	ZIP:ation being requestedFrequ	Physician     Physician     Name:     Address:     City:     Phone:     TIN:     NPI:		Retail Ph     Other:     State:     Fax:     PIN:	ZIP:
F. DIAGNOSIS INFORMATION						
Primary ICD Code:						
G. CLINICAL INFORMATION - For Initiation Requests (clinic			s <u>entirety</u> for all p	ecertification requ	lests.	
Note:       Inflectra, Entyvio, Remicade, and Simponi Aria are the preferred products for MA plans. For MAPD plans, Inflectra, Entyvio, and Remicade are preferred for ulcerative colitis and Enbrel, Humira, Kevzara, Otezla, 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 Inflectra (infliximab-dyyb) 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)         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) 🔲 Xeljanz/Xeljanz XR (tofacitinib)						

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – Rea	nuired clinical information must be complete	ad in its entirety for all precertif	ication requests
☐ Yes       No       Will Inflectra (infliximab-dyyb) be         ☐ Yes       No       Has the patient been tested for biologic therapy?		-	
	est 🔲 interferon-gamma assay (IGRA) 📋	chest x-ray	
	est: positive negative unknown ve latent or active TB? latent active		
	ill TB treatment be started before initiation of	of therapy with Inflectra (inflixin	nab-dvvb)?
Ankylosing Spondylitis and Other Spondyloart			
Please select which of the following applies to the Yes No Is there evidence that the disea	patient: Ankylosing spondylitis Oth se is active?	er spondyloarthropathy	
Yes No Is there evidence of inflammato	•		
Yes No Has the patient had an ineffective Please provide the names and NSAID #1:		nti-inflammatory drugs (NSAID	s)?
NSAID #2:			
Behcet's Disease			
Behcet's Uveitis	ig tiled		
Yes No Is the disease refractory?			
Chronic Cutaneous/Pulmonary sarcoidosis	tomatic despite treatment with steroids?		
Please provide the daily dose o	f steroids: Dose:mg		
Please select: azathioprine	tomatic despite treatment with immunosupp	pressants?	
Crohn's Disease			
Yes No Does the patient have a diagnost Please indicate how long the patient have a diagnost Please indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the patient have a diagnost place indicate how long the place in		rohn'a diagona	
☐ Yes ☐ No Does the patient have a diagnost		onn's disease.	
Please indicate the severity of t	he patient's disease: 🗌 mild 🛛 moderate		
	nt have a documented diagnosis of active C all signs/symptoms that apply:	ronn's disease?	
	pain arthritis bleeding diarrhe	a 🔲 internal fistulae 🔲 inte	stinal obstruction
	perianal disease 🗌 spondylitis 🗋 w		
Yes No Have the Crohi	n's disease symptoms remained active des  ds?	pite treatment with 6-mercapto	purine, azathioprine,
	all medications that apply: 🗌 6-mercaptop	urine 🔲 azathioprine	
	oids- please identify: 🗌 prednisone 🛛 hy	drocortisone 🗌 methylpredni	solone 🗌 Other:
Hidradenitis Suppurativa Please indicate the stage of hidradenitis suppurat	iva:           Hurlev stage I (mild disease)	Hurley stage II (moderate of	disease)
	Hurley stage III (severe disease)		
Yes No Has the patient completed a tria	al of antibiotics?		
☐ Yes ☐ No Does the patient ☐ Yes ☐ No Was the treatm	nt have a contraindication to oral antibiotics	<i>!</i>	
Immune Checkpoint Inhibitor- Induced Toxiciti			
Please indicate therapy used:	es		
CTLA-4			
Please select drug: ipilimumab Other:			
Please select drug: I nivolumab pembro	olizumab 🔲 Other:		
DPD-L1			
Please select drug: 🗌 atezolizumab 🔲 avelumab 🔲 durvalumab 🔲 Other:			
Please explain:			
Yes No Do the immune checkpoint inhib PD-1/PD-L1 (e.g., atezolizumati	pitor-induced toxicities persist despite disco p, ipilimumab, nivolumab, pembrolizumab)?		nt inhibitors that target CTLA-4 or



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
	<i>tinued)</i> – Required clinical information must be	e completed in its <u>entirety</u> for all	precertification requests.	
Please indicate the toxicity (check		enviolities does the nationt have?		
	immune checkpoint inhibitor-induced cardiac t ythmias i impaired ventricular function			
	verity of the immune checkpoint inhibitor-induc		e 🗌 severe	
Please indicate which	of the following symptoms the patient exhibits:	: 🗌 7 or more stools per day ov	er baseline 🔲 ileus 🔲 fever 🗌 None	
	ne patient been treated with corticosteroids? e indicate the corticosteroid name:			
	e patient show improvement after 48 hours of	corticosteroids?		
Elevated serum creatinine/acute				
Please indicate the severity of				
Severe (creatinine	greater than 3 times baseline or greater than	4 mg/dL)		
Life-threatening (c	reatinine greater than 6 times baseline; dialysi	s indicated)		
□ None of the above				
Yes No Has the patie	nt been treated with corticosteroids?			
$\square$ Ves $\square$ No. Did the creati	ite the name and length of therapy: Name: nine level remain greater than 2 to 3 times abo	Lengtr	1: Less than 1 week 1 1 week or greater atment with corticosteroids?	
☐ Inflammatory arthritis	The level remain greater than 2 to 5 times abo			
	ent have refractory or severe disease? 🗌 refra			
	responding to corticosteroids or anti-inflammation	tory agents? 🔲 anti-inflammato	ry agents 🔲 corticosteroids	
Please indicate the severity of	f the disease: 🗌 mild 🔲 moderate 🔲 sever	<b>7</b> 0		
	nt been treated with corticosteroids for pneumo			
Please indica	te the corticosteroid name:			
	nt show improvement after 48 hours of corticos	teroids?		
Juvenile Idiopathic Arthritis (Juver Please indicate the severity of the pa	atient's disease:  mild  moderate  sev	vere		
Yes No Is there evidence				
Yes No Does the patient h	ave clinical documentation of polyarticular juve	enile idiopathic arthritis (JRA)?		
☐ Yes ☐ No Was treatment wit	h Enbrel (etanercept) ineffective?			
Yes No Does the patient h	ave a documented intolerance to Enbrel (etan	ercept)?		
Yes No Does the patient h	ave a documented contraindication to Enbrel (	(etanercept)?		
Noninfectious Uveitis				
Yes No Was the treatmen	e corticosteroid name:			
Yes No Was the treatmen	t with immunosuppressive drugs (e.g., azathio	prine, cyclosporine, or methotre	xate) ineffective?	
Please provide the	e name:			
Yes No Does the patient h	ave a documented intolerance to corticosteroid	ds or immunosuppressive drugs	?	
Please indicate the drug(s) the patient has intolerance to: 🗌 corticosteroids 🔲 immunosuppressive drugs				
<ul> <li>Yes No Does the patient have a documented contraindication to corticosteroids or immunosuppressive drugs?</li> <li>Please indicate the drug(s) the patient has contraindication to: Corticosteroids immunosuppressive drugs</li> </ul>				
Plaque Psoriasis			ppressive drugs	
Please indicate the severity of the pa	atient's disease: 🗌 mild 🔲 moderate 📋 sev	vere		
Yes No Is there evidence	hat the disease is active?			
Yes No Is there clinical do				
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:				
	bdy surface area affected by plaque psoriasis:	— %		
	soriasis involve sensitive areas? <i>If yes</i> , pleas		face 🔲 genitals	
	systemic conventional DMARD(s) (e.g., metho		-	
$\longrightarrow$ Yes $\square$ No Was the trial with systemic conventional DMARD(s) not tolerated?				
	re systemic conventional DMARDs contraindic			
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				
$\sim$ rease creak an that apply. $\Box$ restains (inclusionality, above and it) with overlight (inclusionality) and $\Box$ home UVB $\Box$ None of the above				
Please indicate the	e length of trial: Less than 1 month 1 m			



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G. CLINICAL INFORMATION (continued) – R	equired clinical information must be compl	eted in its entirety for all precerti	fication requests
Psoriatic Arthritis		olou in no <u>onaroty</u> for an procord	
Yes No Is there evidence that the dise			
Yes No Does the patient have <b>axial</b> ps	soriatic arthritis?		
Please provid	ment with 2 or more non-steroidal anti-infla e the names and length of treatment:	ammatory drugs (NSAIDs) ineffe	ctive?
	e the names and length of treatment.		
NSAID #2:			
Yes No Does the patient have <b>non-ax</b>	ial psoriatic arthritis?		
multiple joints		-	set with erosive disease involving
$ \longrightarrow \Box \operatorname{Yes} \Box I$	No Was the treatment with methotrexate in		
	$\longrightarrow$ Yes $\square$ No Was treatment with	nethotrexate not tolerated or contraindicate	
		Vas treatment with another conv	
		Please select: Cyclophosphar	
			quine 🗌 leflunomide
		Sulfasalazine	☐ Other, please explain:
Pyoderma Gangrenosum	nented diagnosis of refrectory nyedorme a	angrana aum?	
Yes No Does the patient have a docur			
<b>Reactive Arthritis (Reiter's syndrome) or Infla</b> Please select which applies to the patient:			s (enteropathic arthritis)
$\Box$ Yes $\Box$ No Was the treatment with metho			
	ment with methotrexate not tolerated?		
	ent have a contraindication to methotrexate	e?	
Yes No Was the treatment with sulfast			
	ment with sulfasalazine not tolerated? ent have a contraindication to sulfasalazine	22	
☐ Tes ☐ No Does the pair			
	ment with non-steroidal anti-inflammatory		
	ent have a contraindication to non-steroida	I anti-inflammatory drugs (NSAI	Ds)?
Retinal Vasculitis	ional DMARD ineffective?		
	nt with a conventional DMARD not tolerate	d or contraindicated? □ not tole	rated Contraindicated
Rheumatoid Arthritis		<b>_</b>	
Please indicate the severity of the patient's rheu		severe	
Yes No Is there evidence that the dise			
Yes No Will the patient be using Inflec		ethotrexate?	
	No Was treatment with methotrexate not t $\Rightarrow$ $\Box$ Yes $\Box$ No Was treatment with a	nother conventional DMARD (otl	
			Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued) – R	aguirad alinical information must be comple	tod in its ontircty for all proportif	instion requests	
G. CLINICAL INFORMATION ( <i>continued)</i> – Re Sarcoidosis	equired clinical information must be comple	eted in its <u>entirely</u> for all precertin	cation requests.	
Yes No Is the disease refractory to con	ticosteroids?			
Ulcerative Colitis				
Yes No Is the patient hospitalized with	active fulminant ulcerative colitis?			
	the patient's ulcerative colitis: mild	moderate 🗌 severe		
	nce that the disease is active? refractory to immunosuppression with cortion	costoroids (o.g., hydrocortisono	mothylprodpisologo, prodpisopo)?	
	No Does the patient require continuous im			
	methylprednisolone, prednisone)?			
	$\rightarrow$ Name and dose: Name:			
	Please indicate the route:  Oral	IV		
Name and d	ose: Name:	Dose:		
Please indic	ate the route: 🗌 Oral 🔲 IV			
	t with immunosuppressant agent (e.g., aza lo   Was treatment with immunosuppressa			
	or contraindicated?	ni ageni (e.g., azatnophne, e-m	sicaptopullie) not tolerated	
	$\rightarrow$ Please select: $\Box$ not tolerated $\Box$ cor			
Please selec	t: 🗌 6-mercaptopurine 🔲 azathioprine			
☐ Yes ☐ No Was treatmen	t with 5-aminosalicylic acid agents (e.g., ba	alsalazide, mesalamine, sulfasal	azine) ineffective?	
	No Was treatment with 5-aminosalicylic ac			
	not tolerated or contraindicated?			
	$\rightarrow$ Please select: $\Box$ not tolerated $\Box$ cont: $\Box$ Colazal (balsalazide) $\Box$ Ariso, Asa		owasa, Canasa (mesalamine)	
	Azulfidine (sulfasalazine) Other,			
		· · ·		
Please select the symptoms the patient exhibit: 🗌 more than 10 stools per day 🗌 continuous bleeding 🔲 abdominal pain				
For Continuation of Thereny (clinical docume	distension acute, severe toxic symptoms, including fever and anorexia			
For Continuation of Therapy (clinical docume Please indicate the length of time on Inflectra (in				
☐ Yes ☐ No Is this continuation request a r		flectra (infliximab-dvvb)?		
Yes No Will Inflectra (infliximab-dyyb)			)s (e.g., adalimumab, certolizumab)?	
Yes No Is there clinical documentation				
□ Yes       □ No       Is there clinical documentation supporting disease improvement?         □ Yes       □ No       Does the patient have any risk factors for TB?				
$\square$ The subscription boost the patient have any fisk factors for TB? $\square$ Yes $\square$ No Has the patient had a TB test within the past year?				
(check all that apply): PPD test interferon-gamma assay (IGRA) ichest x-ray				
Please enter the results of the TB test: Dositive Dinegative unknown				
Yes No Has the patient received Inflectra (infliximab-dyyb) within the past 6 months?				
the previous infusion?				
$\square$ Yes $\square$ No	Could the adverse reaction be managed	I through pre-medication in the h	ome or office setting?	
For Crohn's disease, Juvenile idiopathic arthritis, Plaque psoriasis, Rheumatoid arthritis, Ulcerative colitis only:				
Please indicate the severity of the disease at bas	seline (pretreatment with Inflectra (inflixima	b-dyyb)): 🗋 mild 📋 moderate		
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.				
insurance act, which is a chine and subjects s		ə.		

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