

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

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(All fields must be completed and legible for precertification review.) / ☐ Start of treatment: Start date

For Ohio MMP: FAX: 1-855-734-9389

PHONE: 1-855-364-0974 For other lines of business: Please use other form.

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

Please indicate:	☐ Start of treatment: Start date☐ Continuation of therapy: Date					See section	G below.
Precertification R	• •	e or last treatment		 Phone:		Fax:	
A. PATIENT INFO							
First Name:		L	ast Name:			DOB:	
Address:			ity:			State:	ZIP:
Home Phone:	Work Phone:		-	Phone:		Email:	
Current Weight:				lergies:			
B. INSURANCE IN			omo	norgios.			
	#:	Does nat	ient have other	coverage? \Box Ye	es 🗌 No		
	,		Does patient have other coverage?				
Insured:							
C. PRESCRIBER I	NFORMATION						
First Name:		Last Nam	ne:		(Check C	ne): 🔲 M.D. 🗀] D.O. 🗌 N.P. 🗌 P.A.
Address:		•		City:		State:	ZIP:
Phone:	Fax:	St Lic #:		NPI #:	DEA #:	UI	PIN:
Provider Email:	1	Office Contact I			Phone:		
D. DISPENSING P	ROVIDER/ADMINISTRATION INF	ORMATION					
Center Na Home Infusion Agency N Administration Address: City: Phone: TIN: NPI:	sion Center Phone: ame: Center Phone: lame: code(s) (CPT): State: Fax: PIN: DRMATION – Please select the me	ZIP:edication being re	quested	Phone:	acy	Other: State: Fax: PIN:	ZIP:
	flectra (infliximab-dyyb) Dose: _					HCPCS	Code:
	FORMATION – Please indicate pri						
Primary ICD Code: Secondary ICD Code:							
	DRMATION – Required clinical info			entirety for all precertifi	cation requ	ests.	
Note: Inflectra, Enpreferred for Preferred pr	tests (clinical documentation rec tyvio, Remicade, and Simponi A r ulcerative colitis and Enbrel, He oducts vary based on indication Has the patient had prior therapy w Has the patient had a trial and failur ☐ Enbrel (etanercept) ☐ Humira	ria are the prefer umira, Kevzara, (i. ith Inflectra (inflixi re, intolerance, or	red products for the products for the product of th	Skyrizi, and Xeljanz/X in the last 365 days? In to any of the following	(eljanz XR :	are preferred for that apply)	or other indications.
Please explain if th diagnosis (select a	☐ Skyrizi (risankizumab-rzaa) ☐ ere are any other medical reason(Xeljanz/Xejlanz X s) that the patient (adalimumab)	(R (tofacitinib) cannot use any	of the following preferro	ed products	when indicated	for the patient's

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.						
Yes No Will Inflectra (infliximab-dyyb) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)' 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						
	fill TB treatment be started before initiation of	of therapy with Inflectra (inflixin	nab-dyyb)?			
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?						
	ve response to two or more non-steroidal and length of treatment:	nti-inflammatory drugs (NSAID	s)?			
NSAID #2: Behcet's Disease						
Yes No Is the disease refractory to corticosteroids or immunosuppressive drugs?						
Behcet's Uveitis						
Chronic Cutaneous/Pulmonary sarcoidosis ☐ Yes ☐ No Has the patient remained symp	☐ Yes ☐ No Has the patient remained symptomatic despite treatment with steroids?					
Please provide the daily dose of steroids: Dose:mg Yes No Has the patient remained symptomatic despite treatment with immunosuppressants? Please select: azathioprine cyclophosphamide methotrexate Other, please explain:						
Crohn's Disease ☐ Yes ☐ No Does the patient have a diagnosis of fistulizing Crohn's disease? ☐ Please indicate how long the patient has been diagnosed with fistulizing Crohn's disease: ☐ Yes ☐ No Does the patient have a diagnosis of Crohn's disease?						
Please indicate the severity of the patient's disease: mild moderate severe Yes No Does the patient have a documented diagnosis of active Crohn's disease? Please select all signs/symptoms that apply:						
□ abdominal pain □ arthritis □ bleeding □ diarrhea □ internal fistulae □ intestinal obstruction □ megacolon □ perianal disease □ spondylitis □ weight loss □ None of the above □ Yes □ No Have the Crohn's disease symptoms remained active despite treatment with 6-mercaptopurine, azathioprine,						
☐ corticostere	ds? all medications that apply: ☐ 6-mercaptopoolids- please identify: ☐ prednisone ☐ hyo		isolone			
Hidradenitis Suppurativa Please indicate the stage of hidradenitis suppurativa: ☐ Hurley stage I (mild disease) ☐ Hurley stage II (moderate disease)						
☐ Yes ☐ No Has the patient completed a tria	☐ Hurley stage III (severe disease) al of antibiotics?	Unknown	130030)			
☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐						
Immune Checkpoint Inhibitor- Induced Toxicities Please indicate therapy used: ☐ CTLA-4						
Please select drug: 🗌 ipilimumab 🔲 Other:						
☐ PD-1 Please select drug: ☐ nivolumab ☐ pembrolizumab ☐ Other: ☐ PD-L1						
Please select drug: atezolizumab avelumab durvalumab Other: Other Please explain:						
Yes No Do the immune checkpoint inhib	oitor-induced toxicities persist despite discorto, ipilimumab, nivolumab, pembrolizumab)?	ntinuation of immune checkpoi	nt inhibitors that target CTLA-4 or			

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Patient First	Name	•	Patient Last Name	Patient Phone	Patient DOB
G. CLINICA	AL INF	ORMATION (continued) – Re	equired clinical information must be comple	eted in its <u>entirety</u> for all precertifi	cation requests.
Please indi	cate t	he toxicity (check all that app	oly:)		
☐ Cardiac			eckpoint inhibitor-induced cardiac toxicities		
			impaired ventricular function myocard		
☐ Colitis			immune checkpoint inhibitor-induced colitis		
	FIE:	Yes \square No. Has the natient he	ing symptoms the patient exhibits: 7 or en treated with corticosteroids?	more stools per day over baselii	le 🗌 lieus 🔲 level 🔲 Norie
	ͳ		e corticosteroid name:		
			ow improvement after 48 hours of corticost	eroids?	
□ Flevated		n creatinine/acute renal failure	on improvement and to heard or contiduot	oroldo.	
		cate the severity of the disease	:		
		,	າ 3 times baseline or greater than 4 mg/dL))	
		,	eater than 6 times baseline; dialysis indicate		
		None of the above	ator than a times baseline, diaryole indicat	54)	
			to d with continent and 2		
Ч 1	es 🔲	No Has the patient been treat	and length of therapy: Name:	Length: □ Less	than 1 week
ПУ	es 🗆	No Did the creatinine level re	main greater than 2 to 3 times above base	line after 1 week of treatment wit	h corticosteroids?
☐ Inflamma			main grouter than 2 to 0 times above base	mie aner i week er neament wi	in definitional in the second
			ractory or severe disease? 🗌 refractory dis	sease severe disease	
□ Ye	es 🗌	No Is the patient responding to	to corticosteroids or anti-inflammatory ager	nts? 🗌 anti-inflammatory agents	☐ corticosteroids
☐ Pneumo					
			: mild moderate severe		
			ted with corticosteroids for pneumonitis?		
	о П	Please indicate the cortice	osteroid name:overnment after 48 hours of corticosteroids?		
Juvenile Idi	ອຣ ∐ onath	ic Arthritis (Juvenile Rheuma	atoid Arthritis)		
			se: mild moderate severe		
☐ Yes ☐		Is there evidence that the disea			
☐ Yes ☐		Does the patient have clinical of	documentation of polyarticular juvenile idio	pathic arthritis (JRA)?	
☐ Yes ☐		Was treatment with Enbrel (eta		paulio arailiao (oro i).	
		•	• •		
☐ Yes ☐		•	nented intolerance to Enbrel (etanercept)?	1)0	
☐ Yes ☐		•	nented contraindication to Enbrel (etanerce	ept)?	
Noninfectio		⁄eเนร Was the treatment with cortico	storoids inoffactive?		
T 163 D			oid name:		
		Trough mandate the control to	ord frame.		
☐ Yes ☐	No	Was the treatment with immun	osuppressive drugs (e.g., azathioprine, cyc	closporine, or methotrexate) inef	fective?
	\longrightarrow	Please provide the name:			
	NI.	December of the second second			
			nented intolerance to corticosteroids or imr		70
	Please indicate the drug(s) the patient has intolerance to: corticosteroids immunosuppressive drugs Yes No Does the patient have a documented contraindication to corticosteroids or immunosuppressive drugs?				js
			patient has contraindication to: corticos		drugs
Plaque Pso					9-
			se: mild moderate severe		
☐ Yes ☐	No	Is there evidence that the disea	ase is active?		
☐ Yes ☐	No	Is there clinical documentation	of chronic disease?		
☐ Yes ☐	No	Is the patient a candidate for s	ystemic therapy or phototherapy?		
			y ☐ systemic therapy ☐ phototherapy a	and systemic therapy	
Please prov		e patient's Psoriasis Area and			
			area affected by plaque psoriasis:%		
☐ Yes ☐	No	Does the plaque psoriasis invo	olve sensitive areas? If yes, please select:	☐ hands ☐ feet ☐ face ☐	genitals
☐ Yes ☐	No	Was the trial with systemic cor	nventional DMARD(s) (e.g., methotrexate,	acetretin, or cyclosporine) ineffec	tive?
T T		-	vith systemic conventional DMARD(s) not t		
			conventional DMARDs contraindicated?		
			cyclosporine methotrexate myco	ophenolate	re
☐ Yes ☐		Was the trial with phototherapy		_	
I			vith phototherapy not tolerated?		
		☐ Yes ☐ No Is phototherap	* · · · · · · · · · · · · · · · · · · ·		
			•	UVA light (PUVA) ☐ UVB with	coal tar or dithranol
	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				
			ial: Less than 1 month 1 month 5		ıter
				_ =on grot	



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G. CLINICAL INFORMATION (continued) - R	equired clinical information must be o	completed in its <u>entirety</u> for all prece	rtification requests.
Psoriatic Arthritis			
Yes No Is there evidence that the dise			
Yes No Does the patient have axial ps			
	ment with 2 or more non-steroidal and		fective?
	le the names and length of treatment:		
NSAID #1:			
NSAID #2: ☐ Yes ☐ No Does the patient have non-ax			
Tes No Does the patient have non-ax	ent have severe disease at presentat	ion defined as severe disability at o	unset with erosive disease involving
multiple joints		ion, defined as severe disability at c	miset with erosive disease involving
	No Was the treatment with methotrex	cate ineffective?	
, – –		nt with methotrexate not tolerated or	contraindicated?
	> Please selection	ct: not tolerated contraindica	ted
		No Was treatment with another cor	
		→ Please select: ☐ cyclophosph	
			oquine leflunomide
Book down and Community of the Community		∐ sulfasalazine	Other, please explain:
Pyoderma Gangrenosum			
Yes No Does the patient have a docur			
Reactive Arthritis (Reiter's syndrome) or Infla			the foundation of the state of
Please select which applies to the patient: re	` , , -	iniiammatory bowei disease artini	ius (enteropatnic arthritis)
Yes No Was the treatment with metho	ment with methotrexate not tolerated	2	
· = =	ent have a contraindication to methot		
Yes No Was the treatment with sulfas		icato:	
	ment with sulfasalazine not tolerated	?	
	ent have a contraindication to sulfasa		
☐ Yes ☐ No Was the treatment with non-st			
☐ Yes ☐ No Was the treat			
	ent have a contraindication to non-ste		AIDs)?
· ·			
Retinal Vasculitis			
Yes No Was treatment with a convent			
	nt with a conventional DMARD not tol	erated or contraindicated?	lerated
Rheumatoid Arthritis Please indicate the severity of the patient's rheu	motoid arthritis: mild moderat	a Dagwara	
Yes No Is there evidence that the dise		e 🖂 severe	
Yes No Will the patient be using Inflect		vith methotrexate?	
Yes No Was treatmen			
	No Was treatment with methotrexate	e not tolerated or contraindicated?	not tolerated contraindicated
	→ ☐ Yes ☐ No Was treatment v		
	└────────────────────────────────────	ີ່ azathioprine hydroxychloroqເ	ine 🗌 leflunomide 📋 sulfasalazine

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.						
Sarcoidosis		noted in ito <u>originary</u> for all pro-	commodulori roquocic.			
☐ Yes ☐ No Is the disease refractory to co	rticosteroids?					
Ulcerative Colitis						
☐ Yes ☐ No Is the patient hospitalized with						
	the patient's ulcerative colitis: mild	☐ moderate ☐ severe				
	nce that the disease is active?					
	refractory to immunosuppression with cor No Does the patient require continuous in					
	methylprednisolone, prednisone)?	minunosuppression with con-	costerolas (e.g., flyarocortisorie,			
	→ Name and dose: Name:	Dose:				
	Please indicate the route: ☐ Oral					
Name and a	lose: Name:	Doso:				
	eate the route:	Dose				
	nt with immunosuppressant agent (e.g., az					
	No Was treatment with immunosuppress	ant agent (e.g., azathioprine,	6-mercaptopurine) not tolerated			
	or contraindicated? → Please select: □ not tolerated □ co	ontraindicated				
	ct: 6-mercaptopurine azathioprine					
	nt with 5-aminosalicylic acid agents (e.g., b					
	No Was treatment with 5-aminosalicylic a not tolerated or contraindicated?	acid agents (e.g., balsalazide	, mesalamine, sulfasalazine)			
	→ Please select: ☐ not tolerated ☐ co	ontraindicated				
	ct: Colazal (balsalazide) Ariso, As		sa. Rowasa, Canasa (mesalamine)			
	☐ Azulfidine (sulfasalazine) ☐ Othe		· · · · · · · · · · · · · · · · · · ·			
→ Please select the symptoms t	he patient exhibit: more than 10 stools	e, severe toxic symptoms, inc	=			
For Continuation of Therapy (clinical docume		e, severe toxic symptoms, in	studing lever and anorexia			
Please indicate the length of time on Inflectra (ir						
☐ Yes ☐ No Is this continuation request a result of the patient receiving samples of Inflectra (infliximab-dyyb)?						
Yes No Will Inflectra (infliximab-dyyb) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)?						
☐ Yes ☐ No Is there clinical documentation supporting disease stability?						
☐ 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 Infle	ctra (infliximab-dyyb) within the past 6 mor	nths?				
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, Rheumatoid arthritis, Ulcerative colitis only: Please indicate the severity of the disease at baseline (pretreatment with Inflectra (infliximab-dyyb)): mild moderate severe						
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Require			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.