

Please indicate:

MEDICARE FORM

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

Page 1 of 5

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

☐ Start of treatment: Start date / /

For Illinois MMP: 1-855-320-8445 FAX: **PHONE**: 1-866-600-2139 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. See section G below.

	e of last treatment						
y:			Phone:		Fax		
	Last N	ame:			DOB:		
	City:				State:	ZIP:	
Work Phone:	·	Cell F	Phone:		Email:		
kgs Height:	inches or	cms Al	llergies:				
<u> </u>		-					
	Does natient h	ave other	coverage?	□ Ves □ No	1		
			Ū				
ON	-						
	Last Name:			(Check	One):). 🗌 D.O. 🗌 N.P. 🔲	P.A.
			City:		State:	ZIP:	
Fax:	St Lic #:		NPI #:	DEA #:		UPIN:	
	Office Contact Name	e:		Phone:			
DMINISTRATION INF							
Phone: Phone: Phone: State: Fax:	ZIP:		Physician's Specialty F Name: Address: City: Phone: TIN:	s Office Pharmacy	Retail F	ZIP:	
PIN:							
– Please select the me	edication being reques	ted					
	edication being reques	ted	ncy:		HCP	CS Code:	
– Please select the me kimab-dyyb) Dose: I – Please indicate prin	edication being reques	ted Freque ecify any c	ther where applic	able.			
– Please select the me	edication being reques	ted Freque ecify any c	ther where applic	able.			
– Please select the me kimab-dyyb) Dose: I – Please indicate prin	edication being reques mary ICD Code and sp recondary ICD Code: _	ted Freque pecify any c	ther where applic	able. Other ICD	Code:		
– Please select the me kimab-dyyb) Dose: I – Please indicate prin S	edication being reques mary ICD Code and sp secondary ICD Code: _ rmation must be comp	ted Freque pecify any colleted in its	ther where applic	able. Other ICD	Code:		
	Work Phone: kgs Height: N DN Fax: DMINISTRATION INF Physician's Office Phone: Phone: State:	Last N City: Work Phone: kgs Height: inches or Does patient h If yes, provide Insured: DN Last Name: Fax: St Lic #: Office Contact Name DMINISTRATION INFORMATION Physician's Office Phone: Phone: State: ZIP:	Last Name: City: Work Phone: kgs Height:inches orcms All N Does patient have other of the provide ID#:insured: Insured: St Lic #: Office Contact Name: DMINISTRATION INFORMATION Physician's Office Phone: Phone: Phone:	Last Name: City:	Last Name: City: Work Phone: Cell Phone: Allergies: N	Last Name: DOB:	Last Name: DOB: State: ZIP:

Continued on next page



Inflectra® (infliximab-dyyb) Injectable Medication Precertification Request

Page 2 of 5

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

For Illinois MMP:

FAX: 1-855-320-8445 **PHONE:** 1-866-600-2139

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. See section G.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued) - Re	quired clinical information must be complete	ed in its <u>entirety</u> for all precertifi	cation requests.	
☐ Yes ☐ No Will Inflectra (infliximab-dyyb) b ☐ Yes ☐ No Has the patient been tested for biologic therapy? (check all that apply): ☐ PPD to Please enter results of the TB to	e used concomitantly with apremilast, tofaci	tinib, or other biologic DMARD y (IGRAs) or chest x-ray withir	s (e.g., adalimumab, certolizumab)?	
If latent TB, ☐ Yes ☐ No W	/ill TB treatment be started before initiation of	of therapy with Inflectra (inflixin	nab-dyyb)?	
Ankylosing Spondylitis and Other Spondyloar Please select which of the following applies to the Yes No Is there evidence that the disea Is there evidence of inflammato Has the patient had an ineffect Please provide the names and NSAID #1:	e patient: Ankylosing spondylitis Otherse is active? Ory disease? Over response to two or more non-steroidal and the steroid of the stero		s)?	
→ Please indicate: ☐ corticostere	ticosteroids or immunosuppressive drugs? oids			
Behcet's Uveitis ☐ Yes ☐ No Is the disease refractory?				
Chronic Cutaneous/Pulmonary sarcoidosis				
☐ Yes ☐ No Has the patient remained symptone → Please provide the daily dose of the patient remained symptone ☐ Yes ☐ No Has the patient remained symptone ☐ Yes ☐ Yes ☐ No Has the patient remained symptone ☐ Yes ☐ Y	of steroids: Dose:mg			
Crohn's Disease Yes No Does the patient have a diagnomal Please indicate how long the pure yes No Does the patient have a diagnomal Please indicate how long the pure yes No Does the patient have a diagnomal Please indicate how long the pure yes No Does the patient have a diagnomal Please indicate how long the patien	sis of fistulizing Crohn's disease? atient has been diagnosed with fistulizing Cr			
Please indicate the severity of ☐ Yes ☐ No Does the patie → Please select	the patient's disease: ☐ mild ☐ moderate nt have a documented diagnosis of active C all signs/symptoms that apply:	rohn's disease?	ette et et en etter	
☐ megacolor	pain arthritis bleeding diarrhe n perianal disease spondylitis whi's disease symptoms remained active desids?	eight loss 🔲 None of the above	ve	
	all medications that apply: 6-mercaptoputoids-please identify: prednisone hydroxids-please identify:		aslana 🗖 Othor:	
Hidradenitis Suppurativa	olds- please identity. preditisorie into		solone	
Please indicate the stage of hidradenitis suppura	☐ Hurley stage III (severe disease)	☐ Hurley stage II (moderate o	tisease)	
Yes No Does the patient completed a title of the patient completed	nt have a contraindication to oral antibiotics	?		
Immune Checkpoint Inhibitor- Induced Toxicities Please indicate therapy used:				
☐ CTLA-4 Please select drug: ☐ ipilimumab ☐ Other:				
│	olizumab 🗌 Other:			
Please select drug: ☐ atezolizumab ☐ ave	lumab 🔲 durvalumab 🔲 Other:			
Please explain: Yes No Do the immune checkpoint inhi PD-1/PD-L1 (e.g., atezolizumal	bitor-induced toxicities persist despite discorto, ipilimumab, nivolumab, pembrolizumab)?	ntinuation of immune checkpoi	nt inhibitors that target CTLA-4 or	

Continued on next page



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

Page 3 of 5
(All fields must be completed and legible for precertification review.)

For Illinois MMP: **FAX:** 1-855-320-8445 **PHONE**: 1-866-600-2139 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.

See section G.

Patient First	Name	Patient Last Name	Patient Phone	Patient DOB
		equired clinical information must be comple	eted in its <u>entirety</u> for all precertif	ication requests.
	cate the toxicity (check all that ap			
Cardiac	Please select: arrhythmias	eckpoint inhibitor-induced cardiac toxicities impaired ventricular function myocard	itis pericarditis	
Colitis	Please indicate which of the follow	immune checkpoint inhibitor-induced coliticing symptoms the patient exhibits: 7 or		
		e corticosteroid name:	_	
1_	·	ow improvement after 48 hours of corticost	eroids?	
	serum creatinine/acute renal failure			
Pleas		n 3 times baseline or greater than 4 mg/dL		
	☐ Life-threatening (creatinine green☐ None of the above	eater than 6 times baseline; dialysis indicat	ed)	
☐ Ye	es	ted with corticosteroids?		
	Please indicate the name	and length of therapy: Name: main greater than 2 to 3 times above base	Length: Les	s than 1 week 1 week or greater
☐ Inflamma	ntory arthritis			in concosteroids?
		ractory or severe disease?		s 🗌 corticosteroids
Pneumor		o D mild D moderate D covers		
☐ Ye	e indicate the severity of the diseases S No Has the patient been trea Please indicate the cortic	ted with corticosteroids for pneumonitis?		
		osteroid fiame		
Juvenile Idio	opathic Arthritis (Juvenile Rheum	atoid Arthritis)		
		ase: mild moderate severe		
	No Is there evidence that the dise		nothic outhritic / IDANS	
		documentation of polyarticular juvenile idio	patriic artifitis (JRA)?	
	No Was treatment with Enbrel (et			
		mented intolerance to Enbrel (etanercept)?	nt\2	
Noninfection		mented contraindication to Enbrel (etanerce	ερι) <i>?</i>	
☐ Yes ☐	No Was the treatment with cortico	osteroids ineffective? oid name:		
	/ I loade maidate the controlater	old flame.		
Yes 🗆	No Was the treatment with immur Please provide the name:	nosuppressive drugs (e.g., azathioprine, cy	closporine, or methotrexate) inet	ffective?
☐ Yes ☐	No Does the patient have a docur	mented intolerance to corticosteroids or imr	nunosuppressive drugs?	
	Please indicate the drug(s) the	e patient has intolerance to: 🔲 corticostero	ids immunosuppressive dru	ıgs
Yes 🗆	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 Psoi	riasis			3
		ase: mild moderate severe		
Yes				
Yes				
☐ Yes ☐		systemic therapy or phototherapy?	and avatamia therany	
Please prov	ide the patient's Psoriasis Area and	y ☐ systemic therapy ☐ phototherapy a Severity Index (PASI) score:	and systemic therapy	
	•	area affected by plaque psoriasis:%		
		olve sensitive areas? <i>If yes</i> , please select:	☐ hands ☐ feet ☐ face ☐] genitals
P Yes P		nventional DMARD(s) (e.g., methotrexate, with systemic conventional DMARD(s) not		ctive?
		conventional DMARDs contraindicated?	ioloratou :	
		cyclosporine methotrexate mycc	ophenolate	ve
☐ Yes ☐	No Was the trial with phototherap	y ineffective?		
	→ ☐ Yes ☐ No Was the trial v			
	☐ Yes ☐ No Is photothera			
		Psoralens (methoxsalen, trioxsalen) with		h coal tar or dithranol
		☐ UVB (standard or narrow band) ☐ Hon		
	Please indicate the length of t	rial: 🗌 Less than 1 month 🔲 1 month 📋	」2 months ☐ 3 months or gre	ater



Inflectra® (infliximab-dyyb) Injectable Medication Precertification Request

Page 4 of 5

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

For Illinois MMP:

FAX: 1-855-320-8445 **PHONE:** 1-866-600-2139

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. See section G.

Patient First Name	Patient Last Name	Patient Phone	Patient DOB
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 dames October 1997		∐ 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		revate:	
	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

Continued on next page



Inflectra® (infliximab-dyyb) Injectable Medication Precertification Request

Page 5 of 5

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

For Illinois MMP:

FAX: 1-855-320-8445 **PHONE:** 1-866-600-2139

For other lines of business: Please use other form.

Note: Inflectra is preferred for I

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

Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued) – R	equired clinical information mu	st be completed in its entirety for all pre	ecertification requests.	
Sarcoidosis	oquirou omitour intermatieri ma	ot 20 00 mp. ot 00 mm to <u>0.11 m ot 7</u> m. pro	ooramoanon roquocic.	
☐ Yes ☐ No Is the disease refractory to co	rticosteroids?			
Ulcerative Colitis				
Yes No Is the patient hospitalized with				
	nce that the disease is active?	☐ mild ☐ moderate ☐ severe		
		on with corticosteroids (e.g., hydrocortis	sone, methylprednisolone, prednisone)?	
	No Does the patient require co	ontinuous immunosuppression with cort		
	methylprednisolone, predr	nisone)? Dose:		
	Please indicate the route:			
	r lease indicate the route.			
> Name and o	lose: Name:	Dose:		
Please indic	ate the route:			
☐ Yes ☐ No. Was treatmen	nt with immunosuppressant age	ent (e.g., azathioprine, 6-mercaptopurin	e) ineffective?	
		nosuppressant agent (e.g., azathioprine		
	or contraindicated?	🗖		
	→ Please select: ☐ not toler ct: ☐ 6-mercaptopurine ☐ az			
/ Tidase selection	zi. 🔲 o-mercaptoparine 🗀 az	eatmoprine		
		ents (e.g., balsalazide, mesalamine, su		
│	No Was treatment with 5-amir not tolerated or contrainding	nosalicylic acid agents (e.g., balsalazide	e, mesalamine, sulfasalazine)	
	→ Please select: ☐ not toler			
		Ariso, Asacal, Delzicol, Lialda, Penta	sa, Rowasa, Canasa (mesalamine)	
	☐ Azulfidine (sulfasalazine)		
Please select the symptoms to	he nationt exhibit: \(\sim \) more tha	n 10 stools per day 🔲 continuous ble	eding	
/ Floude select the symptome t		n acute, severe toxic symptoms, in		
For Continuation of Therapy (clinical docume	ntation required for all reque	ests):	•	
Please indicate the length of time on Inflectra (ir				
Yes No Is this continuation request a			AADDa (a.e. adaliesumada aastaliesumada)	
☐ Yes ☐ No Will Inflectra (infliximab-dyyb)		remilast, tolacitinib, or other biologic Di	MARDS (e.g., adailmumab, 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?				
├────────────────────────────────────				
☐ Yes ☐ No Has the patient received Inflectra (infliximab-dyyb) 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 infusion?				
For Crohn's disease, Juvenile idiopathic arth				
Please indicate the severity of the disease at ba			•	
H. ACKNOWLEDGEMENT				
Request Completed By (Signature Require			Date:/	
Any person who knowingly files a request for		a medical procedure or service with th		
insurance company by providing materially insurance act, which is a crime and subjects	false information or conceal	s material information for the purpo		

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