

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

Page 1 of 5

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

Please indicate: Start of treatment: Start date / /  Continuation of therapy: Date of last treatment / /				plans varies based on indication. See section G below.						
Precertification Re						ne:		Fax:		
A. PATIENT INFOR	MATION									
First Name:				Last Name:				DOB:		
Address:				City:				State:	ZIP:	
Home Phone:	,	Nork Phone:		Ce	Il Phone:			Email:		
Current Weight:	lbs or kgs	Height:	inches or	cms	Allergies:					
B. INSURANCE INF		<u> </u>			J					
Aetna Member ID #:	:		Does p	atient have oth	er coverage?	☐ Yes	□No			
Group #:						Carrier	r Name: _			
Insured:			Insured	d:						
C. PRESCRIBER IN	IFORMATION									
First Name:			Last N	ame:			(Check	One): 🔲 M.D	D. 🗌 D.O. 🗌 N	N.P. □ P.A.
Address:					City:			State:	ZIP:	
Phone:	Fax:		St Lic #	<b>#</b> :	NPI #:		DEA #:		UPIN:	
Provider Email:	•	(	Office Contac	ct Name:	•		Phone:		•	
D. DISPENSING PR	ROVIDER/ADMINIST	RATION INFO	RMATION							
Home Infusion ( Agency Na Administration of Address: City: Phone: TIN: NPI:	ed    Physion Center    F me: Center    F ame: code(s) (CPT):  RMATION - Please	State: Fax: PIN:	ZIP:	requested	□ Special  Name: Address: City: Phone: TIN: NPI:	ian's Office Ity Pharma	cy	Retail P Other: _  State: Fax: _ PIN: _	ZIP:	
Request is for: Infl								HCP	CS Code:	
F. DIAGNOSIS INFO			-					0 - 1 -		
Primary ICD Code: _						O				
G. CLINICAL INFO					its <u>entirety</u> for all	I precertifica	ation requ	iests.		
Tremfya and Z	vio, Remicade, Sin icade, and unbrand Xeljanz/Xeljanz XR as the patient had pri	nponi Aria, and ed infliximab a are preferred for therapy with rial and failure,	d unbranded are preferred or other ind Inflectra (infl intolerance, dalimumab)	I infliximab are d for ulcerative ications. Prefe iximab-dyyb) w or contraindicat  Kevzara (s.	e colitis and Enkerred products vithin the last 365 tion to any of the arilumab)	brel, Humir vary based days? following? tezla (apren	a, Kevza on indic (select al nilast)	ra, Otezla, R ation. I that apply) ] Rinvoq (upa	invoq, Skyrizi	
Please explain if the diagnosis (select all	re are any other med that apply)   Enbrel (etanercept)	dical reason(s) t	that the patie	ent cannot use a	any of the followir	ng preferred tezla (apren	d product	s when indica	ated for the pati	ient's
I 1	Skyrizi (risankizuma	an-rzaa) IISta	elara (listekii	numah) IITra	emtya (duselkum	ıanı IIX≏	ilanz/Xali	anz XR (tofac	ntinih)	

Continued on next page

For Medicare Advantage Part B: Phone: <u>1-866-503-0857</u> (TTY: <u>711</u>)

Note: Inflectra is preferred for MA

FAX: 1-844-268-7263
For other lines of business:

Please use other form.



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

Page 2 of 5

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

For Medicare Advantage Part B: Phone: <u>1-866-503-0857</u> (TTY: <u>711</u>)

1-844-268-7263

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	9	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INF	ORMATION (continued) – Red	quired clinical information must be complete	ed in its <u>entirety</u> for all precertif	cation 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?				
$\Big  \; \Big  \; \longrightarrow \;$	—————————————————————————————————————				
	If positive, Does the patient has	ve latent or active TB? ☐ latent ☐ active ill TB treatment be started before initiation of		nah-dvvh\?	
Ankylosing Spon	dylitis and Other Spondyloart		or therapy with himootia (himali	ab dyyb).	
Please select whi		patient: Ankylosing spondylitis Oth	er spondyloarthropathy		
☐ Yes ☐ No	Is there evidence of inflammato	ry disease?			
		ve response to two or more non-steroidal a	nti-inflammatory drugs (NSAID	s)?	
<b> </b>	Please provide the names and NSAID #1:				
Bahastia Disessa					
Behcet's Disease		icosteroids or immunosuppressive drugs?			
		oids  immunosuppressive drugs			
Behcet's Uveitis		-			
	Is the disease refractory?				
	us/Pulmonary sarcoidosis  Has the patient remained symp	tomatic despite treatment with steroids?			
I T - \	Diagon provide the deily does a	fataraida. Daga ma			
Yes No	Has the patient remained symp Please select: ☐ azathioprine	tomatic despite treatment with immunosupp  Cyclophosphamide methotrexate	oressants? ☐ Other, please explain:		
Crohn's Disease	Does the notiont have a diagnos	ois of fictulizing Crahn's discoss?			
		sis of fistulizing Crohn's disease? atient has been diagnosed with fistulizing Cr	rohn's disease		
☐ Yes ☐ No	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?				
$ \hspace{.05cm} \hspace{.05cm} \longrightarrow \hspace{.05cm}  $		he patient's disease:  mild moderate			
		nt have a documented diagnosis of active C	crohn's disease?		
	☐ abdominal	all signs/symptoms that apply: pain ☐ arthritis ☐ bleeding ☐ diarrhe	a □ internal fistulae □ inte	stinal obstruction	
		perianal disease spondylitis w			
		n's disease symptoms remained active desp			
		all medications that apply:   6-mercaptop			
Hidradenitis Sup		oids- please identify:   prednisone hy	drocortisone	solone U Other:	
		iva:	☐ Hurley stage II (moderate	disease)	
		☐ Hurley stage III (severe disease)	• • •		
☐ Yes ☐ No	Has the patient completed a tria	I of antibiotics?			
	☐ Yes ☐ No Does the patien	nt have a contraindication to oral antibiotics	?		
$ $ $\longrightarrow$	☐ Yes ☐ No Was the treatm	ent with antibiotics ineffective?			
Immune Checkpo	oint Inhibitor- Induced Toxicition	es			
CTLA-4					
Please select drug: ☐ ipilimumab ☐ Other:					
☐ PD-1 Please select drug: ☐ nivolumab ☐ pembrolizumab ☐ Other: ☐ PD-L1					
Please select drug: atezolizumab avelumab Other: 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)?				

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 Medicare Advantage Part B: Phone: <u>1-866-503-0857</u> (TTY: <u>711</u>)

FAX: <u>1-844-268-7263</u> 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	equired clinical information must be comp	leted in its entirety for all precertif	ication requests		
		теква III ко <u>станску</u> гоган ртвостан	iodion roquocio.		
Please indicate the toxicity (check all that apply:)  □ Cardiac  Which life-threatening immune checkpoint inhibitor-induced cardiac toxicities does the patient have?  Please select: □ arrhythmias □ impaired ventricular function □ myocarditis □ pericarditis  Please indicate the severity of the immune checkpoint inhibitor-induced colitis: □ mild □ moderate □ severe  Please indicate which of the following symptoms the patient exhibits: □ 7 or more stools per day over baseline □ ileus □ fever □ None  □ Yes □ No Has the patient been treated with corticosteroids?  Please indicate the corticosteroid name: □					
	ow improvement after 48 hours of cortico	steroids?			
☐ Elevated serum creatinine/acute renal failure Please indicate the severity of the disease: ☐ Severe (creatinine greater than 3 times baseline or greater than 4 mg/dL) ☐ Life-threatening (creatinine greater than 6 times baseline; dialysis indicated)					
☐ None of the above ☐ Yes ☐ No Has the patient been trea	ted with corticosteroids?				
Please indicate the name    Yes   No Did the creatinine level re   Inflammatory arthritis   Yes   No Does the patient have refile     Yes   No Is the patient responding     Pneumonitis   Please indicate the severity of the disease     Yes   No Has the patient been treated     Please indicate the corticular     Yes   No Did the patient show improve     Juvenile Idiopathic Arthritis (Juvenile Rheuman     Please indicate the severity of the patient's disease     Yes   No Is there evidence that the disease     Yes   No Does the patient have clinical of the patient of the patient of the patient's disease     Yes   No Does the patient have clinical of the patient	and length of therapy: Name:	eline after 1 week of treatment widisease  severe disease ents?  anti-inflammatory agents	th corticosteroids?		
Noninfectious Uveitis  Yes No Was the treatment with cortico  Please indicate the corticoster	oid name:				
Yes No Was the treatment with immun	osuppressive drugs (e.g., azathioprine, c	yclosporine, or methotrexate) inef	ffective?		
Yes □ No Does the patient have a documented intolerance to corticosteroids or immunosuppressive drugs?  Please indicate the drug(s) the patient has intolerance to: □ corticosteroids □ immunosuppressive drugs  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 Psoriasis					
Please indicate the severity of the patient's diseated and Yes No Is there evidence that the diseated and Yes No.					
Yes No Is there clinical documentation					
Yes No Is the patient a candidate for s  Please select: phototherapy Please provide the patient's Psoriasis Area and Please indicate the percentage of body surface a  Yes No Does the plaque psoriasis invo	y ☐ systemic therapy ☐ phototherapy Severity Index (PASI) score: area affected by plaque psoriasis:	/ <sub>6</sub>	] genitals		
☐ Yes ☐ No Are systemic	with systemic conventional DMARD(s) no conventional DMARDs contraindicated?    cyclosporine   methotrexate   my	t tolerated?			
Yes ☐ No Was the trial v☐ Yes ☐ No Is phototherap	vith phototherapy not tolerated?	th 11\/∆ light /D1\\/∆\ □ 11\/D#	h coal tar or dithranol		
	☐ Psoralens (methoxsalen, trioxsalen) will ☐ UVB (standard or narrow band) ☐ Ho ial: ☐ Less than 1 month ☐ 1 month	ome UVB   None of the above			



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

Page 4 of 5

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

For Medicare Advantage Part B: Phone: <u>1-866-503-0857</u> (TTY: <u>711</u>)

FAX: <u>1-844-268-7263</u>

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		
			05 0		
G. CLINICAL INFORMATION (continued) – R	equired clinical information must be comp	leted in its <u>entirety</u> for all pre	certification requests.		
Psoriatic Arthritis  ☐ Yes ☐ No Is there evidence that the dise	age is active?				
Yes No Does the patient have <b>axial</b> ps		ammatan, druga (NCAIDa) ir	ooffootius?		
	ment with 2 or more non-steroidal anti-infl le the names and length of treatment:	animatory drugs (NSAIDS) ii	lenective?		
	e the names and length of treatment.				
NSAID #2:					
☐ Yes ☐ No Does the patient have <b>non-ax</b>					
Yes No Does the pati	ent have severe disease at presentation,	defined as severe disability a	at onset with erosive disease involving		
multiple joints		noffootive?			
→ Li Yes Li	No Was the treatment with methotrexate i  → ☐ Yes ☐ No Was treatment wit		or contraindicated?		
		not tolerated    contraind			
			conventional DMARD ineffective?		
		Please select:   cyclophos			
	·		loroquine  leflunomide		
		☐ sulfasalaz	ine  Other, please explain:		
Pyoderma Gangrenosum					
☐ Yes ☐ No Does the patient have a docur	mented diagnosis of refractory pyoderma	gangrenosum?			
Reactive Arthritis (Reiter's syndrome) or Infla	mmatory Bowel Disease Arthritis (Ente	eropathic Arthritis)			
Please select which applies to the patient:   re		flammatory bowel disease ar	thritis (enteropathic arthritis)		
Yes No Was the treatment with method					
	ment with methotrexate not tolerated?				
•	ent have a contraindication to methotrexa	te?			
Yes No Was the treatment with sulfasalazine ineffective?					
Yes No Was the treatment with sulfasalazine not tolerated?					
☐ Yes ☐ No Does the patient have a contraindication to sulfasalazine?					
│    │ Yes    │ No					
Yes No Does the patient have a contraindication to non-steroidal anti-inflammatory drugs (NSAIDs)?					
Please provide the name:		ai anti-iiliaminatory urugs (i	isalbs)!		
Retinal Vasculitis					
☐ Yes ☐ No Was treatment with a convent	ional DMARD ineffective?				
— ☐ Yes ☐ No Was treatmer	nt with a conventional DMARD not tolerate	ed or contraindicated? 🔲 no	t tolerated		
Rheumatoid Arthritis					
Please indicate the severity of the patient's rheumatoid arthritis:   mild   moderate   severe					
Yes No Is there evidence that the disease is active?					
Yes No Will the patient be using Inflectra (infliximab-dyyb) in combination with methotrexate?					
Yes No Was treatment with methotrexate ineffective?					
├────────────────────────────────────					
			oguine		

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 Medicare Advantage Part B: Phone: <u>1-866-503-0857</u> (TTY: <u>711</u>)

FAX: <u>1-844-268-7263</u>

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		
O OLINIOAL INCORMATION (			6		
<ul> <li>G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.</li> <li>Sarcoidosis</li> </ul>					
Yes No Is the disease refractory to con	ticosteroids?				
Ulcerative Colitis					
☐ Yes ☐ No Is the patient hospitalized with					
Please indicate the severity of	the patient's ulcerative colitis:   mild	moderate  severe			
☐ Yes ☐ No Is there evide		innataraida (a.g. budranartiana	mathylproduicalana praduicana)2		
	refractory to immunosuppression with corti No Does the patient require continuous in				
	methylprednisolone, prednisone)?				
	→ Name and dose: Name:				
	Please indicate the route:   Oral	] IV			
Name and d	ose: Name:	Dose:			
	ate the route:				
	t with immunosuppressant agent (e.g., aza No Was treatment with immunosuppressa				
	or contraindicated?	ant agent (c.g., azatmopime, o-m	icroaptopuline) not tolerated		
	→ Please select: ☐ not tolerated ☐ co				
> Please select	ct:	cyclosporine			
☐ Yes ☐ No Was treatmen	t with 5-aminosalicylic acid agents (e.g., b	alsalazide. mesalamine. sulfasa	alazine) ineffective?		
	No Was treatment with 5-aminosalicylic a				
	not tolerated or contraindicated?				
Please selec	→ Please select: ☐ not tolerated ☐ co t: ☐ Colazal (balsalazide) ☐ Ariso, Asa		Rowasa Canasa (mesalamine)		
7 1 15456 56166	☐ Azulfidine (sulfasalazine) ☐ Other		towasa, Sanasa (mesalamme)		
> Please select the symptoms t	he patient exhibit: more than 10 stools		g 🔲 abdominal pain		
	☐ distension ☐ acute	e, severe toxic symptoms, includ	ing fever and anorexia		
For Continuation of Therapy (clinical docume	_				
Please indicate the length of time on Inflectra (in	77 /	floatra (infliximah duuh)?			
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?					
└────────────────────────────────────					
Please enter the results of the TB test: ☐ positive ☐ negative ☐ unknown ☐ 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 in		d there was an add a three to the	h		
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	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.					

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