

MEDICARE FORM

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

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

Virginia (HMO D-SNP) 1-833-280-5224 FAX: PHONE: 1-855-463-0933

For other lines of business: Please use other form.

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

Please indicate:	Start of treatment: Star			1			aries based on indication. tion G below.
Precertification Re	quested By:					Fax	
A. PATIENT INFOR							
First Name:			Last Name:			DOB:	
Address:			City:			State:	ZIP:
Home Phone:	Work P	hone:	Cell	Phone:		Email:	
Current Weight:	lbs_orkgs_Hei	ght: inches or	cms	Allergies:		1	
B. INSURANCE INF	ORMATION	-		-			
Aetna Member ID #	:	Does p	patient have othe	er coverage?	🗌 Yes 🗌 No		
Group #:		lf yes,	If yes, provide ID#: Carrier Name:				
Insured:		Insured	Insured:				
C. PRESCRIBER IN	FORMATION						
First Name:		Last Na	ame:		(Check	-	D. 🗌 D.O. 🗌 N.P. 🗌 P.A.
Address:				City:	I	State:	ZIP:
Phone:	Fax:	St Lic #		NPI #:	DEA #:		UPIN:
Provider Email:		Office Contac	ct Name:		Phone:		
Place of Administr Self-administer Outpatient Infus Center Na Home Infusion of Agency Na Administration of Address: City: Phone: TIN: NPI: E. PRODUCT INFO Request is for: Infl F. DIAGNOSIS INFO	ed Physician's sion Center Phone: me: Center Phone: ame: code(s) (CPT):	Office ZIP: the medication being se: ate primary ICD Code Secondary ICD	requested Frequested and specify any Code:	Physician Specialty Name: Address: City: Phone: Phone: NPI: uency: / other where apple	icable. Other ICD	Retail I Other: State: Fax: PIN: HCF Code:	
For Initiation Reque	ests (clinical documentation	on required for all re	equests):				
preferred for Preferred pro Yes No Ha Yes No Ha Please explain if the diagnosis (select all	yvio, Remicade, and Simp ulcerative colitis and Enb ducts vary based on indic as the patient had prior there as the patient had a trial and Enbrel (etanercept) Skyrizi (risankizumab-rzaa are are any other medical re that apply) Enbrel (etanercept) Kyrizi (risankizumab-rzaa	rel, Humira, Kevzara cation. apy with Inflectra (infl d failure, intolerance, umira (adalimumab))	a, Otezla, Rinvo liximab-dyyb) wit or contraindicati C Kevzara (sa z XR (tofacitinib) ent cannot use ar C Kevzara (sa	q, Skyrizi, and Xe thin the last 365 da on to any of the fo rilumab) ☐ Otez ny of the following rilumab) ☐ Otez	eljanz/Xeľjanz XF ays? llowing? (select a la (apremilast) [preferred product	t are preferr I that apply)] Rinvoq (up s when indic	ed for other indications. adacitinib) ated for the patient's

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – Rea	guired clinical information must be complete	ed 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? → (check all that apply): □ PPD te	e used concomitantly with apremilast, tofac	itinib, or other biologic DMARE y (IGRAs) or chest x-ray within	os (e.g., adalimumab, certolizumab)?
If positive, Does the patient ha	ve latent or active TB? I latent active	of therapy with Inflectra (inflixin	nab-dyyb)?
Ankylosing Spondylitis and Other Spondyloart Please select which of the following applies to the Yes No Is there evidence that the disea	hropathies patient:		
Yes No Is there evidence of inflammato	ry disease?		
Yes No Has the patient had an ineffective Please provide the names and NSAID #1:	length of treatment:	nti-inflammatory drugs (NSAID	s)?
NSAID #2: Behcet's Disease			
Yes No Is the disease refractory to cort Please indicate: Corticoster	icosteroids or immunosuppressive drugs? ids immunosuppressive drugs ig tried:		
Behcet's Uveitis			
Chronic Cutaneous/Pulmonary sarcoidosis	f steroids: Dose:mg tomatic despite treatment with immunosupp		
Crohn's Disease	cyclophosphamide methotrexate	Other, please explain:	
Yes No Does the patient have a diagnost Please indicate how long the patient have a diagnost Please indicate how long the patient have been been been been been been been be	atient has been diagnosed with fistulizing C	rohn's disease:	
Yes No Does the patient have a diagnost Please indicate the severity of t	sis of Crohn's disease? he patient's disease: 🗌 mild 🛛 moderate	severe	
Yes No Does the patier	nt have a documented diagnosis of active C		
	all signs/symptoms that apply: pain arthritis bleeding diarrhe	a 🔲 internal fistulae 🔲 inte	stinal obstruction
🖵 Yes 🗌 No 🛛 Have the Crohi	n ☐ perianal disease ☐ spondylitis ☐ w n's disease symptoms remained active des		
	ds? all medications that apply: ☐ 6-mercaptop oids- please identify: ☐ prednisone		isolono 🗖 Othor:
Hidradenitis Suppurativa			
Please indicate the stage of hidradenitis suppurat	Hurley stage III (severe disease)	 Hurley stage II (moderate of Unknown 	Jisease)
	nt have a contraindication to oral antibiotics	?	
Yes No Was the treatm			
Immune Checkpoint Inhibitor- Induced Toxicitie Please indicate therapy used:	es		
Please select drug: ipilimumab Other: D-1			
Please select drug: I nivolumab pembro	blizumab 🗌 Other:		
 PD-L1 Please select drug: atezolizumab avel Other Please explain: 	lumab 🔲 durvalumab 🔲 Other:		
Yes No Do the immune checkpoint inhit	pitor-induced toxicities persist despite disco p. ipilimumab, nivolumab, pembrolizumab)?		nt inhibitors that target CTLA-4 or



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G CLINICAL INFORMATION (continu	<i>led)</i> – Required clinical information must b	a completed in its entirety for all r	procertification requests	
Please indicate the toxicity (check all		e completed in its <u>entirety</u> for all p	recentification requests.	
		toxicities does the natient have?		
 □ Cardiac Which life-threatening immune checkpoint inhibitor-induced cardiac toxicities does the patient have? Please select: □ arrhythmias □ impaired ventricular function □ myocarditis □ pericarditis □ Colitis 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: 				
	atient show improvement after 48 hours of			
Elevated serum creatinine/acute ren Please indicate the severity of the	al failure e disease:			
	eater than 3 times baseline or greater than			
☐ Life-threatening (crea	tinine greater than 6 times baseline; dialys	is indicated)		
☐ Yes ☐ No Has the patient b	een treated with corticosteroids?			
\rightarrow Please indicate t	he name and length of therapy: Name:	Length:	Less than 1 week 1 week or greater ment with corticosteroids?	
☐ Yes ☐ No Did the creatinine	evel remain greater than 2 to 3 times abo	ove baseline after 1 week of treati	ment with corticosteroids?	
	have refractory or severe disease?	actory disease 🔲 severe diseas	se	
	ponding to corticosteroids or anti-inflamma	itory agents? 🗌 anti-inflammatory	y agents □ corticosteroids	
Pneumonitis	diagona — mild — moderate — agus	ro.		
Yes No Has the patient b	e disease:	ionitis?		
Please indicate t	he corticosteroid name:	atoroida?		
Juvenile Idiopathic Arthritis (Juvenile	Reumatoid Arthritis)	steroids?		
	nt's disease: 🗌 mild 📋 moderate 📋 se	vere		
☐ Yes ☐ No Is there evidence that	the disease is active?			
Yes No Does the patient have	e clinical documentation of polyarticular juv	enile idiopathic arthritis (JRA)?		
Yes No Was treatment with E	nbrel (etanercept) ineffective?			
☐ Yes ☐ No Does the patient have	e a documented intolerance to Enbrel (etan	nercept)?		
	a documented contraindication to Enbrel	(etanercept)?		
Noninfectious Uveitis	h corticosteroids ineffective?			
	orticosteroid name:			
Yes No Was the treatment with Please provide the na	th immunosuppressive drugs (e.g., azathio me:	prine, cyclosporine, or methotrex	ate) ineffective?	
Yes No Does the patient have	a documented intolerance to corticosteroi	ids or immunosuppressive drugs?		
└────────────────────────────────────				
Please indicate the drug(s) the patient has contraindication to: Corticosteroids Controsteroids				
Plaque Psoriasis				
Please indicate the severity of the patient Yes No Is there evidence that	nt's disease: mild moderate	vere		
	nentation 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:				
	surface area affected by plaque psoriasis:			
	iasis involve sensitive areas? <i>If yes</i> , pleas		-	
	temic conventional DMARD(s) (e.g., metho the trial with systemic conventional DMAR) ineffective?	
Yes No Are s	ystemic conventional DMARDs contraindic	cated?		
	tretin cyclosporine methotrexate	mycophenolate None of	the above	
Yes No Was the trial with pho				
	the trial with phototherapy not tolerated?			
	☐ Yes ☐ No Is phototherapy contraindicated?			
Please check all that apply: Please check all that apply: Please check all that apply: VSB (standard or narrow band) Home UVB None of the above				
Please indicate the le	ngth of trial: \Box Less than 1 month \Box 1 n			



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continued) – Re	equired clinical information must be compl	leted in its <u>entirety</u> for all precertit	fication requests.
Please provide NSAID #1:		ammatory drugs (NSAIDs) ineffe	ctive?
\square Yes \square No Does the patient have non-ax			
Yes No Does the patient multiple joints	ent have severe disease at presentation, c ? No Was the treatment with methotrexate in —> ☐ Yes ☐ No Was treatment with —> Please select: ☐ ☐ Yes ☐ No W	neffective? n methotrexate not tolerated or co not tolerated	ontraindicated? d entional DMARD ineffective?
Pyoderma Gangrenosum			
Yes No Does the patient have a docum	nented diagnosis of refractory pyoderma g	gangrenosum?	
Reactive Arthritis (Reiter's syndrome) or Infla			
☐ Yes ☐ No Does the patie ☐ Yes ☐ No Was the treatment with sulfasa ☐ Yes ☐ No Was the treatment with sulfasa ☐ Yes ☐ No Was the treatment with sulfasa ☐ Yes ☐ No Was the treatment with sulfasa ☐ Yes ☐ No Was the treatment with non-step ☐ Yes ☐ No Was the treatment with non-step ☐ Yes ☐ No Was the treatment with non-step	trexate ineffective? ment with methotrexate not tolerated? ent have a contraindication to methotrexat alazine ineffective? ment with sulfasalazine not tolerated? ent have a contraindication to sulfasalazin eroidal anti-inflammatory drugs (NSAIDs) ment with non-steroidal anti-inflammatory	e? e? ineffective? drugs (NSAIDs) not tolerated?	
	ent have a contraindication to non-steroida	, , ,	Ds)?
Retinal Vasculitis			
☐ Yes ☐ No Was treatment with a conventi → ☐ Yes ☐ No Was treatment Rheumatoid Arthritis	t with a conventional DMARD not tolerate		rated 🔲 contraindicated
Please indicate the severity of the patient's rheur		severe	
□ Yes □ No Will the patient be using Inflect □ Yes □ No Was treatment □ Yes □ No Was treatment □ Yes □ Yes □ Yes	tra (infliximab-dyyb) in combination with m it with methotrexate ineffective? No Was treatment with methotrexate not \Rightarrow \Box Yes \Box No Was treatment with a	tolerated or contraindicated?	



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G. CLINICAL INFORMATION (continued) – R	equired clinical information must be	completed in its entirety for all p	recertification requests.	
Sarcoidosis		eenipieted in the <u>entitety</u> for an p		
☐ Yes ☐ No Is the disease refractory to co	rticosteroids?			
Ulcerative Colitis	active fulminant ulcerative colitis?			
	the patient's ulcerative colitis:	ild 🔲 moderate 🔲 severe		
	nce that the disease is active?			
	No Does the patient require continu		tisone, methylprednisolone, prednisone)? orticosteroids (e.g., hydrocortisone,	
	methylprednisolone, prednison	e)?		
	→ Name and dose: Name: Please indicate the route: □ 0			
	lose: Name:	Dose:		
Please indic	ate the route:			
	nt with immunosuppressant agent (e			
	No Was treatment with immunosup or contraindicated?	opressant agent (e.g., azathiopri	ne, 6-mercaptopurine) not tolerated	
	\rightarrow Please select: \Box not tolerated			
Please selection	ct: 🗌 6-mercaptopurine 🔲 azathio	oprine 🗌 cyclosporine		
	nt with 5-aminosalicylic acid agents			
	No Was treatment with 5-aminosal not tolerated or contraindicated		de, mesalamine, sulfasalazine)	
	\rightarrow Please select: \Box not tolerated			
	ct: 🗌 Colazal (balsalazide) 🗌 Ari	so, Asacal, Delzicol, Lialda, Pen	tasa, Rowasa, Canasa (mesalamine)	
	Azulfidine (sulfasalazine)	Other, please explain:		
Please select the symptoms t	he patient exhibit: 🗌 more than 10	stools per day 🔲 continuous b	leeding 🔲 abdominal pain	
☐ distension ☐ acute, severe toxic symptoms, including fever and anorexia				
For Continuation of Therapy (clinical docume Please indicate the length of time on Inflectra (ir				
Yes ☐ No Is this continuation request a l		es of Inflectra (infliximab-dyyb)?		
Yes No Will Inflectra (infliximab-dyyb)	be used concomitantly with apremil			
□ 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				
	the results of the TB test: D positiv			
Yes No Has the patient received Inflect	ctra (infliximab-dyyb) within the past	6 months?		
The previous i		/or potentially life-threatening ad	verse event that occurred during or following	
	o Could the adverse reaction be m	anaged through pre-medication	in the home or office setting?	
For Crohn's disease, Juvenile idiopathic arth				
Please indicate the severity of the disease at ba	seline (pretreatment with inflectra (i	niliximab-dyyb)): 🗋 mild 📋 mo		
H. ACKNOWLEDGEMENT				
Request Completed By (Signature Require		dieel weeeduwe en een die oordi	Date: / /	
Any person who knowingly files a request for insurance company by providing materially insurance act, which is a crime and subjects	false information or conceals ma	aterial information for the purp		

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