

MEDICARE FORM

Renflexis® (infliximab-abda) Injectable **Medication Precertification Request**

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(All fields must be completed and legible for precertification review.) Start of treatment: Start date / _/ Please indicate: Continuation of therapy: Date of last treatment / /

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

For other lines of business: Please use other form.

Note: Renflexis is non-preferred for select indications on MAPD plans. Preferred products vary based on indication. Renflexis is not subject to step therapy on MA plans or for ulcerative colitis on MAPD plans. See section G below.

Precertification Requested	Ву:				Phone):		Fax:	
A. PATIENT INFORMATION									
First Name:				Last	Name:				
Address:				City:				State:	ZIP:
Home Phone:		Worl	k Phone:				Cell Phone:		
DOB:	Allergies:	•					E-mail:		
Current Weight:	lbs or	kgs	Height:		inches o	or	cms		
B. INSURANCE INFORMATIC	N	-	_						
Aetna Member ID #:			Does patient have o	ther of	coverage?	ПΥ	′es 🗌 No		
Group #:			If yes, provide ID#:						
Insured:			Insured:						
C. PRESCRIBER INFORMATI	ON								
First Name:			Last Name:				(Check One): 🗌 M.D. 🗌 D	.O. 🗌 N.P. 🗌 P.A.
Address:			•	C	City:			State:	ZIP:
Phone:	Fax:		St Lic #:	Ν	NPI #:		DEA #:	UP	IN:
Provider Email:		Offi	ice Contact Name:				Phone:	1	
D. DISPENSING PROVIDER/A	DMINISTRATION IN	NFORM/	ATION						
Place of Administration: Self-administered Outpatient Infusion Center Center Name: Home Infusion Center Agency Name: Administration code(s) (Cl Address: City: Phone: TIN: NPI: E. PRODUCT INFORMATION	Phone: PT): State: Fax: PIN:	Z	ZIP:		Name: Address: City: Phone: TIN:	's Off	fice macy	C Retail Pharma	
Request is for: Renflexis (in		ose:		Freq	uency:			HCPCS Co	de:
F. DIAGNOSIS INFORMATION				-				_	
Primary ICD Code:								Code:	
G. CLINICAL INFORMATION – Required clinical information must be completed in its <u>entirety</u> for all precertification requests. <u>For Initiation Requests (clinical documentation required for all requests):</u> Note: Renflexis is non-preferred for select indications on MAPD plans. Enbrel, Humira, Kevzara, Otezla, Rinvoq, Skyrizi, and Xeljanz/Xeljanz XR are the preferred products. Preferred products vary based on indication. Renflexis is not subject to step therapy on MA plans or for ulcerative colitis on MAPD plans.									
 □ Yes □ No Has the patient had prior therapy with Renflexis (infliximab-abda) 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) □ Skyrizi (risankizumab-rzaa) □ Xeljanz/Xeljanz XR (tofacitinib) 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 (co	ntinued) – Required clinical information	on must be completed in its <u>entirety</u> for	all precertification requests.	
			ogic DMARDs (e.g., adalimumab, certolizumab)?	
		erferon-release assay (IGRAs) or chest	x-ray within 6 months of initiation a	
biologic therapy (check all that ar	ç oply): □ PPD test □ interferon-gamn	na assav (IGRA) 🗍 chest x-rav		
	ults of the TB test: positive neg			
	s the patient have latent or active TB?			
		ted before initiation of therapy with Rer	flexis (infliximab-abda)?	
Ankylosing Spondylitis and Othe	r Spondyloarthropathies			
		spondylitis D Other spondyloarthrop	athy	
Yes No Is there evidence				
Yes No Is there evidence				
	had an ineπective response to two or n he names and length of treatment:	nore non-steroidal anti-inflammatory dr	igs (NSAIDs)?	
NSAID #1:				
Behcet's Disease				
	efractory to corticosteroids or immunos	uppressive drugs?		
\square Please indicate:	corticosteroids immunosuppre	ssive drugs		
	he name of drug tried:			
Behcet's Uveitis				
Yes No Is the disease re				
Chronic Cutaneous/Pulmonary S	arcoloosis remained symptomatic despite treatme	nt with storoids?		
	he daily dose of steroids: Dose:			
Yes No Has the patient I	emained symptomatic despite treatme	ent with immunosuppressants?		
Please select:] azathioprine 🔲 cyclophosphamide	🗌 methotrexate 🛛 Other, please e	kplain:	
Crohn's Disease				
	have a diagnosis of fistulizing Crohn's			
	now long the patient has been diagnos			
	have a diagnosis of Crohn's disease? the severity of the patient's disease:			
	Does the patient have a documented of			
	Please select all signs/symptoms that	-		
		oleeding 🔲 diarrhea 🔲 internal fistu	ae 🔲 intestinal obstruction	
	🗌 megacolon 🔲 perianal disease	🗌 spondylitis 🔲 weight loss 🗌 none	of the above	
		emained active despite treatment with	6-mercaptopurine, azathioprine,	
	or corticosteroids?	oly: 🔲 6-mercaptopurine 🛛 azathiop	ine	
/		prednisone hydrocortisone n		
Hidradenitis Suppurativa				
	enitis suppurativa: Hurley stage I	(mild disease)	(moderate disease)	
Yes INo Has the patient of	_ , ,			
	Does the patient have a contraindication	on to oral antibiotics?		
	Was the treatment with antibiotics inef			
Immune Checkpoint Inhibitor- In	duced Toxicities			
Please indicate therapy used:				
CTLA-4: Please select drug:				
		Other:		
Other, please explain:		valumab 🔲 Other:		
	checkpoint inhibitor-induced toxicities r	persist despite discontinuation of immu	ne checkpoint inhibitors that target CTLA-4 or	
	., atezolizumab, ipilimumab, nivolumal		is showpoint initiations that target of En-4 of	



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued	0 – Required clinical information must be	completed in its entirety for all pr	ecertification requests	
		completed in its <u>entirety</u> for all pr	ecertification requests.	
Please indicate the toxicity (check all the	<u>at apply):</u>			
Cardiac	knoint inhibitor induced cordine tovicities	does the nationt have?		
•	kpoint inhibitor-induced cardiac toxicities			
	npaired ventricular function 🛛 myocard			
Colitis	muna abadynaint inhibitar induaad aalitik	y 🗆 mild 🗖 moderate 🗖 equi		
	nmune checkpoint inhibitor-induced colitis			
	g symptoms the patient exhibits:			
	v improvement after 48 hours of corticost		IIe	
Elevated serum creatinine/acute renal fa				
Please indicate the severity of the dis				
	n 3 times baseline or greater than 4 mg/d			
	eater than 6 times baseline; dialysis indica			
\square None of the above				
\Box Yes \Box No Has the patient be	een treated with corticosteroids?			
		Length. [☐ Less than 1 week ☐ 1 week or greater	
	e level remain greater than 2 to 3 times at		-	
☐ Inflammatory arthritis				
	ve refractory or severe disease? 🔲 refra	ctory disease 🛛 severe disease		
	nding to corticosteroids or anti-inflammato			
		,		
_	sease: 🗌 mild 🔲 moderate 🔲 severe			
-	een treated with corticosteroids for pneur			
	ne corticosteroid name:			
	now improvement after 48 hours of cortico			
Juvenile Idiopathic Arthritis (Juvenile Rh				
Please indicate the severity of the patient's		ere		
Yes No Is there evidence that the				
Yes No Does the patient have cli	nical documentation of polyarticular juve	nile idiopathic arthritis (JRA)?		
Yes No Was treatment with Enbr				
Yes No Does the patient have a c		rcept)?		
Yes No Does the patient have a c	documented contraindication to Enbrel (e	etanercept)?		
Noninfectious Uveitis				
Yes No Was the treatment with c	corticosteroids ineffective?			
Please indicate the corticosteroid name:				
☐ Yes ☐ No Was the treatment with ir	mmunosuppressive drugs (e.g., azathiop	rine, cyclosporine, or methotrexat	e) ineffective?	
Please provide the name				
☐ 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				
☐ 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 Psoriasis				
Please indicate the severity of the patient's disease: 🗌 mild 🔲 moderate 🔲 severe				
Yes No Is there evidence that the	e disease is active?			
Yes No Is there clinical documen	tation of chronic disease?			
Yes No Is the patient a candidate for systemic therapy or phototherapy?				
Please select: D phototherapy D systemic therapy D phototherapy and systemic therapy				
Please provide the patient's Psoriasis Area and Severity Index (PASI) score:				
Please indicate the percentage of body surface area affected by plaque psoriasis:%				
☐ Yes ☐ No Does the plaque psoriasis involve sensitive areas? <i>If yes</i> , please select: ☐ hands ☐ feet ☐ face ☐ genitals				
Yes No Was the trial with systemic conventional DMARD(s) (e.g., methotrexate, acetretin, or cyclosporine) ineffective?				
\square Yes \square No Was the trial with systemic conventional DMARD(s) not tolerated?				
	Yes No Are systemic conventional DMARDs contraindicated?			
Please select: 🗌 acitretin 📋 cyclosporine 📋 methotrexate 📋 mycophenolate 📋 None of the above				



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G. CLINICAL INFORMATION (continued) – Re		ted in its <u>entirety</u> for all precertific	ation requests.		
Yes No Was the trial with phototherapy	•				
Please check all that apply:	Psoralens (methoxsalen, trioxsalen) with	LIVA light (PLIVA)			
	UVB with coal tar or dithranol				
-	UVB (standard or narrow-band)				
] Home UVB				
	None of the above				
Please indicate the length of tr	ial: 🗌 Less than 1 month 📋 1 month 📋	2 months 3 months or great	ier		
Psoriatic Arthritis					
☐ Yes ☐ No Is there evidence that the dise	ase is active?				
Yes No Does the patient have axial ps					
	ment with 2 or more non-steroidal anti-inflar	nmatory drugs (NSAIDs) ineffect	ve?		
	e the names and length of treatment:				
NSAID #1:					
Yes No Does the patient have non-ax	ial psoriatic arthritis?				
	ent have severe disease at presentation, de	fined as severe disability at onse	t with erosive disease involving		
multiple joints					
	No Was the treatment with methotrexate ine				
	\longrightarrow Yes \Box No Was treatment with		itraindicated?		
		not tolerated Contraindicated			
		as treatment with another conver			
	$ \longrightarrow P $	ease select: 🗌 cyclophosphami			
		hydroxychloroqu			
Pyoderma Gangrenosum			Other, please explain:		
Yes No Does the patient have a docum	nented diagnosis of refractory pyoderma ga	ingrenosum?			
Reactive Arthritis (Reiter's syndrome) or Infla		•			
Please select which applies to the patient:			(enteropathic arthritis)		
Yes No Was the treatment with metho			,		
└──── ◯ Yes □ No Was the treat	\square Yes \square No Was the treatment with methotrexate not tolerated?				
	Yes No Does the patient have a contraindication to methotrexate?				
□ Yes □ No Was the treatment with sulfasalazine ineffective?					
	ment with sulfasalazine not tolerated?				
Yes No Does the patie	ent have a contraindication to sulfasalazine	?			
Yes □ No Was the treatment with non-ste	eroidal anti-inflammatory drugs (NSAIDs) in	offective?			
	ment with non-steroidal anti-inflammatory d				
	ent have a contraindication to non-steroidal		s)?		
Please provide the name:					
Retinal Vasculitis					
Yes No Was treatment with a conventi	ional DMARD ineffective?				
	t with a conventional DMARD not tolerated	or contraindicated? not tolera	ted Contraindicated		
Rheumatoid Arthritis					
Please indicate the severity of the patient's rheur		severe			
Yes No Is there evidence that the dise		athetroveto?			
Yes No Will the patient be using Renfle		iethotfexate?			
└────────────────────────────────────		lerated or contraindicated?	at tolerated		
└────────────────────────────────────					
Please select: azathioprine hydroxychloroquine leftunomide leftunomide leftunomide leftunomide leftunomide sulfasalazine					
		, , , ,			

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) – R	equired clinical information must be completed	tod in its ontiraty for all procertif	ication requests		
Sarcoidosis		eled in its <u>entirety</u> for all precertif	ication requests.		
Yes No Is the disease refractory to con	rticosteroids?				
Ulcerative Colitis					
☐ Yes ☐ No Is the patient hospitalized with	active fulminant ulcerative colitis?				
	the patient's ulcerative colitis: mild	moderate 🔲 severe			
Yes No Is there evide	nce that the disease is active?				
	refractory to immunosuppression with cortion				
	No Does the patient require continuous im methylprednisolone, prednisone)?	imunosuppression with corticost	eroids (e.g., hydrocortisone,		
	\rightarrow Name and dose: Name:	Dose:			
	Please indicate the route: Oral] IV			
	lose: Name:	Dose:			
· · · · · · · · · · · · · · · · · · ·	cate the route:				
	nt with immunosuppressant agent (e.g., aza	,			
	No Was treatment with immunosuppressa or contraindicated?	ant agent (e.g., azathioprine, 6-m	iercaptopurine) not tolerated		
	$ ightarrow$ Please select: \Box not tolerated \Box co	ontraindicated			
Please select	ct: 🗌 6-mercaptopurine 🔲 azathioprine 🛽 [_ cyclosporine			
	nt with 5-aminosalicylic acid agents (e.g., ba				
$ \qquad \qquad$	No Was treatment with 5-aminosalicylic a	cid agents (e.g., balsalazide, me	salamine, sulfasalazine)		
	not tolerated or contraindicated?				
	\rightarrow Please select: \Box not tolerated \Box co ct: \Box Colazal (balsalazide) \Box Ariso, Asa		owaga Capaga (magalamina)		
	Ariso, Asa		Swasa, Canasa (mesalamme)		
\rightarrow Please select the symptoms the	ne patient exhibit: mere than 10 stools p		abdominal pain		
, · · · · · · · · · · · · · · · · · · ·		, severe toxic symptoms, includi			
For Continuation of Therapy (clinical docume	ntation required for all requests):				
Please indicate the length of time on Renflexis (
Yes No Is this continuation request a result of the patient receiving samples of Renflexis (infliximab-abda)?					
Yes No Will Renflexis (infliximab-abda) 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?					
└────────────────────────────────────					
(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 Renflexis (infliximab-abda) 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 Crohp's disease Invention in the home or office setting?					
For Crohn's disease, Juvenile idiopathic arthritis, Plaque psoriasis, and Rheumatoid arthritis, Ulcerative colitis only: Please indicate the severity of the disease at baseline (pretreatment with Renflexis (infliximab-abda)): mild moderate severe					
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Require	ed):		Date: / /		
Any person who knowingly files a request for					
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.