

Renflexis® (infliximab-abda) Injectable Medication Precertification Request

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

Please indicate:	Start of treatment: Start date // /	_				
	☐ Continuation of therapy: Date of last treatment _	/	'	1		

For Ohio MMP:

FAX: 1-855-734-9389 **PHONE:** 1-855-364-0974

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 Secur	on G below.
Precertification Re	quested By:				Phone:		Fax	
A. PATIENT INFORM	MATION							
First Name:				Last	Name:			
Address:				City	:		State:	ZIP:
Home Phone:		Wor	k Phone:			Cell Phone:		
DOB:	Allergies:					E-mail:		
Current Weight:	lbs or	kgs	Height:		inches or	cms	3	
B. INSURANCE INFO	ORMATION							
Aetna Member ID #	:		Does patient have of	other	coverage?	Yes □ No		
Group #:			If yes, provide ID#: Carrier Name:					
Insured:			Insured:					
C. PRESCRIBER INI	FORMATION							
First Name:			Last Name:			(Check One	e): 🔲 M.D.	☐ D.O. ☐ N.P. ☐ P.A.
Address:				(City:		State:	ZIP:
Phone:	Fax:		St Lic #:	١	NPI #:	DEA #:		UPIN:
Provider Email:	'	Off	ice Contact Name:			Phone:		
D. DISPENSING PRO	OVIDER/ADMINISTRATIO	N INFORM	ATION					
Center Nan Home Infusion C Agency Nan Administration co Address: City: Phone:	d Physician's ion Center Phone: ne: Phone: me: State State Fax: PIN:	:2	ZIP:		Phone:	ffice rmacy	Retail P Other _ Other _ State: Fax: _ PIN: _	ZIP:
Request is for: Ren	nflexis (infliximab-abda)	: Dose:	_	Freq	luency:		НСРО	CS Code:
F. DIAGNOSIS INFO	RMATION - Please indica	ate primary I	CD Code and specify	any c	other where applicable	e		
Primary ICD Code:		Secon	dary ICD Code:			Other ICD (Code:	
G. CLINICAL INFOR	RMATION – Required clinic	al information	on must be completed	in its	entirety for all precer	tification reque	ests.	
For Initiation Reques	sts (clinical documentation	on required	for all requests):					
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? 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 (continued					
☐ Yes ☐ No Will Renflexis (infliximate of the state of t			DMARDs (e.g., adalimumab, certolizumab)? y within 6 months of initiation a		
	PPD test 🔲 interferon-gamma assay (IGRA) ☐ chest x-ray			
	he TB test: ☐ positive ☐ negative ☐				
	tient have latent or active TB? ☐ latent				
	No Will TB treatment be started before	initiation of therapy with Renflexis	; (infliximab-abda)?		
Ankylosing Spondylitis and Other Spon	-				
Please select which of the following applie		S Uther spondyloarthropathy			
Yes ☐ No Is there evidence that th☐ Yes ☐ No Is there evidence of infla					
Yes No Has the patient had an i		teroidal anti-inflammatory drugs (NSAIDs)?		
Please provide the name		tereraan ama mmammater, arage (
Behcet's Disease					
	to corticosteroids or immunosuppressive				
	costeroids immunosuppressive drug				
	e of drug tried:				
Behcet's Uveitis ☐ Yes ☐ No Is the disease refractory	12				
Chronic Cutaneous/Pulmonary Sarcoide					
☐ Yes ☐ No Has the patient remaine		roids?			
Please provide the daily					
Yes No Has the patient remaine					
· ·	oprine 🗌 cyclophosphamide 🔲 metho	otrexate U Other, please explain	1:		
Crohn's Disease	diagnosis of fictulizing Crobn's diagnosis				
Yes No Does the patient have a	g the patient has been diagnosed with fis	tulizina Crohn's disease.			
Yes No Does the patient have a		tuizing Croffit's disease.			
	erity of the patient's disease: mild	moderate ☐ severe			
	e patient have a documented diagnosis of				
	e select all signs/symptoms that apply:				
	dominal pain 🔲 arthritis 🔲 bleeding [
	gacolon 🔲 perianal disease 🔲 spondy				
	e Crohn's disease symptoms remained a costeroids?	active despite treatment with 6-me	rcaptopurine, azathioprine,		
	e check all medications that apply: 🔲 6-m		_		
	ticosteroids- please identify: 🗌 prednisor	ne	Iprednisolone		
Hidradenitis Suppurativa	uppurativa. D Hurlay ataga I (mild diaga		Jorges diagona)		
Please indicate the stage of hidradenitis st	☐ Hurley stage III (severe d		erate disease)		
Yes No Has the patient complete	ed a trial of antibiotics? e patient have a contraindication to oral a	antihi ation?			
Yes No Was the	e treatment with antibiotics ineffective?	anubioucs?			
Immune Checkpoint Inhibitor- Induced					
Please indicate therapy used:	Oxicities				
☐ CTLA-4: Please select drug: ☐ ipilimu	ımab □ Other:				
☐ PD-1: Please select drug: ☐ nivolu	mab pembrolizumab Other:				
☐ PD-L1: Please select drug: ☐ atezolizumab ☐ avelumab ☐ durvalumab ☐ Other:					
Other, please explain:					
Yes No Do the immune checkpo PD-1/PD-L1 (e.g., atezo	oint inhibitor-induced toxicities persist des lizumab, ipilimumab, nivolumab, pembrol	pite discontinuation of immune chizumab)?	eckpoint inhibitors that target CTLA-4 or		

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued	d) – Required clinical information must be	completed in its entirety for all pr	ecertification requests			
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Please indicate the toxicity (check all that apply):						
☐ Cardiac						
	expoint inhibitor-induced cardiac toxicities	does the patient have?				
_	mpaired ventricular function 🔲 myocard					
Colitis	_ ,	_,				
_	mmune checkpoint inhibitor-induced colitis	s: 🗌 mild 📗 moderate 🔲 seve	ere			
Please indicate which of the following	ng symptoms the patient exhibits: 7 or	more stools per day over baseline	e 🗌 ileus 🔲 fever 🔲 None			
☐ Yes ☐ No Has the patient bee	en treated with corticosteroids? <i>If yes,</i> plea	ase indicate the corticosteroid na	me:			
	w improvement after 48 hours of corticost	eroids?				
☐ Elevated serum creatinine/acute renal						
Please indicate the severity of the d						
	n 3 times baseline or greater than 4 mg/d					
	eater than 6 times baseline; dialysis indica	ated)				
□ None of the above	soon trooted with continuatoraids?					
Yes No Has the patient b		Longth: [☐ Less than 1 week ☐ 1 week or greater			
	e level remain greater than 2 to 3 times at					
☐ Inflammatory arthritis	Flover remain greater than 2 to 5 times at	bove baseline after 1 week of trea	then with corticosteroids:			
	ave refractory or severe disease? 🗌 refra	ctory disease				
	nding to corticosteroids or anti-inflammato					
☐ Pneumonitis	-					
	lisease: 🗌 mild 🔲 moderate 🔲 severe					
	peen treated with corticosteroids for pneur					
	he corticosteroid name:					
· ·	how improvement after 48 hours of cortico	osteroids?				
Juvenile Idiopathic Arthritis (Juvenile R						
Please indicate the severity of the patient's		ere				
Yes No Is there evidence that the		nile idionathie arthritic (IDA)?				
☐ Yes ☐ No Does the patient have conditional or Does the Does t		niie idiopatnic artifitis (JRA)?				
Yes No Does the patient have a		ercent)?				
☐ Yes ☐ No Does the patient have a	· ·	• •				
Noninfectious Uveitis	`	1 /				
	☐ Yes ☐ No Was the treatment with corticosteroids ineffective?					
Please indicate the corticosteroid name:						
	Yes No Was the treatment with immunosuppressive drugs (e.g., azathioprine, cyclosporine, or methotrexate) ineffective?					
Please provide the name	e:					
☐ Yes ☐ No Does the patient have a documented intolerance to corticosteroids or immunosuppressive drugs?						
	g(s) the patient has intolerance to: corti					
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		ere				
Yes No Is there clinical document						
│						
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?						
	e trial with systemic conventional DMARD					
	temic conventional DMARDs contraindica					
Please select: ☐ acitret	tin 🗌 cyclosporine 🔲 methotrexate 📋	☐ mycophenolate ☐ None of the	e above			



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB				
C. CLINICAL INFORMATION (southwest). D		And in the autimate for all non-conti	6-4:				
G. CLINICAL INFORMATION (continued) – R		eted in its <u>entirety</u> for all precerti	ication requests.				
1 7 7	│						
Yes No Is phototherap							
	☐ Psoralens (methoxsalen, trioxsalen) with	UVA light (PUVA)					
_	UVB with coal tar or dithranol						
	UVB (standard or narrow-band)						
	☐ Home UVB ☐ None of the above						
	rial: Less than 1 month 1 month	2 months 3 months or gre	ater				
Psoriatic Arthritis		_					
☐ Yes ☐ No Is there evidence that the dise	ease is active?						
Yes No Does the patient have axial ps							
· ·	ment with 2 or more non-steroidal anti-inflar	mmatory drugs (NSAIDs) ineffe	ctive?				
·	e the names and length of treatment:						
<u> </u>							
☐Yes ☐ No Does the patient have non-ax							
	ent have severe disease at presentation, de	efined as severe disability at on	set with erosive disease involving				
multiple joints	·? No Was the treatment with methotrexate inc	offoctive?					
	→ ☐ Yes ☐ No Was treatment with		ontraindicated?				
		not tolerated					
		as treatment with another conv					
	└─── > PI	ease select: cyclophosphar					
			quine leflunomide leflunomide leflunomide				
Pyoderma Gangrenosum							
☐ Yes ☐ No Does the patient have a docur	nented diagnosis of refractory pyoderma ga	angrenosum?					
Reactive Arthritis (Reiter's syndrome) or Infla							
Please select which applies to the patient: re		ammatory bowel disease arthriti	s (enteropathic arthritis)				
Yes No Was the treatment with methotrexate ineffective?							
 Yes ☐ 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 pati	ent have a contraindication to sulfasalazine	?					
☐ Yes ☐ No Was the treatment with non-st	eroidal anti-inflammatory druge (NSAIDe) in	neffective?					
	ment with non-steroidal anti-inflammatory d						
	ent have a contraindication to non-steroidal		Ds)?				
Please provide the name:							
Retinal Vasculitis	to a LDMADD to effect the O						
☐ Yes ☐ No Was treatment with a convent	ional DMARD ineffective <i>?</i> nt with a conventional DMARD not tolerated	or contraindicated? \square not tole	rated				
Rheumatoid Arthritis	it with a conventional blunkto not tolerated	or contraindicated: I not tole	Tated Gontramdicated				
Please indicate the severity of the patient's rheu	matoid arthritis: mild moderate	severe					
Yes No Is there evidence that the dise							
Yes No Will the patient be using Renflexis (infliximab-abda) in combination with methotrexate?							
Yes No Was treatmer		plerated or contraindicated?	not tolerated. Contraindicated				
Yes No Was treatment with methotrexate not tolerated or contraindicated? not tolerated contraindicated tolerated or conventional DMARD (other than methotrexate) ineffective?							
		•	e leflunomide sulfasalazine				

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) – R Sarcoidosis	equired clinical information must be	completed in its <u>entirety</u> for all prece	ertification requests.			
Yes No Is the disease refractory to co	rticosteroide?					
Ulcerative Colitis	rticosteroids :					
Yes No Is the patient hospitalized with	active fulminant ulcerative colitis?					
	f the patient's ulcerative colitis: m	ild ☐ moderate ☐ severe				
	nce that the disease is active?					
☐ Yes ☐ No Is the patient	refractory to immunosuppression wi	th corticosteroids (e.g., hydrocortiso	ne, methylprednisolone, prednisone)?			
	methylprednisolone, prednisone		, -			
		Dose:				
	Please indicate the route: C					
	dose: Name:	Dose:				
	cate the route: Oral IV	a conthibution Construction	in affa ativa 2			
		.g., azathioprine, 6-mercaptopurine) opressant agent (e.g., azathioprine, (
	or contraindicated?	opressant agent (e.g., azatmopine, i	o-mercaptopulme) not tolerated			
	→ Please select: ☐ not tolerated					
	ct: 🗌 6-mercaptopurine 🔲 azathiop					
		e.g., balsalazide, mesalamine, sulfa				
│		icylic acid agents (e.g., balsalazide,	mesalamine, sulfasalazine)			
	not tolerated or contraindicated → Please select: ☐ not tolerated					
		ontraindicated so, Asacal, Delzicol, Lialda, Pentasa	Rowasa Canasa (mesalamine)			
	☐ Azulfidine (sulfasalazine) ☐	Other, please explain:	<u> </u>			
Please select the symptoms the		stools per day	=			
For Continuation of Thorany (clinical document	-	acute, severe toxic symptoms, incli	uding lever and anorexia			
For Continuation of Therapy (clinical docume						
Please indicate the length of time on Renflexis (infliximab-abda): Yes \[\subseteq No \] 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?						
☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐ ☐						
Please enter	the results of the TB test: positive	e	-,			
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						
Yes No Does the pati the previous		or potentially life-threatening advers	e event that occurred during or following			
		anaged through pre-medication in th	e 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)):						
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Require	ed):		Date: /			
Any person who knowingly files a request for	r authorization of coverage of a m	edical procedure or service with t	he intent to injure, defraud or deceive			
any insurance company by providing materia	-		e of misleading, commits a fraudulent			
insurance act which is a crime and subjects	such person to criminal and civil p	enalties				

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