

Renflexis® (infliximab-abda) Injectable Medication Precertification Request

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

	,		3	'		,
Please indicate:	☐ Start of treatment: Start date	1	1			
	☐ Continuation of therapy: Date of	last tre	atment	1	1	_

For Illinois MMP: FAX: 1-855-320-8445

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

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.

							See Section	on o below.
Precertification	Requested By:				Phone:		Fax:	:
A. PATIENT INFO	RMATION							
First Name:				Last N	lame:			
Address:				City:			State:	ZIP:
Home Phone:		Work F	Phone:			Cell Phone:		
DOB:	Allergies:					E-mail:		
Current Weight:	lbs or	kgs	Height: _		inches or	cms		
B. INSURANCE I	NFORMATION							
Aetna Member II	O #:		oes patient have o	ther c	overage?	es No		
Group #:			If yes, provide ID#: Carrier Name:					
Insured:			Insured:					
C. PRESCRIBER	INFORMATION							
First Name:		L	ast Name:			(Check One	:):	☐ D.O. ☐ N.P. ☐ P.A.
Address:		•		Ci	ty:		State:	ZIP:
Phone:	Fax:	S	St Lic #:	NI	기 #:	DEA #:		UPIN:
Provider Email:	,	Office	Contact Name:			Phone:		
D. DISPENSING F	PROVIDER/ADMINISTRATIO	N INFORMAT	TON					
Center N Home Infusior Agency N Administration Address: City: Phone:	ered Physician's usion Center Phone: lame: n Center Phone: Name: n code(s) (CPT): State Fax: PIN:	ZIF	D:		Phone:	fice macy	Retail Pl	ZIP:
Request is for: R	Renflexis (infliximab-abda)	: Dose:		Frequ	ency:		НСРС	S Code:
F. DIAGNOSIS IN	FORMATION – Please indica	ite primary ICE	Code and specify a	any oth	er where applicable	e		
Primary ICD Code	e:	Seconda	ry ICD Code:			Other ICD 0	Code:	
G. CLINICAL INFO	ORMATION – Required clinic	al information	must be completed i	in its <u>e</u>	ntirety for all precert	ification reque	sts.	
	uests (clinical documentation							V II W II 17
	s non-preferred for select in ducts. Preferred products v							
☐ 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 (continued) – Required clinical information must be	completed in its <u>entirety</u> for all pr	ecertification requests.		
Yes No Has the patient been test biologic therapy?	ed for TB with a PPD test, interferon-rele	ease assay (IGRAs) or chest x-ray	DMARDs (e.g., adalimumab, certolizumab)? y within 6 months of initiation a		
	PPD test interferon-gamma assay (I				
	le TB test: ☐ positive ☐ negative ☐ lient have latent or active TB? ☐ latent				
	No Will TB treatment be started before		(inflivimab abda)?		
Ankylosing Spondylitis and Other Spond		illidation of therapy with Nermexis	s (IIIIIIAIIII ab-abda) :		
Please select which of the following applies	•	□ Other spondyloarthropathy			
☐ Yes ☐ No Is there evidence that the		Guier spenayisanun spanny			
☐ Yes ☐ No Is there evidence of inflat					
Yes No Has the patient had an ir	•	teroidal anti-inflammatory drugs (NSAIDs)?		
> Please provide the name		, ,	•		
NSAID #1:					
NSAID1 #2:					
Behcet's Disease					
	to corticosteroids or immunosuppressive				
	costeroids immunosuppressive drugs of drug tried:				
Behcet's Uveitis	or drug tried.				
Yes No Is the disease refractory	2				
Chronic Cutaneous/Pulmonary Sarcoido					
☐ Yes ☐ No Has the patient remained		roids?			
Please provide the daily					
Yes No Has the patient remained					
	oprine 🗌 cyclophosphamide 🔲 metho	trexate U Other, please explain	1:		
Crohn's Disease	diamanda of fictulizione Occidente discussione				
Yes No Does the patient have a	diagnosis of fistulizing Cronn's disease? I the patient has been diagnosed with fist	tulizing Crobn's discoss:			
Yes No Does the patient have a		duizing Cronin's disease.			
	rity of the patient's disease: mild	moderate □ severe			
	e patient have a documented diagnosis o				
	select all signs/symptoms that apply:				
☐ abdominal pain ☐ arthritis ☐ bleeding ☐ diarrhea ☐ internal fistulae ☐ intestinal obstruction					
☐ meg	jacolon 🔲 perianal disease 🔲 spondyl	litis	ne above		
	e Crohn's disease symptoms remained a osteroids?	ctive despite treatment with 6-me	rcaptopurine, azathioprine,		
Please	check all medications that apply: ☐ 6-m	ercaptopurine 🔲 azathioprine			
cort	icosteroids- please identify: 🗌 prednisor	ne hydrocortisone methy	Iprednisolone		
Hidradenitis Suppurativa	_	_			
Please indicate the stage of hidradenitis su	ppurativa: Hurley stage I (mild disea Hurley stage III (severe d		erate disease)		
Yes No Has the patient complete					
	e patient have a contraindication to oral a	antibiotics?			
. — —	treatment with antibiotics ineffective?				
Immune Checkpoint Inhibitor- Induced T	oxicities				
Please indicate therapy used:	mah				
☐ CTLA-4: Please select drug: ☐ ipilimur☐ PD-1: Please select drug: ☐ nivolun	nab Dombrolizumah DOthor:				
☐ PD-1: Please select drug: ☐ nivolumab ☐ pembrolizumab ☐ 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					
☐ Yes ☐ No Do the immune checkpoi	nt inhibitor-induced toxicities persist des izumab, ipilimumab, nivolumab, pembroli		eckpoint inhibitors that target CTLA-4 or		

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C. CLINICAL INFORMATION (continued	d) Required clinical information must be	completed in its entirety for all pr	coortification requests			
G. CLINICAL INFORMATION (continued		completed in its <u>entirety</u> for all pro	ecerunication requests.			
Please indicate the toxicity (check all the	<u>lat apply):</u>					
Cardiac	expoint inhibitor-induced cardiac toxicities	does the nationt have?				
_	mpaired ventricular function					
Colitis	inpaired ventificular function	itis 🔲 pericarditis				
_	mmune checkpoint inhibitor-induced colitis	s:□ mild □ moderate □ seve	are			
1	ng symptoms the patient exhibits: 7 or					
	en treated with corticosteroids? <i>If yes,</i> plea					
	w improvement after 48 hours of corticoste					
☐ Elevated serum creatinine/acute renal f	failure					
Please indicate the severity of the d	isease:					
☐ Severe (creatinine greater tha	n 3 times baseline or greater than 4 mg/d	L)				
☐ Life-threatening (creatinine gre	eater than 6 times baseline; dialysis indica	ated)				
☐ None of the above						
☐ Yes ☐ No Has the patient b	een treated with corticosteroids?					
			Less than 1 week 1 week or greater			
	e level remain greater than 2 to 3 times ab	pove baseline after 1 week of trea	tment with corticosteroids?			
☐ Inflammatory arthritis						
<u> </u>	ave refractory or severe disease? refract	•				
1	nding to corticosteroids or anti-inflammato	ory agents? anti-inflammatory	agents corticosteroids			
Please indicate the severity of the d	isease: mild moderate severe					
	peen treated with corticosteroids for pneun					
	the corticosteroid name:	nonius:				
	how improvement after 48 hours of cortice	osteroids?				
Juvenile Idiopathic Arthritis (Juvenile R		301010100				
Please indicate the severity of the patient's		ere				
☐ Yes ☐ No Is there evidence that th						
☐ Yes ☐ No Does the patient have cl		nile idiopathic arthritis (JRA)?				
Yes No Was treatment with Enb		, ,				
☐ Yes ☐ No Does the patient have a	documented intolerance to Enbrel (etane	rcept)?				
☐ Yes ☐ No Does the patient have a	documented contraindication to Enbrel (e	etanercept)?				
Noninfectious Uveitis						
☐ Yes ☐ No Was the treatment with o						
Please indicate the corti	Please indicate the corticosteroid name:					
Yes No Was the treatment with i		rine, cyclosporine, or methotrexat	e) ineffective?			
Please provide the name	e:					
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		ere				
☐ Yes ☐ No Is there clinical documentation 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:						
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?						
☐ Yes ☐ 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) – R		eted in its <u>entirety</u> for all precertif	ication requests.		
Yes No Was the trial with phototherap					
☐ Yes ☐ No Was the trial No ☐ Yes ☐ No Is phototherap					
	oy contramulcated? ☐ Psoralens (methoxsalen, trioxsalen) with	LIVA light (PLIVA)			
_	UVB with coal tar or dithranol	rova light (rova)			
-	UVB (standard or narrow-band)				
	☐ Home UVB				
	☐ None of the above				
Please indicate the length of to	rial: 🗌 Less than 1 month 🔲 1 month 📋	2 months 3 months or great	ater		
Psoriatic Arthritis					
Yes No Is there evidence that the dise					
Yes No Does the patient have axial ps		(NOAID -) : #	the O		
	ment with 2 or more non-steroidal anti-infla	immatory drugs (NSAIDs) ineffec	tive?		
·	e the names and length of treatment:				
NSAID #1: NSAID #2:					
Yes No Does the patient have non-ax					
Yes No Does the pati	ent have severe disease at presentation, d	efined as severe disability at ons	et with erosive disease involving		
multiple joints		offortivo?			
Tes 🗇	No Was the treatment with methotrexate in → □ Yes □ No Was treatment with		entraindicated?		
		not tolerated contraindicated			
		as treatment with another conve			
		lease select: 🔲 cyclophosphan			
			uine 🗌 leflunomide		
		☐ sulfasalazine	Other, please explain:		
Pyoderma Gangrenosum					
Yes No Does the patient have a docur		•			
Reactive Arthritis (Reiter's syndrome) or Infla			(antaranathia arthritia)		
Please select which applies to the patient: ☐ re ☐ Yes ☐ No Was the treatment with metho		ammatory bower disease artificis	(enteropatine artifitis)		
T T	ment with methotrexate not tolerated?				
☐ Yes ☐ No Does the patient have a contraindication to methotrexate?					
Yes No Was the treatment with sulfasalazine ineffective?					
Yes No Was the treatment with sulfasalazine not tolerated?					
☐ Yes ☐ No Does the pati	ent have a contraindication to sulfasalazine	e?			
Yes No Was the treatment with non-st					
	ment with non-steroidal anti-inflammatory o ent have a contraindication to non-steroida		2012		
Please provide the name:		il anti-illianimatory drugs (NSAIL	JS)!		
Retinal Vasculitis					
Yes No Was treatment with a convent	ional DMARD ineffective?				
	nt with a conventional DMARD not tolerated	d or contraindicated? 🔲 not toler	rated contraindicated		
Rheumatoid Arthritis					
Please indicate the severity of the patient's rheu		severe			
Yes No Is there evidence that the dise					
Yes No Will the patient be using Renfl		nethotrexate?			
Yes No Was treatmer		alarated or contraindicated a	not tolorated. antroindicated		
Yes No Was treatment with methotrexate not tolerated or contraindicated? not tolerated contraindicated tolerated or conventional DMARD (other than methotrexate) ineffective?					
Please select: azathioprine hydroxychloroquine leflunomide sulfasalazine					
	/5455 55.555 🗀 424	,, a. a., yaa. yaani			

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) – R	equired clinical information must be co	mpleted in its entirety for all	nrecertification requests		
Sarcoidosis	oquilou omiliour milermation muct be ec	inplotod in to <u>ottaroty</u> for dir	processinous requests.		
☐ Yes ☐ No Is the disease refractory to co	rticosteroids?				
Ulcerative Colitis					
Yes No Is the patient hospitalized with					
	the patient's ulcerative colitis: mild				
	nce that the disease is active?	corticosteroids (e.g. hydroci	ortisone, methylprednisolone, prednisone)?		
	No Does the patient require continuor	, -	• • • • • • • • • • • • • • • • • • • •		
		• • • • • • • • • • • • • • • • • • • •	, -		
	Name and dose: Name:	Dose:			
Name and	Please indicate the route: U Ora	I ∐ IV			
Name and (nose: Name:	Dose:			
☐ Yes ☐ No. Was treatmen	nt with immunosuppressant agent (e.g.	azathioprine 6-mercaptopu	rine) ineffective?		
	No Was treatment with immunosupp				
	or contraindicated?	_	· · · · ·		
	→ Please select: ☐ not tolerated [
	ct:		sulfasalazine) ineffective?		
	No Was treatment with 5-aminosalic				
	not tolerated or contraindicated?	9 (9,	,		
	→ Please select: ☐ not tolerated [
> Please sele			ntasa, Rowasa, Canasa (mesalamine)		
No. 2012 and and the community was the	☐ Azulfidine (sulfasalazine) ☐ C		blanding.		
Please select the symptoms to	ne patient exhibit: more than 10 sto		oleeding		
For Continuation of Therapy (clinical docume		oute, severe toxic symptoms	, moldaring lever and anorexia		
Please indicate the length of time on Renflexis (
Yes No Is this continuation request a	result of the patient receiving samples				
Yes No Will Renflexis (infliximab-abda		st, tofacitinib, or other biolog	c 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 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 Crohn's disease, Juvenile idiopathic arthritis, Plaque psoriasis, and Rheumatoid arthritis, Ulcerative colitis only:					
Please indicate the severity of the disease at ba					
H. ACKNOWLEDGEMENT		<i>"</i> – –			
THE POSITION OF THE POSITION O					
Request Completed By (Signature Require	ed):		Date: /		
Any person who knowingly files a request fo					
any insurance company by providing materia			rpose of misleading, commits a fraudulent		
insurance act, which is a crime and subjects	such person to criminal and civil per	alties.			

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