

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	- /	<u> </u>
Precertification Re	equested By:		Phone:

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

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

Duo o o utifi ti	Democrated Des				DI			on G below.
	Requested By:				Phone:			•
A. PATIENT INFO	DRMATION			1 1	Maria			
First Name:					Name:		1	
Address:		1		City:		Т	State:	ZIP:
Home Phone:		Work	Phone:			Cell Phone:		
DOB:	Allergies:					E-mail:		
Current Weight: _	lbs or	kgs	Height:		inches or	cms	;	
B. INSURANCE I	NFORMATION							
Aetna Member II	D #:		Does patient have o	other	coverage?	Yes 🗌 No		
Group #:			If yes, provide ID#: Carrier Name:					
Insured:			Insured:					
C. PRESCRIBER	INFORMATION							
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:	'	Offic	ce Contact Name:			Phone:		
D. DISPENSING	PROVIDER/ADMINISTRATIO	N INFORMA	ATION					
Center N Home Infusio Agency Administration Address: City: Phone: TIN: NPI: E. PRODUCT INF	ered Physician's fusion Center Phone: Name: n Center Phone: Name: n code(s) (CPT): State Fax: PIN:	Z	IP:	_	Phone: TIN: NPI:	ffice rmacy	Retail P Other _ State: Fax: _ PIN: _	ZIP:
_	Renflexis (infliximab-abda)				-		HCPC	S Code:
	NFORMATION – Please indica							
•	e:		<u> </u>				·	
G. CLINICAL INF	ORMATION – Required clinic	al informatio	n must be completed	in its	entirety for all precer	tification reque	sts.	
Note: Renflexis i the preferred pro MAPD plans. Yes No Yes No Please explain if t diagnosis (select a	Has the patient had prior therathes the patient had a trial and Enbrel (etanercept) How Skyrizi (risankizumab-rzaa) Chere are any other medical reall that apply).	adications of arry based of apy with Rend failure, intolumira (adalin) Xeljan: ason(s) that all all all and a son(s) that all all all and a son(s)	n MAPD plans. Enbron indication. Renfle flexis (infliximab-abdaterance, or contraindic numab)	xis is a) with cation (sarile nib) e any	not subject to step nin the last 365 days to any of the followin umab)	o therapy on N ? ng? (select all the premilast)	that apply) Rinvoq (upa when indica	for ulcerative colitis on adacitinib)



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued	d) – Required clinical information must be	e completed in its <u>entirety</u> for all pi	recertification requests.	
☐ Yes ☐ No Will Renflexis (infliximab	o-abda) be used concomitantly with apren	nilast, tofacitinib, or other biologic	DMARDs (e.g., adalimumab, certolizumab)?	
	PPD test interferon-gamma assay (
	he TB test: ☐ positive ☐ negative ☐ tient have latent or active TB? ☐ latent			
· — —	No Will TB treatment be started before	initiation of therapy with Renflexis	s (infliximab-abda)?	
Ankylosing Spondylitis and Other Spon Please select which of the following applie		s Other enondyloarthronathy		
Yes No Is there evidence that the		o Guier spondyloartinopatily		
☐ Yes ☐ No Is there evidence of infla				
Yes No Has the patient had an i	•	teroidal anti-inflammatory drugs (NSAIDs)?	
Please provide the name				
NSAID #1:				
NSAID1 #2:				
Behcet's Disease				
Yes No Is the disease refractory	v to corticosteroids or immunosuppressive icosteroids □ immunosuppressive drug			
	e of drug tried:			
Behcet's Uveitis				
☐ Yes ☐ No Is the disease refractory	<i>(</i> ?			
Chronic Cutaneous/Pulmonary Sarcoide	osis			
☐ Yes ☐ No Has the patient remaine	d symptomatic despite treatment with ste	roids?		
Please provide the daily				
Yes No Has the patient remaine	oprine		n·	
Crohn's Disease	opinie cyclophosphamide metric	otresate		
Yes No Does the patient have a	diagnosis of fistulizing Crohn's disease?			
	g the patient has been diagnosed with fis	tulizing Crohn's disease:		
☐ Yes ☐ No Does the patient have a		J		
Please indicate the seve	erity of the patient's disease: mild	moderate severe		
	e patient have a documented diagnosis o	of active Crohn's disease?		
	e select all signs/symptoms that apply:			
	dominal pain			
	gacolon perianal disease spondy	=		
	e Crohn's disease symptoms remained a costeroids?	active despite treatment with 6-me	ercaptopurine, azathioprine,	
	e check all medications that apply: \Box 6-n		_	
	ticosteroids- please identify: 🗌 prednisor	ne	Iprednisolone	
Hidradenitis Suppurativa	uppurativa.	and)	Jarata diaggas)	
Please indicate the stage of hidradenitis st	☐ Hurley stage III (severe o		derate disease)	
Yes No Has the patient complete		and the first of the second		
	e patient have a contraindication to oral a e treatment with antibiotics ineffective?	antibiotics?		
Immune Checkpoint Inhibitor- Induced				
Please indicate therapy used:	Toxicities			
☐ CTLA-4: Please select drug: ☐ ipilimu	ımah □ 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	oint inhibitor-induced toxicities persist des dizumab, ipilimumab, nivolumab, pembro		neckpoint inhibitors that target CTLA-4 or	

Continued on next page



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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				
Please indicate the toxicity (check all the		остіріосов її ко <u>отклосу</u> гог вії рі					
☐ Cardiac	iat appryji						
	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 corti	costeroid name:						
☐ Yes ☐ No Was the treatment with		rine, cyclosporine, or methotrexat	e) ineffective?				
Please provide the name	e:						
Yes No Does the patient have a							
	g(s) the patient has intolerance to: corti						
Yes No Does the patient have a	documented contraindication to corticost	eroids or immunosuppressive dru	gs?				
_	Please indicate the drug(s) the patient has contraindication to: corticosteroids immunosuppressive drugs						
•	Plaque Psoriasis						
Please indicate the severity of the patient's Yes No Is there evidence that the		ere					
☐ Yes ☐ No — Is there evidence that the ☐ 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?							
└────────────────────────────────────							
	temic conventional DMARDs contraindica						
Please select: ☐ acitretin ☐ cyclosporine ☐ methotrexate ☐ mycophenolate ☐ None of the above							



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
C. CLINICAL INFORMATION (constituted)		A in its				
G. CLINICAL INFORMATION (continued) – Re ☐ Yes ☐ No Was the trial with phototherapy		ried in its <u>entirety</u> for all precertific	ation requests.			
Yes No Was the trial vitin phototherapy						
☐ 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					
	ial: Less than 1 month 1 month	1.2 months ☐ 3 months or great	ter			
Psoriatic Arthritis		, e. g. e.				
☐ Yes ☐ No Is there evidence that the dise	ase is active?					
Yes No Does the patient have axial ps	oriatic arthritis?					
	ment with 2 or more non-steroidal anti-inflar	mmatory drugs (NSAIDs) ineffect	ive?			
1	e the names and length of treatment:					
Yes No Does the patient have non-ax	ial psoriatic arthritis?					
1 T - ·	ent have severe disease at presentation, de	efined as severe disability at onse	et with erosive disease involving			
multiple joints		,	3			
└────────────────────────────────────	No Was the treatment with methotrexate inc					
_	→ ☐ Yes ☐ No Was treatment with		ıtraindicated?			
		not tolerated	ational DMARD ineffective?			
		ease select: cyclophosphami				
	,	☐ hydroxychloroqu				
		☐ sulfasalazine ☐	Other, please explain:			
Pyoderma Gangrenosum						
Yes No Does the patient have a docum		~				
Reactive Arthritis (Reiter's syndrome) or Infla Please select which applies to the patient: rea			(antoronathic arthritis)			
Yes No Was the treatment with metho		illillatory bower disease artifitis	(enteropatine artifitis)			
	ment with methotrexate not tolerated?					
	ent 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	neffective?				
	☐ Yes ☐ No Was the treatment with non-steroidal anti-inflammatory drugs (NSAIDs) ineffective? ☐ Yes ☐ No Was the treatment with non-steroidal anti-inflammatory drugs (NSAIDs) not tolerated?					
☐ Yes ☐ No Does the patient have a contraindication to non-steroidal anti-inflammatory drugs (NSAIDs)?						
Please provide the name:						
Retinal Vasculitis						
Yes No Was treatment with a conventi		ar contraindicated? not talors	stad			
☐ Yes ☐ No Was treatment with a conventional DMARD not tolerated or contraindicated? ☐ not tolerated ☐ contraindicated ☐ Rheumatoid Arthritis						
Please indicate the severity of the patient's rheumatoid arthritis: mild moderate severe						
☐ Yes ☐ No Is there evidence that the dise						
☐ Yes ☐ No Will the patient be using Renfle		nethotrexate?				
Yes No Was treatment with methotrexate ineffective?						
☐ Yes ☐ No Was treatment with methotrexate not tolerated or contraindicated? ☐ not tolerated ☐ contraindicated ☐ with another conventional DMARD (other than methotrexate) ineffective?						
Please select: azathioprine hydroxychloroquine leflunomide sulfasalazine						

Continued on next page



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
C. CLINICAL INFORMATION (southwest) B		stad in its autinotes for all monocouti	6 4:			
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Sarcoidosis						
☐ Yes ☐ No Is the disease refractory to con	rticosteroids?					
Ulcerative Colitis						
☐ Yes ☐ No Is the patient hospitalized with	active fulminant ulcerative colitis?					
I I	the patient's ulcerative colitis: mild	moderate severe				
<u> </u>	nce that the disease is active?					
	refractory to immunosuppression with corti No Does the patient require continuous im					
	methylprednisolone, prednisone)?		croids (c.g., flydrocortisone,			
	→ Name and dose: Name:	Dose:				
	Please indicate the route: Oral] IV				
Name and d	lose: Name: ate the route:	Dose:				
	ate the route: Orai IV it with immunosuppressant agent (e.g., aza		effective?			
	No Was treatment with immunosuppressa					
	or contraindicated?		,			
	→ Please select: ☐ not tolerated ☐ co					
	ct: 6-mercaptopurine azathioprine [lazina) ineffective?			
	nt with 5-aminosalicylic acid agents (e.g., ba No Was treatment with 5-aminosalicylic a		The state of the s			
	not tolerated or contraindicated?	old agents (e.g., baloalazide, m	Journal of Sandodiazino)			
	→ Please select: ☐ not tolerated ☐ co					
> Please select	ct: 🔲 Colazal (balsalazide) 🔲 Ariso, Asa		· · ·			
	☐ Azulfidine (sulfasalazine) ☐ Other,					
Please select the symptoms tr	ne patient exhibit:	per day ⊔ continuous bleeding , severe toxic symptoms, includi	· — ·			
For Continuation of Therapy (clinical docume		, severe toxic symptoms, includi	ng lever and anorexia			
Please indicate the length of time on Renflexis (i						
☐ Yes ☐ No Is this continuation request a r	result of the patient receiving samples of Re	enflexis (infliximab-abda)?				
Yes No Will Renflexis (infliximab-abda		facitinib, or other biologic DMAF	RDs (e.g., adalimumab, certolizumab)?			
☐ Yes ☐ No Is there clinical documentation						
☐ 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						
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 baseline (pretreatment with Renflexis (infliximab-abda)): mild moderate severe						
H. ACKNOWLEDGEMENT						
			Dato: / /			
Request Completed By (Signature Require		procedure or semiles with the	Date: /			
Any person who knowingly files a request for	<u> </u>	•				
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.