

Remicade® (infliximab) Injectable **Medication Precertification Request**

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

Virginia (HMO D-SNP) FAX: 1-833-280-5224 **PHONE**: 1-855-463-0933 For other lines of business:

Please use other form.

Note: Remicade is preferred for MA plans. Preferred status for MAPD plans varies based on

	☐ Start of treatment: Start date ☐ Continuation of therapy: Da		1		indication. See section G below.
	uested By:	·			Fax:
A. PATIENT INFORM					
First Name:			Last Name:		
Address:			City:	St	ate: ZIP:
Home Phone:		Work Phone:		Cell Phone:	
DOB:	Allergies:			Email:	
		gs Height:	inches or	r cms	
B. INSURANCE INFO		ngo ricigiti.			
		Does patient have	other coverage?	Yes □ No	
			If yes, provide ID#: Carrier Name: _		
Insured:		Insured:			
C. PRESCRIBER IN	FORMATION				
First Name:		Last Name:		(Check One)	: M.D. D.O. N.P. P.A.
Address:			City:	St	ate: ZIP:
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	UPIN:
Provider Email:		Office Contact Name:		Phone:	<u> </u>
D. DISPENSING PRO	OVIDER/ADMINISTRATION IN	FORMATION			
Place of Administra	ation:		Dispensing Provide	er/Pharmacy:	
☐ Self-administered	d Physician's Office		☐ Physician's Offic	ce 🔲 Retail Phari	macy
	on Center Phone:		☐ Specialty Pharn	nacy 🔲 Mail Order	
	ne:		Other:		
	enter Phone: me:		Name:		
☐ Administration co	ode(s) (CPT):		Address:		
Address:					ZIP:
	State:		_ ·		:
	Fax:				:
	PIN:		NPI:		·
NPI:	MATION Discussion of the second		INFI.		
	RMATION – Please select the m				HODOS Codos
	icade (infliximab) Dose:	<u>'</u>			HCPCS Code:
	RMATION – Please indicate pr				
-					
	MATION – Required clinical info		d in its <u>entirety</u> for all pr	ecertification requests	
•	sts (clinical documentation re				
Note: Remicade, Inflectra, Entyvio, and Simponi Aria are the preferred products for MA plans. For MAPD plans, Remicade, Inflectra, and Entyvio are preferred for ulcerative colitis and Enbrel, Humira, Kevzara, Otezla, Rinvoq, Skyrizi, and Xeljanz/Xeljanz XR are preferred for other indications. Preferred products vary based on indication.					
Yes No Has the patient had prior therapy with Remicade (infliximab) 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)					
Yes No Will Remicade (infliximab) be used concomitantly with apremilast, tofacitinib, or other biologic DMARDs (e.g., adalimumab, certolizumab)? Has the patient been tested for TB with a PPD test, interferon-release assay (IGRAs) or chest x-ray within 6 months of initiation a biologic therapy?					
└────────────────────────────────────					
If positive, Does the patient have latent or active TB? ☐ latent ☐ active					
•	atent TB, Yes No Will T			with Remicade (inflixim	nab)?



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Council Au Int OSEMATION (continued) Resulted clinical information must be completed in its antiroby for all present scation requests. Analysiosing Spondylitis and Other Spondylocarthropathy	Patient First N	lame	Patient Last Name	Patient Phone	Patient DOB	
Presses select with of the following applies to the patient. Ankylosing spondylitis Other spondyloarthropathy Presses select without of the following applies to the patient. Ankylosing spondylitis Other spondyloarthropathy Presses No Is there evidence inflammatory disease?	G. CLINICAL	INFORMATION (continued) - Re	l equired clinical information must be c	completed in its entirety for all	precertification requests.	
Please select which of the following applies to the patient Ankylosing spondylitis Other spondyloarthropathy Yes No Is there evidence that the disease is active?	î e			ompiotod in ito <u>ontiroty</u> for all j	procertimodical requests.	
Please provide the names and length of treatment: NSAID #2	Please select Yes N Yes N	e select which of the following applies to the patient: Ankylosing spondylitis Other spondyloarthropathy S No Is there evidence that the disease is active? S No Is there evidence of inflammatory disease?				
Select's Disease Is the disease refractory to corticosteroids or immunosuppressive drugs	Yes Lin	Please provide the names and length of treatment: NSAID #1:				
Please indicate onticosteroids Immunosuppressive drugs Please provide the name of drug tried: Please provide the daily dose of steroids: Pose:	Behcet's Dise	·				
Yes No No Is the disease refractory?	☐ Yes ☐ N	Yes ☐ No Is the disease refractory to corticosteroids or immunosuppressive drugs? Please indicate: ☐ corticosteroids ☐ immunosuppressive drugs				
Cront's Disease Yes No						
Yes No						
Please provide the daily dose of steroids: Dose:mg Yes No						
Please select azathioprine cyclophosphamide methotrexate Other, please explain: Crohn's Disease? Yes No Does the patient have a diagnosis of fistulizing Crohn's disease? Please indicate how long the patient has been diagnosed with fistulizing Crohn's disease? Please indicate the severity of the patient shade adaptosis of Crohn's disease? Please indicate the severity of the patient stage adocumented diagnosis of active Crohn's disease? Please select all signs/symptoms that apply: abdominal pain antitritis bleeding diarrhea internal fistulae intestinal obstruction megacolon perianal diseases spondylitis weight loss None of the above megacolon perianal diseases spondylitis weight loss None of the above megacolon perianal disease spondylitis weight loss None of the above megacolon perianal disease spondylitis weight loss None of the above megacolon perianal disease spondylitis weight loss None of the above megacolon perianal disease spondylitis weight loss None of the above megacolon perianal disease spondylitis weight loss None of the above megacolon perianal disease spondylitis weight loss None of the above megacolon perianal disease		→ Please provide the daily dose	of steroids: Dose:mg			
Yes No Does the patient have a diagnosis of flatulizing Crohn's disease: Please indicate how long the patient has been diagnosed with fistulizing Crohn's disease: No Does the patient have a diagnosis of Crohn's disease: Please indicate the severity of the patient's disease: Mo Does the patient have a documented diagnosis of active Crohn's disease? Please indicate the severity of the patient have a documented diagnosis of active Crohn's disease? Please select all signs/symptoms that apply: Bodominal pain arthritis bleeding diarrhea internal fistulae		$ ightarrow$ Please select: \square azathioprine			iin:	
Yes			onin of fintulizing Crobn's discoso?			
Yes No Does the patient have a diagnosis of Crohn's disease? mild moderate severe Please indicate the severity of the patient's disease? Please select all signs/symptoms that apply: abdominal pain arthritis bleeding diarrhea internal fistulae intestinal obstruction megacolon perianal disease spondylitis weight loss None of the above megacolon meg				izing Crohn's disease:		
Ves	☐ Yes ☐ N	lo Does the patient have a diagno	osis of Crohn's disease?			
Please select all signs/symptoms that apply:						
abdominal pain arthritis bleeding diarrhea internal fistulae intestinal obstruction megacolon perianal disease spondylitis weight loss None of the above or corticosteroids? Please check all medications that apply: 6-mercaptopurine azathioprine or corticosteroids? Please check all medications that apply: 6-mercaptopurine azathioprine or corticosteroids Please check all medications that apply: 6-mercaptopurine azathioprine or corticosteroids Please indicate the stage of hidradenitis suppurativa: Hurley stage (mild disease) Hurley stage Il (moderate disease) Hurley stage Il (moderate disease) Please indicate the stage of hidradenitis suppurativa: Hurley stage (mild disease) Unknown Please indicate the stage of hidradenitis suppurativa: Hurley stage (mild disease) Unknown Please indicate the stage of hidradenitis suppurativa: Hurley stage (mild disease) Unknown Please indicate the stage of hidradenitis suppurativa: Please stage Unknown Please indicate the duration of the medication to oral antibiotics? Please indicate the duration of the medication trial: Less than 1 month 1 month 1 month 2 months 3 months (90 days) or greater Please indicate therapy used: CTLA-4 Please select drug: ipilimumab Other: Please select drug: ipilimumab Other: Please select drug: atezolizumab avelumab Other: Please edicate drug: atezolizumab avelumab Other: Please indicate the toxicity, (check all that apply): All that apply: Please indicate the toxicity, (check all that apply): Please indicate the toxicity, (check all that apply): Please indicate which of the following symptoms the patient exhibits: 7 or more stools per day over baseline ileus fever None Please indicate which of the following symptoms the patient exhibits: 7 or more stools per day over baseline ileus fever None Please indicate the corticosteroids Please indicate the corticosteroids Please indicate the				active Cronn's disease?		
Yes No Have the Crohn's disease symptoms remained active despite treatment with 6-mercaptopurine, azathioprine, or corticosteroids?				diarrhea	☐ intestinal obstruction	
cr corticosteroids?						
Please check all medications that apply: 6-mercaptopurine azathioprine corticosteroids- please identify: prednisone methylprednisolone Other: Hidradenitis Suppurativa Hurley stage I (mild disease) Hurley stage II (moderate disease) Hurley stage II (m				tive despite treatment with 6-m	nercaptopurine, azathioprine,	
dotation				rcantonurine		
Hidradenitis Suppurativa Please indicate the stage of hidradenitis suppurativa: Hurley stage I (mild disease) Hurley stage II (moderate disease) Hurley stage III (moderate di						
Hurley stage III (severe disease) Unknown Yes No Does the patient completed a trial of antibiotics? Yes No Does the patient have a contraindication to oral antibiotics? Yes No Was the treatment with antibiotics ineffective? Yes No Was the treatment with antibiotics ineffective? Please indicate the duration of the medication trial: Less than 1 month 1 month 2 months 3 months (90 days) or greater 2 months 3 months (90 days) or greater 2 months 3 months (90 days) or greater 3 months (90 days) or greater 4 please select drug: ipilimumab Other: PD-1 Please select drug: nivolumab pembrolizumab Other: PD-1 Please select drug: nivolumab avelumab durvalumab 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 PD-1/PD-L1 (e.g., atezolizumab, ipilimumab, nivolumab, pembrolizumab)? Please indicate the toxicity, (check all that apply): 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. mid moderate severe Please indicate which of the following symptoms the patient exhibits: 7 or more stools per day over baseline ileus fever None Please indicate the severity of the immune checkpoint inhibitor-induced colitis. mid moderate severe Please indicate the teoxicity at the immune checkpoint inhibitor-induced colitis. mid moderate severe Please indicate the teoxicity at the immune checkpoint inhibitor-induced colitis. mid moderate severe Please indicate the corticosteroid anne: Please indicate the corticosteroid anne: Please indicate the corticosteroid anne: Please indicate the corticoste		Suppurativa		-		
Yes No Does the patient have a contraindication to oral antibiotics? Yes No Was the treatment with antibiotics ineffective? Please indicate the duration of the medication trial: Less than 1 month 1 month 2 months 3 months (90 days) or greater 3 months (9			☐ Hurley stage III (severe dise		derate disease)	
Yes				tibiotics?		
Immune Checkpoint Inhibitor-Induced Toxicities 2 months 3 months (90 days) or greater		→ ☐ Yes ☐ No Was the treate	ment with antibiotics ineffective?			
Immune Checkpoint Inhibitor-Induced Toxicities Please indicate therapy used: CTLA-4						
Please indicate therapy used: CTLA-4 Please select drug: ipilimumab Other:	Immune Ched	cknoint Inhibitor-Induced Toxicit	ies	☐ 2 months ☐ 3 months (9	00 days) or greater	
Please select drug: ipilimumab Other: PD-1 Please select drug: nivolumab pembrolizumab Other: PD-L1 Please select drug: atezolizumab avelumab durvalumab Other: Please select drug: atezolizumab avelumab other: Please explain: Please explain: Please explain: Please explain: Please explain: Please indicate the toxicity, (check all that apply): Please indicate the toxicity, (check all that apply): Cardiac Which life-threatening immune checkpoint inhibitor-induced cardiac toxicities does the patient have? Please select: arrhythmias impaired ventricular function myocarditis pericarditis 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 corticosteroids?						
PD-1 Please select drug:	□ CTLA-4					
Please select drug: nivolumab pembrolizumab Other: PD-L1 Please select drug: atezolizumab avelumab durvalumab Other: Other Please explain: Yes No						
□ PD-L1 Please select drug: □ atezolizumab □ avelumab □ 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 PD-1/PD-L1 (e.g., atezolizumab, ipilimumab, nivolumab, pembrolizumab)? Please indicate the toxicity, (check all that apply): □ □ 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?						
Other Please explain: ☐ Yes ☐ No Do the immune checkpoint inhibitor-induced toxicities persist despite discontinuation of immune checkpoint inhibitors that target CTLA-4 or PD-1/PD-L1 (e.g., atezolizumab, ipilimumab, nivolumab, pembrolizumab)? Please indicate the toxicity, (check all that apply): ☐ 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: ☐ Please Indicate The	□ PD-L1					
☐ Yes No Do the immune checkpoint inhibitor-induced toxicities persist despite discontinuation of immune checkpoint inhibitors that target CTLA-4 or PD-1/PD-L1 (e.g., atezolizumab, ipilimumab, nivolumab, pembrolizumab)? Please indicate the toxicity, (check all that apply): Which life-threatening immune checkpoint inhibitor-induced cardiac toxicities does the patient have? ☐ Cardiac Which life-threatening immune checkpoint inhibitor-induced cardiac toxicities does the patient have? ☐ Please select: ☐ arrhythmias ☐ impaired ventricular function ☐ myocarditis ☐ 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: ☐ Indicate the c	☐ Other					
□ 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: □	Yes No Do the immune checkpoint inhibitor-induced toxicities persist despite discontinuation of immune checkpoint inhibitors that target CTLA-4 or					
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:						
Colitis Please indicate the severity of the immune checkpoint inhibitor-induced colitis.	☐ Cardiac					
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:	☐ Colitis				te 🗌 severe	
Please indicate the corticosteroid name:		Please indicate which of the following symptoms the patient exhibits: 🗌 7 or more stools per day over baseline 🗌 ileus 🔲 fever 🔲 None				
Yes No Did the patient show improvement after 48 hours of corticosteroids?						
		Yes No Did the patient sh	ow improvement after 48 hours of co	rticosteroids?		



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Patient F	irst Name	е	Patient Last Name	Patient Phone	Patient DOB
			equired clinical information must be comple	ted in its <u>entirety</u> for all precertifi	cation requests.
		he toxicity, (check all that ap	oly):		
_		m creatinine/acute renal failure			
		icate the severity of the disease			
			es baseline or greater than 4 mg/dL)		
		5 \	an 6 times baseline; dialysis indicated)		
	_	of the above	ested with cortinectoroids?		
		No Has the patient been tree → Please indicate the part	me and length of therapy: Name:	Length: □ Les	s than 1 week
	□ Yes [No Did the creatinine level	me and length of therapy: Name: remain greater than 2 to 3 times above bas	eline after 1 week of treatment v	vith corticosteroids?
	nmatory		g		
	☐ Yes [☐ No Does the patient have re	efractory or severe disease? 🔲 refractory o	lisease	
	☐ Yes [☐ No Is the patient responding	g to corticosteroids or anti-inflammatory age	ents? 🗌 anti-inflammatory agen	ts Corticosteroids
☐ Pneu					
			se: mild moderate severe		
	ا Yes ا		eated with corticosteroids for pneumonitis?		
	□ Yes [No Did the patient show im	provement after 48 hours of corticosteroids	7	
		nic Arthritis (Juvenile Rheum		•	
			ase: mild moderate severe		
			documentation of polyarticular juvenile idio	pathic arthritis (JRA)?	
		Is there evidence that the dise		,	
☐ Yes	☐ No	Was treatment with Enbrel (eta	anercept) ineffective?		
			nented intolerance to Enbrel (etanercept)?		
☐ Yes	☐ No	Does the patient have a docur	nented contraindication to Enbrel (etanerce	pt)?	
Noninfed					
☐ Yes	☐ No	Was the treatment with cortico	steroids ineffective?		
	\longrightarrow	Please indicate the corticoster	oid name:		
□Yes	□ No	Was the treatment with immur	osuppressive drugs (e.g., azathioprine, cyc	closporine or methotrexate) inef	fective?
7		Please provide the name:			
∏ Yes	☐ No 〔	Does the patient have a docur	nented intolerance to corticosteroids or imner patient has intolerance to: corticosteroi	nunosuppressive drugs?	
		Please indicate the drug(s) the	e patient has intolerance to:	ds Immunosuppressive druge?	gs
Tes			e patient has contraindication to: corticosteroids o		2 drugs
Plaque F			patient has contrained and to. Governood	inimunosappressive	, di ago
Please in	ndicate th	ne severity of the patient's disea	ase: mild moderate severe		
		Is there evidence that the dise			
	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 plague 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: ☐ acetretin ☐ cyclosporine ☐ methotrexate ☐ mycophenolate ☐ None of the above					
Yes No Was the trial with phototherapy ineffective?					
☐ Yes ☐ No Was the trial with phototherapy not tolerated?					
☐ Yes ☐ No Is phototherapy contraindicated? Please check all that apply: ☐ 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	2 months	ater
Ì				onaloomonalo or grea	

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) - R	equired clinical information must be comp	leted in its <u>entirety</u> for all prece	ertification requests.		
Psoriatic Arthritis					
☐ Yes ☐ No Is there evidence that the dise	ease is active?				
☐ Yes ☐ No Does the patient have axial ps	soriatic arthritis?				
	ment with 2 or more non-steroidal anti-infl	ammatory drugs (NSAIDs) ine	ffective?		
	e the names and length of treatment:				
NSAID #1:					
NSAID #2:					
Yes No Does the patient have non-ax	ial psoriatic arthritis? ent have severe disease at presentation,	defined as sovere disability at	anast with arasiva diagona involving		
multiple joints		defined as severe disability at	onset with erosive disease involving		
	No Was the treatment with methotrexate i	neffective?			
	→ ☐ Yes ☐ No Was treatment wit	h methotrexate not tolerated o	contraindicated?		
		not tolerated 🔲 contraindica			
			nventional DMARD ineffective?		
		Please select:	<u> </u>		
			roquine leflunomide		
Pyoderma Gangrenosum		☐ suiiasaiazin	e Dther, please explain:		
Yes No Does the patient have a docur	mented diagnosis of refractory pyoderma	rangrenosum?			
Reactive Arthritis (Reiter's syndrome) or Infla	,,,				
Please select which applies to the patient:	•	. ,	ritis (enteronathic arthritis)		
Yes No Was the treatment with metho		lammatory bower discuss artif	mas (orneropauno aramas)		
	ment with methotrexate not tolerated?				
	ent have a contraindication to methotrexa	te?			
☐ Yes ☐ No Was the treatment with sulfas					
	ment with sulfasalazine not tolerated?				
☐ Yes ☐ No Does the patient have a contraindication to sulfasalazine?					
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 convent	ional DMARD ineffective?				
	nt with a conventional DMARD not tolerate	ed or contraindicated? 🔲 not t	olerated		
Rheumatoid Arthritis					
Please indicate the severity of the patient's rheumatoid arthritis: mild moderate severe					
☐ Yes ☐ No Is there evidence that the disease is active?					
Yes No Will the patient be using Remicade (infliximab) in combination with methotrexate?					
Yes No Was treatment with methotrexate ineffective?					
☐ Yes ☐ No Was treatment with methotrexate not tolerated or contraindicated? ☐ not tolerated ☐ contraindicated ☐ Contraindicated ☐ Contraindicated ☐ Contraindicated ☐ Contraindicated					
			iner tnan metnotrexate) ineπective? ine □ leflunomide □ sulfasalazine		
Sarcoidosis	i lease sciect. 🔲 aza	ппортпе Шпучгохуоппогоди			
Vos No. Is the disease refractory to con	ticostoroids?				

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB			
G. CLINICAL INFORMATION (continued) – Required clinical information mus	t be completed in its <u>entirety</u> for all p	precertification requests.			
Ulcerative Colitis ☐ Yes ☐ No ☐ Is the patient hospitalized with active fulminant ulcerative colitis? ☐ Please indicate the severity of the patient's ulcerative colitis: ☐ mild ☐ moderate ☐ severe ☐ Yes ☐ No ☐ Is there evidence that the disease is active? ☐ Yes ☐ No ☐ Is the patient refractory to immunosuppression with corticosteroids (e.g., hydrocortisone, methylprednisolone, prednisone)? ☐ Yes ☐ No ☐ Dose: ☐ Do						
Name a	and dose: Name: Oral	Dose:				
Yes Please	Yes No Was treatment with immunosuppressant agent (e.g., azathioprine, 6-mercaptopurine) ineffective? Yes No Was treatment with immunosuppressant agent (e.g., azathioprine, 6-mercaptopurine) not tolerated or contraindicated? Please select: not tolerated contraindicated Please select: 6-mercaptopurine azathioprine cyclosporine					
Yes No Was treatment with 5-aminosalicylic acid agents (e.g., balsalazide, mesalamine, sulfasalazine) ineffective? Yes No Was treatment with 5-aminosalicylic acid agents (e.g., balsalazide, mesalamine, sulfasalazine) not tolerated or contraindicated? Please select: □ not tolerated □ contraindicated Please select: □ Colazal (balsalazide) □ Ariso, Asacal, Delzicol, Lialda, Pentasa, Rowasa, Canasa (mesalamine) □ Azulfidine (sulfasalazine) □ Other, please explain: □ Please select the symptoms the patient exhibit: □ more than 10 stools per day □ continuous bleeding □ abdominal pain						
	☐ distension ☐ acute, severe toxic symptoms, including fever and anorexia					
For Continuation of Therapy (clinical doc Please indicate the length of time on Remid		<u>sts):</u>				
Yes No Is this continuation request a result of the patient receiving samples of Remicade (infliximab)? Yes No Is there clinical documentation supporting disease stability? Yes No No Is there clinical documentation supporting disease improvement? Yes No No Is there clinical documentation supporting disease improvement? Yes No No Is there clinical documentation supporting disease improvement? Yes No No Is there clinical documentation supporting disease improvement? Yes No No Has the patient had a TB test within the past year? Yes No No Has the patient had a TB test within the past year? Yes No No Has the patient received Remicade (infliximab) within the past 6 months? Yes No No No No No No No N						
For Crohn's disease, Juvenile idiopathic Please indicate the severity of the disease						
H. ACKNOWLEDGEMENT						
Request Completed By (Signature Red	quired):		Date: //			
	ially false information or conceals	material information for the pure	n the intent to injure, defraud or deceive any rpose of misleading, commits a fraudulent			

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