

Page 1 of 10

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

Aetna Precertification Notification Phone: 1-866-752-7021 (TTY: 711)

FAX: <u>1-888-267-3277</u>

For Medicare Advantage Part B: Please Use Medicare Request Form

ontinuation of therapy: Dat	te of last treatment _	/	1			
• •	<u> </u>	·	Phone:		Fay:	
					1 dx	
Ж		Last Name	a·			
			.		State:	ZIP:
W 1 51	0 11 101	City.	000		+	ZIP.
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	inches or	_ cms	Allergies:			
			_			
				er Name: _		
	Insured:					
If yes, provide ID #:	_	Medicaid:	Yes N	lo If yes, p	orovide ID #:	
ATION						
	Last Name:			(Check On	e): 🔲 M.D. 🔲 🗅).O. 🗌 N.P. 🗌 P.A.
		City:			State:	ZIP:
Fax:	St Lic #:	NPI #	# :	DEA #:	UF	PIN:
	Office Contact Nar	me:			Phone:	
☐ Dermatologist ☐ Gast	troenterologist 🔲 R	Rheumatolo	gist 🗌 Other	· ,		
ER/ADMINISTRATION INFOR	RMATION					
☐ Physician's Office enter Phone:		 Nar	Physician's Offi Specialty Pharn ne:	ce nacy	Retail Pharm Other	nacy
		Aut				
) (CPT):		— Pho	one:		Fax:	
) (CPT):		— Pho	one:		Fax:	
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ON e (infliximab) Dose:		Pho TIN	one: : _ Frequency: _		Fax:	
ON e (infliximab) Dose: TION – Please indicate primar	y ICD Code and specify	Pho TIN	one: : _ Frequency: _ /here applicable.		Fax: PIN:	
ON e (infliximab) Dose: TION – Please indicate primar Seco	y ICD Code and specify ondary ICD Code:	Pho TIN	one: l: Frequency: _ /here applicable.	Other ICD (Fax:PIN:	
ON e (infliximab) Dose: TION – Please indicate primar	y ICD Code and specify ondary ICD Code:	Pho TIN	one: l: Frequency: _ /here applicable.	Other ICD (Fax:PIN:	
	Work Phone: s or kgs Height: _ NTION If yes, provide ID #: ATION Fax: Dermatologist Gast R/ADMINISTRATION INFOR	Work Phone: Sorkgs Height:inches or NTION Does patient have If yes, provide ID# Insured: Insured: ATION Last Name: Fax: St Lic #: Office Contact Name Dermatologist Gastroenterologist Fax: Gastroenterologist Fax: Fax: Gastroenterologist Fax: Fax: Fax: Fax: Fax: Fax: Fax: Fax:	Last Name City: Work Phone: Cell Phone: Gorkgs Height:inches orcms ATION Does patient have other cover lf yes, provide ID#:lnsured: Insured: If yes, provide ID #: Medicaid: ATION Last Name: City: Fax: St Lic #: NPI # Office Contact Name: Dermatologist Gastroenterologist Rheumatologist	Last Name: City: DOB: Sorkgs	Last Name: City: Work Phone: DOB: DOB:	Last Name: State: State: Work Phone: Cell Phone: DOB: E-mail: State: Work Phone: Cell Phone: DOB: E-mail: State: DOB: E-mail: State: DOB: E-mail: State: Stat



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Aetna Precertification Notification Phone: <u>1-866-752-7021</u> (TTY: <u>711</u>)

FAX: <u>1-888-267-3277</u>

For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB
Fallent First Name	Falletit Last Name	Fatient Frione	
G. CLINICAL INFORMATION (continue	ed) – Required clinical information must be	completed in its entirety for all prece	ertification requests.
	(a)qu	<u> </u>	, another oquests.
interver severe	an outpatient hospital setting? patient experienced an adverse event with tions (e.g., acetaminophen, steroids, diphe adverse event (anaphylaxis, anaphylactoid ately after an infusion?	enhydramine, fluids, other pre-medic	cations or slowing of infusion rate) or a
☐ Yes ☐ No Does th	patient developed antibodies to infliximable patient have severe venous access issuent hospital setting?		
☐ Yes ☐ No Does the the infu	e patient have significant behavioral issues sion therapy AND the patient does not hav	e access to a caregiver?	ment that would impact the safety of
☐ Yes ☐ No Is the p patient's manage	□ F	le respiratory, cardiovascular, or rena or predispose the patient to a severe te medical personnel and equipment	adverse event that cannot be t?
For Initiation Requests (clinical docum	entation required for all requests):		
Yes No Is the requested quantit literature (e.g., Microme Please select: Suppr	ested for initiation of treatment at a higher different supported by the manufacturer's prescribed by the manufacturer's prescribed by the manufacturer's prescribing information in the control by the manufacturer's prescribing information in the control by dosing guidelines found in the control by the manufacturer's diagnosis?	oing information or dosing guidelines treatment guidelines)? ormation quency supported by the manufactur mpendia or current literature	found in the compendia or current
Acute graft versus host disease			
compendia, current treations of the compendia current treations. Is the requested drug by the compension of the compensi	ty supported by dosing guidelines found in atment guidelines)? eing prescribed by or in consultation with a nced an inadequate response to systemic e patient experienced an intolerance to cors No Does the patient have a contraine	n oncologist or hematologist? corticosteroids? ticosteroids?	e.g., Micromedex DrugDex, NCCN
Ankylosing spondylitis and Non-radio			
Please select which of the following app Yes No Is the requested drug be	, 2 and 6:Please indicate mainte lies to the patient: ☐ Active ankylosing sp eing prescribed by or in consultation with a	ondylitis (AS) Active non-radiog rheumatologist?	raphic axial spondyloarthritis (nr-axSpA)
indicated for active anky	eived or is currently receiving a biologic (e ylosing spondylitis or active non-radiograph e patient experienced an inadequate respo	nic axial spondyloarthritis? onse with at least TWO nonsteroidal	
☐ Yes ☐ No Has the patient had an	intolerance or contraindication to at least ineffective response, contraindication, or in d failed treatment with Avsola (infliximab-a	ntolerance to Simponi Aria?	e to a documented intolerable adverse
event (e.g., rash, nause yes \sum \text{Yes \subseteq} No Was the second of the	ea, vomiting)? Please indicate: ☐ Avsola he adverse event unexpected and not attrit	☐ Inflectra buted to the active ingredient as des	
Behçet's disease	n adverse reaction for both the brand and b	nosimiai medicalion)!	
☐ Yes ☐ No Is the requested quantite compendia, current treated		·	e.g., Micromedex DrugDex, NCCN
Yes No Has the patient ever red (excluding receiving the	eing prescribed by or in consultation with a ceived or is currently receiving Otezla or a drug via samples or a manufacturer's pati e patient had an inadequate response to a cine, systemic glucocorticoids, azathioprine	biologic (e.g., Humira) indicated for t ent assistance program)? t least one nonbiologic medication fo	-



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For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.					
Crohn's disease Please indicate loading dose at weeks 0,	2 and 6: Please indicate mainter	nance dose: frequency	:weeks		
For under 18 years of age only: Yes No Does the prescriber record done?	gnize that a dose above 5 mg per kg is a l	higher dose and the prescriber confi	rms that appropriate monitoring will be		
dose of	e prescribed dose exceed an induction do 10 mg per kg thereafter?	se of 10 mg per kg at week 0, week	2, and week 6, and a maintenance		
All requests: ☐ Yes ☐ No Has the patient been diaç	gnosed with moderately to severely active	Crohn's disease (CD)?			
Yes No Is the requested drug bei	· ,	gastroenterologist?			
Yes No Is the patient 18 years of		- to discation on total control of Fortist	. 0		
	patient had an ineffective response, contraction treatment with Avada (infliving b. a)	•			
Yes No Has the patient tried and	a, vomiting)? Please indicate: 🗌 Avsola		e to a documented intolerable adverse		
` • •	adverse event unexpected and not attrib		ribed in the prescribing information		
	own adverse reaction for both the brand a	_	nibed in the presenting information		
Hidradenitis suppurativa					
Yes No Is the requested quantity compendia, current treati		he compendia or current literature (e	e.g., Micromedex DrugDex, NCCN		
☐ Yes ☐ No Has the patient been diag	gnosed with severe, refractory hidradenitis				
Yes No Is the requested drug bei					
	eceiving the drug via samples or a manufa	cturer's patient assistance program)	?		
	patient experienced an inadequate respor	ntolerance to oral antibiotics?			
☐ Yes ☐ No Has the patient tried and	☐ Yes ☐ No Does the patient				
event (e.g., rash, nausea	, vomiting)? Please indicate: Avsola	☐ Inflectra			
	adverse event unexpected and not attributive dverse reaction for both the brand and bio		ibed in the prescribing information (i.e.,		
Immune checkpoint inhibitor (e.g., CTL					
Yes No Is the requested quantity compendia, current treatr	ment guidelines)?		e.g., Micromedex DrugDex, NCCN		
Yes No Is the requested drug bei					
Yes No Has the patient experience					
	patient experienced an intolerance to cort No Does the patient have a contraind				
/ 🗆 100	Yes No Does the patient have		colitis?		
Immune checkpoint inhibitor (e.g., CTL	, —				
Yes No Is the requested quantity compendia, current treatr		the compendia or current literature (e	e.g., Micromedex DrugDex, NCCN		
☐ Yes ☐ No Does the patient have se	evere disease?				
☐ Yes ☐ No Is the requested drug be					
Yes No Has the patient experien					
	patient experienced an intolerance to corti No Does the patient have a contraind				



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For Medicare Advantage Part B: Please Use Medicare Request Form

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 prece	rtification requests.		
Plaque psoriasis Please indicate loading dose at weeks 0, 2	2 and 6: Please indicate mainte	nance dose: frequency	weeks		
Yes No Has the patient been diag	nosed with moderate to severe plaque pa	soriasis?	weeks		
Yes No Is the requested drug bein					
☐ Yes ☐ No Has the patient ever recei					
	t of moderate to severe plaque psoriasis	(excluding receiving the drug via sar	nples or a manufacturer's patient		
assistance program)? ☐ Yes ☐ No Are crucia	al body areas (e.g., hands, feet, face, ned	ck scaln genitals/groin intertrigingu	s areas) affected?		
	dicate the percentage of body surface are				
	natient experienced an inadequate respon plogic treatment with methotrexate, cyclo		erapy (e.g., UVB, PUVA) or		
└─── Yes [No Does the patient have a clinical re and acitretin?	eason to avoid pharmacologic treatm	ent with methotrexate, cyclosporine		
	Please indicate clinical reason to a				
		r other chronic liver disease Bre			
	☐ Risk of treatment-related toxici	ity 🔲 Significant comorbidity prohib	ce or adverse event Hypersensitivity ts use of systemic agents (e.g., liver or		
☐ Yes ☐ No Has the patient had an ine		uncontrolled hypertension)	er, piease explairi.		
Yes No Has the patient tried and f	ailed treatment with Avsola (infliximab-ax	xxq) or Inflectra (infliximab-dyyb) due	to a documented intolerable adverse		
	vomiting)? Please indicate: Avsolated Avsolate		ihed in the prescribing information (i.e.		
known ad	lverse reaction for both the brand and bio	osimilar medication)?	ibed in the prescribing information (i.e.,		
Psoriatic arthritis with or without co-exis					
Please indicate loading dose at weeks 0, 2 Please indicate which of the following appl		nance dose: trequency:	weeks		
☐ WITH co-existent plaque psoriasis	noo to the patient.				
	s being treated as the primary diagnosis?	•			
Please go to plaque p Yes No Has the patient been diag		12			
☐ Yes ☐ No Is the requested drug bein	ng prescribed by or in consultation with a	rheumatologist or dermatologist?			
☐ Yes ☐ No Has the patient ever recei	ved or is currently receiving a biologic (e.	.g., Humira) or targeted synthetic dru			
indicated for active psorial	tic arthritis (excluding receiving the drug patient have mild to moderate disease?	via samples or a manufacturer's pati	ent assistance program)?		
	□ No Does the patient have severe dis	sease?			
•	patient have enthesitis or predominantly				
Yes No Has the patient had an inadequate response to methotrexate, leflunomide, or another conventional synthetic					
		stered at an adequate dose and dur			
Yes \square No Has the patient had an intolerance to methotrexate, leflunomide, or another conventional					
synthetic drug (e.g., sulfasalazine)? ☐ Yes ☐ No Does the patient have a contraindication to methotrexate or leflunomide?					
		$ ightarrow$ \square Yes \square No Does the patient			
	\downarrow		thetic drug (e.g., sulfasalazine)?		
	•	ate the contraindication:	3 (3 /		
			holic liver disease or other chronic liver		
disease ☐ Drug interaction ☐ Risk of treatment-related toxicity					
☐ Pregnancy or currently planning pregnancy ☐ Breastfeeding					
☐ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension) ☐ Hypersensitivity					
		intolerance or adverse event	,		
☐ Other:					
Yes No Has the patient had an ine					
Yes No Has the patient tried and f			to a documented intolerable adverse		
	, vomiting)? Please indicate: ☐ Avsola adverse event unexpected and not attribu		ibod in the prescribing information (i.e.		
	lverse reaction for both the brand and bic		ibed in the prescribing information (i.e.,		



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For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORM	MATION <i>(continued)</i> – Required clinical info	prmation must be completed in its entirety	for all precertification requests.
Pyoderma gangrenos	<u> </u>		· ·
Yes No Is the		idelines found in the compendia or curren	nt literature (e.g., Micromedex DrugDex, NCCN
	e requested drug being prescribed by or in c	consultation with a dermatologist?	
☐ Yes ☐ No Hast	he patient ever received or is currently rece	iving a biologic (e.g., Humira) indicated fo	or the treatment of pyoderma gangrenosum (excluding
	ving the drug via samples or a manufacturer		1,7 0 0 (0
			or immunosuppressive therapy (e.g., cyclosporine,
	cyclosporine, m	ycophenolate mofetil)?	oids and immunosuppressive therapy (e.g.,
		Does the patient have a contraindication (e.g., cyclosporine, mycophenolate mofet	to corticosteroids and immunosuppressive therapy til)?
Reactive arthritis			
		idelines found in the compendia or currer	nt literature (e.g., Micromedex DrugDex, NCCN
	pendia, current treatment guidelines)?	11.11.11.11.11.11	
	requested drug being prescribed by or in co		the treatment of reactive arthritis (excluding receiving
	rug via samples or a manufacturer's patient		the healthern of reactive artiflus (excluding receiving
	es 🔲 No Has the patient experienced an ir	nadequate response after at least 3 montl	
	than or equal to 15 mg per week	0 mg twice daily or maximally tolerated do	ose, or b) methodiexate at a dose greater
		experienced an intolerance to sulfasalazir	ne and methotrexate?
		Does the patient have a contraindication obstruction)?	to sulfasalazine (e.g., porphyria, intestinal or urinary
	\vdash	☐ Yes ☐ No Does the patient have a	a contraindication to methotrexate?
		Please indicate the contraindication:	
		☐ Clinical diagnosis of alcohol use of	disorder, alcoholic liver disease or other chronic liver
		disease ☐ Drug interaction ☐ Risk	•
		Pregnancy or currently planning p	• , _
		_ • • • • • • • • • • • • • • • • • • •	use of systemic agents (e.g., liver or kidney disease,
		blood dyscrasias, uncontrolled hyper	, _ ,,
		☐ History of intolerance or adverse of	event



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11011e. 1-000-732-7021 (111. /

AX: <u>1-888-267-3277</u>

For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued)) – Required clinical information must be a	completed in its entirety for all pr	ecertification requests	
	required similar mismation must be	ompieted in its <u>entirety</u> for all pr	ecertinodian requests.	
Rheumatoid arthritis Please indicate loading dose at weeks 0, 2	2 and 6: Please indicate mainter	aansa dasa: fragua	nev: wooks	
Yes No Has the patient been diag			ncy:weeks	
☐ Yes ☐ No Is the requested drug being				
☐ Yes ☐ No Has the patient ever recei			drug (e.g. Rinyog Xelianz) indicated	
-	/ active rheumatoid arthritis (excluding red			
program)?	active medinatoid artifilis (excluding rec	serving the drug via samples of a	i manufacturer s patient assistance	
	patient meet either of the following: a) the	e natient was tested for the rheu	matoid factor (RF) biomarker and	
	omarker test was positive, or b) the patie			
	anti-CCP biomarker test was positive?			
└── ☐ Yes	☐ No Has the patient been tested for all	I of the following biomarkers: a)	rheumatoid factor (RF), b) anti-cyclic	
			ind/or erythrocyte sedimentation rate (ESR)?	
	uested medication being prescribed in co			
	dicate a clinical reason for the patient to r			
	cal diagnosis of alcohol use dis <u>or</u> der, alco		_	
	of treatment-related toxicity Pregnar			
	ficant comorbidity prohibits use of system		sease, blood dyscrasias, uncontrolled	
	nsion) 🗌 Hypersensitivity 🗎 History of i	ntolerance or adverse event		
Othe		- Park - Land		
Yes	No Does the patient have other reas	on or no clinical reason not to us	se methotrexate or letiunomide?	
V Ves	☐ No. Has the nationt experienced an in	nadequate response after at leas	st 3 months of treatment with methotrexate	
	at a dose greater than or equal to	15 mg ner week?	of months of treatment with methotrexate	
	Yes \(\subseteq \text{No Has the patient e}	xperienced an intolerance to meth	notrexate?	
	T Yes □ No	Does the patient have a contrain the contraindication:	ndication to methotrexate?	
	─────────────────────────────────────	he contraindication:		
	☐ Clinical dia	agnosis of alcohol use disorder,	alcoholic liver disease or other chronic liver	
	disease 🗌 🛭	Orug interaction 🔲 Risk of treatr	ment-related toxicity	
	<u> </u>	y or currently planning pregnanc	, <u> </u>	
	☐ Significan	t comorbidity prohibits use of sys	stemic agents (e.g., liver or kidney disease,	
	blood dyscras	sias, uncontrolled hypertension)	☐ Hypersensitivity	
		intolerance or adverse event		
. <u> </u>	Other:			
	s the requested medication being prescri			
\hookrightarrow	Please indicate a clinical reason for the p			
	Clinical diagnosis of alcohol use di			
☐ Drug interaction ☐ Risk of treatment-related toxicity ☐ Pregnancy or currently planning pregnancy				
☐ Breastfeeding ☐ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease,				
		ension) Hypersensitivity	History of intolerance or adverse event	
	Other:			
	☐ No clinical reason not to use metho			
☐ Yes ☐ No Has the patient had an inc			described and accommon described and describ	
☐ Yes ☐ No Has the patient tried and			due to a documented intolerable adverse	
event (e.g., rash, nausea, vomiting)? Please indicate: Avsola Inflectra				
Yes No Was the adverse event unexpected and not attributed to the active ingredient as described in the prescribing information (i.e.,				
known ac	dverse reaction for both the brand and bio	isimilar medication)?		



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION	(continued) – Required clinical information	on must be completed in its entirety	for all precertification requests.
Sarcoidosis			
		es found in the compendia or curren	t literature (e.g., Micromedex DrugDex, NCCN
	urrent treatment guidelines)?	eation with a dormatalogist or nulmon	alogist?
	ed drug being prescribed by or in consult		essive therapy (e.g., azathioprine, methotrexate)?
\square res \square No rias the patien	Has the patient experienced an intoler	ance to corticosteroids and immuno	suppressive therapy (e.g., azathioprine, methodexate):
7 L 188 T	methotrexate?		, app. 2001/2 a.e.apy (e.g., a_aa.e.epe,
		a contraindication to corticosteroids	and immunosuppressive therapy (e.g., azathioprine,
	methotrexate?		and minutescappings and apply (e.g., a_aamepinie,
Takayasu's arteritis	motifoli chate.		
	ed quantity supported by dosing quideling	es found in the compendia or curren	t literature (e.g., Micromedex DrugDex, NCCN
compendia. ci	urrent treatment guidelines)?	os lodina in the compendid of current	micratare (e.g., Micromedex Bragbex, 140014
	nt been diagnosed with refractory Takaya	su's arteritis?	
	ed drug being prescribed by or in consult		
		vith corticosteroids or immunosuppre	essive therapy (e.g., methotrexate, azathioprine,
mycophenola			
└── ☐ Yes ☐ No	Has the patient experienced an intoler		suppressive therapy (e.g., methotrexate,
	azathioprine, mycophenolate mofetil)?		
	Yes No Does the patient have		and immunosuppressive therapy (e.g.,
Ulcerative colitis	methotrexate, azatnio	prine, mycophenolate mofetil)?	
	at weeks 0, 2 and 6: Please indi	cate maintenance dose:	frequency: weeks
For under 18 years of age only			
☐ Yes ☐ No Does the pres	criber recognize that a dose above 5 mg	per kg is a higher dose and the pres	scriber confirms that appropriate monitoring
will be done?			
	cribed dose exceed an induction dose of	f 10 mg per kg at week 0, week 2, ar	d week 6, and a maintenance dose of 10 mg per kg
thereafter? All requests:			
	nt been diagnosed with moderately to sev	verely active ulcerative colitis (LIC)?	
	ed drug being prescribed by or in consult		
☐ Yes ☐ No Is the pateint	,	3	
	Has the patient had an ineffective resp	oonse, contraindication, or intoleranc	e to Entyvio?
☐ Yes ☐ No Has the patier	nt tried and failed treatment with Avsola (infliximab-axxq) or Inflectra (inflixima	ab-dyyb) due to a documented intolerable adverse
		ate: 🗌 Avsola 🔲 Inflectra	
└────────────────────────────────────			ent as described in the prescribing information (i.e.,
	known adverse reaction for both the bi	rand and biosimilar medication)?	
Uveitis			UII I D D NOON
		es found in the compendia or curren	t literature (e.g., Micromedex DrugDex, NCCN
	urrent treatment guidelines)? ed drug being prescribed by or in consult	ration with an anhthalmologist or rha	matelogist?
☐ Yes ☐ No Has the natier	at ever received or currently receiving a h	oiologic (e.g. Humira) indicated for th	ne treatment of uveitis (excluding receiving the drug
	r a manufacturer's patient assistance pro		to deathern of dvoids (oxoldding receiving the drug
→ □ Yes □ No	Has the patient experienced an inade	quate response with corticosteroids	or immunosuppressive therapy (e.g., methotrexate,
	azathioprine, mycophenolate mofetil)?)	
	$ egthinspace igspace \square$ Yes $\ \square$ No $\ $ Has the patient exper	ienced an intolerance to corticostero	ids and immunosuppressive therapy (e.g.,
	methotrexate, azathio	prine, mycophenolate mofetil)?	
			to corticosteroids and immunosuppressive therapy
□ Ves □ No. Has the nation	, <u>-</u>	methotrexate, azathioprine, mycoph	enolate molettl)? ab-dyyb) due to a documented intolerable adverse
T	•	te: Avsola Inflectra (milixima	b-dyyb) due to a documented intolerable adverse
			ent as described in the prescribing information (i.e.,
> □ 100 □ M	known adverse reaction for both the bi		s as assorbed in the prosonoring information (i.e.,



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Patient First Name	Patient Last Name	Patient Phone	Patient DOB
G. CLINICAL INFORMATION (continu	ued) – Required clinical information must	be completed in its <u>entirety</u> for all precer	tification requests.
For Continuation Requests (clinical d	locumentation required):		
Please indicate maintenance dose:	weeks		
		amples or a manufacturer's patient assist	
		scribing information or dosing guidelines f	ound in the compendia or current
	nedex DrugDex, NCCN compendia, curre ported by the manufacturer's prescribing		
		l frequency supported by the manufacture	er's prescribing information for the
/ _	patient's diagnosis?	quee, eupperiou e, a.euu.u.u.u	. o processing innormation for the
	pported by dosing guidelines found in the		
•	Yes No Is the supporting informati		
For Crohn's disease, Ulcerative colit	is, Rheumatoid arthritis, Ankylosing s	spondylitis, Non-radiographic axial spo	ndylitis, Psoriatic arthritis,
Plaque psoriasis, Hidradenitis suppu	ırativa, or Uveitis only:		
☐ Yes ☐ No Has the patient tried a	and failed treatment with Avsola (inflixima	ab-axxq) or Inflectra (infliximab-dyyb) due	to a documented intolerable adverse
	isea, vomiting)? Please indicate: 🗌 Av		
		ttributed to the active ingredient as descri	bed in the prescribing information (i.e.,
Know Acute graft versus host disease	n adverse reaction for both the brand an	d biosimilar medication)?	
	being prescribed by or in consultation wi	th an oncologist or hematologist?	
	ienced an inadequate response to system		
	the patient have an intolerance or contra		
Ankylosing spondylitis and Non-radi			
		g spondylitis (AS) 🔲 Active non-radiogra	phic axial spondyloarthritis (nr-axSpA)
	being prescribed by or in consultation wi		
	ved or maintained a positive clinical respo lition since starting treatment with the req	onse as evidenced by low disease activity	or improvement in signs and
	of the following the patient has experien		
		g., morning stiffness)	ve
Behcet's disease		, –	
☐ Yes ☐ No Is the requested drug	being prescribed by or in consultation wit	th a rheumatologist?	
☐ Yes ☐ No Has the patient achiev	ed or maintained a positive clinical respo	onse as evidenced by low disease activity	or improvement in signs and
symptoms of the cond	lition since starting treatment with the req	uested drug?	
Crohn's disease			
	diagnosed with moderately to severely ac		
	being prescribed by or in consultation wi	th a gastroenterologist?	
For under 18 years of age only:	ecognize that a dose above 5 mg per kg	is a higher dose and the prescriber confin	ms that appropriate monitoring will
be done?	scognize that a dose above 5 mg per kg i	is a higher dose and the prescriber comin	ns that appropriate monitoring will
For 18 years of age or older only:			
Please select which applies to this req	uest: Dosage decrease Dosage ir	ncrease 🗌 Continued therapy on current	dose
	for an adult patient following loss of resp		
☐ Yes ☐ No Does the patient requi	ire a dose above 5 mg per kg due to loss	s of response at the current dose?	
All requests:			
Yes No Does the prescribed d			
Yes No Has the patient achiev		45 185 1	45.46
		tive clinical response as evidenced by lov	disease activity or improvement in
	and symptoms of the condition since sta	ining treatment with the requested drug? in dosing regimen due to the patient not	achieving an adequate clinical response
	at the current dose?	e in dosing regimen due to the patient not	achieving an adequate clinical response
Pleas		ent has experienced an improvement in f	rom baseline:
		a ☐ body weight ☐ abdominal mass	
		computed tomography enterography (CT	
		ent on a disease activity scoring tool (e.g	
Index	([CDAI] score)		•
<u>If non</u>	<u>ne of the above applies</u> : ☐ Yes ☐ No	Is this request for an increase in dosing re	egimen due to the patient not achieving
		an adequate clinical response at the curr	ent dose?
Hidradenitis suppurativa	dia	- mikin	
	diagnosed with severe, refractory hidrade being prescribed by or in consultation wi		
		onse as evidenced by low disease activity	or improvement in signs and symptoms
of the condition since	starting treatment with the requested dru	ug?	, , , , , , , , ,
	the patient has experienced since starting		
		uced formation of new sinus tracts and sc	
		in pain from baseline reduction in sup	opuration from baseline
	ses from baseline improvement in qu		
	assessment tool from baseline 🔲 none	ב טו נווכ מטטעכ	



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☐ Yes ☐ No Is the requested drug being prescribed by or in consultation with a rheumatologist?

symptoms of the condition since starting treatment with the requested drug?

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

Aetna Precertification Notification

Phone: <u>1-866-752-7021</u> (TTY: <u>711</u>) FAX: <u>1-888-267-3277</u>

For Medicare Advantage Part B: Please Use Medicare Request Form

Patient DOB Patient First Name Patient Last Name Patient Phone G. CLINICAL INFORMATION (continued) - Required clinical information must be completed in its entirety for all precertification requests Immune checkpoint inhibitor (e.g., CTLA-4, PD-L1 inhibitor) toxicity Yes No Is the requested drug being prescribed by or in consultation with an oncologist or hematologist? \square Yes \square No \square Has the patient experienced an inadequate response to corticosteroids? → ☐ Yes ☐ No Has the patient experienced an intolerance to corticosteroids? → ☐ Yes ☐ No Does the patient have a contraindication to corticosteroids? \longrightarrow \square Yes $\dot{\square}$ No Does the patient have cardiac toxicity? Immune checkpoint inhibitor (e.g., CTLA-4, PD-L1 inhibitor) toxicity- (Immunotherapy arthritis) Yes No Is the requested drug being prescribed by or in consultation with an oncologist or hematologist? ☐ Yes ☐ No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug? Plaque psoriasis or Psoriatic arthritis WITH co-existent plaque psoriasis Yes No Has the patient been diagnosed with moderate to severe plaque psoriasis? Yes No Is the requested drug being prescribed by or in consultation with a dermatologist? Yes No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug? ☐ Yes ☐ No Has the patient experienced a reduction in body surface area (BSA) affected from baseline? → 🔲 Yes 🗍 No Has the patient experienced an improvement in signs and symptoms of the condition from baseline (e.g., itching, redness, flaking, scaling, burning, cracking, pain)? Psoriatic arthritis WITHOUT co-existent plaque psoriasis ☐ Yes ☐ No Is the requested drug being prescribed by or in consultation with a rheumatologist or dermatologist? Yes No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug? Please indicate which of the following the patient has experienced an improvement in from baseline: □ number of swollen joints □ number of tender joints □ dactylitis □ enthesitis □ axial disease □ skin and/or nail involvement ☐ none of the above Pyoderma gangrenosum ☐ Yes ☐ No Is the requested drug being prescribed by or in consultation with a dermatologist? Yes No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug? Reactive arthritis Yes No Is the requested drug being prescribed by or in consultation with a rheumatologist? ☐ Yes ☐ No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and symptoms of the condition (e.g., tender joint count, swollen joint count, or pain) since starting treatment with the requested drug? Rheumatoid arthritis ☐ Yes ☐ No Has the patient been diagnosed with moderately to severely active rheumatoid arthritis (RA)? Yes No Is the requested drug being prescribed by or in consultation with a rheumatologist? ☐ Yes ☐ No Is this a request for a change in dosing regimen? ☐ Yes ☐ No Does the patient require a dose above 3 mg per kg due to an incomplete response at the current dose? Yes No Does the patient require dosing more frequent than every 8 weeks due to an incomplete response at the current dosing frequency? ☐ Yes ☐ No For 18 years of age or older only: Is the requested drug for an adult patient with incomplete response? ☐ Yes ☐ No Does the prescribed dose exceed 10 mg per kg? ☐ Yes ☐ No Has the patient achieved or maintained a positive clinical response since starting treatment with the requested drug? ⇒ ☐ Yes ☐ No Has the patient experienced substantial disease activity improvement (e.g., at least 20% from baseline) in tender joint count, swollen joint count, pain, or disability? ⇒ ☐ Yes ☐ No Is this a request for an increase in dosing regimen due to the patient not achieving an adequate clinical. response at the current dose or frequency? Sarcoidosis ☐ Yes ☐ No Is the requested drug being prescribed by or in consultation with a dermatologist or pulmonologist? ☐ Yes ☐ No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and Symptoms of the condition since starting treatment with the requested drug? Takayasu's arteritis ☐ Yes ☐ No Has the patient been diagnosed with refractory Takayasu's arteritis?

Yes No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and



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

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For Medicare Advantage Part B: Please Use Medicare Request Form

Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G. CLINICAL INFORMATION (continued)	 Required clinical information must be 	completed in its entirety for all prece	rtification requests.		
Ulcerative colitis:					
☐ Yes ☐ No Has the patient been diag	nosed with moderately to severely active	e ulcerative colitis (UC)?			
Yes No Is the requested drug being		gastroenterologist?			
Yes No Has the patient achieved		. But I			
	patient achieved or maintained a positive I symptoms of the condition since startin				
	he following the patient experienced an i				
	P) ☐ fecal calprotectin (FC) ☐ appea				
,	ce enterography (MRE), or intestinal ultra		computed temography emerography		
, ,. ,	ease activity scoring tool (e.g., Ulcerative	_ 0 ,	[UCEIS]. Mayo Score)		
☐ none of the above	, 3 (3,-	,	1 , , ,		
For under 18 years of age only:					
☐ Yes ☐ No Does the prescriber recog	Jnize that a dose above 5 mg per kg is a	higher dose and the prescriber confi	rms that appropriate monitoring		
will be done?					
For 18 years of age or older only: ☐ Yes ☐ No Was the patient on a dose	e exceeding 5 mg per kg as a pediatric p	atient and is continuing that dose into	adulthood?		
	prescriber recognize that a dose above				
monitoring will be done?					
Uveitis			_		
Yes No Is the requested drug bein					
Yes No Has the patient achieved or maintained a positive clinical response as evidenced by low disease activity or improvement in signs and					
	symptoms of the condition since starting treatment with the requested drug?				
Please indicate which of the following the patient has experienced since starting treatment with the requested drug: ☐ reduced frequency of recurrence compared to baseline ☐ decreased reliance on topical corticosteroids					
☐ zero anterior chamber inflammation or reduction in anterior chamber inflammation compared to baseline ☐ none of the above					
·					
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Required): Date:/					
Any person who knowingly files a reque	est for authorization of coverage of a r	nedical procedure or service with t	he intent to injure defraud or deceive		
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive 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 subj			g,		

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