

Avsola™ (infliximab-axxq) Injectable Medication Precertification Request

Page 1 of 5

(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: Avsola is non-preferred. Preferred products vary based on indication and plan type.

Please indicate: Start	·				See section G	below.
☐ Conti	nuation of therapy: Date o	of last treatment	<u> </u>			
Precertification Requested	Ву:		Phone:		Fax:	
A. PATIENT INFORMATION						
First Name:			Last Name:			
Address:			City:		State:	ZIP:
Home Phone:	Worl	k Phone:		Cell Phone:		
DOB:	Allergies:			E-mail:		
Current Weight:	lbs orkgs	Height: _	inches or	cms	i	
B. INSURANCE INFORMATIO		ı				
Aetna Member ID #:		Does patient have o	_	☐ Yes ☐ No		
Group #:			(Carrier Name:		
Insured:		Insured:	_			
Medicare: ☐ Yes ☐ No If		_	Medicaid: Yes	☐ No If yes, pr	ovide ID #:	
C. PRESCRIBER INFORMATION	ON	L t N		(0)	\	
First Name:		Last Name:	la:	(Check One	1	O.
Address:	<u></u>	T	City:	1	State:	ZIP:
Phone:	Fax:	St Lic #:	NPI #:	DEA #:	UPI	IN:
Provider E-mail:		Office Contact Nam			Phone:	
Specialty (Check one):	<u> </u>		neumatologist 🔲 O	ther:		
D. DISPENSING PROVIDER/A	DMINISTRATION INFORM	ATION				
Place of Administration:	_				y: Patient Select	
	☐ Physician's Office		-		☐ Retail Pharma	-
Outpatient Infusion Center Center Name:				-	Other	
	Phone:					
Agency Name: Address:						
☐ Administration code(s) (CPT					State:	
Address:	04-4-	710			Fax:	
City:Phone:					PIN:	
TIN:			— NPI:			
NPI:			<u> </u>			
E. PRODUCT INFORMATION						
Request is for: Avsola (inflix	kimab-axxq) Dose:		Frequenc	cy:		
F. DIAGNOSIS INFORMATION	I – Please indicate primary l	CD Code and specify	any other where applica	able.		
Primary ICD Code:	Second	dary ICD Code:		Other ICD C	Code:	
G. CLINICAL INFORMATION – Required clinical information must be completed in its entirety for all precertification requests.						
For All Requests (clinical documentation required for all requests):						
Note: Avsola is non-preferred. The preferred products for MA plans are Entyvio, Inflectra, Remicade, and Simponi Aria. For MAPD plans, Inflectra, Entyvio, and Remicade, are preferred for ulcerative colitis and Enbrel, Humira, Kevzara, Otezla, Rinvog, 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 Avsola (infliximab-axxq) 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)						
☐ Entyvio (vedolizumab) ☐ Inflectra (infliximab-dyyb) ☐ Remicade (infliximab) ☐ Simponi Aria (golimumab) ☐ 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 the apply)						
☐ Entyvio (vedolizumab) ☐ Inflectra (infliximab-dyyb) ☐ Remicade (infliximab) ☐ Simponi Aria (golimumab)						
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Page 2 of 5

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Patient First Name	Patient Last Name	Patient Phone	Patient DOB	
G. CLINICAL INFORMATION (continued)				
Please explain if there are any other medical diagnosis (select all the apply)	al reason(s) that the patient cannot use a	any of the following preferred proc	lucts when indicated for the patient's	
• • • • • • • • • • • • • • • • • • • •	☐ Humira (adalimumab) ☐ Kevzara (s	arilumab) 🔲 Otezla (apremilast)) ☐ Rinyog (upadacitinib)	
	rzaa) 🔲 Xeljanz/Xeljanz XR (tofacitinib			
-				
Yes No Will the requested drug be (e.g., Olumiant, Xeljanz)?	used in combination with any other biolo	gic or targeted synthetic disease-	modifying anti-rheumatic drug (DMARD)	
☐ Yes ☐ No Has the patient received a				
		, interferon-release assay (IGRA)	or chest x-ray within 6 months of initiating	
a biologic → (Check al	merapy <i>?</i> I that apply):	gamma assay (IGRA) 🔲 chest :	x-rav	
Please er	nter the results of the TB test: positive	negative 🔲 unknown		
	e, Does the patient have latent or active			
ii iatent i	B , ☐ Yes ☐ No Has treatment for late	eatment initiated		
	patient have risk factors for TB?	_	•	
☐ Yes ☐	No Has the patient been tested for tul	berculosis (TB) within the previou	s 12 months?	
	→ (Check all that apply): ☐ PPD tes Please enter the results of the TB			
	> If positive, Does the patient have			
			(TB) infection been initiated or completed?	
For Initiation Requests:	Ple	ase select: treatment initiated	☐ treatment completed	
Ankylosing spondylitis or axial spondylo	parthritis			
Please select which of the following applies	to the patient: Active ankylosing spo		ndyloarthritis	
Yes No Has the patient previously			dal auti inflammatam dunina (NCAIDa) au	
	patient experienced an inadequate respontolerance or contraindication to at least		dal anti-inflammatory drugs (NSAIDs), or	
Please indicate the preferred alternatives for	or ankylosing spondylitis (AS) or axial spo		ffective, not tolerated, or are contraindicated:	
☐ Cosentyx ☐ Enbrel ☐ Humira ☐ R	temicade 🔲 Simponi Aria			
Behçet's syndrome	to the end of the treets	ment of Reheat's discours?		
☐ Yes ☐ No Has the patient received O → ☐ Yes ☐ No Has the			on for Behcet's disease (e.g., colchicine.	
Crohn's disease				
Yes No Has the patient been diagnosed with moderately to severely active Crohn's disease (CD)?				
☐ Yes ☐ No Does the patient have fistulizing Crohn's disease? ☐ Yes ☐ No Has the patient previously received a biologic indicated for moderately to severely active Crohn's disease?				
	☐ No Has the patient tried and had an			
T	Yes No Does the par	tient have a contraindication or in	tolerance to at least one conventional	
	therapy option	on (e.g.,azathioprine [Azasan, Imu	uran], budesonide [Entocort EC],	
			hol], methylprednisolone [Solu-Medrol], one, sulfasalazine [Azulfidine, Sulfazine],	
	rifaximin [Xif	axan], tacrolimus)?	•	
→ Please select: ☐ Sulfasalazine (Azulfidine, Sulfazine) ☐ Metronidazole (Flagyl) ☐ Ciprofloxacin				
(Cipro) ☐ Prednisone ☐ Budesonide (Entocort EC) ☐ Azathioprine (Azasan, Imuran) ☐ Mercaptopurine (Purinethol) ☐ Methotrexate ☐ Methylprednisolone (Solu-Medrol)				
☐ Rifaximin (Xifaxan) ☐ Tacrolimus				
Please indicate the preferred alternatives for Crohn's disease that have been ineffective, not tolerated, or are contraindicated:				
☐ Humira ☐ Entyvio ☐ Remicade ☐ Stelara (intravenous formulation)				
Granulomatosis with polyangiitis (Wegener's granulomatosis)				
Yes No Has the patient experienced an inadequate response with corticosteroids or immunosuppressive therapy (e.g., cyclophosphamide, azathioprine, methotrexate, or mycophenolate mofetil)?				
azatnioprine, methotrexate, or mycophenolate moretti)? ———————————————————————————————————				
azathioprine, methotrexate, or mycophenolate mofetil)?				
Yes No Does the patient have a contraindication to corticosteroids and immunosuppressive therapy (e.g., cyclophosphamide, azathioprine, methotrexate, or mycophenolate mofetil)?				
	(c.g., cyolophosphaniue, aza	annopinio, incurouexate, or myco	priorioiate moretiny:	



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Page 3 of 5

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G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests.				
Hidradenitis suppurativa				
☐ Yes ☐ No Has the patient been diagn	osed with severe, refractory hidradenitis	suppurativa?		
☐ Yes ☐ No Has the patient previously				
	patient experienced an inadequate resp			
Y (es \(\sum \) No Has the patient experienced	an intolerable adverse effect to of patient have a contraindication to		
☐ Yes ☐ No Has the patient had an inef			oral artiblotics :	
Juvenile idiopathic arthritis	,			
☐ Yes ☐ No Has the patient previously i	received a biologic indicated for juvenile	idiopathic arthritis?		
	patient experienced an inadequate respo			
			nt with corticosteroids (e.g., prednisone,	
тетлую ☐ Yes ☐ No Has the patient had an inef			east 3 months of treatment with leflunomide	
Yes No Has the patient had an inef	•			
Immune checkpoint inhibitor toxicity	receive respondes, some amaication or mix	noralise to Elibroi.		
Yes No Has the patient experience	d an inadequate response to corticoster	oids?		
Yes No Does the				
Plaque psoriasis				
☐ Yes ☐ No Has the patient been diagn	osed with chronic, severe plaque psoria	sis?		
☐ Yes ☐ No Has the patient previously i				
	oody surface area (BSA) affected (prior t	o starting the requested medication	on)?	
Please select: ☐ Less th		hands feet face neck scalp de	nitals/groin, intertriginous areas) affected?	
	than or equal to 3% of BSA	Tiarrae, reet, raee, rieek, eearp, ge	maio, grom, mora ignicae areae, anosiea.	
	patient experienced an inadequate respo		totherapy (e.g., UVB, PUVA) or	
pharmac	ologic treatment with methotrexate, cycl	osporine or acitretin?		
Yes	Yes No Does the patient have a clinical reason to avoid pharmacologic treatment with methotrexate, cyclosporine and acitretin?			
		nt have severe psoriasis that warr	ants a biologic DMARD as first-line	
therapy (i.e. at least 10% of the body surface area (BSA) or crucial body areas (e.g., hands,				
		k, scalp, genitals/groin, intertrigino	,	
			Alcoholism, alcoholic liver disease or	
other chronic liver disease Breastfeeding Cannot be used due to risk of treatment-related toxicity				
☐ Drug interaction with traditional systemic agent ☐ Pregnancy or planning pregnancy ☐ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled				
hypertension)				
☐ Other reason to avoid pharmacologic treatment				
Yes No Does the patient have severe psoriasis that warrants a biologic DMARD as first-line				
therapy (i.e. at least 10% of the body surface area (BSA) or crucial body areas (e.g.,				
hands, feet, face, neck, scalp, genitals/groin, intertriginous areas) are affected)? Please indicate the preferred alternatives for plaque psoriasis that have been ineffective, not tolerated, or are contraindicated:				
☐ Humira ☐ Ilumya ☐ Otezla ☐ Remicade ☐ Skyrizi ☐ Stelara ☐ Taltz ☐ Tremfya				
Psoriatic arthritis				
☐ Yes ☐ No Has the patient been diagnosed with active psoriatic arthritis (PsA)?				
Please indicate the preferred alternatives for psoriatic arthritis that have been ineffective, not tolerated, or are contraindicated:				
☐ Cosentyx ☐ Enbrel ☐ Humira ☐ Otezla ☐ Remicade ☐ Simponi Aria				
Pyoderma gangrenosum				
Yes No Has the patient previously received a biologic medication indicated for the treatment of pyoderma gangrenosum?				
Yes No Has the patient experienced an inadequate response with corticosteroids or immunosuppressive therapy (e.g., cyclosporine or mycophenolate mofetil)?				
	□ No Has the patient experienced a	n intolerance to corticosteroids an	d immunosuppressive therapy (e.g.,	
	cyclosporine or mycophenolate	e mofetil)?		
☐ Yes ☐ No Does the patient have a contraindication to corticosteroids and immunosuppressive therapy (e.g., cyclosporine mycophenolate mofetil)?				
	tnerapy (e.g	., cyclosporine mycopnenolate mo	nem):	

Continued on next page



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Page 4 of 5

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G. CLINICAL INFORMATION (continued) – Re	equired clinical information must be comple	eted in its <u>entirety</u> for all precertific	cation requests.		
Reactive arthritis					
Yes No Has the patient previously received a biologic medication indicated for the treatment of reactive arthritis? Has the patient experienced an inadequate response after at least 3 months of treatment with methotrexate titrated 20 mg per week? Yes No Has the patient experienced intolerance to methotrexate? Yes No Does the patient have a contraindication to methotrexate? Please indicate the contraindication: ☐ History of intolerance or adverse event ☐ Alcoholism, alcoholic liver disease ☐ Elevated liver transaminases ☐ Interstitial pneumonitis or clinically significant pulmonary fibrosis ☐ Renal impairment ☐ Pregnancy or planning pregnancy ☐ Blood dyscrasias (e.g., thrombocytopenia, leukopenia, significant anemia) ☐ Myelodysplasia ☐ Hypersensitivity ☐ Significant drug interaction ☐ Other					
Rheumatoid arthritis			, caner		
☐ Yes ☐ No Has the patient been diagnosed with moderately to severely active rheumatoid arthritis (RA)? ☐ Yes ☐ No Has the patient previously received a biologic or targeted synthetic disease modifying drug (e.g., Xeljanz) indicated for moderately to severely active rheumatoid arthritis? ☐ Yes ☐ No Is the requested medication being prescribed in combination with methotrexate or leflunomide? ☐ Please indicate a clinical reason for the patient to not use methotrexate or leflunomide: ☐ History of intolerance or adverse event ☐ Alcoholism, alcoholic liver disease or other chronic liver disease ☐ Elevated liver transaminases ☐ Interstitial					
. ☐ Breastfeed	pneumonitis or clinically significant pulmonary fibrosis Renal impairment Pregnancy or planning pregnancy Breastfeeding Blood dyscrasias (e.g., thrombocytopenia, leukopenia, significant anemia) Myelodysplasia				
☐ Hypersensitivity ☐ Significant drug interaction ☐ Yes ☐ No Does the patient have other reason or no clinical reason not to use methotrexate or leflunomide? ✓ Please explain:					
☐ ☐ Yes ☐ No	Has the patient experienced an inadequ		ths of treatment with the		
	methotrexate dose greater than or equa → ☐ Yes ☐ No Has the patient experie		,		
		es the patient have a contraindica			
	Ple	ase indicate the contraindication:			
		History of intolerance or adverse			
☐ Alcoholism, alcoholic liver disease or other chronic liver disease ☐ Elevated liver transaminases ☐ Interstitial pneumonitis or clinically significant pulmonary fibrosis ☐ Renal impairment ☐ Pregnancy or planning pregnancy ☐ Breastfeeding ☐ Blood dyscrasias (e.g., thrombocytopenia, leukopenia, significant anemia) ☐ Myelodysplasia ☐ Hypersensitivity ☐ Significant drug interaction ☐ Other ☐ No clinical reason not to use methotrexate or leflunomide					
Yes No Is the requested medication being prescribed in combination with methotrexate or leflunomide?					
Please indicate a clinical reason for the patient to not use methotrexate or leflunomide: Please indicate a clinical reason for the patient to not use methotrexate or leflunomide: Interstitial event Pregnancy or planning pregnancy Breastfeeding Blood dyscrasias (e.g., thrombocytopenia, leukopenia, significant anemia) Myelodysplasia Hypersensitivity Significant drug interaction Other No clinical reason not to use methotrexate or leflunomide					
Please indicate the preferred alternatives for rheumatoid arthritis have been ineffective, not tolerated, or are contraindicated: □ Enbrel □ Humira □ Kevzara □ Orencia □ Remicade □ Rinvoq □ Simponi Aria □ Xeljanz/Xeljanz XR Sarcoidosis					
Sarcoldosis Yes No Has the patient experienced an inadequate response with corticosteroids or immunosuppressive therapy?					
Yes No Does the patient experienced an inadequate response with conticosteroids of immunosuppressive therapy? Yes No Does the patient have a contraindication to corticosteroids and immunosuppressive therapy?					
Takayasu's arteritis					
☐ Yes ☐ No Has the patient experienced an inadequate response with corticosteroids or immunosuppressive therapy (e.g., methotrexate, azathioprine, or mycophenolate mofetil)?					
☐ Yes ☐ No Has the patient experienced an intolerance to corticosteroids and immunosuppressive therapy (e.g., methotrexate, azathioprine, or mycophenolate mofetil)?					
Yes No Does the patient have a contraindication to corticosteroids and immunosuppressive therapy (e.g., methotrexate, azathioprine, or mycophenolate mofetil)?					

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G. CLINICAL INFORMATION (continu	<i>red)</i> – Required clinical information	must be completed in its entirety	for all precertification requests.	
Ulcerative colitis ☐ Yes ☐ No Has the patient been diag	acced with moderately to acversly act	tive ulcorative colitic (LIC)?		
	patient been hospitalized for fulminar		s bleeding, severe toxic symptoms	
	g fever and anorexia)?	it dicerative contis (e.g., continuous	blocding, severe toxic symptoms,	
☐ Yes ☐ No Has the patient previously received a biologic or targeted synthetic disease modifying drug (e.g., Xeljanz) indicated for				
	tely to severely active ulcerative colities		, , , ,	
└── ☐ Yes	s ☐ No Has the patient tried and ha	d an inadequate response to at lea	st one conventional therapy option?	
			ntolerance to at least one conventional	
			muran], corticosteroid [e.g., budesonide,	
			dnisolone, prednisone, cyclosporine Pentasa, Canasa, Rowasa], mercaptopurine	
		ol], sulfasalazine, tacrolimus [Progr		
		nitis only])?	1	
			steroid (e.g., budesonide [Entocort, Uceris],	
			hthylprednisolone [Medrol, Solu-Medrol],	
			ne (e.g., Apriso, Asacol, Lialda, Pentas, Canasa, Tacrolimus (Prograf) Metronidazole	
	(Flagyl) or Ciprofloxacin (
Please indicate the preferred alternatives f			aindicated:	
☐ Humira ☐ Entyvio ☐ Remicade ☐ Xeljanz ☐ Stelara (intravenous formulation)				
Uveitis				
Yes No Has the patient previously				
	patient experienced an inadequate re prine, or mycophenolate mofetil)?	esponse with corticosteroids or imm	nunosuppressive therapy (e.g., methotrexate,	
☐ Yes ☐ No Has the patient experienced an intolerance to corticosteroids and immunosuppressive therapy (e.g.,				
methotrexate, azathioprine, or mycophenolate mofetil)?				
Yes No Does the patient have a contraindication to corticosteroids and immunosuppressive				
therapy (e.g., methotrexate, azathioprine, or mycophenolate mofetil)?				
Yes No Has the patient had an ineffective response, contraindication or intolerance to Humira?				
For Continuation Requests:				
☐ Yes ☐ No Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program? ☐ Yes ☐ No Has the patient achieved or maintained positive clinical response as evidenced by low disease activity or improvement in signs and symptoms				
since starting treatment with the requested drug?				
H. ACKNOWLEDGEMENT	ar are requested drug.			
H. ACKNOWLEDGEMENT				
Request Completed By (Signature Re	quired):		Date: //	
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 subjects such person to criminal and civil penalties.				

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