

adalimumab-aacf Prior Authorization Request Form Caterpillar Prescription Drug Benefit Phone: 877-228-7909 Fax: 800-424-7640



Instructions: Please fill out all applicable sections completely and legibly. Attach any additional documentation that is important for the review (e.g., chart notes or lab data, to support the authorization request). Information contained in this form is Protected Health Information under HIPAA.

MEMBER INFORMATION	
LAST NAME:	FIRST NAME:
PHONE NUMBER:	DATE OF BIRTH:
STREET ADDRESS:	
CITY:	STATE: ZIP CODE:
PATIENT INSURANCE ID NUMBER:	

MALE	FEMALE	HEIGHT (IN/CM):	WEIGHT (LB/KG):	ALLERGIES:

IF YOU ARE NOT THE PATIENT OR THE PRESCRIBER, YOU WILL NEED TO SUBMIT A PHI DISCLOSURE AUTHORIZATION FORM WITH THIS REQUEST WHICH CAN BE FOUND AT THE FOLLOWING LINK: <u>https://magellanrx.com/member/external/commercial/common/doc/en-us/phi_disclosure_authorization.pdf</u>

PATIENT'S AUTHORIZED REPRESENTATIVE (IF APPLICABLE): ______AUTHORIZED REPRESENTATIVE'S PHONE NUMBER: ______

PRESCRIBER INFORMATION			
LAST NAME:	FIRST NAME:		
PRESCRIBER SPECIALTY:	EMAIL ADDRESS:		
NPI NUMBER:	DEA NUMBER:		
PHONE NUMBER:	FAX NUMBER:		
STREET ADDRESS:			
CITY:	STATE: ZIP CODE:		
REQUESTOR (if different than prescriber):	OFFICE CONTACT PERSON:		

MEDICATION OR MEDICAL DISPENSING INFORMATION				
MEDICATION NAME:				
DOSE/STRENGTH:	FREQUENCY:	LENGTH OF THERAPY/REFILLS:	QUANTITY:	
NEW THERAPY		IF RENEWAL: DATE THERAPY INITIATED:		
DURATION OF THERAPY (SPECIFIC DATES):				

Continued on next page.

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adalimumab-aacf

Prior Authorization Request Form





MEMBER'S LAST NAME:	MEMBER'S FIRST NAME:		
1. HAS THE PATIENT TRIED ANY OTHER	MEDICATIONS FOR THIS CONDITION?	YES (if yes, complete below) NO	
MEDICATION/THERAPY (SPECIFY DRUG NAME AND DOSAGE):	DURATION OF THERAPY (SPECIFY DATES):	RESPONSE/REASON FOR FAILURE/ALLERGY:	
2. LIST DIAGNOSES:		ICD-10:	
) Code(s):		
	PLEASE PROVIDE ALL RELEVANT CLINICA	AL INFORMATION TO SUPPORT A	
PRIOR AUTHORIZATION.			
Clinical Information: Will adalimumab-aacf be used in combination with another biologic response modifier [such as but not limited to:Kineret (anakinra), Rituxan (rituximab), Remicade (infliximab), Orencia (abatacept), Cimzia (certolizumab pegol), Enbrel (etanercept), Simponi (golimumab), Actemra (tocilizumab)], Xeljanz (tofacitinib), Rinvoq(upadacitinib), Olumiant(baricitinib), Cibinqo(abrocitinib)? Yes No Select if the requested medication is prescribed by the following specialist: Dermatologist Gastroenterologist Ophthalmologist Rheumatologist			
□ Yes □ No	r the following: failure to at least TWO (2) non-steroida ied:		
Are non-steroidal anti-inflammatory ag documentation.	gents (NSAIDs) contraindicated in this p	atient? 🗆 Yes 🗆 No Please provide	
Has the patient had a trial of methotre	xate? Yes No Please provide docum	nentation.	
For <u>Crohn's disease</u> , also answer the for Select if the patient has had a trial of the 5-ASA/mesalamine 6-mercaptopurine Azathioprine	•		

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MANAGEMENT





Glucocorticoid therapy

Methotrexate

*Please provide supporting chart notes for verification

Is there clinical rationale explaining why the patient cannot try glucocorticoid therapy, methotrexate, azathioprine, 6-mercaptopurine or 5-ASA/mesalamine?*
Ves
No*Please provide supporting documentation.

For <u>hidradenitis suppurativa</u>, also answer the following: Is the medication being used prior to surgery? Yes
No

For juvenile idiopathic arthritis (JIA), also answer the following:

Has the patient had a trial and inadequate response to therapy with oral disease modifying anti-rheumatic agents (DMARDs) [e.g., methotrexate, azathioprine (Imuran), auranofin (Ridaura), hydroxychloroquine (Plaquenil), sulfasalazine (Azulfidine), leflunomide (Arava)]? \Box Yes \Box No**Please submit chart documentation on therapies the patient has tried*.

Does the patient have chronic liver disease such as chronic alcohol abuse/alcoholism, chronic hepatitis, fatty liver, nonalcoholic steatohepatitis/NASH, or elevated liverenzymes?
□ Yes □ No Please provide documentation.

Is there clinical rationale explaining why the patient cannot try a DMARD?*
Yes
No *Please provide supporting documentation.

For <u>non-infectious uveitis</u>, also answer the following: Does the patient have isolated anterior uveitis? □ Yes □ No Please provide documentation.

For plaque psoriasis, also answer the following:

Does the patient have plaques covering at least 10% of their body surface area (BSA) or < 10% of BSA, but with involvement of palms, soles, head and neck, or genitalia which causes disruption of normal activities?
Yes
No Please provide documentation.

Select if patient has had previous treatment failure with any of the following:

□ Acitretin □ Methotrexate

□ Cyclosporine □ Phototherapy

*Please submit what therapies the patient has tried.

Is there clinical rationale explaining why the patient cannot try any of the following: methotrexate, cyclosporine, acitretin, or phototherapy?*
Yes
No
Please provide supporting documentation.

For psoriatic arthritis, also answer the following:

Has the patient had a trial and inadequate response to therapy with oral disease modifying anti-rheumatic agents (DMARDs) [e.g., methotrexate, azathioprine (Imuran), auranofin (Ridaura), hydroxychloroquine (Plaquenil), penicillamine (Cuprimine), sulfasalazine (Azulfidine), leflunomide (Arava)]?* \Box Yes \Box No *Please submit chart documentation on therapies the patient has tried.

Does the patient have chronic liver disease such as chronic hepatitis, fatty liver, nonalcoholic steatohepatitis/NASH, or elevated liverenzymes?
Que Yes
Que No Please provide documentation.







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Is there clinical rationale explaining why the patient cannot try a DMARD?* Yes No <i>Please provide supporting documentation</i>
For pyoderma gangrenosum, also answer the following:
Has the patient had a trial of glucocorticoid therapy? Yes No
Please provide supporting chart notes.
Select if the patient has had a trial of the following systemic therapies: Azathioprine Dapsone Nicotine Chlorambucil Hyperbaric oxygen Tacrolimus Cyclophosphamide Intravenous immune globulin Thalidomide Cyclosporine Mycophenolate
Is there clinical rationale explaining why the patient cannot try corticosteroids and one additional systemic therapy (such as cyclosporine, mycophenolate, dapsone, azathioprine, etc.)?* Yes Doe No *Please provide supporting documentation.
For requirements in arthritic also answer the following:
For <u>rheumatoid arthritis,</u> also answer the following: Has the patient had a trial with methotrexate or another oral disease modifying anti-rheumatic agent (DMARD)
such as azathioprine (Imuran), auranofin (Ridaura), hydroxychloroquine (Plaquenil), sulfasalazine (Azulfidine), or
leflunomide (Arava)? 🗆 Yes 🗆 No
*Please submit chart documentation on therapies the patient has tried.
Does the patient have chronic liver disease such as chronic alcohol abuse/alcoholism chronic hepatitis, fatty liver, nonalcoholic steatohepatitis/NASH, or elevated liver enzymes? Yes No Please provide documentation.
Is the patient unable to take a non-biologic DMARD due to patient is a male of fatherhood potential or a female of childbearing potential? Yes I No Please provide documentation.
If "no" to the above question, provide the rationale explaining why the patient cannot take the prerequisite non- biologic DMARDs:
For <u>ulcerative colitis</u> , also answer the following:
Select if the patient has tried and failed at least one of the following therapies:
□ Corticosteroids
Azathioprine
G-mercaptopurine
*Please provide supporting chart notes.
Is there clinical rationale explaining why the patient cannot try corticosteroids, azathioprine, or 6- mercaptopurine?* Yes No
*Please provide supporting documentation.
Reauthorization:
If this is a reauthorization request, answer the following questions:
Select if the requested medication is prescribed by the following specialist:
□ Dermatologist
□ Gastroenterologist
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Ophthalmologist

Rheumatologist

For all <u>indications except non-infectious uveitis</u>, also answer the following: Is the patient continuing to have a positive clinical response and remission of disease with continued use of Humira?* Yes I No

*Please provide documentation supporting this information.

For <u>non-infectious uveitis</u>, also answer the following:

Has the patient 's response been evaluated at a recent office visit (i.e., occurring after previous date of approval)?* □ Yes □ No

*Please provide progress notes/chart notes from the patient's ophthalmologist or rheumatologist supporting this information.

Has the patient had a positive response to therapy?
□ Yes
□ No Please provide documentation.

Are there any other comments, diagnoses, symptoms, medications tried or failed, and/or any other information the physician feels is important to this review?

Please note: Not all drugs/diagnosis are covered on all plans. This request may be denied unless all required information is received.

ATTESTATION: I attest the information provided is true and accurate to the best of my knowledge. I understand that the Health Plan, insurer, Medical Group or its designees may perform a routine audit and request the medical information necessary to verify the accuracy of the information reported on this form.

Prescriber Signature or Electronic I.D. Verification:

Date:

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FAX THIS FORM TO: 800-424-7640

MAIL REQUESTS TO: Magellan Rx Management Prior Authorization Program

Attn: CP - 4201 P.O. Box 64811 St. Paul, MN 55164-0811



