

Tremfya HMSACOM - Prior Authorization Request

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. **Please respond below and fax this form to CVS Caremark toll-free at 1-866-237-5512**. If you have questions regarding the prior authorization, please contact CVS Caremark at **1-808-254-4414**. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect[®] 1-800-237-2767.

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Patient's Name:	Date:
Patient's ID:	Patient's Date of Birth:
Patient's Phone Number:	
Physician's Name:	
Specialty:	NPI#:
Physician Office Telephone:	Physician Office Fax:

Approvals may be subject to dosing limits in accordance with FDA-approved labeling, accepted compendia, and/or evidence-based practice guidelines.

Additional Demographic Information:

Patient Weight: _		kg
Patient Height: _	ft	inches

Indicate where the drug is being dispensed:

- □ Office □ Outpatient Hospital □ Ambulatory Surgical □ Inpatient Hospital
- □ Off Campus Outpatient Hospital □ Urgent Care □ Emergency Room □ Birthing Center
- □ Military Facility □ Skilled Nursing Facility □ Nursing Facility □ Hospice
- □ Inpatient Psychiatric □ Psychiatric Residential Treatment □ End Stage Renal Facility
- □ Psychiatric Facility □ Pharmacy □ Other

Indicate where the drug is being administered:

- □ Ambulatory surgical □ Home □ Inpatient Hospital
- □ Office □ Outpatient Hospital □ Pharmacy

What is the ICD-10 code?

What product is being requested?
Tremfya IV
Tremfya SQ

Send completed form to: CVS Caremark Specialty Programs. Fax: 1-866-237-5512

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Criteria Ouestions:

1. Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic drug (e.g., Olumiant, Otezla, Xeljanz) for the same indication?

 \square Yes, *Continue to #2*

 \square No, Continue to #2

2. Has the patient ever received (including current utilizers) a biologic (e.g., Humira) or targeted synthetic drug (e.g., Olumiant, Xeljanz) associated with an increased risk of tuberculosis?

□ Yes, Continue to #9

 \square No, Continue to #3

3. Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [TST], interferon-release assay [IGRA]) within 6 months of initiating therapy?

□ Yes, Continue to #4

 \square No. Continue to #4

4. What were the results of the tuberculosis (TB) test?

D Positive for TB. *Continue to #5*

□ Negative for TB, *Continue to #9*

Unknown, No Further Questions

5. Which of the following applies to the patient?

D Patient has latent TB and treatment for latent TB has been initiated, Continue to #6

□ Patient has latent TB and treatment for latent TB has been completed, *Continue to #6*

□ Patient has latent TB and treatment for latent TB has not been initiated, *Continue to #6*

□ Patient has active TB, *Continue to #6*

Indication

- 9. What is the diagnosis?
- □ Plaque psoriasis, *Continue to #100*
- D Psoriatic arthritis WITH co-existent plaque psoriasis, *Continue to #10*
- □ Psoriatic arthritis, *Continue to #200*
- Ulcerative colitis. *Continue to #250*
- **Other**, No Further Questions

10. Is the requested drug being prescribed by or in consultation with a rheumatologist or dermatologist?

- □ Yes, Continue to #11
- \square No. Continue to #11
- 11. What is the primary diagnosis being treated?
- □ Psoriatic arthritis, *Continue to #201*
- □ Plaque psoriasis, *Continue to #101*

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Plaque Psoriasis

100. Is the requested drug being prescribed by or in consultation with a dermatologist?

□ Yes, Continue to #101

□ No. Continue to #101

101. Has the patient been diagnosed with moderate to severe plaque psoriasis?

 \square Yes, Continue to #102

 \square No. Continue to #102

102. Is the patient an adult?

□ Yes, Continue to #103

□ No, Continue to #103

Continuation of Therapy

103. Is this request for continuation of therapy with the requested drug?

□ Yes, Continue to #104

 \square No. Continue to #108

104. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?

□ Yes, Continue to #108

 \square No. Continue to #105

□ Unknown, *Continue to #108*

105. 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, Continue to #106 □ No, Continue to #106

106. Has the patient experienced a reduction in body surface areas (BSA) affected from baseline?

D Yes, No Further Questions \square No. Continue to #107

107. Has the patient experienced an improvement in signs and symptoms of the condition from baseline (e.g., itching, redness, flaking, scaling, burning, cracking, pain)?

□ Yes, No Further Questions

□ No, No Further Questions

Initial Therapy

Prior treatment with another biologic or targeted synthetic drug

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108. Has the patient ever received or is currently receiving a biologic (e.g., Humira) or targeted synthetic drug (e.g., Sotyktu, Otezla) indicated for the treatment of moderate to severe plaque psoriasis (excluding receiving the drug via samples or a manufacturer's patient assistance program)?

D Yes, No Further Questions

□ No, Continue to #109

Requirements regarding prior therapy

109. Are crucial body areas (e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas) affected?

□ Yes, No Further Ouestions

 \square No. Continue to #110

110. Is the percentage of body surface area (BSA) affected (prior to starting the requested medication) less then 3%?

□ Yes, Continue to #111

□ No, Continue to #111

111. What is the percentage of body surface area (BSA) affected (prior to starting the requested medication)?

Greater than or equal to 3% to less than 10% of BSA, *Continue to #112*

Greater than or equal to 10% of BSA, No Further Questions

112. Has the patient experienced an inadequate response, or has an intolerance to phototherapy (e.g., UVB, PUVA) or pharmacologic treatment with methotrexate, cyclosporine or acitretin?

T Yes, No Further Ouestions

□ No, Continue to #113

113. Does the patient have a clinical reason to avoid pharmacologic treatment with methotrexate, cyclosporine and acitretin?

 \square Yes, *Continue to #114*

 \square No. Continue to #114

114. Please indicate clinical reason to avoid pharmacologic treatment with methotrexate, cyclosporine, and acitretin

Clinical diagnosis of alcohol use disorder, alcoholic liver disease or other chronic liver disease, *No Further* Questions

Drug interaction, No Further Questions

Risk of treatment-related toxicity, *No Further Questions*

D Pregnancy or currently planning pregnancy, No Further Questions

□ Breastfeeding, No Further Questions

□ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension), No Further Questions

□ Hypersensitivity, No Further Questions

History of intolerance or adverse event, No Further Questions

Other, No Further Questions

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Psoriatic Arthritis

200. Is the requested drug being prescribed by or in consultation with a rheumatologist or dermatologist?

□ Yes, Continue to #201

 \square No. Continue to #201

201. Is the patient an adult?

 \Box Yes, Continue to #202

 \square No. Continue to #202

Continuation of Therapy

202. Is this request for continuation of therapy with the requested drug?

□ Yes, Continue to #203

 \square No, Continue to #210

203. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?

 \square Yes, Continue to #210

 \square No. Continue to #204

□ Unknown, Continue to #210

204. 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, Continue to #205

□ No, Continue to #205

205. Has the patient experienced improvement in any of the following from baseline?

D Number of swollen joints, No Further Questions

□ Number of tender joints, No Further Questions

Dactylitis, No Further Questions

D Enthesitis, No Further Questions

Skin and/or nail involvement, No Further Questions

□ Functional status, *No Further Questions*

C-reactive protein (CRP), No Further Questions

□ None of the above, *No Further Questions*

Initial Therapy

210. Has the patient been diagnosed with active psoriatic arthritis (PsA)?

 \square Yes, Continue to #211

 \square No, Continue to #211

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211. Has the patient ever received or is currently receiving a biologic (e.g., Humira) or targeted synthetic drug (e.g., Rinvoq, Otezla) indicated for the treatment of active psoriatic arthritis (excluding receiving the drug via samples or a manufacturer's patient assistance program)?

□ Yes, No Further Questions

 \square No, Continue to #212

212. What is the patient's disease severity?

□ Mild to moderate, *Continue to #213*

Severe, No Further Questions

213. Does the patient have enthesitis?

□ Yes, No Further Questions

 \square No. Continue to #214

214. Has the patient had an inadequate response to methotrexate, leflunomide, or another conventional synthetic drug (e.g., sulfasalazine) administered at an adequate dose and duration?

□ Yes, No Further Questions

□ No, Continue to #215

215. Has the patient had an intolerance to methotrexate, leflunomide, or another conventional synthetic drug (e.g., sulfasalazine)?

□ Yes, No Further Questions

 \square No, Continue to #216

216. Does the patient have a contraindication to methotrexate or leflunomide?

 \Box Yes. Continue to #217

 \square No. Continue to #218

217. Please indicate the contraindication to methotrexate or leflunomide

Clinical diagnosis of alcohol use disorder, alcoholic liver disease, or other chronic liver disease, *No Further* Questions

Drug interaction, No Further Questions

I Risk of treatment-related toxicity, *No Further Questions*

D Pregnancy or currently planning pregnancy, No Further Questions

□ Breastfeeding, No Further Ouestions

□ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension), No Further Questions

□ Hypersensitivity, No Further Questions

History of intolerance or adverse event, No Further Questions

□ None of the above, *No Further Questions*

218. Does the patient have a contraindication to another conventional synthetic drug (e.g., sulfasalazine)?

D Yes, No Further Questions

□ No, No Further Questions

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Ulcerative colitis

250. Has the patient been diagnosed with moderately to severely active ulcerative colitis?

□ Yes, Continue to #251

□ No, Continue to #251

251. Is the patient an adult?

□ Yes, Continue to #252

 \square No. Continue to #252

252. Is the requested drug being prescribed by or in consultation with a gastroenterologist?

□ Yes, Continue to #253

□ No, Continue to #253

253. Which of the following applies to this request for the requested drug?

Initiation of the intravenous (IV) loading dose, No Further Questions

□ Initiation of the subcutaneous (SQ) maintenance dose, No Further Questions

Continuation of the subcutaneous (SQ) maintenance dose, *Continue to #254*

254. Has the patient achieved or maintained remission OR achieved or maintained a positive clinical response to treatment as evidenced by low disease activity or improvement in signs and symptoms of the condition since starting treatment with the requested drug?

Tyes, achieved or maintained remission, *No Further Questions*

□ Yes, achieved or maintained a positive clinical response, *Continue to #255*

□ No or none of the above, *Continue to #255*

255. Which of the following has the patient experienced and improvement in from baseline?

□ Stool frequency, No Further Questions

□ Rectal bleeding, *No Further Questions*

Urgency of defecation, No Further Questions

C-reactive protein (CRP), No Further Questions

□ Fecal calprotectin (FC), No Further Questions

□ Appearance of the mucosa on endoscopy, computed tomography enterography (CTE), magnetic resonance enterography (MRE), or intestinal ultrasound, No Further Questions

□ Improvement on a disease activity scoring tool (e.g., Ulcerative Colitis Endoscopic Index of Severity [UCEIS], Mayo Score), No Further Questions

□ None of the above, *No Further Questions*

I attest that this information is accurate and true, and that documentation supporting this information is available for review if requested by CVS Caremark or the benefit plan sponsor.

Prescriber or Authorized Signature

Date (mm/dd/yy)

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