

Skyrizi

CareFirst 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 1-855-330-1720**. If you have questions regarding the prior authorization, please contact CVS Caremark at **1-888-877-0518**. 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:	
Physician's Name:			
Specialty:		NPI#:	
Physician Office Telephone:		Physician Office Fax:	
Referring Provider Info: 🗆 Same as Re	equesting Provid	ler	
Name:		NPI#:	
Fax:		Phone:	
<u>Rendering</u> Provider Info: □ Same as Re Name:	_		
Fax:		Phone:	
accepted comp Required Demographic Information:	oendia, and/or ev	vidence-based practice guidelines.	
Patient Weight:	kg		
Patient Height:			
Please indicate the place of service for the	requested drug:		
☐ Ambulatory Surgical	☐ Home	Off Campus Outpatient Hospital	
On Campus Outpatient Hospital	Office	☐ Pharmacy	
What is the ICD-10 code?			

Send completed form to: Case Review Unit CVS Caremark Specialty Programs Fax: 1-855-330-1720

Psoriasis Enhanced SGM 4179-A Criteria Questions:
Is the diagnosis moderate or severe plaque psoriasis?
Sting to Sharing SCM 2047 A Criteria Occasion 1
□ No, Skip to Skyrizi SGM 3047-A Criteria Question 1
1. What is the patient's age? Indicate in years. ☐ 18 years of age or older, Continue to 2 ☐ Less than 18 years of age, Skip to Skyrizi SGM 2047-A Criteria Question 1
 2. What is the diagnosis? ☐ Plaque psoriasis, Continue to 3 ☐ Plaque psoriasis with co-existing psoriatic arthritis, Skip to Skyrizi SGM 2047-A Criteria Question 1 ☐ Other, please specify:, Skip to Skyrizi SGM 2047-A Criteria Question 1
3. Is the request for Sotyktu? ☐ Yes, Continue to 4 ☐ No, Continue to 5
4. Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic drug (e.g., Otezla)? ☐ Yes, <i>Continue to 7</i> ☐ No, <i>Continue to 7</i>
5. Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic drug (e.g., Otezla, Sotyktu) for the same indication? ☐ Yes, Continue to 6 ☐ No, Continue to 6
6. What is the requested medication?
☐ Otezla, <i>Continue to 11</i>
☐ Other, please specify:, <i>Continue to 7</i> 7. 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, <i>Continue to 12</i> ☐ No, <i>Continue to 8</i>
8. Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [TST], interferon-release assay [IGRA]) within 12 months of initiating therapy? Tyes, Continue to 9 No, Continue to 11
9. What were the results of the TB test? □ Positive for TB, Continue to 10 □ Negative for TB, Continue to 12 □ Unknown, No further questions
10. Which of the following applies to the patient? ☐ Patient has latent TB and treatment for latent TB has been initiated, <i>Continue to 12</i> ☐ Patient has latent TB and treatment for latent TB has been completed, <i>Continue to 12</i>

☐ Patient has latent TB and treatment for latent TB has ☐ Patient has active TB, <i>Continue to 12</i>	not been initiated, Continue to 12
11. What is the severity of the disease? ☐ Mild plaque psoriasis, <i>Skip to Skyrizi SGM 2047-A C.</i> ☐ Moderate plaque psoriasis, <i>Continue to 13</i> ☐ Severe plaque psoriasis, <i>Continue to 13</i>	riteria Question 1
12. Has the patient been diagnosed with moderate to sev ☐ Yes, <i>Continue to 13</i> ☐ No, <i>Continue to 13</i>	ere plaque psoriasis?
13. Is the requested drug prescribed by or in consultation ☐ Yes, <i>Continue to 14</i> ☐ No, <i>Continue to 14</i>	n with a dermatologist?
 14. Is this request for continuation of therapy with the reapplicable)? ☐ Yes, Continue to 15 ☐ No, Continue to 21 	equested drug or a biosimilar of the requested drug (if
15. Is the patient currently receiving the requested drug samples or a manufacturer's patient assistance program? ☐ Yes, Continue to 21 ☐ No, Continue to 16 ☐ Unknown, Continue to 21	or a biosimilar of the requested drug (if applicable) through
16. Has the patient achieved or maintained a positive cli improvement in signs and symptoms of the condition sir the requested drug? ☐ Yes, <i>Continue to 17</i> ☐ No, <i>Continue to 17</i>	nical response as evidenced by low disease activity or nce starting treatment with the requested drug or a biosimilar of
ACTION REQUIRED: Attach supporting chart note(s) involvement of BSA percent. ☐ Less than or equal to 3%	n body surface area (BSA) percent? Indicate in percentage. or medical record documentation for current psoriasis ACTION REQUIRED: Submit supporting documentation, Skip IN REQUIRED: Submit supporting documentation. Continue to
BSA from baseline. □ Less than 75% BSA improvement	or medical record documentation for percent improvement of
documentation, <i>Continue to 19</i> Greater than or equal to 75% BSA improvement supporting documentation, <i>Skip to Skyrizi SGM 2047-A</i>	% ACTION REQUIRED: Submit Criteria Question 51

19. What is the patient's percent reduction in the Psoria		
score reduction in percentage. <i>ACTION REQUIRED</i> :	Attach supportin	ig chart note(s) or medical record documentation
for percent reduction of PASI score from baseline. Greater than or equal to 75% reduction	0/	ACTION PEOUIPED: Submit supporting
documentation, Skip to Skyrizi SGM 2047-A Criteria Q		ACTION REQUIRED. Submit supporting
☐ Greater than or equal to 50% and less than 75% red	juesiion 31	0/ ACTION DECLUDED.
	uction	% ACTION REQUIRED:
Submit supporting documentation, <i>Continue to 20</i>	0/ C	20
☐ Less than 50% reduction	_%, Continue to	20
20. What is the patient's Dermatology Life Quality Ind <i>REQUIRED</i> : Attach supporting chart note(s) or medic (DLQI) score. ☐ Less than or equal to 5	cal record docume	entation for Dermatology Life Quality Index
	ACTION KE	CIRED . Submit supporting documentation,
Skip to Skyrizi SGM 2047-A Criteria Question 51 Greater than 5, No feet	further questions	
21. Has the patient received or is currently receiving a Otezla) within the past 120 days indicated for the treat the drug via samples or a manufacturer's patient assistantes, medical record documentation, or claims history ☐ Yes, <i>Skip to Skyrizi SGM 2047-A Criteria Question</i> ☐ No, <i>Continue to 22</i>	ment of moderate ance program)? As supporting previous	e to severe plaque psoriasis (excluding receiving ACTION REQUIRED: If Yes, please attach chart
22. Is the percentage of body surface area (BSA) affect ☐ Yes, <i>Continue to 23</i> ☐ No, <i>Continue to 23</i>	ted (prior to starti	ing the requested medication) less than 3%?
23. What is the percentage of body surface area (BSA) percentage. <i>ACTION REQUIRED</i> : Attach supporting (BSA) affected.		
☐ Greater than or equal to 3% but less than 10%		% ACTION REQUIRED: Submit
supporting documentation, Continue to 24		
☐ Greater than or equal to 10%	% ACTION	REOUIRED : Submit supporting documentation
Continue to 33		
24. What is the patient's Psoriasis Area Severity Index <i>REQUIRED</i> : Attach supporting chart note(s) or medic score.	cal record docume	entation of Psoriasis Area Severity Index (PASI)
☐ Greater than or equal to 10	ACTION	REQUIRED: Submit supporting
documentation, Continue to 26		
	ON REQUIRED:	: Submit supporting documentation, Continue to
25	-	
25. Is the affected area severe at localized sites and ass of distress (e.g., nail disease or involvement of high-inflexures and genitals)? <i>ACTION REQUIRED</i> : If Yes, documentation of affected area(s) with significant func	npact and difficul , please attach sup	lt-to-treat sites such as face, scalp, palms, soles, pporting chart notes or medical record

☐ Yes, Continue to 33 ☐ No, Continue to 33
26. Has the patient had an inadequate response at the maximum tolerated dose to a medium to super-high potency topical corticosteroid therapy for a duration of at least 4 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart notes, medical record documentation, or claims history of all prior and current use of treatment regimens for topical corticosteroid therapy, including dosage, duration, and response to therapy. ☐ Yes, <i>Continue to 33</i> ☐ No, <i>Continue to 27</i>
27. Has the patient had an inadequate response at the maximum tolerated dose to a topical calcineurin inhibitor therapy for a duration of at least 8 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for topical calcineurin inhibitor therapy, including dosage, duration, and response to therapy. ☐ Yes, <i>Continue to 33</i> ☐ No, <i>Continue to 28</i>
28. Has the patient had an inadequate response at the maximum tolerated dose to a topical vitamin D analog therapy for a duration of at least 12 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for topical vitamin D analog therapy, including dosage, duration, and response to therapy. Tyes, <i>Continue to 33</i> No, <i>Continue to 29</i>
29. Has the patient had an inadequate response at the maximum tolerated dose to a topical retinoid therapy for a duration of at least 12 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for topical retinoid therapy, including dosage, duration, and response to therapy. Tyes, <i>Continue to 33</i> No, <i>Continue to 30</i>
30. Has the patient had an inadequate response at the maximum tolerated dose to a topical aryl hydrocarbon receptor agonist therapy for a duration of at least 12 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for topical aryl hydrocarbon receptor agonist therapy, including dosage, duration, and response to therapy. ☐ Yes, <i>Continue to 33</i> ☐ No, <i>Continue to 31</i>
31. Has the patient had an inadequate response at the maximum tolerated dose to a topical phosphodiesterase 4 inhibitor therapy for a duration of at least 8 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for topical phosphodiesterase 4 inhibitor therapy, including dosage, duration, and response to therapy. Yes, <i>Continue to 33</i> No, <i>Continue to 32</i>
32. Are crucial body areas (e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas) affected? <i>ACTION REQUIRED</i> : If yes, please attach chart notes or medical record documentation of affected areas. ¬ Yes, <i>Continue to 33</i> ¬ No, <i>Continue to 33</i>

33. Has the patient had a trial of phototherapy (e.g., UVB, PUVA) for a duration of at least 3 months? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s) or medical record documentation for phototherapy, including dosage, duration, and response to therapy. ☐ Yes, <i>Continue to 35</i> ☐ No, <i>Continue to 34</i>
34. Does the patient meet any of the following criteria: a) the patient has experienced an intolerable adverse event with phototherapy, b) the patient has a clinical reason to avoid phototherapy, or c) the patient does not have access to phototherapy? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous treatments tried (if applicable), including duration and response to therapy. If therapy is not advisable, please attach documentation of clinical reason to avoid therapy. 35
☐ Yes, clinical reason to avoid phototherapy <i>ACTION REQUIRED</i> : Submit supporting documentation, <i>Continue to 35</i> ☐ Yes, does not have access to phototherapy <i>ACTION REQUIRED</i> : Submit supporting documentation, <i>Continue to 35</i> ☐ None of the above, <i>Continue to 35</i> ☐ None of the above, <i>Continue to 35</i> ☐
35. Has the patient had a trial of methotrexate at a dose of at least 25 mg/week or at the maximum tolerated dose for a duration of at least 3 months? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for methotrexate, including dosage, duration, and response to therapy. ☐ Yes, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> ☐ No, <i>Continue to 36</i>
36. Has the patient had a trial of cyclosporine at a dose of at least 5 mg/kg/day or at the maximum tolerated dose for a duration of at least 6 weeks? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for cyclosporine, including dosage, duration, and response to therapy. The Yes, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> No, <i>Continue to 37</i>
37. Has the patient had a trial of acitretin at a dose of at least 50 mg/day or at the maximum tolerated dose for a duration of at least 3 months? <i>ACTION REQUIRED</i> : If Yes, please attach supporting chart note(s), medical record documentation, or claims history of all prior and current use of treatment regimens for acitretin, including dosage, duration, and response to therapy. □ Yes, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> □ No, <i>Continue to 38</i>
38. Does the patient have a clinical reason to avoid systemic pharmacologic treatment with methotrexate, cyclosporine, and acitretin? <i>ACTION REQUIRED</i> : Please attach documentation of clinical reason to avoid therapy. ☐ Yes, <i>Continue to 39</i> ☐ No, <i>Continue to 39</i>
39. Please indicate the 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, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> ☐ Drug interaction, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> ☐ Risk of treatment-related toxicity, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> ☐ Pregnancy or currently planning pregnancy, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> ☐ Breastfeeding, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i>

□ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension), <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> □ Hypersensitivity, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> □ History of intolerance or adverse event, <i>Skip to Skyrizi SGM 2047-A Criteria Question 51</i> □ Other, please specify, <i>No Further Questions</i>
Skyrizi SGM 2047-A Criteria Questions: 1. Will the requested drug be used in combination with any other biologic (e.g., Humira) or targeted synthetic drug (e.g., Otezla, Xeljanz) for the same indication? ☐ Yes, Continue to 2 ☐ 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 6 ☐ No, Continue to 3
3. Has the patient had a tuberculosis (TB) test (e.g., tuberculosis skin test [TST], interferon-release assay [IGRA]) within 12 months of initiating therapy? ☐ Yes, Continue to 4 ☐ No, Continue to 4
4. What were the results of the TB test? ☐ Positive for TB, Continue to 5 ☐ Negative for TB, Continue to 6 ☐ Unknown, No further questions
5. Which of the following applies to the patient? ☐ Patient has latent TB and treatment for latent TB has been initiated, <i>Continue to 6</i> ☐ Patient has latent TB and treatment for latent TB has been completed, <i>Continue to 6</i> ☐ Patient has latent TB and treatment for latent TB has not been initiated, <i>Continue to 6</i> ☐ Patient has active TB, <i>Continue to 6</i>
6. What is the diagnosis? ☐ Plaque psoriasis, Continue to 10 ☐ Psoriatic arthritis WITH co-existent plaque psoriasis, Continue to 7 ☐ Psoriatic arthritis, Continue to 25 ☐ Crohn's disease, Continue to 40 ☐ Ulcerative colitis, Continue to 45 ☐ Other please specify No further questions
Other, please specify

☐ Yes, Continue to 8 ☐ No, Continue to 8
8. Is the requested drug being prescribed by or in consultation with a rheumatologist or dermatologist? ☐ Yes, <i>Continue to 9</i> ☐ No, <i>Continue to 9</i>
 9. What is the primary diagnosis being treated? ☐ Psoriatic arthritis, <i>Continue to 27</i> ☐ Plaque psoriasis, <i>Continue to 12</i>
 10. Is the patient an adult (18 years of age or older)? ☐ Yes, Continue to 11 ☐ No, Continue to 11
 11. Is the requested drug being prescribed by or in consultation with a dermatologist? ☐ Yes, Continue to 12 ☐ No, Continue to 12
 12. Has the patient been diagnosed with moderate to severe plaque psoriasis? ☐ Yes, Continue to 13 ☐ No, Continue to 13
 13. Is this request for continuation of therapy with the requested drug? ☐ Yes, Continue to 14 ☐ No, Continue to 18
14. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?
☐ Yes, Continue to 18
□ No, Continue to 15
☐ Unknown, Continue to 18
15. 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, <i>Continue to 16</i> ☐ No, <i>Continue to 16</i>
16. Has the patient experienced a reduction in body surface area (BSA) affected from baseline? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes or medical record documentation of decreased body surface area affected. <i>ACTION REQUIRED</i> : Submit supporting documentation ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 17</i>
17. Has the patient experienced an improvement in signs and symptoms of the condition from baseline (e.g., itching, redness, flaking, scaling, burning, cracking, pain)? <i>ACTION REQUIRED</i> : If Yes, please attach chart

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notes or medical record documentation of improvement in signs and symptoms.

☐ Yes, Continue to 50 ☐ No, Continue to 50
18. 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)? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried. ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 19</i>
19. Are crucial body areas (e.g., hands, feet, face, neck, scalp, genitals/groin, intertriginous areas) affected? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes or medical record documentation of affected areas. ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 20</i>
20. Is the percentage of body surface area (BSA) affected (prior to starting the requested medication) less than 3%? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>Continue to 21</i>
21. What is the percentage of body surface area (BSA) affected (prior to starting the requested medication)? Indicate percentage. <i>ACTION REQUIRED</i> : Please attach chart notes or medical record documentation of body surface area affected.
☐ Greater than or equal to 3% to less than 10% of BSA
Submit supporting documentation, Continue to 22 Greater than or equal to 10% of BSAACTION REQUIRED: Submit supporting documentation, Continue to 50
22. 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? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy. Yes, <i>Continue to 50</i> No, <i>Continue to 23</i>
23. Does the patient have a clinical reason to avoid pharmacologic treatment with methotrexate, cyclosporine, and acitretin? <i>ACTION REQUIRED</i> : If Yes, please attach documentation of clinical reason to avoid therapy. ☐ Yes, <i>Continue to 24</i> ☐ No, <i>Continue to 24</i>
24. Please indicate the 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, <i>Continue to 50</i>
☐ Drug interaction, Continue to 50
☐ Risk of treatment-related toxicity, <i>Continue to 50</i>
☐ Pregnancy or currently planning pregnancy, Continue to 50
☐ Breastfeeding, Continue to 50

☐ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension), <i>Continue to 50</i>
☐ Hypersensitivity, Continue to 50
☐ History of intolerance or adverse event, <i>Continue to 50</i>
□ Other, please specify, Continue to 50
25. Is the patient an adult (18 years of age or older)? ☐ Yes, Continue to 26 ☐ No, Continue to 26
26. Is the requested drug being prescribed by or in consultation with a rheumatologist or dermatologist? ☐ Yes, <i>Continue to 27</i> ☐ No, <i>Continue to 27</i>
27. Is this request for continuation of therapy with the requested drug? ☐ Yes, Continue to 28 ☐ No, Continue to 31
28. Is the patient currently receiving the requested drug through samples or a manufacturer's patient assistance program?
☐ Yes, Continue to 31
□ No, Continue to 29
☐ Unknown, Continue to 31
29. 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 30 ☐ No, Continue to 30
30. Which of the following has the patient experienced an improvement in from baseline? <i>ACTION REQUIRED</i> : Please attach chart notes or medical record documentation supporting positive clinical response.
□ Number of swollen joints ACTION REQUIRED: Submit supporting documentation, Continue to 50
□ Number of tender joints ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Dactylitis ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Enthesitis ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Skin and/or nail involvement ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Functional status ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ C-reactive protein (CRP) ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ None of the above, <i>Continue to 50</i>
31. Has the patient been diagnosed with active psoriatic arthritis (PsA)? ☐ Yes, Continue to 32 ☐ No, Continue to 32

32. Has the patient ever received or is currently receiving a biologic (e.g., Humira) or targeted synthetic drug (e.g., Rinvoq, Otezla) that is indicated for the treatment of active psoriatic arthritis (excluding receiving the drug via samples or a manufacturer's patient assistance program)? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried. ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 33</i>
33. What is the patient's disease severity?
☐ Mild to moderate, <i>Continue to 34</i>
☐ Severe, Continue to 50
34. Does the patient have enthesitis? ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 35</i>
35. 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? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy. ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 36</i>
36. Has the patient had an intolerance to methotrexate, leflunomide, or another conventional synthetic drug (e.g., sulfasalazine)? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes, medical record documentation, or claims history supporting previous medications tried, including response to therapy. ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 37</i>
37. Does the patient have a contraindication to methotrexate or leflunomide? <i>ACTION REQUIRED</i> : If Yes, please attach documentation of clinical reason to avoid therapy. ☐ Yes, <i>Continue to 39</i> ☐ No, <i>Continue to 38</i>
38. Does the patient have a contraindication to another conventional synthetic drug (e.g., sulfasalazine)? <i>ACTION REQUIRED</i> : If Yes, please attach documentation of clinical reason to avoid therapy. ☐ Yes, <i>Continue to 50</i> ☐ No, <i>Continue to 50</i>
39. Please indicate the contraindication methotrexate or leflunomide.
\square Clinical diagnosis of alcohol use disorder, alcoholic liver disease, or other chronic liver disease, <i>Continue to 50</i>
☐ Drug interaction, Continue to 50
☐ Risk of treatment-related toxicity, <i>Continue to 50</i>
☐ Pregnancy or currently planning pregnancy, <i>Continue to 50</i>
☐ Breastfeeding, <i>Continue to 50</i> ☐ Significant comorbidity prohibits use of systemic agents (e.g., liver or kidney disease, blood dyscrasias, uncontrolled hypertension), <i>Continue to 50</i>

☐ Hypersensitivity, <i>Continue to 50</i>	
☐ History of intolerance or adverse ev	vent, Continue to 50
☐ Other, please specify	
40. Has the patient been diagnosed wit ☐ Yes, Continue to 41 ☐ No, Continue to 41	th moderately to severely active Crohn's disease (CD)?
41. Is the requested drug being prescri ☐ Yes, Continue to 42 ☐ No, Continue to 42	bed by or in consultation with a gastroenterologist?
42. Which of the following applies to	this request for the requested drug?
☐ Initiation of the intravenous (IV) lo	ading dose, Continue to 50
\square Initiation of the subcutaneous (SQ)	maintenance dose, Continue to 50
☐ Continuation of the subcutaneous (S	SQ) maintenance dose, Continue to 43
evidenced by low disease activity or in with the requested drug? ACTION RE documentation of remission.	nined remission OR achieved or maintained a positive clinical response as improvement in signs and symptoms of the condition since starting treatment EQUIRED: If Yes, please attach chart notes or medical record sion ACTION REQUIRED: Submit supporting documentation, Continue to
50	to in the second supporting documentation, community
☐ Yes, achieved or maintained a posit	tive clinical response, Continue to 44
☐ None of the above, <i>No further quest</i>	•
	tient experienced improvement in from baseline? <i>ACTION REQUIRED</i> : ecord documentation supporting positive clinical response.
☐ Abdominal pain or tenderness <i>ACT</i>	TION REQUIRED: Submit supporting documentation, Continue to 50
☐ Diarrhea ACTION REQUIRED: Sa	ubmit supporting documentation, Continue to 50
☐ Body weight ACTION REQUIRED	D: Submit supporting documentation, Continue to 50
☐ Abdominal mass ACTION REQUI	(RED : Submit supporting documentation, Continue to 50
☐ Appearance of the mucosa on endos	: Submit supporting documentation, Continue to 50 scopy, computed tomography enterography (CTE), magnetic resonance asound ACTION REQUIRED: Submit supporting documentation,
	scoring tool (e.g., Crohn's Disease Activity Index [CDAI] score) <i>ACTION umentation, Continue to 50</i>
☐ None of the above, <i>Continue to 50</i>	
45. Has the patient been diagnosed wit ☐ Yes, Continue to 46 ☐ No, Continue to 46	th moderately to severely active ulcerative colitis (UC)?

46. Is the requested drug being prescribed by or in consultation with a gastroenterologist? ☐ Yes, Continue to 47 ☐ No, Continue to 47
47. Which of the following applies to this request for the requested drug?
☐ Initiation of the intravenous (IV) loading dose, <i>Continue to 50</i>
☐ Initiation of the subcutaneous (SQ) maintenance dose, <i>Continue to 50</i>
☐ Continuation of the subcutaneous (SQ) maintenance dose, <i>Continue to 48</i>
48. Has the patient achieved or maintained remission OR 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? <i>ACTION REQUIRED</i> : If Yes, please attach chart notes or medical record documentation of remission.
☐ Yes, achieved or maintained remission <i>ACTION REQUIRED</i> : Submit supporting documentation, Continue to 50
☐ Yes, achieved or maintained a positive clinical response, <i>Continue to 49</i>
☐ None of the above, <i>No further questions</i>
49. Which of the following has the patient experienced improvement in from baseline? <i>ACTION REQUIRED</i> : Please attach chart notes or medical record documentation supporting positive clinical response.
☐ Stool frequency ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Rectal bleeding ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Urgency of defecation ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ C-reactive protein (CRP) ACTION REQUIRED: Submit supporting documentation, Continue to 50
☐ Fecal calprotectin (FC) <i>ACTION REQUIRED</i> : Submit supporting documentation, Continue to 50 ☐ Appearance of the mucosa on endoscopy, computed tomography enterography (CTE), magnetic resonance enterography (MRE), or intestinal ultrasound <i>ACTION REQUIRED</i> : Submit supporting documentation, Continue to 50
☐ Improvement on a disease activity scoring tool (e.g., Ulcerative Colitis Endoscopic Index of Severity [UCEIS], Mayo Score) <i>ACTION REQUIRED</i> : Submit supporting documentation, Continue to 50
☐ None of the above, <i>Continue to 50</i>
50. What is the diagnosis?
☐ Plaque Psoriasis, <i>Continue to 51</i>
☐ Psoriatic arthritis WITH co-existent plaque psoriasis, <i>Continue to 51</i>
☐ Psoriatic arthritis, <i>Continue to 51</i>
☐ Crohn's disease, Continue to 56
☐ Ulcerative colitis, <i>Continue to 57</i>
51. Is the patient currently receiving the requested drug? ☐ Yes, Continue to 52 ☐ No, Continue to 54

52. Does the prescribed dose exceed 150 mg? ☐ Yes, Continue to 53 ☐ No, Continue to 53
53. Is the prescribed frequency for the maintenance dose more frequent than one dose every 12 weeks? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
54. Does the prescribed dose exceed a loading dose of 150 mg at weeks 0 and 4, and a maintenance dose of 150 mg thereafter? ☐ Yes, Continue to 55 ☐ No, Continue to 55
55. Is the prescribed frequency for the maintenance dose more frequent than one dose every 12 weeks? ☐ Yes, <i>No Further Questions</i> ☐ No, <i>No Further Questions</i>
56. Which of the following applies to this request for the requested drug?
☐ Initiation of the intravenous (IV) loading dose, <i>Continue to 58</i>
☐ Initiation of the subcutaneous (SQ) maintenance dose, <i>Continue to 62</i>
☐ Continuation of the subcutaneous (SQ) maintenance dose, <i>Continue to 62</i>
57. Which of the following applies to this request for the requested drug?
☐ Initiation of the intravenous (IV) loading dose, <i>Continue to 60</i>
☐ Initiation of the subcutaneous (SQ) maintenance dose, <i>Continue to 62</i>
☐ Continuation of the subcutaneous (SQ) maintenance dose, <i>Continue to 62</i>
58. Does the prescribed dose exceed a loading dose of 600 mg at weeks 0, 4, and 8, and a maintenance dose of 360 mg thereafter? ☐ Yes, Continue to 59 ☐ No, Continue to 59
59. Is the prescribed frequency starting at week 12 for the maintenance dose more frequent than one dose every 8 weeks? ☐ Yes, No Further Questions ☐ No, No Further Questions
60. Does the prescribed dose exceed a loading dose of 1200 mg at weeks 0, 4, and 8, and a maintenance dose of 360 mg thereafter? Yes, <i>Continue to 61</i> No, <i>Continue to 61</i>
61. Is the prescribed frequency starting at week 12 for the maintenance dose more frequent than one dose every 8 weeks?

☐ Yes, No Further Questions	
□ No, No Further Questions	
62. Does the prescribed dose exceed 360 mg?	
☐ Yes, Continue to 63 ☐ No, Continue to 63	
63. Is the prescribed frequency for the maintenance dose more frequent than one Yes, <i>No Further Questions</i>	e dose every 8 weeks?
□ No, No Further Questions	
I attest that this information is accurate and true, and that documentation and the same and the same are the same and the same are the	11 0
K	
Prescriber or Authorized Signature	Date (mm/dd/yy)