

# SUBMITTING A LETTER OF MEDICAL NECESSITY

#### INDICATIONS<sup>1</sup>

**Plaque Psoriasis:** SKYRIZI is indicated for the treatment of moderate to severe plaque psoriasis in adults who are candidates for systemic therapy or phototherapy.

Psoriatic Arthritis: SKYRIZI is indicated for the treatment of active psoriatic arthritis in adults.

**Crohn's Disease**: SKYRIZI is indicated for the treatment of moderately to severely active Crohn's disease in adults.

**Ulcerative Colitis:** SKYRIZI is indicated for the treatment of moderately to severely active ulcerative colitis in adults.

#### **SAFETY CONSIDERATIONS**<sup>1</sup>

SKYRIZI is contraindicated in patients with a history of serious hypersensitivity reaction to risankizumab-rzaa or any of its excipients. Serious hypersensitivity reactions, including anaphylaxis, have been reported with use of SKYRIZI. If a serious hypersensitivity reaction occurs, discontinue SKYRIZI and initiate appropriate therapy immediately. SKYRIZI may increase the risk of infection. Instruct patients to report signs or symptoms of clinically important infection during treatment. Should such an infection occur, discontinue SKYRIZI until infection resolves. Evaluate patients for tuberculosis infection prior to initiating treatment with SKYRIZI. Drug-induced liver injury was reported in a patient with Crohn's disease during induction dosing of SKYRIZI. For the treatment of Crohn's disease and ulcerative colitis, evaluate liver enzymes and bilirubin at baseline and during induction (12 weeks). Interrupt treatment with SKYRIZI if drug-induced liver injury is suspected, until this diagnosis is excluded. Avoid use of live vaccines in SKYRIZI patients.



Please see additional Important Safety Information on page 4.

Please click here for full Prescribing Information.

### SUBMITTING A LETTER OF MEDICAL NECESSITY



#### You may need to provide a letter of medical necessity (LMN) if:

- Your patient's claim was denied and you are submitting an appeal letter
- You are requesting a formulary exception or tiering exception to get access for your patient



#### Make sure you have the following for an efficient submission of your LMN:

- Patient's insurance policy/ID number
- · Case ID number if a decision has already been rendered
- Patient's full name, plan identification number, and date of birth
- A brief medical history, including diagnosis, allergies, existing comorbidities, and International Classification of Disease (ICD) code(s)
- Clinical support for your recommendation
- Your office contact information

For support in person or over the phone, call an Access Specialist at 1.877.COMPLETE (1.877.266.7538)

Please see <u>Indications</u> and <u>Important Safety Information</u> on page 4. Please click here for full <u>Prescribing Information</u>.





## SAMPLE LETTER OF MEDICAL NECESSITY

Ask the payer whether a specific form is required to help establish medical necessity. Follow up with the payer if your office does not receive notification of the decision in a timely manner.

Date:	
[Payer Name]	
[Payer Address]	
[Appeals Department]	
Re: [Patient Name]	
[Policy ID/Group Number]	
Date of Service:	
To whom it may concern:	
My name is [Name]	and [Board-certified medical specialty] [NPI] writing on behalf or
my patient, [Patient name]	, to request coverage for [product, dosage, and frequency]
[Patient name]	has been under my care for 0 months for the treatment of [disease or symptoms]
[Product name]	nedical necessity because, after working with [Patient name] is the best treatment for this patient, and it's important that a formulary
exception be made.	is the best treatment for this patient, and it's important that a formulary
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This information is presented for informational purposes only and is not intended to provide reimbursement or legal advice. Providers are encouraged to contact third-party payers for specific information about their coverage policies. For more information, please call an Access Specialist at 1.877.COMPLETE (1.877.266.7538).

Digital version available at CompletePro.com and SkyriziHCP.com

Please see <u>Indications</u> and <u>Important Safety Information</u> on page 4. Please click here for full <u>Prescribing Information</u>.





## INDICATIONS AND IMPORTANT SAFETY INFORMATION FOR SKYRIZI® (risankizumab-rzaa)

#### INDICATIONS<sup>1</sup>

**Plaque Psoriasis:** SKYRIZI is indicated for the treatment of moderate to severe plaque psoriasis in adults who are candidates for systemic therapy or phototherapy.

**Psoriatic Arthritis:** SKYRIZI is indicated for the treatment of active psoriatic arthritis in adults.

**Crohn's Disease:** SKYRIZI is indicated for the treatment of moderately to severely active Crohn's disease in adults.

**Ulcerative Colitis:** SKYRIZI is indicated for the treatment of moderately to severely active ulcerative colitis in adults.

## IMPORTANT SAFETY INFORMATION<sup>1</sup> Hypersensitivity Reactions

SKYRIZI® (risankizumab-rzaa) is contraindicated in patients with a history of serious hypersensitivity reaction to risankizumab-rzaa or any of the excipients. Serious hypersensitivity reactions, including anaphylaxis, have been reported with the use of SKYRIZI. If a serious hypersensitivity reaction occurs, discontinue SKYRIZI and initiate appropriate therapy immediately.

#### Infection

SKYRIZI may increase the risk of infection. Do not initiate treatment with SKYRIZI in patients with a clinically important active infection until it resolves or is adequately treated.

In patients with a chronic infection or a history of recurrent infection, consider the risks and benefits prior to prescribing SKYRIZI. Instruct patients to seek medical advice if signs or symptoms of clinically important infection occur. If a patient develops such an infection or is not responding to standard therapy, closely monitor and discontinue SKYRIZI until the infection resolves.

#### **Tuberculosis (TB)**

Prior to initiating treatment with SKYRIZI, evaluate for TB infection and consider treatment in patients with latent or active TB for whom an adequate course of treatment cannot be confirmed. Monitor patients for signs and symptoms of active TB during and after SKYRIZI treatment. Do not administer SKYRIZI to patients with active TB.

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#### Hepatotoxicity in Treatment of Inflammatory Bowel Disease

Drug-induced liver injury was reported in a patient with Crohn's disease who was hospitalized for a rash during induction dosing of SKYRIZI. For the treatment of Crohn's disease and ulcerative colitis, evaluate liver enzymes and bilirubin at baseline and during induction (12 weeks); monitor thereafter according to routine patient management. Consider an alternate treatment for patients with evidence of liver cirrhosis. Interrupt treatment if drug-induced liver injury is suspected, until this diagnosis is excluded. Instruct your patient to seek immediate medical attention if they experience symptoms suggestive of hepatic dysfunction.

#### **Administration of Vaccines**

Avoid use of live vaccines in patients treated with SKYRIZI. Medications that interact with the immune system may increase the risk of infection following administration of live vaccines. Prior to initiating SKYRIZI, complete all age-appropriate vaccinations according to current immunization guidelines.

#### **Adverse Reactions**

Most common (≥1%) adverse reactions associated with SKYRIZI in plaque psoriasis and psoriatic arthritis include upper respiratory infections, headache, fatigue, injection site reactions, and tinea infections.

In psoriatic arthritis phase 3 trials, the incidence of hepatic events was higher with SKYRIZI compared to placebo. Most common (>3%) adverse reactions associated with SKYRIZI in Crohn's disease are upper respiratory infections, headache, and arthralgia in induction, and arthralgia, abdominal pain, injection site reactions, anemia, pyrexia, back pain, arthropathy, and urinary tract infection in maintenance. Most common (≥3%) adverse reactions associated with SKYRIZI in ulcerative colitis are arthralgia in induction, and arthralgia, pyrexia, injection site reactions, and rash in maintenance.

**Lipid Elevations**: Increases from baseline and increases relative to placebo were observed at Week 4 and remained stable to Week 12 in patients treated with SKYRIZI in Crohn's disease. Lipid elevations observed in patients with ulcerative colitis were similar to those in Crohn's disease.

**Dosage Forms and Strengths:** SKYRIZI (risankizumab-rzaa) is available in a 150 mg/mL prefilled syringe and pen, a 600 mg/10 mL single-dose vial for intravenous infusion, and a 180 mg/1.2 mL or 360 mg/2.4 mL single-dose prefilled cartridge with on-body injector.

Reference: 1. SKYRIZI [package insert]. North Chicago, IL: AbbVie Inc.

