

SOTYKTU 360 SUPPORT ACCESS & REIMBURSEMENT GUIDE SIMPLY COMMITTED TO PATIENT SUPPORT



SOTYKTU 360 SUPPORT Overview

Simply committed to patient support

START



Enroll Patients With the Start Form

See slide **7** for additional details on how to complete and submit the SOTYKTU Start Form.



SOTYKTU Free Trial Offer*

Patients new to SOTYKTU may be eligible to receive a **30-day** free trial in the mail.* See slide **13** for more details.



Benefits Investigation

SOTYKTU 360 SUPPORT can help conduct a benefits investigation and share the results with your office and the patient. See slide 15 for more details.



Prior Authorization (PA) & Appeals Support

If a PA or appeal is needed, SOTYKTU 360 SUPPORT can communicate requirements to your office and can check the status once submitted. See slide 17 for more details



Additional Resources

See slide **26** for information for your patients.

STAY



SOTYKTU Bridge Program*

Eligible, commercially insured patients may access SOTYKTU for up to 3 years if there is a delay or denial during the insurance coverage determination process. See slide **21** for more details.

SAVE



SOTYKTU Co-Pay Assistance Program*

Eligible, commercially insured patients may pay as little as \$0 every month for SOTYKTU with the SOTYKTU Co-Pay Assistance Card. See slide 23 for more details.



Third Party Referrals

SOTYKTU 360 SUPPORT may be able to help identify possible independent financial support options for patients with affordability concerns. See slide 24 for more details.



^{*}Please click here to review Program Terms and Conditions.

The accurate completion of reimbursement or coverage-related documentation is the responsibility of the healthcare provider and patient. Bristol-Myers Squibb and its agents make no guarantee regarding reimbursement for any service or item.

SOTYKU (deucravacitinib) Indication and Important Safety Information



INDICATION

SOTYKTU™ (deucravacitinib) is indicated for the treatment of moderate-to-severe plaque psoriasis in adults who are candidates for systemic therapy or phototherapy. <u>Limitations of Use:</u>

SOTYKTU is not recommended for use in combination with other potent immunosuppressants.

IMPORTANT SAFETY INFORMATION

CONTRAINDICATIONS

SOTYKTU is contraindicated in patients with a history of hypersensitivity reaction to deucravacitinib or to any of the excipients in SOTYKTU.

WARNINGS AND PRECAUTIONS

Hypersensitivity: Hypersensitivity reactions such as angioedema have been reported. If a clinically significant hypersensitivity reaction occurs, institute appropriate therapy and discontinue SOTYKTU.

Infections: SOTYKTU may increase the risk of infections. Serious infections have been reported in patients with psoriasis who received SOTYKTU. The most common serious infections reported with SOTYKTU included pneumonia and COVID-19. Avoid use of SOTYKTU in patients with an active or serious infection. Consider the risks and benefits of treatment prior to initiating SOTYKTU in patients:

- · with chronic or recurrent infection
- · who have been exposed to tuberculosis
- with a history of a serious or an opportunistic infection
- · with underlying conditions that may predispose them to infection.

Closely monitor patients for the development of signs and symptoms of infection during and after treatment. A patient who develops a new infection during treatment should undergo prompt and complete diagnostic testing, have appropriate antimicrobial therapy initiated and be closely monitored. Interrupt SOTYKTU if a patient develops a serious infection. Do not resume SOTYKTU until the infection resolves or is adequately treated.

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SOTYKU (deucravacitinib) Indication and Important Safety Information (cont'd)



WARNINGS AND PRECAUTIONS (cont'd)

Viral Reactivation

Herpes virus reactivation (e.g., herpes zoster, herpes simplex) was reported in clinical trials with SOTYKTU. Through Week 16, herpes simplex infections were reported in 17 patients (6.8 per 100 patient-years) treated with SOTYKTU, and 1 patient (0.8 per 100 patient-years) treated with placebo. Multidermatomal herpes zoster was reported in an immunocompetent patient. During PSO-1, PSO-2, and the open-label extension trial, the majority of patients who reported events of herpes zoster while receiving SOTYKTU were under 50 years of age. The impact of SOTYKTU on chronic viral hepatitis reactivation is unknown. Consider viral hepatitis screening and monitoring for reactivation in accordance with clinical guidelines before starting and during therapy with SOTYKTU. If signs of reactivation occur, consult a hepatitis specialist. SOTYKTU is not recommended for use in patients with active hepatitis B or hepatitis C.

Tuberculosis (TB): In clinical trials, of 4 patients with latent TB who were treated with SOTYKTU and received appropriate TB prophylaxis, no patients developed active TB (during the mean follow-up of 34 weeks). One patient, who did not have latent TB, developed active TB after receiving 54 weeks of SOTYKTU. Evaluate patients for latent and active TB infection prior to initiating treatment with SOTYKTU. Do not administer SOTYKTU to patients with active TB. Initiate treatment of latent TB prior to administering SOTYKTU. Consider anti-TB therapy prior to initiation of SOTYKTU in patients with a past history of latent or active TB in whom an adequate course of treatment cannot be confirmed. Monitor patients for signs and symptoms of active TB during treatment.

Malignancy including Lymphomas: Malignancies, including lymphomas, were observed in clinical trials with SOTYKTU. Consider the benefits and risks for the individual patient prior to initiating or continuing therapy with SOTYKTU, particularly in patients with a known malignancy (other than a successfully treated non-melanoma skin cancer) and patients who develop a malignancy when on treatment with SOTYKTU.

Rhabdomyolysis and Elevated CPK: Treatment with SOTYKTU was associated with an increased incidence of asymptomatic creatine phosphokinase (CPK) elevation and rhabdomyolysis compared to placebo. Discontinue SOTYKTU if markedly elevated CPK levels occur or myopathy is diagnosed or suspected. Instruct patients to promptly report unexplained muscle pain, tenderness or weakness, particularly if accompanied by malaise or fever.

Laboratory Abnormalities: Treatment with SOTYKTU was associated with increases in triglyceride levels. Periodically evaluate serum triglycerides according to clinical guidelines during treatment. SOTYKTU treatment was associated with an increase in the incidence of liver enzyme elevation compared to placebo. Evaluate liver enzymes at baseline and thereafter in patients with known or suspected liver disease according to routine management. If treatment-related increases in liver enzymes occur and drug-induced liver injury is suspected, interrupt SOTYKTU until a diagnosis of liver injury is excluded.

Continued on next slide



SOTYKU (deucravacitinib) Indication and Important Safety Information (cont'd)



WARNINGS AND PRECAUTIONS (cont'd)

Immunizations: Prior to initiating therapy with SOTYKTU, consider completion of all age-appropriate immunizations according to current immunization guidelines including prophylactic herpes zoster vaccination. Avoid use of live vaccines in patients treated with SOTYKTU. The response to live or non-live vaccines has not been evaluated.

Potential Risks Related to JAK Inhibition: It is not known whether tyrosine kinase 2 (TYK2) inhibition may be associated with the observed or potential adverse reactions of Janus Kinase (JAK) inhibition. In a large, randomized, postmarketing safety trial of a JAK inhibitor in rheumatoid arthritis (RA), patients 50 years of age and older with at least one cardiovascular risk factor, higher rates of all-cause mortality, including sudden cardiovascular death, major adverse cardiovascular events, overall thrombosis, deep venous thrombosis, pulmonary embolism, and malignancies (excluding non-melanoma skin cancer) were observed in patients treated with the JAK inhibitor compared to those treated with TNF blockers. SOTYKTU is not approved for use in RA.

ADVERSE REACTIONS

Most common adverse reactions (≥1% of patients on SOTYKTU and more frequently than with placebo) include upper respiratory infections, blood creatine phosphokinase increased, herpes simplex, mouth ulcers, folliculitis and acne.

SPECIFIC POPULATIONS

Pregnancy: Available data from case reports on SOTYKTU use during pregnancy are insufficient to evaluate a drug-associated risk of major birth defects, miscarriage, or adverse maternal or fetal outcomes. Report pregnancies to the Bristol-Myers Squibb Company's Adverse Event reporting line at 1-800-721-5072.

Lactation: There are no data on the presence of SOTYKTU in human milk, the effects on the breastfed infant, or the effects on milk production. SOTYKTU is present in rat milk. When a drug is present in animal milk, it is likely that the drug will be present in human milk. The developmental and health benefits of breastfeeding should be considered along with the mother's clinical need for SOTYKTU and any potential adverse effects on the breastfed infant from SOTYKTU or from the underlying maternal condition.

Hepatic Impairment: SOTYKTU is not recommended for use in patients with severe hepatic impairment.

SOTYKTU is available in 6 mg tablets.

Please see U.S. Full Prescribing Information, including Medication Guide, for SOTYKTU



It begins with your SOTYKTU 360 SUPPORT team

Your SOTYKTU 360 SUPPORT team:





DEDICATED SOTYKTU Support Coordinators*

- Assist with benefits investigations and prior authorizations (PAs)
- Explain insurance coverage and pharmacy benefits
- Help eligible patients access SOTYKTU if it isn't initially covered by commercial or private insurance via the SOTYKTU Bridge Program[†]
- Enroll eligible patients into the SOTYKTU Co-Pay Assistance Program[†]
- Identify and coordinate with a specialty pharmacy to arrange shipments
- Provide patients with information about other available resources

Available to you and your patients from 8 AM to 11 PM ET, Monday through Friday, at **1-888-SOTYKTU** (768-9588).



Access and Reimbursement Managers (ARMs)

- Educate on SOTYKTU 360 SUPPORT after a prescribing decision has been made
- Assist with patient access challenges
- Provide timely responses to access and reimbursement questions
- · Share knowledge regarding local access landscape

Contact your ARM for general access or reimbursement support questions and to schedule an office visit.



^{*}SOTYKTU Support Coordinators can provide general information about SOTYKTU but cannot provide medical advice.

[†]Please click here to review Program Terms and Conditions.

The accurate completion of reimbursement or coverage-related documentation is the responsibility of the healthcare provider and patient. Bristol-Myers Squibb and its agents make no guarantee regarding reimbursement for any service or item.

Enrolling your patients in SOTYKTU 360 SUPPORT

Get started in 3 steps. To enroll your patient, assist them in completing the Start Form





SELECT

Complete the Start Form and, if applicable, **select** the following options:

- SOTYKTU Free Trial Offer*
- Maintenance Dose
- SOTYKTU Bridge Program*†



SIGN

- After you sign, ensure that patients sign the Patient Authorization
- Eligible, commercially insured patients can enroll in the SOTYKTU Co-Pay Assistance* Program by signing in the designated area



SUBMIT

• Fax the fully completed and signed Start Form to 1-888-381-0029 or submit through CoverMyMeds.com. To get started, create an account on CoverMyMeds.com

NOW IT'S OUR TURN

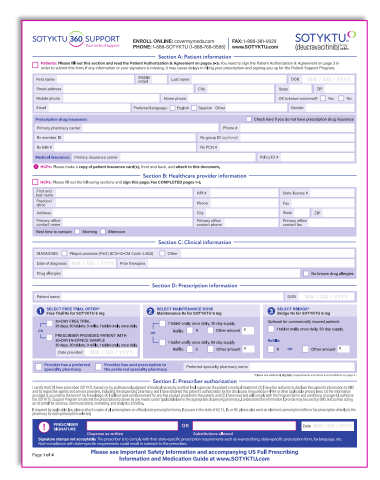
After submitting the Start Form, let your patients know that their SOTYKTU Support Coordinator will be in touch to help patients access prescribed medication.



[†]For patients eligible for the SOTYKTU Bridge Program, submit a Prior Authorization (PA) to the patient's payer as soon as possible. If the PA is denied, submit an appeal, exception, and/or Letter of Medical Necessity within 90 days or per payer requirements.

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SOTYKTU (360) SUPPORT





Steps to complete the SOTYKTU Start Form



FIRST

Complete Patient Information in Section A on page 1 of the form

SECOND

Complete the HCP information in Sections B-E on page 1 of the form

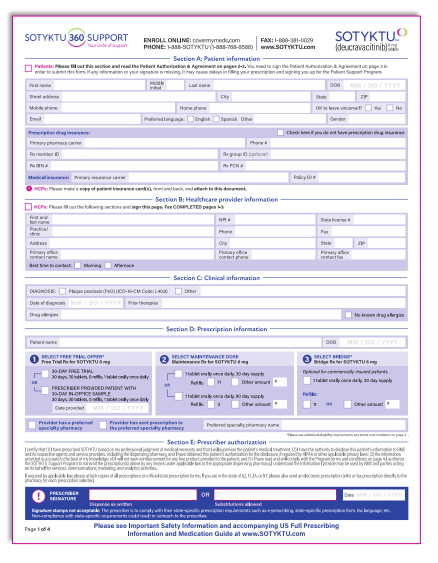
THIRD

Have the patient sign and date the bottom of the Patient Authorization Agreement (PAA) on page 3



All sections marked with a pink exclamation point must be filled out for the form to be processed.

NOTE: Once you have completed these 3 sections, you have fully completed the Start Form and are ready to submit.





Completing the SOTYKTU Start Form

PATIENT INFORMATION

Fill Out the Start Form on Page 1, Section A

The Basics

As you fill out the form, be sure to include:

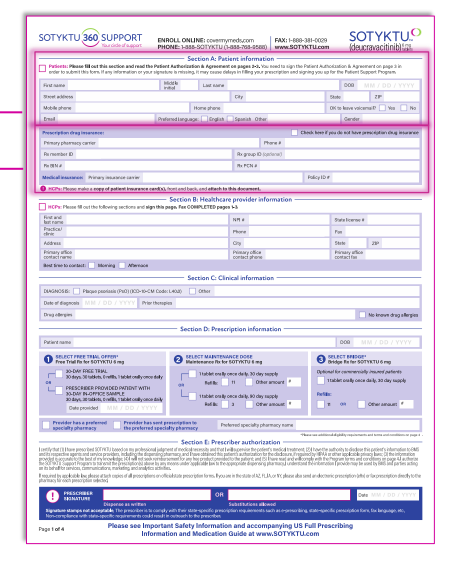
- Name and date of birth: The patient's full name and date of birth are required for processing
- Phone number: Mobile or home phone number is required to allow us to contact patients with additional questions or notifications
- Email: Patients can receive communications and important updates about product shipments through email

Prescription Drug Insurance

 Insurance: Filling in the right insurance carrier and policy number is required to determine whether the patient may be approved for SOTYKTU and covered for the cost of therapy

Important Note for You and Your Patients:

 All patients should read: Patient Authorization and Agreement (PAA) (pages 2-3)





TO HOME

Completing the SOTYKTU Start Form (cont'd)

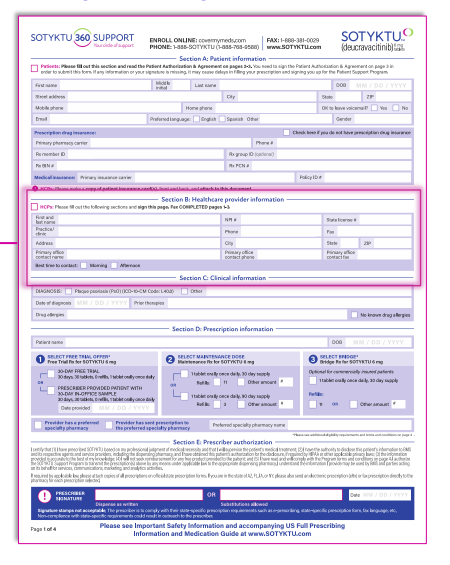
HCP INFORMATION

Fill Out the Start Form on Page 1, Sections B-E

Prescriber Information

As you fill out the form, be sure to include:

Your Name, address, and contact information: Just as with the
patient, it's required to have your full name and phone number so we can
contact you directly as we process the Start Form; also, be sure to
include the address of your office, your NPI number, and your State
Medical License number







Completing the SOTYKTU Start Form (cont'd)

Treatment Information

- **Diagnosis:** It is required to identify the patient's diagnosis
- SOTYKTU Free Trial Offer*: The SOTYKTU Free Trial Offer for SOTYKTU includes a 30-day supply of SOTYKTU that is available at no cost to eligible patients through SOTYKTU 360 SUPPORT. See the SOTYKTU Free Trial Offer full terms and conditions on page 4 of the Start Form
- Maintenance Dose: 30-day and 90-day supplies of the SOTYKTU maintenance dose can be ordered using this form. Just check the appropriate box and indicate the number of refills you may want to prescribe. You may also indicate if there is a preferred Specialty Pharmacy to use
- SOTYKTU Bridge Program*: Patients experiencing a delay or denial with coverage may be eligible for the SOTYKTU Bridge Program. Eligible patients with commercial or private insurance may be able to receive SOTYKTU free of charge for up to 3 years while awaiting coverage. See the SOTYKTU Bridge Program full terms and conditions on page 4 of the Start Form

Prescriber Authorization

Signature: Be sure to sign the form when you are finished. Your signature is required in order for the Start Form to be processed





The accurate completion of reimbursement or coverage-related documentation is the responsibility of the healthcare provider and patient. Bristol-Myers Squibb and its agents make no guarantee regarding reimbursement for any service or item.





TO HOME

Completing the SOTYKTU Start Form (cont'd)

PATIENT SIGNATURES

Obtain Signature and Date on Page 3

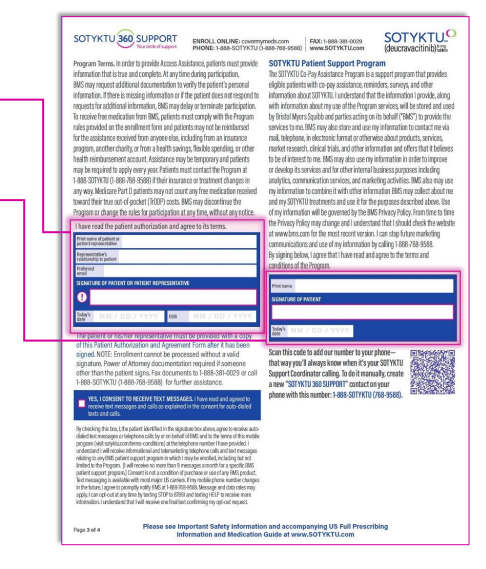
Patient Authorization

The patient's signature and date are required. If patients prefer to fill out the form electronically, they can visit <u>SOTYKTU.com/esign</u> to provide an electronic signature.

SOTYKTU Co-Pay Assistance* Program Enrollment

Patients may choose to enroll in the SOTYKTU Co-Pay Assistance Program, if they are eligible, by signing in the designated area. Eligible, commercially insured patients may pay as little as \$0 per month for SOTYKTU.

Once the patient has signed the form, you should provide them with a **photocopy** of the signature page as well as page 4 with the program terms and conditions. Be sure to keep the **original** signature page for your office, as you will need it for your submission.







TO HOME

^{*}Please click here to review Program Terms and Conditions.

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SOTYKTU Free Trial Offer

For eligible patients getting started on SOTYKTU





First-time eligible patients may receive a 30-day free trial in the mail*

Complete and submit the Start Form through CoverMyMeds.com or fax it to 1-888-381-0029 for the 30-day SOTYKTU Free Trial Offer

ELIGIBILITY REQUIREMENTS

To be eligible for the SOTYKTU Free Trial Offer for SOTYKTU (deucravacitinib)*:

- Patients must be new patients who have not previously received a sample or filled a prescription for SOTYKTU
- Patient must have a valid 30-day prescription for SOTYKTU for an on-label indication
- Patients are 18 years of age or older
- Patients are residents of the United States or a US Territory





Coverage and Access to SOTYKTU

Support during the coverage determination process

After your patient is enrolled, their SOTYKTU Support Coordinator will:





FAX A SUMMARY OF BENEFITS FORM TO YOUR OFFICE AFTER COVERAGE IS VERIFIED

- Summary of Benefits form includes SOTYKTU out-of-pocket costs, expected coverage, PA requirements, and co-pay and bridge eligibility
- Results will be faxed to your office



PROVIDE PAYER-SPECIFIC PA FORMS AND ASSIST WITH APPEALS, IF NEEDED:

Access template letters are available on <u>SOTYKTUHCP.com</u>

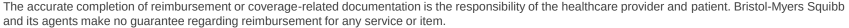


CALL YOUR PATIENTS TO HELP THEM UNDERSTAND:

- Information about their insurance coverage and out-of-pocket costs for SOTYKTU
- If they may be eligible for the SOTYKTU Co-Pay Assistance Program*
- Whether there may be other programs or resources to assist them with treatment access



^{*}Please click here to review Program Terms and Conditions.





Key areas of the Summary of Benefits form

SOTYKTU 360 SUPPORT during the coverage determination process





PATIENT ELIGIBILITY FOR SOTYKTU CO-PAY ASSISTANCE* AND/OR SOTYKTU BRIDGE PROGRAM* WILL BE INCLUDED ON THE SUMMARY OF BENEFITS FORM, IF APPLICABLE

PRESCRIPTION COVERAGE MAY BE REPORTED AS: COVERED, NOT COVERED, OR UNDISCLOSED

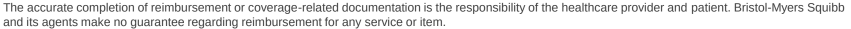
- 2
- If covered, the report will include the patient's:
 - Deductible
 - Co-Pay and frequency of payment and/or
 - Coinsurance with an out-of-pocket maximum
- If not covered, or if a PA or formulary exception request is denied by the plan, it can be appealed;
 - Refer to the template letter of appeal on <u>SOTYKTUHCP.com</u>
 - If an NDC block is in place, a formulary exception letter may be submitted to request its removal.
 Refer to the template letter of formulary exception on <u>SOTYKTUHCP.com</u>
 - Additional payer-specific documents may be sent along with the Summary of Benefits form, if applicable

3

THE AUTHORIZATION INFORMATION REQUIREMENTS SECTION WILL INDICATE IF A PRIOR AUTHORIZATION (PA) IS REQUIRED:

- If a PA is required, the authorization number and how to submit the PA will be included
- Refer to the template letter of medical necessity and formulary exception letter on <u>SOTYKTUHCP.com</u> for potential next steps

^{*}Please click here to review Program Terms and Conditions.





SOTYKTU 360 Support SOTYKTU 360 SUPPORT Phone: 1-888-SOTYKTU (1-888-768-9588) Fax: 1-888-381-0029 Hours: Monday through Friday, 8 AM - 8 PM ET FROM: SOTYKTU 360 Support PATIENT NAME PATIENT DATE OF BIRTH PATIENT CASE ID: SUBJECT: Summary of Benefits PROVIDER INFORMATION Prescriber: PHARMACY INSURANCE COVERAGE INFORMATIO Policy Effective Date (MM/DD/YYYY) □ Covered ☐ Not Covered Co-Pay (30 day): □ Undisclosed Coverage Details/Comments: AUTHORIZATION INFORMATION/REQUIREMENTS Fax PA results to SOTYKTU 360 Support at 1-888-381-0029 Inis accument is provided for information purposed only and is not intended to provide i SOTYKTU 360 Support do not guarantee payer relimbursement for product treatment an werranties, expressed or implied, as to the accuracy or completeness of the information

SOTYKTU prior authorization and appeal checklist



The checklist below highlights items and information that may help support a prior authorization (PA) decision from a patient's health insurance plan. Be sure to review the insurer's guidelines for obtaining a PA, as these can differ by insurer, the medication being prescribed, and other factors.

- Review the health plan's PA submission options
- Complete a PA request form (if required by patient's health plan)
- Provide documentation supporting the treatment decision
- Include patient prescription insurance information:
 Copy the front and back of the prescription insurance card
- Confirm the health plan received the PA request
- Ocument the PA approval number and duration
- When you receive PA determination, please fax to your SOTYKTU Support Coordinator

covermymeds®

CoverMyMeds offers electronic PA support.

An electronic PA is available for submitting and tracking prior authorizations.

To learn more, visit CoverMyMeds.com or call 1-800-705-9613



For additional information or assistance, please contact your SOTYKTU Support Coordinator at 1-888-SOTYKTU (768-9588)
8 AM to 11 PM ET, M-F



Supporting Information and Access Template Letters



The following supporting information may be included within or accompany your communications to request coverage for SOTYKTU.

Clinical information to support the treatment decision, for example (if applicable):

- Percentage of body surface area affected
- sPGA score and/or PASI score
- Body location of plaques including pictures of plaque severity if possible
- Date of patient diagnosis
- ICD-10 code
- Previous treatment(s) of plaque psoriasis, duration, patient's response and reason(s) for discontinuation

Additional supporting information:

- **Prescribing Information**
- Journal articles or clinical guidelines

and its agents make no guarantee regarding reimbursement for any service or item.



Download access letter templates



- Formulary exception letter
- Letter of appeal
- Letter of medical necessity
- Letter of reverification

The example templates above may be used to support requests for access to SOTYKTU. Information provided in the templates are for informational purposes for patients who have been prescribed SOTYKTU. Templates are not intended to substitute for a prescriber's independent, clinical decision-making. Completed letters must be submitted by the prescriber on the prescriber's letterhead, along with any relevant medical records.



^{*}Abbreviations: PA, Prior Authorization; sPGA, static Physicians Global Assessment; PASI, Psoriasis Area and Severity Index; IGA, Investigator's Global Assessment The accurate completion of reimbursement or coverage-related documentation is the responsibility of the healthcare provider and patient. Bristol-Myers Squibb

Formulary exceptions overview



A formulary exception may be requested:

- To obtain access to a product that has been prescribed for a patient but is not included on a plan's formulary, or
- For removal of a utilization management requirement for a formulary product, such as:



Step Therapy (ST) / Step Edit (ED)

A payer process in which patients must first try one therapy before they are permitted to "step" to another drug



Prior Authorization (PA)

A process through which a request for provisional affirmation of coverage is submitted to the insurance provider for review before the item or service is furnished to the beneficiary and before the claim is submitted for processing



Quantity Limit (QL)

The highest amount of a prescription drug that can be given to your patient by their pharmacy in a specified period of time (for example, 30 tablets per month)



For additional information or assistance, please contact your SOTYKTU Support Coordinator at 1-888-SOTYKTU (768-9588)

8 AM to 11 PM ET, M-F



Handling appeals



If coverage for SOTYKTU is denied, the treating HCP or patient may submit an appeal. The process flow below highlights some options for pursuing an appeal:



Plan Guidelines

Refer to the health plan's specific guidelines for appeals, as the plan may have multiple levels of appeals with different requirements



Targeted Response

The appeal letter should provide the prescriber's clinical rationale as to why the product preferred on the insurer's formulary is not appropriate



For additional information or assistance, please contact your SOTYKTU Support Coordinator at 1-888-SOTYKTU (768-9588)

8 AM to 11 PM ET, M-F



SOTYKTU Bridge Program

For commercially insured patients taking SOTYKTU who are denied or experience a delay in coverage





If denied coverage or experiencing a delay in coverage, commercially insured patients may be eligible to receive SOTYKTU at no cost for up to 3 years while awaiting a coverage determination*

The program begins when coverage is delayed after the initial submission of prior authorization (PA). If PA is denied, you must file an appeal within 90 days or per the payer requirements to keep your patient in the program.

ELIGIBILITY REQUIREMENTS

To be eligible for the SOTYKTU Bridge Program for SOTYKTU (deucravacitinib):

- A SOTYKTU prescription for an FDA-approved use
- Commercial insurance with coverage
- Submitting a Prior Authorization (PA) within 90 days of SOTYKTU Bridge Program enrollment
- Submitting an Appeal/Exception/Letter of Medical Necessity (LMN) to challenge PA payer outcome within 90 days or per payer guidelines of PA outcome if coverage is denied
- Program requires a periodic check of your insurance coverage status to confirm your continued eligibility, including, but not limited to the annual reverification process.
 Program is available until your commercial insurance covers your medication for up to 36 months (dispensed in 30-day prescriptions). Up to 12 months coverage for residents in Massachusetts, Minnesota, and Rhode Island
- · A signed Patient Authorization and Agreement (PAA) is on file
- US residents only
- SOTYKTU Bridge Program is not available to patients who have prescription insurance coverage through Medicare, Medicaid, or any other federal or state program



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Patient Financial Support

SOTYKTU Co-Pay Assistance Program





Commercially insured patients may **pay as little as \$0 every month** for SOTYKTU with the SOTYKTU Co-Pay Assistance Card*

To enroll in the SOTYKTU Co-Pay Assistance Program, eligible patients can call their SOTYKTU Support Coordinator at 1-888-SOTYKTU (768-9588), self-enroll at SOTYKTUCoPaySignup.com, or sign the designated area on the Start Form. If a patient is uninsured or underinsured, their SOTYKTU Support Coordinator can discuss what options may be available to them

ELIGIBILITY REQUIREMENTS

- Patients must have commercial (private) insurance, but their coverage does not cover the full cost of the prescription. Co-pay assistance is not valid where the entire cost of the prescription is reimbursed by insurance
- Patients are not eligible if they have prescription insurance coverage through a state or federal healthcare program, including but not limited to Medicare, Medicaid, Medigap, CHAMPUS, TRICARE, Veterans Affairs (VA), or Department of Defense (DOD) programs; patients who move from commercial to state or federal healthcare program insurance will no longer be eligible
- Cash-paying patients are not eligible for co-pay assistance
- · Patients must be 18 years of age or older
- Patients must live in the United States or United States territories
- Eligible patients with an activated co-pay card and a valid prescription may pay as little as \$0 per 30-day supply; monthly and annual maximum program benefits apply and may vary from patient to patient, depending on the terms of a patient's prescription drug plan and based on factors determined solely by Bristol-Myers Squibb

CO-PAY ACTIVATION OPTIONS

- Signature on page 3 of the Start Form: Direct activation by patient
- <u>SOTYKTUCoPaySignup.com</u>: Direct activation by patient
- 1-888-SOTYKTU (768-9588): Direct activation by patient, or the SOTYKTU Support Coordinator may contact patient before submitting the prescription to the specialty pharmacy



SOTYKTU 360 SUPPORT can help identify potential financial assistance programs for uninsured and underinsured patients





Referrals to independent charitable foundations that may be able to assist eligible uninsured or underinsured patients who have an established financial hardship



Referrals to Low-Income Subsidy Program (Medicare Extra Help)

It is important to note that charitable foundations and the Low-Income Subsidy Program are independent from Bristol Myers Squibb and have their own eligibility criteria and evaluation process. Bristol Myers Squibb cannot guarantee that a patient will receive assistance.





Additional Resources

Get committed support

Have questions or need assistance? There are 3 ways to get support:





your Access and Reimbursement
Manager for access and reimbursement
assistance or to schedule a conversation



1-888-SOTYKTU (768-9588)

8 AM to 11 PM ET • Monday to Friday to speak with your SOTYKTU Support Coordinator



VISIT

SOTYKTUHCP.com for information and resources, including the enrollment form, to help your patients with access to SOTYKTU



SOTYKTU select product information



SOTYKTU prescriptions can be filled at any specialty pharmacy*

For the SOTYKTU Free Trial Offer and SOTYKTU Bridge Program, please enroll eligible patients in SOTYKTU 360 SUPPORT through CoverMyMeds.com or fax: 1-888-381-0029[†]

- Indication: SOTYKTU is indicated for the treatment of adults with moderate-to-severe plaque psoriasis who are candidates for systemic therapy or phototherapy
- NDC 0003-0895-11 30 ct tablets
- ICD-10 Code: L40.0
- The dosage strength of SOTYKTU is 6 mg in tablet form. The treatment schedule is one tablet of SOTYKTU, once daily
- SOTYKTU should be stored at 68°F to 77°F (20°C to 25°C) in the original container or blister pack





 $^{{}^*\}text{Please be aware that some payers mandate a specific specialty pharmacy for SOTYKTU}.$

[†]Please click here to review Program Terms and Conditions.

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SOTYKTU 360 SUPPORT Your circle of support



