

Access and Reimbursement Guide for Healthcare Providers

Information on Distribution, Patient Support, Coverage, and Access

Enroll your patients at time of prescription to support the patient experience and access to programs



Please see the Important Safety Information on pages 3-4 and click here to see the full Prescribing Information for AYVAKIT.

Blueprint Medicines is pleased to provide this information to help you and your office staff navigate coverage and access for AYVAKIT[®] (avapritinib). It is not intended to supersede any individual payer guidance and/or processes. Please be sure to check directly with each patient's insurance for any specific requirements needed to help obtain coverage and access. This document is presented for informational purposes only and does not guarantee reimbursement.

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Product Information

INDICATION

AYVAKIT® (avapritinib) is indicated for the treatment of adult patients with:

Unresectable or Metastatic Gastrointestinal Stromal Tumor (GIST)

harboring a platelet-derived growth factor receptor alpha (PDGFRA) exon 18 mutation, including PDGFRA D842V mutations

Advanced SM (AdvSM)

including patients with aggressive systemic mastocytosis (ASM), systemic mastocytosis with an associated hematological neoplasm (SM-AHN), and mast cell leukemia (MCL)

<u>Limitations of Use:</u> AYVAKIT is not recommended for the treatment of patients with AdvSM with platelet counts of <50 X 10°/L

Indolent Systematic Mastocytosis (ISM)

Limitations of Use: AYVAKIT is not recommended for the treatment of patients with ISM with platelet counts of <50 x 10⁹/L

IMPORTANT SAFETY INFORMATION

There are no contraindications for AYVAKIT.

Intracranial Hemorrhage—Serious intracranial hemorrhage (ICH) may occur with AYVAKIT treatment; fatal events occurred in <1% of patients. Overall, ICH (e.g, subdural hematoma, ICH, and cerebral hemorrhage) occurred in 2.9% of 749 patients who received AYVAKIT in clinical trials. In GIST patients, ICH occurred in 3 of 267 patients (1.1%) and two (0.7%) of the events were Grade \geq 3 and resulted in discontinuation. In AdvSM patients who received AYVAKIT at 200 mg daily, ICH occurred in 2 of 75 patients (2.7%) who had platelet counts \geq 50 x 10⁹/L prior to initiation of therapy and in 3 of 80 patients (3.8%) regardless of platelet counts. In ISM patients, no events of ICH occurred in the 246 patients who received any dose of AYVAKIT in the PIONEER study.

Monitor patients closely for risk factors of ICH which may include history of vascular aneurysm, ICH or cerebrovascular accident within the prior year, concomitant use of anticoagulant drugs, or thrombocytopenia.

Symptoms of ICH may include headache, nausea, vomiting, vision changes, or altered mental status. Advise patients to seek immediate medical attention for signs or symptoms of ICH.

Permanently discontinue AYVAKIT if ICH of any grade occurs. In AdvSM patients, a platelet count must be performed prior to initiating therapy. AYVAKIT is not recommended in AdvSM patients with platelet counts <50 x 10⁹/L. Following treatment initiation, platelet counts must be performed every 2 weeks for the first 8 weeks. After 8 weeks of treatment, monitor platelet counts every 2 weeks or as clinically indicated based on platelet counts. Manage platelet counts of <50 x 10⁹/L by treatment interruption or dose reduction.

Cognitive Effects—Cognitive adverse reactions can occur in patients receiving AYVAKIT and occurred in 33% of 995 patients overall in patients who received AYVAKIT in clinical trials including: 41% of 601 GIST patients (5% were Grade ≥3), 28% of 148 AdvSM patients (3% were Grade ≥3), and 7.8% of patients with ISM who received AYVAKIT + best supportive care (BSC) versus 7.0% of patients who received placebo + BSC (<1% were Grade 3). Depending on the severity and indication, withhold AYVAKIT and then resume at same dose or at a reduced dose upon improvement, or permanently discontinue.

Photosensitivity—AYVAKIT may cause photosensitivity reactions. In all patients treated with AYVAKIT in clinical trials (n=1049), photosensitivity reactions occurred in 2.5% of patients. Advise patients to limit direct ultraviolet exposure during treatment with AYVAKIT and for one week after discontinuation of treatment.

Embryo-Fetal Toxicity—AYVAKIT can cause fetal harm when administered to a pregnant woman. Advise pregnant women of the potential risk to a fetus. Advise females and males of reproductive potential to use an effective contraception during treatment with AYVAKIT and for 6 weeks after the final dose of AYVAKIT. Advise women not to breastfeed during treatment with AYVAKIT and for 2 weeks following the final dose.

IMPORTANT SAFETY INFORMATION (CONT.)

Adverse Reactions—The most common adverse reactions (>20%) in patients with unresectable or metastatic GIST were edema, nausea, fatigue/asthenia, cognitive impairment, vomiting, decreased appetite, diarrhea, increased lacrimation, abdominal pain, constipation, rash, dizziness, and hair color changes.

The most common adverse reactions (>20%) in patients with AdvSM were edema, diarrhea, nausea, and fatigue/asthenia.

The most common adverse reactions (>10%) in patients with ISM were eye edema, dizziness, peripheral edema, and flushing.

Drug Interactions—Avoid coadministration of AYVAKIT with strong or moderate CYP3A inhibitors. If coadministration with a moderate CYP3A inhibitor cannot be avoided in patients with GIST or AdvSM, reduce dose of AYVAKIT. Avoid coadministration of AYVAKIT with strong or moderate CYP3A inducers.

To report suspected adverse reactions, contact Blueprint Medicines Corporation at 1-888-258-7768 or FDA at 1-800-FDA-1088 or **www.fda.gov/medwatch**.

AYVAKIT is available in 25 mg, 50 mg, 100 mg, 200 mg and 300 mg tablets.

DOSING & ADMINISTRATION

Recommended Administration (Section 2.1)*

Administer AYVAKIT[®] orally on an empty stomach, at least 1 hour before or 2 hours after a meal. Do not make up for a missed dose within 8 hours of the next scheduled dose. Do not repeat dose if vomiting occurs after AYVAKIT but continue with the next scheduled dose.

GIST Harboring PDGFRA Exon 18 Mutations (Section 2.2)*

Select patients for treatment with AYVAKIT based on the presence of a PDGFRA exon 18 mutation. An FDA-approved test for the detection of exon 18 mutations is not currently available.

The recommended dosage of AYVAKIT is 300 mg orally once daily in patients with GIST. Continue treatment until disease progression or unacceptable toxicity.

Advanced Systemic Mastocytosis (Section 2.3)*

The recommended dosage of AYVAKIT is 200 mg orally once daily in adult patients with AdvSM. Continue treatment until disease progression or unacceptable toxicity.

Indolent Systemic Mastocytosis (Section 2.4)*

The recommended dosage of AYVAKIT is 25 mg orally once daily in patients with ISM.

Dosage Modifications for Adverse Reactions (Section 2.5)*

The recommended dosage reductions and modifications for adverse reactions are provided in the tables below.

Recommended Dosage Reductions for AYVAKIT for Adverse Reactions			
Dose Reduction Level	Dosage in patients with GIST ⁺	Dosage in patients with AdvSM [‡]	
First dose reduction	200 mg once daily	100 mg once daily	
Second dose reduction	100 mg once daily	50 mg once daily	
Third dose reduction	-	25 mg once daily	

⁺Permanently discontinue AYVAKIT in patients with GIST who are unable to tolerate a dose of 100 mg once daily.

[‡]Permanently discontinue AYVAKIT in patients with AdvSM who are unable to tolerate a dose of 25 mg once daily.

*AYVAKIT® (avapritinib) Prescribing Information. Blueprint Medicines Corporation; Cambridge, MA.

Recommended Dosage Modifications for AYVAKIT [®] for Adverse Reactions			
Adverse Reaction	Severity*	Dosage Modification	
Patients with GIST or AdvSM			
Intracranial Hemorrhage	Any grade	Permanently discontinue AYVAKIT.	
	Grade 1	Continue AYVAKIT at same dose or reduced dose or withhold until improvement to baseline or resolution. Resume at same dose or reduced dose.	
Cognitive Effects	Grade 2 or Grade 3	Withhold AYVAKIT until improvement to baseline, Grade 1, or resolution. Resume at same dose or reduced dose.	
	Grade 4	Permanently discontinue AYVAKIT.	
Other	Grade 3 or Grade 4	Withhold AYVAKIT until improvement to less than or equal to Grade 2. Resume at same dose or reduced dose, as clinically appropriate.	
Patients with AdvSM			
Thrombocytopenia	<50 × 10º/L	Interrupt AYVAKIT until platelet count is ≥50 × 10°/L, then resume at reduced dose. If platelet counts do not recover above 50 × 10°/L, consider platelet support.	

*Severity as defined by the National Cancer Institute Common Terminology Criteria for Adverse Events version 5.0.

Concomitant Use of Strong and Moderate CYP3A Inhibitors (Section 2.6)⁺

Avoid concomitant use of AYVAKIT with strong or moderate CYP3A inhibitors. If concomitant use with a moderate CYP3A inhibitor cannot be avoided, the starting dosage of AYVAKIT is as follows:

- GIST: 100 mg orally once daily
- AdvSM: 50 mg orally once daily

For ISM, avoid concomitant use of AYVAKIT with strong or moderate CYP3A inhibitors.

Dosage Modifications for Severe Hepatic Impairment (Section 2.7)*

A modified starting dosage of AYVAKIT is recommended for patients with severe hepatic impairment (Child-Pugh Class C):

- GIST: 200 mg orally once daily
- AdvSM: 100 mg orally once daily
- ISM: 25 mg orally every other day

⁺AYVAKIT[®] (avapritinib) Prescribing Information. Blueprint Medicines Corporation; Cambridge, MA.

ORDERING INFORMATION

AYVAKIT® (avapritinib) is available through a select network of specialty pharmacies and specialty distributors.

Specialty Pharmacy Provider Network

To prescribe AYVAKIT, please complete the YourBlueprint[®] Enrollment Form and fax it to 1-866-370-3082, or send your patient's prescription to one of the authorized specialty pharmacies listed below.

Biologics

Phone: 1-800-850-4306 Fax: 1-800-823-4506 ePrescribe: Biologics or NPI# 1487640314

Onco360

Phone: 1-877-662-6633 Fax: 1-877-662-6355 ePrescribe: Oncomed DBA Onco360 or NPI# 1679618151

PANTHERx Rare Pharmacy

Phone: 1-833-918-2015 Fax: 1-855-246-3986 ePrescribe: PANTHERx Rare Pharmacy or NPI #1659762524

AYVAKIT Product Information

AYVAKIT tablets are supplied in 5 dosage strengths:

Specialty Distribution Network

The following specialty distributors are authorized to drop-ship AYVAKIT to qualified accounts.

Physician Dispensing Offices

Cardinal Health

Specialty Distribution Phone: 1-855-855-0708 Email: <u>GMB-SPD-Specialty</u> @cardinalhealth.com

McKesson Specialty Health

Phone: 1-855-477-9800 Email: <u>MSH.CustomerCare-MSPL</u> @mckesson.com

Oncology Supply

Phone: 1-800-633-7555 Email: service@oncologysupply.com

Institutions/Hospitals

ASD Healthcare Phone: 1-800-746-6273 Email: <u>service@asdhealthcare.com</u>

Cardinal Health

Specialty Distribution Phone: 1-855-855-0708 Email: <u>GMB-SPD-Specialty</u> @cardinalhealth.com

McKesson Plasma and Biologics Phone: 1-877-625-2566 Email: MPBOrders@mckesson.com

Blueprint Medicines does not endorse the use of any particular specialty pharmacy or specialty distributor listed above and makes no representation or guarantee of services or coverage of any product. This list is current as of 3/2024 and may be updated from time to time.

Dosage Strength	300 mg	200 mg	100 mg	50 mg	25 mg
NDC Codes	10-digit code:	10-digit code:	10-digit code:	10-digit code:	10-digit code:
	72064-130-30	72064-120-30	72064-110-30	72064-150-30	72064-125-30
	11-digit code:	11-digit code:	11-digit code:	11-digit code:	11-digit code:
	72064-0130-30	72064-0120-30	72064-0110-30	72064-0150-30	72064-0125-30
Description	300 mg, capsule-	200 mg, capsule-	100 mg, round, white	50 mg, round, white	25 mg, round, white
	shaped, white film-	shaped, white film-	film-coated tablet,	film-coated tablet with	film-coated tablet with
	coated tablet, printed	coated tablet, printed	printed with blue ink	debossed text. One side	debossed text. One side
	with blue ink "BLU" on	with blue ink "BLU" on	"BLU" on one side and	reads "BLU" and the	reads "BLU" and the
	one side and "300" on	one side and "200" on	"100" on the other side;	other side reads "50";	other side reads "25";
	the other side; available	the other side; available	available in bottles of 30	available in bottles of 30	available in bottles of 30
	in bottles of 30 tablets.	in bottles of 30 tablets.	tablets.	tablets.	tablets.

Please note that splitting or breaking up individual pills is not advised.

The **blue** zero converts the 10-digit NDC code to the 11-digit NDC code. Some payers may require each NDC to be listed on the claim. Payer requirements regarding the use of NDCs may vary. Electronic data exchange generally requires use of the 11-digit NDC.

Storage: Store AYVAKIT at controlled room temperature 20 °C to 25 °C (68 °F to 77 °F); excursions are permitted between 15 °C and 30 °C (59 °F and 86 °F) [see USP Controlled Room Temperature].

PATIENT SUPPORT WITH YourBlueprint® NAVIGATING THE APPROVAL PROCESS DENIALS & APPEALS

Patient Support with YourBlueprint®

Enroll your patients at time of prescription to support the patient experience and access to programs



YourBlueprint is a patient support program designed with your patients in mind. YourBlueprint assists eligible patients throughout many aspects of treatment by providing a variety of support along the treatment journey.

Resources to assist your patients with financial needs



CO-PAY ASSISTANCE

Eligible patients with commercial insurance may be able to reduce their out-of-pocket costs (co-pay, coinsurance, or deductible) to as little as \$0 per fill up to an annual maximum of \$25,000. Please contact YourBlueprint to learn more about eligibility



PATIENT ASSISTANCE PROGRAM (PAP)

Eligible patients with no insurance, limited coverage, or unaffordable out-of-pocket costs may be able to receive their medication at no cost

Resources to ensure continued access

O COVERAGE INTERRUPTION

A no-cost, limited supply in the event of a patient experiencing a temporary lapse in coverage while on therapy



DOSE EXCHANGE

Allows patients whose healthcare provider (HCP) recommends a dose modification to exchange their remaining medication for the new dose at no cost

Resources to help your patients rapidly access treatment once prescribed and while coverage is being confirmed



QUICKSTART

A no-cost, limited supply in the event of an insurance coverage delay



REIMBURSEMENT SUPPORT AND RESOURCES

Benefits verification and resources related to prior authorizations, appeals, and formulary exceptions provided by Blueprint Medicines

Resource to support your patients once treatment has begun

PSYCHOSOCIAL PATIENT SUPPORT CALLS For those who opt in to the program

Learn more about accessing each of the programs in the following pages

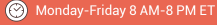
If you are interested in enrolling your patient in co-pay assistance, go to **PORTAL.TRIALCARD.COM/YOURBLUEPRINT**

Context 1-888-BLUPRNT (1-888-258-7768)



Fax: 1-866-370-3082

info@yourblueprint.com



www.YourBlueprint.com

Please see the Important Safety Information on <u>page 3-4</u> and click here to see the full <u>Prescribing Information</u> for AYVAKIT (avapritinib).

To access support services for your patient, fill out the YourBlueprint® enrollment form

- Be sure to completely fill in the <u>enrollment form</u>, including patient signatures and HCP signature. If the patient is unable to sign in person, the patient can submit their signature through DocuSign <u>here</u>
- Submitting the enrollment form to YourBlueprint at the time of prescribing will enable the YourBlueprint team to proactively support your patient's access needs



click here to see the full Prescribing Information for AYVAKIT (avapritinib).

DENIALS & APPEALS

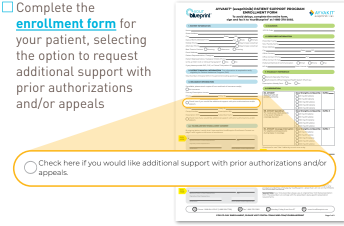
Supporting Access to Treatment



What is it?

YourBlueprint[®] will work with the patient's insurance plan to determine the path to access and communicate with you the requirements for coverage, including the correct form to submit, the supporting documentation to provide, and where to send it

What do we need from you?



Ensure patient's insurance information is completed on the form



What is it?

Should the patient's coverage determination be delayed more than 5 business days from the date your office submits the PA to the payer, YourBlueprint will provide eligible patients with up to a 60-day limited supply of nocost medication pending a final coverage determination or, if needed, a PAP eligibility determination may be made

What do we need from you?

- Complete the <u>enrollment form</u> for your patient, selecting the QuickStart prescription in section 8B of the enrollment form
- Provide YourBlueprint with the PA submission date with the enrollment form

Context Phone: 1-888-BLUPRNT (1-888-258-7768)

Fax: 1-866-370-3082

int com

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FOR CO-PAY ENROLLMENT, PLEASE VISIT PORTAL.TRIALCARD.COM/YOURBLUEPRINT

Please see the Important Safety Information on <u>page 3-4</u> and click here to see the full <u>Prescribing Information</u> for AYVAKIT (avapritinib).



Financial Needs

CO-PAY ASSISTANCE

What is it?

For eligible patients enrolled in co-pay assistance who have commercial insurance, YourBlueprint® will assist with their out-of-pocket expenses, and patients can pay as little as \$0 for their Blueprint Medicines therapy up to an annual maximum of \$25,000. Please contact YourBlueprint to learn more about eligibility

What do we need from you?

- Enroll your patient via the online portal here
- Once enrolled, adjudication information will be assigned to your patient and you can adjudicate the claim using your pharmacy system

Your Medically Integrated Dispensing (MID) pharmacy must be contracted with our co-pay processor to adjudicate claims. Contact your Blueprint Medicines Area Business Manager for more information.



PATIENT ASSISTANCE PROGRAM (PAP)

What is it?

Patients with no insurance, no coverage for AYVAKIT®, or high out-of-pocket costs, including Medicare Part D, for their Blueprint Medicines therapy may be eligible to receive their therapy at no cost through our noncommercial dispensing pharmacy

What do we need from you?

- Complete the **enrollment form** for your patient, selecting the prescription in section 3 and 8A of the enrollment form
- □ If patient has insurance but no coverage for their therapy, provide YourBlueprint the prior authorization and two (2) subsequent appeal denials with the enrollment form

Please contact YourBlueprint for current eligibility criteria.

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FOR CO-PAY ENROLLMENT, PLEASE VISIT PORTAL.TRIALCARD.COM/YOURBLUEPRINT

Please see the Important Safety Information on page 3-4 and click here to see the full Prescribing Information for AYVAKIT (avapritinib).

Continued Access to Treatment While on Therapy



COVERAGE INTERRUPTION

What is it?

Should the patient experience a temporary lapse in coverage for their therapy, YourBlueprint® will provide eligible patients with a limited supply of no-cost medication. Examples of eligible coverage lapse could be PA expiration or job transition

What do we need from you?

□ Complete the <u>enrollment form</u> for your patient, selecting the Coverage Interruption prescription in section 8C of the enrollment form

🚊 DOSE EXCHANGE

What is it?

Should the patient experience a dose modification while on AYVAKIT[®], the patient may exchange their remaining medication for the new prescribed dose at no cost to them

What do we need from you?

Complete the **Dose Exchange Form** for your patient and submit to the YourBlueprint non-commercial pharmacy for dispensing

ResultPSYCHOSOCIAL PATIENT SUPPORT CALLS

YourBlueprint enrolled patients are given the option to participate in psychosocial support phone calls. These calls are intended to support patients through conversation topics that are focused on various aspects of therapy, such as: how to prepare for upcoming doctors visits, nutrition, workplace accommodations, or talking with their family about diagnosis. The call schedule is dependent on the patient's preference.

All programs are subject to eligibility criteria. For more information, please connect with YourBlueprint for details.

C Phone: 1-888-BLUPRNT (1-888-258-7768) Fax: 1-866-370-3082

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FOR CO-PAY ENROLLMENT, PLEASE VISIT PORTAL.TRIALCARD.COM/YOURBLUEPRINT

Please see the Important Safety Information on <u>page 3-4</u> and click here to see the full <u>Prescribing Information</u> for AYVAKIT (avapritinib).

PATIENT SUPPORT

WITH YourBlueprint[®]

DOSE EXCHANGE PROGRAM

PRODUCT

INFORMATION

At times, a patient's dose may need to be adjusted during the course of treatment, and in order to facilitate this process, the YourBlueprint® Dose Exchange Program is available. If your patient

meets the requirements outlined in the Program Eligibility section below, they may qualify for this program. Please fax the completed and signed form to YourBlueprint.

Please note that the YourBlueprint Dose Exchange Program is facilitated by the YourBlueprint non-commercial pharmacy and not by the pharmacy to which the patient's previous prescription was submitted. For future refills, a new prescription will need to be submitted to the patient's current dispensing pharmacy.

In order to be eligible to participate in the Dose Exchange Program:

- Prescriber must complete the **Dose Exchange Form**
- Patient must reside in the United States or its territories
- Patient must have remaining pills from a current prescription
- Patient must return his or her remaining pills. Instructions for return will be provided with a pre-addressed envelope for the patient to return the unused quantity of previous strength
- Patient must not have already had 2 separate dose adjustments under the YourBlueprint Dose Exchange Program

blueprint	DOSE	KIT® (avapritinib) EXCHANGE FORM	avapritinib tablets
AYVAKIT Dose Exchange Progra	need to be adjusted during the co am is available. If your patient me ease fax the completed and signe	ets the requirements outlined in tl	facilitate this process, the YourBlueprint® he Program Eligibility section below, they
Please note that the YourBluep the pharmacy to which the pati patient's current dispensing ph	ient's previous prescription was su	gram is facilitated by the YourBlue Jbmitted. For future refills, a new p	print non-commercial pharmacy and not by orescription will need to be submitted to the
Patient's Current Pharmacy:	Onco360	O PANTHERx Rare Pharmacy	O Medically Integrated Dispenser (MID)
1. PROGRAM ELIGIBILITY			
In order to be eligible to participate in th	e AYVAKIT Dose Exchange Program:		
Prescriber must complete the AYVAKI	l Dose Exchange Form		
Patient must reside in the United State	es or its territories		
Patient must have remaining pills from	n a current prescription		
	ng pills. Instructions for return will be prov	ided with a pre-addressed envelope for the	patient to return the unused quantity of
previous strength Patient must not have already bad thr	ee (3) separate dose adjustments under th	e YourBlueprint AYVAKIT Dose Exchange P	rocitam -
			-9
2. PATIENT INFORMATION			
Patient Name (First, MI, Last): Patient Phone:	Patient Address:	D0	B (MM/DD/YYYY):
City:		State: ZIP	
_ny		State21P	· · · · · · · · · · · · · · · · · · ·
3. PRESCRIBER INFORMATION			
Prescriber Name (First, MI, Last):			
Practice Name:		Practice Contact:	
Practice Address:			
City:		State:ZIP	
Phone:	Fax	NP	#:
4. AYVAKIT REPLACEMENT PRESCR			
Current AYVAKIT Dose:	New AYVAKIT Dose (No Refills):		
25 mg (30 tablets)	25 mg (30 tablets)		
50 mg (30 tablets)	050 mg (30 tablets)		
000 mg (30 tablets) 200 mg (30 tablets)	0100 mg (30 tablets) 200 mg (30 tablets)		
300 mg (30 tablets)	300 mg (30 tablets)	Directions for use:	
	0		
RE Prescriber's Signature:			Date:
Special Note: If a New York prescriber, pl	ease use an original New York State prescri	iption form. The prescriber is to comply with	the prescriber's state-specific prescription requirements
5. TERMS AND CONDITIONS			
			y to exchange may not exceed 30 tablets per adjustment
	and patient will not submit a claim for reir	mbursement or otherwise seek payment fro ator for a refund or credit	orn any source for the dose exchange product, and the
	annea to aracprint medicinea or na datinas	The second set are interfaced and set in the side of	to any other patient or distributed elsewhere
The prescriber, prescriber's institution, dose exchange product will not be retr		inn. me product provided may not be giver	
The prescriber, prescriber's institution, dose exchange product will not be retr	tended only for the patient listed on this fo	I'm me product provided may not be given	
The prescriber, prescriber's institution, dose exchange product will not be reti- Product provided in this program is int agree to the terms and conditions ou	tended only for the patient listed on this fo	inn, me product provideu may not be giver	Date:
The prescriber, prescriber's institution, dose exchange product will not be retu- Product provided in this program is int agree to the terms and conditions ou NDP Prescriber's Signature	tended only for the patient listed on this fo	AND SIGNED FORM TO 1-866	
The prescriber, prescriber's institution, doze exchange product will not be ret Product provided in this program is int agree to the terms and conditions ou Prescriber's Signature	tended only for the patient listed on this form: PLEASE FAX COMPLETED A	AND SIGNED FORM TO 1-866	-370-3082
The prescriber, prescriber's institution, dose exchange product will not be retu- Product provided in this program is ini lagree to the terms and conditions ou Drescriber's Signature	tended only for the patient listed on this form: PLEASE FAX COMPLETED A	AND SIGNED FORM TO 1-866	-370-3082

🔇 Phone: 1-888-BLUPRNT (1-888-258-7768) Fax: 1-866-370-3082

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FOR CO-PAY ENROLLMENT, PLEASE VISIT PORTAL.TRIALCARD.COM/YOURBLUEPRINT

Please see the Important Safety Information on page 3-4 and click here to see the full Prescribing Information for AYVAKIT (avapritinib).



Navigating the Approval Process

Blueprint Medicines can help with questions you may have about the approval process, including prior authorization (PA) and navigating coverage denials for AYVAKIT® (avapritinib).

Contact YourBlueprint at **1-888-BLUPRNT** (1-888-258-7768) for assistance.

ENROLL YOUR PATIENTS IN YOURBLUEPRINT AT TIME OF PRESCRIPTION TO SUPPORT THE PATIENT EXPERIENCE AND ACCESS TO PROGRAMS

BENEFITS VERIFICATION

YourBlueprint and our network of specialty pharmacies can help conduct benefit verifications

YourBlueprint and our network of specialty pharmacies can conduct a benefits verification to determine a patient's health insurance coverage and out-of-pocket costs. After verifying coverage, we will provide a summary of benefits to you over the phone as well as by fax. The patient can call to review the summary of benefits verbally, and upon request, receive a copy by mail.

Prior authorization requirement? We can help

YourBlueprint and our network of specialty pharmacies can support your patient through the process of managing a prior authorization requirement. Here is what you can expect:

First, we will coordinate with your patient's insurer to gather the prior authorization requirements, including the payer-specific documents We will then contact you to help guide you through the submission process and provide you the necessary documents for you to complete, including a <u>documentation checklist</u> After your office submits the prior authorization request, upon your request, we can track the progress and communicate the status of a prior authorization to you

PRIOR AUTHORIZATION (PA) CHECKLIST

The information shown below may be required by payers to obtain a prior authorization for AYVAKIT[®] (avapritinib), however, individual payers may have their own forms or requirements.

In the case of a prior authorization denial or need for a formulary exception request, detailed information about those processes and documentation requirements are found in the Denials & Appeals section of this guide.



Ensure you complete and submit all fields in the prior authorization form, as required by the payer, which may include:

- Patient's name
- Patient's insurance company and policy number
- Patient's date of birth
- Patient's diagnosis / ICD-10 code(s)
- Derivider details, specialty, contact information, and NPI number
- AYVAKIT NDC, dosage, route of administration, and estimated duration of treatment
- Proof of appropriate diagnosis
- □ Patient's lab results (e.g., platelet counts)
- □ Clinic notes with patient subtype

Even if not part of the prior authorization form, it may also be helpful to include the following:

- □ Full Prescribing Information
- □ Information related to the treatment decision
- □ Clinical practice guidelines

If a prior authorization has been denied, see <u>page 16</u> for an appeals checklist or available for download at <u>www.YourBlueprint.com/HCP</u>

ENSURE YOUR OFFICE STAYS RESPONSIVE TO ALL YOURBLUEPRINT AND SPECIALTY PHARMACY FOLLOW UPS TO ENSURE TIMELY RESPONSES THROUGH THE PA PROCESS

Denials & Appeals

COMMON REASONS FOR COVERAGE DENIALS

Here are some common reasons for coverage denials that may be resolvable through the appeals or formulary exception request processes.



Missing Information Coverage request is missing information or there was a data error



New Drug Not yet reviewed by payer and considered non-formulary



2

3

Prior Authorization Required PA not submitted with coverage request



Insurance Information Patient's insurance changed or coverage has lapsed

If a request for coverage of AYVAKIT[®] (avapritinib) is denied, it may be resolvable through the standard appeals process, which consists of three levels.

1st Level Appeal

Contact payer to request a consideration of the denial. This may include a "peer-to-peer" discussion with the medical reviewer

2nd Level Appeal

At this step, the appeal is typically reviewed by a medical director of the plan to determine whether the request should be accepted within the coverage guidelines

Independent External Review

If attempts to appeal a coverage decision have not been successful, an external review can be conducted by an independent third party to make a binding decision

Patients may also assist with the appeals process.

If a request for coverage of AYVAKIT is denied, patients can contact their employer's benefits administrator or their health plan for additional information on how to appeal the payer's decision or to request an external review.

In some cases, it may be necessary to submit a formulary exception request to the payer. Common processes for commercial payers and Medicare Part D are described in this guide.

AYVAKIT® (avapritinib) APPEALS REQUEST CHECKLIST

If the patient's health plan has not established coverage or has denied coverage for AYVAKIT, it may be necessary to submit an appeal or a formulary exception request.

The information below includes general information, however, individual payers may have their own forms or documentation requirements.

Review the denial letter or notification received

Understand why coverage for AYVAKIT was denied and consider the following common questions:

- □ Has coverage for AYVAKIT been established for patient's condition / diagnosis?
- Did the prior authorization include all information as required by the payer or was information missing? Note: some payers may require confirmation of the diagnosis with associated documentation (e.g., provider attestation, bone marrow biopsy results)
- □ Was the insurance information correct?
- Did the patient's insurance change or coverage lapse?

Initiate the appeals process

Understand the payer's specific process or requirements:

- Use payer-specific forms, if available
- □ Follow payer's instructions on the appeals submission process and filing timelines
- □ Include all required documentation such as
 - Letter of medical necessity
 - Biomarker status
 - Treatment rationale

A sample letter is provided in this guide and available for download at www.YourBlueprint.com/HCP.

The sample letter is provided for information only and supplying the information with requests does not guarantee coverage for AYVAKIT. The information is not intended to substitute for or influence the physician's independent clinical decision.

For use when submitting a PA (see checklist on page 14)

Sample Letter of Medical Necessity	
[Physician Practice Letterhead at the top of the letter] [Date]	[P] [D]
[Name of Insurance company] [Address]	[N: [N:
[City, State Zip Code]	(Ai
	(Ci
Re: [Patient's Name]	
[Patient's Group Policy Number] [Patient's Date of Birth]	Re
	-
To whom it may concern:	De
	Ple
I am writing on behalf of my patient, [Patient's Name], to document the medical necessity for [INSERT PRODUCT] ^w (insert generic name)] and to provide information about my patient's medical history and	de
treatment to justify this therapy and subsequent payment.	
Listed below are [Patient's Name] diagnosis, medical history, treatment plan, and other supporting	l h PR
information which confirm the medical necessity and appropriateness of [INSERT PRODUCT].	ani
Patient's diagnosis, medical history, treatment plan and any other supporting information	INTO [IN
[Include information regarding your patient's diagnosis, such as:	Su
 Brief description of the patient's diagnosis, including the applicable ICD-10 code(s); 	
History with this patient; Previous therapies and results of such therapies;	
 Current treatment plan; and 	
 Other supporting information (e.g., USPI, NCCN guidelines, HCP office-selected clinical notes).] 	Tre
Enclosed in support of this matter are the following documents: [Practice to list the names of each	
de anno anti-arte de la constructione de la construction de la const	
attached]. Based on the above and attached information, I am confident that you will agree that	
attached]. Based on the above and attached information, I am confident that you will agree that [INSERT PRODUCT] is indicated and medically necessary for [Patient's Name].	
document including the package insert and provide a short description of each document being attached). Based on the above and attached information, I am confident that you will agree that [INSERT PRODUCT] is indicated and medically necessary for [Patient's Name]. Please contact me at [Insert phone number of Practice OR Physician] for any additional information that you might need to ensure prompt approval of [INSERT PRODUCT] for [Patient's Name].	cai
attached]. Based on the above and attached information, I am confident that you will agree that [INSERT PRODUCT] is indicated and medically necessary for [Patient's Name]. Please contact me at [Insert phone number of Practice OR Physician] for any additional information that you might need to ensure prompt approval of [INSERT PRODUCT] for [Patient's Name].	cai ap
attached]. Based on the above and attached information, I am confident that you will agree that [NSERT PRODUCT] is indicated and medically necessary for [Patient's Name]. Please contact me at [Insert phone number of Practice OR Physical Or any additional information that you might need to ensure prompt approval of [INSERT PRODUCT] for [Patient's Name]. Sincerely,	cai ap, Sin
attached]. Based on the above and attached information, I am confident that you will agree that [NSERT PRODUCT] is indicated and medically necessary for [Patient's Name]. Please contact me at [Insert phone number of Practice OR Physician] for any additional information that you might need to ensure prompt approval of [INSERT PRODUCT] for [Patient's Name]. Sincerely, [Prescriber's Signature]	Ple car apj Sin (Pr
attached). Based on the above and attached information, I am confident that you will agree that [INSERT PRODUCT] is indicated and medically necessary for [Patient's Name]. Please contact me at [Insert phone number of Precice OR Physician] for any additional information that	car apj Sin

Sample Formulary Exception Request

For use when requesting a coverage exception when a drug is not yet covered on formulary (see page 15)



For use when appealing a coverage denial (see **page 15**)

Sample Letter of Appeal Physician Practice Letterhead at the top of the letter] Date]

Name of Medical Director] Name of Insurance Company] Tity. State Zip Code

e: [Patient's Name, Group Policy Number, Date of Birth] – Letter of Appeal for [INSERT PRODUCTTM generic name]]

Dear [Medical Director Name],

Please consider this letter an appeal of your decision to deny coverage [Insert Denial Reason, if known] or [INSERT PRODUCT] for my patient, [Patient's Name]. I am requesting that you review my patient's lenied claim for coverage and reverse your previous decision.

have included additional information to support my decision to treat my patient with [INSERT RODUCT]. In my clinical judgement, [INSERT PRODUCT] (as you will note from the information below nd attached) is medically necessary and appropriate for [Patient's Name]. This letter includes information on [Patient's Name] medical history, prognoses and my medical rationale for selecting INSERT PRODUCT] to be used.

ummary of Medical History [Patient's Name] is a [Age, Gender]. [He/She] was diagnosed with [Insert description of disease or condition] on [Date]. [Include a biref description of patient's medical history and attach patient's chart notes]. [Include product Package Insert and note that use is within labeled indication].

reatment Rationale Given my patient's medical history, [the lack of response to other medications] and the patient's current condition and prognosis, I strongly believe that the use of [INESRT PRODUCT] for [Patient's Name] is medically necessary and appropriate and coverage should be approved. [Include any relevant clinical guidelines, such as NCCN guidelines]

lease call me or my office staff at [Physician's tele ber OR Pra nber] if I an provide you with any additional information. I look forward to receiving your timely response and pproval for treatment with [INSERT PRODUCT] for [Patient's Name].

incerely.

Prescriber's Signature]

Prescriber's Name] Attachments: Enclose supporting documentation]



Electronic versions of these sample letters are available on www.YourBlueprint.com/HCP

Diagnostic Coding

DIAGNOSIS CODES FOR THE IDENTIFICATION OF ADVANCED SM

Based on the AdvSM indications for AYVAKIT[®] (avapritinib), examples of diagnosis codes that may be appropriate are listed below. The codes provided below are information only. Coding is a clinical decision that can only be made by a provider based on the condition of the patient being treated. The use of the following codes does not suggest or guarantee reimbursement and may not be comprehensive. The codes listed are current as of February 2024 and are subject to change.

ICD-10 Diagnosis Codes

Code	Description
D47.02	Systemic mastocytosis
D47.09	Other mast cell neoplasms of uncertain behavior
C94.30	Mast cell leukemia, not having achieved remission
C94.31	Mast cell leukemia, in remission
C94.32	Mast cell leukemia, in relapse
C96.21	Aggressive systemic mastocytosis

DIAGNOSIS CODES FOR THE IDENTIFICATION OF INDOLENT SYSTEMIC MASTOCYTOSIS

Based on the ISM indications for AYVAKIT, examples of diagnosis codes that may be appropriate are listed below. The codes provided below are information only. Coding is a clinical decision that can only be made by a provider based on the condition of the patient being treated. The use of the following codes does not suggest or guarantee reimbursement and may not be comprehensive. The codes listed are current as of February 2024 and are subject to change.

ICD-10 Diagnosis Codes		
Code	Description	
D47.02	Systemic mastocytosis	
D47.09	Other mast cell neoplasms of uncertain behavior	

KIT, KIT proto-oncogene receptor tyrosine kinase; CPT, correct procedural terminology.

PDGFRA Exon 18 Mutation-Positive Unresectable or Metastatic GIST

DIAGNOSIS CODES FOR THE IDENTIFICATION OF GIST

Based on the PDGFRA Exon 18 Mutation-Positive Unresectable or Metastatic GIST indication for AYVAKIT[®] (avapritinib), examples of diagnosis codes that may be appropriate are listed below. The codes provided below are information only. Coding is a clinical decision that can only be made by a provider based on the condition of the patient being treated. The use of the following codes does not suggest or guarantee reimbursement and may not be comprehensive. The codes listed are current as of February 2024 and are subject to change.

ICD-10 Diagnosis Codes

Code	Description
C49.A0	Gastrointestinal stromal tumor, unspecified site
C49.A1	Gastrointestinal stromal tumor of esophagus
C49.A2	Gastrointestinal stromal tumor of stomach
C49.A3	Gastrointestinal stromal tumor of small intestine
C49.A4	Gastrointestinal stromal tumor of large intestine
C49.A5	Gastrointestinal stromal tumor of rectum
C49.A9	Gastrointestinal stromal tumor of other sites

PDGFRA, platelet-derived growth factor receptor alpha.

Please see the Important Safety Information on pages 3-4 and full Prescribing Information for AYVAKIT.

