

REIMAGINE

RECURRENT LGSOC

DOSING AND SIDE EFFECTS TRACKER

For patients taking
Avmapki Fakzynja Co-Pack

Use this tool to track treatment and record changes to your health during treatment, including any side effects.

Bring this tracker to your appointments to discuss with your healthcare provider.

THE FIRST FDA-APPROVED TREATMENT SPECIFICALLY for women who have low-grade serous ovarian cancer (LGSOC) that has come back (recurrent), **and** LGSOC with an abnormal *KRAS* gene, **and** previously been treated with medicine for their cancer. It is not known if Avmapki Fakzynja Co-Pack is safe and effective in children.

IMPORTANT SAFETY INFORMATION

Avmapki Fakzynja Co-Pack may cause serious side effects, including eye problems, severe skin reactions, liver problems, and muscle problems such as rhabdomyolysis. These are not all the possible side effects of Avmapki Fakzynja Co-Pack. Call your doctor for medical advice about side effects.

Please see additional Important Safety Information throughout this brochure and discuss with your healthcare provider.

Click here for full [Prescribing Information](#) and [Patient Information](#).

 **AVMAPKI® FAKZYNJA® CO-PACK**
(avutometinib capsules; defactinib tablets) 0.8 mg; 200 mg

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Our dedicated team is ready to help

For more information and helpful resources on access and support, call today: **866-351-8372**
8:00 AM to 8:00 PM ET Monday—Friday

Download or print a new copy of this calendar before the start of every new treatment cycle





Your medical history

Before taking Avmapki Fakzynja Co-Pack, tell your healthcare provider about all of your medical conditions, including if you have a history of eye or vision problems, have had severe skin reactions, or have liver problems.

Detail all of your medical conditions here:



Your pregnancy status and plans

Before taking Avmapki Fakzynja Co-Pack, tell your healthcare provider about all of your medical conditions, including if you are pregnant or plan to become pregnant. Avmapki Fakzynja Co-Pack can harm your unborn baby.

Females who are able to become pregnant:

- Your healthcare provider should do a pregnancy test before you start treatment with Avmapki Fakzynja Co-Pack.
- You should use effective birth control (contraception) during treatment and for 1 month after the last dose of Avmapki Fakzynja Co-Pack.
- Tell your healthcare provider right away if you become pregnant or think you may be pregnant during treatment with Avmapki Fakzynja Co-Pack.

Detail all of your questions and concerns about pregnancy here:

Males with a female partner who is able to become pregnant:

- You should use effective birth control (contraception) during treatment and for 4 months after the last dose of Avmapki Fakzynja Co-Pack.
- If you are breastfeeding or plan to breastfeed. It is not known if Avmapki Fakzynja Co-Pack passes into your breast milk. Do not breastfeed during treatment and for 2 weeks after the last dose of Avmapki Fakzynja Co-Pack.

IMPORTANT SAFETY INFORMATION (CONT'D)

Avmapki Fakzynja Co-Pack may cause fertility problems in females and males, which may affect your ability to have children. Talk to your healthcare provider if this is a concern for you.

Please see additional Important Safety Information throughout this brochure and discuss with your healthcare provider.



About the medications you take

Avmapki Fakzynja Co-Pack may affect the way other medicines work, and other medicines may affect how Avmapki Fakzynja Co-Pack works. It's important to tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements.

Especially tell your healthcare provider if you take:

Proton pump inhibitors (PPIs), such as omeprazole, pantoprazole, etc.

H2 receptor antagonists (H2 blockers), such as cimetidine, ranitidine, etc.

A blood thinner called warfarin

Check the applicable boxes above and list all medications you take below, including prescription and over-the-counter medicines, vitamins, and herbal supplements:

Avmapki Fakzynja Co-Pack may cause serious side effects, including eye problems, severe skin reactions, liver problems, and muscle problems (rhabdomyolysis).

Use the following sections to track and monitor all side effects you experience during treatment.



Monitoring your eye health

Eye problems are common with Avmapki Fakzynja Co-Pack and can be severe. Your healthcare provider will send you to see an eye care professional before starting treatment, during treatment, and for any new or worsening eye problems. Tell your healthcare provider if you get any new or worsening eye problems including:

- Changes in vision, such as blurred vision, double vision, or vision loss
- Dry eye(s)
- Eye and eyelid inflammation, including redness, swelling, and itching
- Eye pain
- New or increased floaters (small dark spots or squiggly lines that float across your vision)
- Light hurting eyes or seeing flashes of light

Notes:

**Watch the
eye safety video**



Monitoring your skin health

Skin reactions are common with Avmapki Fakzynja Co-Pack and can be severe or life threatening. Tell your healthcare provider if you develop any new or worsening skin reactions including:

- Skin rash or acne
- Dry skin
- Itching
- Blisters on your skin
- Redness or swelling of your face, hands, or soles of your feet
- Blisters or sores in your mouth
- Peeling of your skin
- Skin infection

Notes:

**Watch the
skin safety video**





Monitoring your liver health

Liver problems are common with Avmapki Fakzynja Co-Pack and can be severe. Treatment with Avmapki Fakzynja Co-Pack can increase the level of aspartate aminotransferase, alanine aminotransferase, and bilirubin in your blood. Your healthcare provider will do blood tests before starting each treatment cycle and during treatment to check your liver function. Tell your healthcare provider if you get any signs and symptoms of severe liver problems including:

- Yellowing of your skin or the white part of your eyes
- Itchy skin
- Feeling very tired
- Flu-like symptoms
- Nausea or vomiting
- Upper right side stomach pain
- Dark or tea colored urine
- Bleeding or bruising more easily than normal

Notes:



Monitoring your muscle health

Avmapki Fakzynja Co-Pack may cause muscle problems (rhabdomyolysis) that can be severe. Treatment with Avmapki Fakzynja Co-Pack can increase the level of an enzyme in your blood called creatine phosphokinase (CPK) and can be a sign of muscle damage. Your healthcare provider will perform blood tests to check your levels of CPK before starting each treatment cycle and during treatment. Tell your healthcare provider if you get any signs or symptoms of increased CPK including:

- Weakness or difficulty moving arms and legs
- Muscle or bone aches or pain that does not go away
- Dark, red colored urine
- Decreased urine output

Notes:



Monitoring your overall health

It's important that you tell your healthcare provider about any problems or discomfort that you may be experiencing during treatment.

 Indicates side effect determined by labwork

The most common side effects of Avmapki Fakzynja Co-Pack include:

- Nausea
- Tiredness
- Increased aspartate aminotransferase 
- Diarrhea
- Swelling in the body (edema)
- Decreased hemoglobin 
- Increased alanine aminotransferase 
- Vomiting
- Increased blood bilirubin 
- Increased triglycerides 
- Stomach-area pain
- Acid reflux
- Increased alkaline phosphatase 
- Decreased platelets 
- Constipation
- Shortness of breath
- Cough
- Urinary tract infection
- Decreased lymphocytes 
- Decreased neutrophils 
- Proteinuria 

Notes:

Tell your healthcare provider if you are experiencing gastrointestinal side effects such as nausea, diarrhea, vomiting, stomach-area pain, acid reflux, or constipation, and detail your symptoms here:

If you are experiencing swelling in the body, also called edema, please detail your symptoms here:

IMPORTANT SAFETY INFORMATION (CONT'D)

Tell your healthcare provider if you have any side effect that bothers you or that does not go away.

These are not all of the possible side effects of Avmapki Fakzynja Co-Pack. Call your doctor for medical advice about side effects.

Please see additional Important Safety Information throughout this brochure and discuss with your healthcare provider.

How should I take Avmapki Fakzynja Co-Pack?

Avmapki:

- On **Weeks 1, 2, and 3**, take 3.2 mg (4 capsules) on **Day 1** and **Day 4**
- On **Week 4**, take a one-week break
- Take each dose at the same time **with food**
- Swallow capsules whole. Do not chew, break or open the capsules
- **Keep refrigerated** in original bottle

3.2 mg (4 capsules)



1 DOSE

Fakzynja:

- On **Weeks 1, 2, and 3**, take 200 mg (1 tablet) **two times a day**
- On **Week 4**, take a one-week break
- Take **with food**
- Swallow tablets whole. Do not chew, break or crush the tablets
- **Keep refrigerated** in original bottle

200 mg (1 tablet)



1 DOSE

Product images shown are not actual size.

Use the table below for monthly treatment guidance.

Avmapki | Fakzynja

	Time	Day 1	Day 2	Day 3	Day 4	Day 5	Day 6	Day 7
Weeks 1, 2, & 3	Morning Same Time				+			
	Evening							
Week 4	ONE-WEEK BREAK. Do not take Avmapki Fakzynja Co-Pack during Week 4.							

TAKE Avmapki Fakzynja Co-Pack exactly as directed by your healthcare provider.

DO NOT change your dose or stop taking Avmapki Fakzynja Co-Pack without talking to your healthcare provider.

IF YOU VOMIT after taking Avmapki capsules or Fakzynja tablets, do not take an additional dose. Take the next scheduled dose as prescribed by your healthcare provider.

IF YOU MISS A DOSE of Avmapki by more than 24 hours or if you miss a dose of Fakzynja by more than 6 hours, skip the missed dose and take the next scheduled dose as prescribed by your healthcare provider. Do not take two doses at the same time to make up for a missed dose.

DISCARD any remaining medication at the end of the 3-week treatment period as instructed by your healthcare provider.

IF YOU DEVELOP CERTAIN SIDE EFFECTS, your healthcare provider may change your dose, temporarily stop, or completely stop your treatment with Avmapki Fakzynja Co-Pack.

IF YOU TAKE ANTACIDS, take Fakzynja tablets 2 hours before or 2 hours after taking the antacid.

Dosing and side effects tracker

Use the calendar below to track your treatment and note when side effects occur.

How to use the calendar

- A Note the cycle number and date when your new treatment begins.
- B Write in the date you take your medication.
- C Check the box each time you take your medication.
- D Record any side effect you experience on each day.

A Cycle # 3 Start date 11 / 1 / 26

B DATE 11 / 1 / 26

C

DAY 01
Avmapki AM ☒
Fakzynja AM ☒ PM ☒

D Side effects: Blurred vision

After noting the side effect on the calendar here, please put the details of each side effect in its respective section of the tracker on the prior pages.

Cycle # _____ Start date _____ / _____ / _____

WEEK 1

DAY 01 DATE / / Avmapki AM ☐ Fakzynja AM ☐ PM ☐	DAY 02 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 03 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 04 DATE / / Avmapki AM ☐ Fakzynja AM ☐ PM ☐	DAY 05 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 06 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 07 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐
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Side effects: _____

WEEK 2

DAY 08 DATE / / Avmapki AM ☐ Fakzynja AM ☐ PM ☐	DAY 09 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 10 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 11 DATE / / Avmapki AM ☐ Fakzynja AM ☐ PM ☐	DAY 12 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 13 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 14 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐
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Side effects: _____

WEEK 3

DAY 15 DATE / / Avmapki AM ☐ Fakzynja AM ☐ PM ☐	DAY 16 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 17 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 18 DATE / / Avmapki AM ☐ Fakzynja AM ☐ PM ☐	DAY 19 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 20 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐	DAY 21 DATE / / NO Avmapki Fakzynja AM ☐ PM ☐
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Side effects: _____

WEEK 4

DAY 22 DATE / / NO Avmapki and NO Fakzynja	DAY 23 DATE / / NO Avmapki and NO Fakzynja	DAY 24 DATE / / NO Avmapki and NO Fakzynja	DAY 25 DATE / / NO Avmapki and NO Fakzynja	DAY 26 DATE / / NO Avmapki and NO Fakzynja	DAY 27 DATE / / NO Avmapki and NO Fakzynja	DAY 28 DATE / / NO Avmapki and NO Fakzynja
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Side effects: _____

Please see additional Important Safety Information throughout this brochure and discuss with your healthcare provider.

Care team information

Fill out this page with the names and contact information of your care team, which may include your medical oncologist, nurse, and any other healthcare professional assisting you during your treatment.

PRIMARY CARE

Name: _____

Email: _____

Phone: _____

MEDICAL ONCOLOGIST

Name: _____

Email: _____

Phone: _____

DERMATOLOGIST

Name: _____

Email: _____

Phone: _____

OPHTHALMOLOGIST

Name: _____

Email: _____

Phone: _____

GYNECOLOGIST

Name: _____

Email: _____

Phone: _____

NURSE

Name: _____

Email: _____

Phone: _____



Please see additional Important Safety Information throughout this brochure and discuss with your healthcare provider.

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Please see additional Important Safety Information throughout this brochure and discuss with your healthcare provider.
Click here for full [Prescribing Information](#) and [Patient Information](#).