

My Milestones"

A free digital companion on the Medisafe® app to support you through your DARZALEX FASPRO® treatment journey

Medisafe is a registered trademark of Medisafe Project, Ltd.



What is DARZALEX FASPRO® (daratumumab and hyaluronidase-fihj)?

DARZALEX FASPRO® is a prescription medicine used to treat adult patients with multiple myeloma:

- in combination with the medicines bortezomib, melphalan, and prednisone in people with newly diagnosed multiple myeloma who cannot receive a type of stem cell transplant that uses their own stem cells (autologous stem cell transplant)
- in combination with the medicines lenalidomide and dexamethasone in people with newly diagnosed multiple myeloma who cannot receive a type of stem cell transplant that uses their own stem cells (autologous stem cell transplant) and in people whose multiple myeloma has come back or did not respond to treatment who have received at least one prior medicine to treat multiple myeloma
- in combination with the medicines bortezomib, thalidomide, and dexamethasone in newly diagnosed people who are eligible to receive a type of stem cell transplant that uses their own stem cells (autologous stem cell transplant)
- in combination with the medicines bortezomib and dexamethasone in people who have received at least one prior medicine to treat multiple myeloma
- in combination with the medicines pomalidomide and dexamethasone in people who have received at least one prior medicine, including lenalidomide and a proteasome inhibitor, to treat multiple myeloma
- in combination with the medicines carfilzomib and dexamethasone in people whose multiple myeloma has come back or did not respond to treatment who have received one to three prior medicines to treat multiple myeloma
- alone in people who have received at least three prior medicines, including a proteasome inhibitor and an immunomodulatory agent, **or** did not respond to a proteasome inhibitor and an immunomodulatory agent It is not known if DARZALEX FASPRO® is safe and effective in children.

SELECT IMPORTANT SAFETY INFORMATION

Do not receive DARZALEX FASPRO® if you have a history of a severe allergic reaction to daratumumab, hyaluronidase, or any of the ingredients in DARZALEX FASPRO®. See page 7 for a complete list of ingredients in DARZALEX FASPRO®.

Please see additional Important Safety Information on pages 5-7. Please see the Product Information in back pocket.

DARZALEX FASPRO® MyMilestones™ resources tailored to your needs

With this program, you can connect with helpful tools and trackers, treatment updates, educational videos, and other resources relevant to your multiple myeloma treatment journey.



Gives you access to support and resources throughout the course of your treatment journey



Celebrates your treatment milestones



Helps you have more meaninaful conversations with your healthcare team about your treatment and how it's affecting you



Provides details so you know what to expect before, during, and after receiving treatment



Delivers reminders for treatments and other medical appointments



Offers access to cost support information to help you pay for DARZALEX FASPRO®

SELECT IMPORTANT SAFETY INFORMATION

- Serious allergic reactions and other severe injection-related reactions. Serious allergic reactions and reactions due to release of certain substances by your body (systemic) that can lead to death can happen with DARZALEX FASPRO®. Your healthcare provider may temporarily stop or completely stop treatment with DARZALEX FASPRO® if you have a serious reaction. Tell your healthcare provider or get medical help right away if you get any of these symptoms during or after an injection of DARZALEX FASPRO®.
- shortness of breath or trouble breathing
- dizziness or lightheadedness (hypotension)
- o cough
- wheezing
- heart beating faster than

- (hypoxia)
- runny or stuffy nose
- headache
- o itching
- o low oxygen in the blood
- throat tightness or irritation
- high blood pressure

Please see additional Important Safety Information on pages 5-7. Please see the Product Information in back pocket.

- eye pain
- o nausea
- vomiting
- o chills
- o fever
- o chest pain
- blurred vision

GET ACCESS FOR FREE!

How to scan the QR code using your phone to access the MyMilestones™ program

- Open the camera app on your phone
- Point the camera at the QR code until a pop-up banner appears on your screen
- Tap on the pop-up to open the Apple App Store or Google Play Store
- Follow the prompts on your phone to download and install the Medisafe app
- Once downloaded, open the **Medisafe App** on your phone



Milestones.

- Tap "Get started"
- Tap "Continue" on the MyMilestones™ screen and follow the prompts for adding DARZALEX FASPRO® reminders to the Medisafe app

Please see Important Safety Information on pages 5-7. Please see the Product Information in back pocket.





Patient Support From Janssen Compass™

Personalized Support. From People Who Care.

Janssen Compass™ is a free, personalized patient support program that provides one-on-one guidance, information, and educational resources to you about your disease. It also may help you understand your insurance coverage and cost support options, as well as tips to help you get started and stay on track with your treatment.

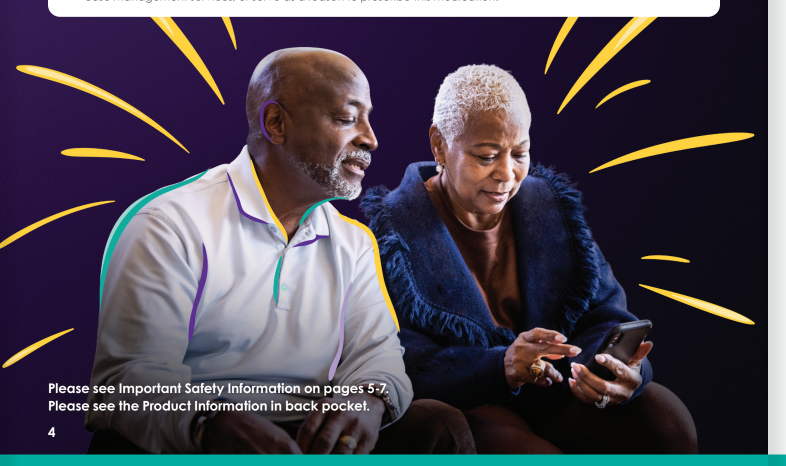


Connect with us!

Your dedicated Janssen Compass[™] Care Navigator is a phone call away! Scan the QR code with your phone or visit www.darzalex.com/faspro/janssen-compass to request your first call and learn more about how Janssen Compass[™] can be here for you.

You can also call us at 844-NAV-1234 (844-628-1234), Monday through Friday, 8:30 AM — 8:30 PM ET

Janssen Compass™ is limited to education for patients about their Janssen therapy, its administration, and/or their disease. It is intended to supplement a patient's understanding of their therapy and is not intended to provide medical advice, replace a treatment plan from the patient's doctor or nurse, provide case management services, or serve as a reason to prescribe this medication.



Important Safety Information for DARZALEX FASPRO®

Do not receive DARZALEX FASPRO® if you have a history of a severe allergic reaction to daratumumab, hyaluronidase, or any of the ingredients in DARZALEX FASPRO®. See below for a complete list of ingredients in DARZALEX FASPRO®.

Before you receive DARZALEX FASPRO®, tell your healthcare provider about all of your medical conditions, including if you:

- have a history of breathing problems
- have had shingles (herpes zoster)
- have ever had or might now have a hepatitis B infection as DARZALEX FASPRO® could cause hepatitis B virus to become active again. Your healthcare provider will check you for signs of this infection before, during, and for some time after treatment with DARZALEX FASPRO®. Tell your healthcare provider right away if you get worsening tiredness or yellowing of your skin or white part of your eyes.
- are pregnant or plan to become pregnant. DARZALEX FASPRO® may harm your unborn baby. Tell your healthcare provider right away if you become pregnant or think that you may be pregnant during treatment with DARZALEX FASPRO®.
- ° Females who are able to become pregnant should use an effective method of birth control (contraception) during treatment and for 3 months after your last dose of DARZALEX FASPRO®. Talk to your healthcare provider about birth control methods that you can use during this time.
- ° Before starting DARZALEX FASPRO® in combination with lenalidomide, thalidomide, or pomalidomide, females and males must agree to the instructions in the lenalidomide, thalidomide, or pomalidomide REMS program.
- The lenalidomide, thalidomide, and pomalidomide REMS have more information about effective methods of birth control, pregnancy testing, and blood donation for females who can become pregnant.
- For males who have female partners who can become pregnant, there is information in the lenalidomide, thalidomide, and pomalidomide REMS about sperm donation and how lenalidomide, thalidomide, and pomalidomide can pass into human semen.
- are breastfeeding or plan to breastfeed. It is not known if DARZALEX FASPRO® passes into your breast milk. You should not breastfeed during treatment with DARZALEX FASPRO®. Talk to your healthcare provider about the best way to feed your baby during treatment with DARZALEX FASPRO®.

Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins, and herbal supplements.

Continued on next page

Important Safety Information for DARZALEX FASPRO® (cont)

How will I receive DARZALEX FASPRO®?

- DARZALEX FASPRO® may be given alone or together with other medicines used to treat multiple myeloma.
- DARZALEX FASPRO® will be given to you by your healthcare provider as an injection under the skin in the stomach area (abdomen).
- DARZALEX FASPRO® is injected over 3 to 5 minutes.
- Your healthcare provider will decide the time between doses as well as how many treatments you will receive.
- Your healthcare provider will give you medicines before each dose of DARZALEX FASPRO® and after each dose of DARZALEX FASPRO® to help reduce the risk of serious allergic reactions and other reactions due to release of certain substances by your body (systemic).

If you miss any appointments, call your healthcare provider as soon as possible to reschedule your appointment.

DARZALEX FASPRO® may cause serious reactions, including:

- Serious allergic reactions and other severe injection-related reactions. Serious allergic reactions and reactions due to release of certain substances by your body (systemic) that can lead to death can happen with DARZALEX FASPRO®. Your healthcare provider may temporarily stop or completely stop treatment with DARZALEX FASPRO® if you have a serious reaction. Tell your healthcare provider or get medical help right away if you get any of these symptoms during or after an injection of DARZALEX FASPRO®.
- shortness of breath low oxygen in the blood o eye pain or trouble breathing (hypoxia) o nausea throat tightness or irritation dizziness or lightheadedness vomiting (hypotension) runny or stuffy nose o chills o cough • headache o fever wheezing o itching o chest pain heart beating faster than high blood pressure blurred vision usual
- **Injection site reactions.** Skin reactions at or near the injection site (local), including injection site reactions, can happen with DARZALEX FASPRO®. Symptoms at the site of injection may include itching, swelling, bruising, pain, rash, bleeding, or redness of the skin. These reactions sometimes happen more than 24 hours after an injection of DARZALEX FASPRO®.



- **Decreases in blood cell counts.** DARZALEX *FASPRO*® can decrease white blood cell counts, which help fight infections, and blood cells called platelets, which help to clot blood. Decreases in blood cell counts are common with DARZALEX *FASPRO*® but can be severe. Your healthcare provider will check your blood cell counts during treatment with DARZALEX *FASPRO*®. Tell your healthcare provider if you develop fever or have signs of bruising or bleeding.
- Changes in blood tests. DARZALEX FASPRO® can affect the results of blood tests to match your blood type. These changes can last for up to 6 months after your final dose of DARZALEX FASPRO®. Your healthcare provider will do blood tests to match your blood type before you start treatment with DARZALEX FASPRO®. Tell all of your healthcare providers that you are being treated with DARZALEX FASPRO® before receiving blood transfusions.

The most common side effects of DARZALEX FASPRO® when used alone include cold-like symptoms (upper respiratory infection) and decreased red blood cell counts.

The most common side effects of DARZALEX FASPRO® used in combination therapy include:

п	$r \cap \cap$	ness	
н	$\Gamma \subset G$	コロニシフ	

nausea

diarrhea

shortness of breath

trouble sleeping

headache

• fever

• cough

muscle spasms

• back pain

vomiting

high blood pressure

 cold-like symptoms (upper respiratory infection) nerve damage causing tingling, numbness, or pain

constipation

• lung infection (pneumonia)

• swollen hands, ankles, or feet

decreased red blood cell counts

These are not all the possible side effects of DARZALEX FASPRO®. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

General information about the safe and effective use of DARZALEX FASPRO®

Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. You can ask your pharmacist or healthcare provider for information about DARZALEX FASPRO® that is written for health professionals.

Active ingredient: daratumumab and hyaluronidase-fihi

Inactive ingredients: L-histidine, L-histidine hydrochloride monohydrate, L-methionine, polysorbate 20, sorbitol, water for injection

Please see the accompanying Product Information.

cp-143282v9

6



My Milestones™

Find multiple myeloma treatment support in the palm of your hand



Treatment reminders



Educational resources



Access to cost support information



Treatment milestone celebrations

Scan the QR code with your phone to install the FREE Medisafe app!



Add DARZALEX FASPRO® to your list of medications in the Medisafe app to gain access to the MyMilestones™ program.
See page 3 for detailed instructions.





Please see Important Safety Information on pages 5-7. Please see the Product Information in back pocket.

Apple® and App Store® are registered trademarks of Apple Inc. Google Play and the Google Play logo are trademarks of Google LLC. © Janssen Biotech, Inc. 2023 04/23 cp-320986v2

