PATIENT INFORMATION

EGRIFTA SV[®] (eh-GRIF-tuh ESS-vee) (tesamorelin) for injection

for subcutaneous use

2 mg vial

Read the Patient Information that comes with EGRIFTA SV before you start to take EGRIFTA SV and each time you get a refill. There may be new information. This leaflet does not take the place of talking to your healthcare provider about your medical condition or your treatment.

What is EGRIFTA SV?

• EGRIFTA SV is an injectable prescription medicine used to reduce the excess stomach-area (abdominal) fat in HIVinfected adult patients with lipodystrophy. EGRIFTA SV is a growth hormone-releasing factor (GHRF).

The long-term safety of EGRIFTA SV on the heart and blood vessels (cardiovascular) is not known.

EGRIFTA SV is not for weight loss management.

It is not known whether taking EGRIFTA SV helps improve how well you take (compliance with) antiretroviral medicines.

It is not known if EGRIFTA SV is safe and effective in children.

EGRIFTA SV is not recommended to be used in children with open or closed bone growth plates (epiphyses).

Who should not use EGRIFTA SV?

Do not use EGRIFTA SV if you:

- have a pituitary gland tumor, have had pituitary gland surgery, have other problems related to your pituitary gland, or have had radiation treatment to your head or a head injury.
- have active cancer. Any previous cancer should be inactive, and any previous cancer treatment should be complete before starting EGRIFTA SV.
- are allergic to tesamorelin or any of the ingredients in EGRIFTA SV. See the end of this leaflet for a complete list of ingredients in EGRIFTA SV.
- are pregnant or plan to become pregnant. EGRIFTA SV can harm your unborn baby. If you become pregnant, stop using EGRIFTA SV and talk with your healthcare provider.

What should I tell my healthcare provider before using EGRIFTA SV?

- Before using EGRIFTA SV, tell your healthcare provider about all of your medical conditions, including if you:
 have or have had cancer.
- have problems with your blood sugar or diabetes. Some people with diabetes who use EGRIFTA SV may develop or may have worsening eye problems.
- have scheduled heart or stomach surgery.
- have breathing problems.
- are breastfeeding or plan to breastfeed. It is not known if EGRIFTA SV passes into your breast milk. The Centers for
 Disease Control and Prevention (CDC) recommends that HIV-infected mothers **not** breastfeed to avoid the risk of passing
 HIV infection to your baby. Talk with your healthcare provider about the best way to feed your baby if you are using
 EGRIFTA SV.

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

How should I use EGRIFTA SV?

- Read the detailed "Instructions for Use" that comes with EGRIFTA SV before you start using it. Your healthcare provider will show you how to inject EGRIFTA SV.
- Use EGRIFTA SV exactly as your healthcare provider tells you to use it.
- Inject EGRIFTA SV under the skin (subcutaneously) of your stomach-area (abdomen).
- Change (rotate) the injection site on your stomach-area with each dose. Do not inject EGRIFTA SV into scar tissue, bruises or your belly button.
- Do not share your EGRIFTA SV syringe or needles with other people, even if the needle has been changed. You may give other people a serious infection or get a serious infection from them.

What are the possible side effects of EGRIFTA SV?

EGRIFTA SV may cause serious side effects, including:

- increase risk of new cancer in HIV positive patients or your cancer coming back (reactivation). Stop using EGRIFTA SV if any cancer symptoms come back.
- increased levels of your insulin-like growth factor-1 (IGF-1). Your healthcare provider will do blood tests to check your IGF-1 levels while you are taking EGRIFTA SV.
- **swelling (fluid retention).** EGRIFTA SV can cause swelling in some parts of your body. Call your healthcare provider if you have swelling, an increase in joint pain or pain or numbness in your hands or wrist (carpal tunnel syndrome). Joint pain and swelling of your arms, hands, legs and feet are common side effects of EGRIFTA SV, but may sometimes be serious.
- **increase in blood sugar (glucose) or diabetes.** Your healthcare provider will check your blood sugar before you start taking EGRIFTA SV and during your treatment with EGRIFTA SV.
- serious allergic reaction. Some people using EGRIFTA SV may have an allergic reaction. Stop using EGRIFTA SV and get emergency medical help right away if you have any of the following symptoms:

	0	a rash over your body	0	hives	0	swelling of your face or throat	
	0	shortness of breath or	0	fast heartbeat	0	feeling of faintness or fainting	
		trouble breathing	0	itching	0	reddening or flushing of the skin	
•	 injection site reactions. Injection site reactions are a common side effect of EGRIFTA SV but may sometimes be serious. Change (rotate) your injection site to help lower your risk for injection site reactions. Call your healthcare provider for medical advice if you have any of the following symptoms around the area of the injection site: redness itching pain 						
	0	irritation	0	bruising or bleeding	0	rash	
	0	swelling					
•	increased risk of death in people who have critical illnesses because of heart or stomach surgery, trauma or serious breathing (respiratory) problems has happened when taking certain amounts of growth hormone.						
The most common side effects of EGRIFTA SV include:							
	•	pain in legs and arms		• mus	cle pain		
These are not all the possible side effects of EGRIFTA SV. For more information, ask your healthcare provider or pharmacist.							
Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.							
You may also report side effects to THERA patient support toll-free at 1-833-23THERA (1-833-238-4372).							
How should I store EGRIFTA SV 2 mg vials, Sterile Water for Injection, syringes and needles?							
•	 You will be given two boxes from the pharmacy when you get your prescription of EGRIFTA SV: Store the 2 mg EGRIFTA SV vials in the Medication Box they come in, at room temperature at 20°C to 25°C (68°F to 77°F). 						
	 Store the Sterile Water for Injection, syringes and needles that come in the Injection Box at room temperature at 20°C to 25°C (68°F to 77°F). 						
•							
•							
•	Throw away any Sterile Water for Injection left in the bottle after use.						
Do not use EGRIFTA SV after the expiration date (EXP) printed on the carton and vial labels.							
Keep EGRIFTA SV and all medicines out of the reach of children.							
General information about the safe and effective use of EGRIFTA SV. Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. Do not use EGRIFTA SV for a condition for which it was not prescribed. Do not give EGRIFTA SV to other people, even if they have the same symptoms you have. It may harm them.							
	You can ask your healthcare provider or pharmacist for information about EGRIFTA SV that is written for health professionals.						
What are the ingredients in EGRIFTA SV? Active ingredient: tesamorelin (as an acetate salt) Inactive ingredients: histidine, hydrochloric acid, mannitol, polysorbate 20, sucrose EGRIFTA SV does not contain any preservative.							
Man	ufacture	ed by Theratechnologies Inc., 2015 Peel Street, S	Suite 110	00, Montréal, Québec, Canada H3A	1T8 US Licen	se No. 2091 for Theratechnologies Inc.	
		formation about EGRIFTA SV, go to www.EGRIF			A (1-833-238	-4372). Revised: 02/2024	