

March 23, 2022

Dear Valued Patient,

You're receiving this update because you have previously been prescribed NATPARA® (parathyroid hormone) for Injection or you are part of our NATPARA® Special Use Program (SUP). Takeda is providing a U.S. regulatory update for NATPARA® (parathyroid hormone) to inform that the Company has received a Complete Response Letter (CRL) from the U.S. Food and Drug Administration (FDA) in response to the Prior Approval Supplement (PAS) submitted in August 2021 to address the potential for rubber particulate formation that led to the U.S. NATPARA recall in September 2019.

The CRL indicates that the FDA has completed its review of the NATPARA PAS and determined that it cannot be approved in its present form. Takeda is evaluating the details of the CRL to determine next steps. In the meantime, we are deeply disappointed to inform the hypoparathyroidism community that NATPARA's commercial return in the U.S. is indefinitely delayed.

Importantly, because there are no U.S. FDA-approved treatment alternatives for chronic hypoparathyroidism patients, Takeda intends to provide patients who are enrolled in the NATPARA SUP with continued access to therapy free of charge, in accordance with regulatory oversight and under the discretion of the FDA, until a commercial product is available.

With the goal of limiting supply interruption for SUP patients, we continue to work on the separate supply challenges surrounding protein particle formation that we have described over the past year. Those challenges are unrelated to the PAS and the recall. It is important to underscore that all product released for patient use continues to meet Takeda's quality standards and the safety profile of NATPARA has not changed. Our Takeda OnePath® Patient Support Managers will continue to work with SUP patients to ensure they have access to their prescribed NATPARA treatment.

We understand and have tremendous empathy for how much the community has been impacted without NATPARA and are disheartened to share this update. Takeda leaders will host a video teleconference with the US hypoparathyroidism community to provide more context regarding this communication in the coming weeks. Please watch for more information about that upcoming event in the NATPARA Updates section of our Takeda US Newsroom at https://www.takeda.com/en-us/newsroom/natpara-updates/. Information should be posted before the end of March. We will also share the information with the HypoPARAthyroidism Association and HypoPara Support & Advocacy.

Cheryl Schwartz

Senior Vice President and Business Unit Head

US Rare Disease Business Unit

Chenge Schwaf

Richard C. Ascroft, RPh, JD

Fichal C. AscrifA

Senior Vice President
Patient and Market Access
US Business Unit

What is NATPARA® (parathyroid hormone) for Injection?

- NATPARA is a prescription parathyroid hormone used with calcium and vitamin D to control low blood calcium (hypocalcemia) in people with low parathyroid hormone blood levels (hypoparathyroidism).
- NATPARA is only for people who do not respond well to treatment with calcium and active forms of vitamin D alone, because it may increase the possible risk of bone cancer (osteosarcoma).
- NATPARA was not studied in people with hypoparathyroidism caused by calcium-sensing receptor mutations.
- NATPARA was not studied in people who get sudden hypoparathyroidism after surgery.
- It is not known if NATPARA is safe and effective for children 18 years of age and younger. NATPARA should not be used in children and young adults whose bones are still growing.

Important Safety Information

What is the most important information I should know about NATPARA? Warning: Possible bone cancer (osteosarcoma).

- During animal drug testing, NATPARA caused some rats to develop a bone cancer called
 osteosarcoma. It is not known if people who take NATPARA will have a higher chance of getting
 osteosarcoma. Tell your doctor right away if you have pain in any areas of your body that does
 not go away, or any new or unusual lumps or swelling under your skin that is tender to touch.
- NATPARA is only available through the NATPARA Risk Evaluation and Mitigation Strategy
 (REMS) Program. The purpose of the NATPARA REMS program is to inform patients about the
 potential risk of osteosarcoma associated with the use of NATPARA. For more information about
 this REMS program, call 1-855-NATPARA (628-7272) or go to www.NATPARAREMS.com.

NATPARA may cause other serious side effects, including:

High blood calcium (hypercalcemia)

NATPARA can cause some people to have a higher blood calcium level than normal.

- Your doctor should check your blood calcium before you start and during your treatment with NATPARA.
- Tell your doctor if you have nausea, vomiting, constipation, low energy, or muscle weakness. These may be signs that you have too much calcium in your blood.

Low blood calcium (hypocalcemia)

- People who stop using or miss a dose of NATPARA may have an increased risk of severe low blood calcium levels.
- Tell your doctor if you have tingling of your lips, tongue, fingers and feet, twitching of face muscles, cramping of feet and hands, seizures, depression, or have problems thinking or remembering.

Who should not use NATPARA?

• **Do not use NATPARA** if you are allergic to parathyroid hormone or any of the ingredients in NATPARA.

What should I tell my healthcare provider before using NATPARA?

Before you start using NATPARA, tell your doctor about all of your medical conditions. Tell
your doctor about all the medicines you take, including prescription and over-the-counter
medicines, vitamins, and herbal supplements.

What are the possible side effects of NATPARA?

- NATPARA may cause serious side effects like allergic (hypersensitivity) reaction, including anaphylaxis. Tell your healthcare provider or get emergency medical help right away if you have any of the following symptoms of an allergic reaction:
- swelling of your face, lips, mouth, or tongue
- itching

breathing problems

- rash
- fainting, dizziness, feeling lightheaded (low blood pressure)
- hives

- fast heartbeat
- The most common side effects of NATPARA include: tingling, tickling, or burning feeling of the skin, low or high blood calcium, headache, nausea, reduced sense of touch or sensation, diarrhea, vomiting, pain in joints, too much calcium in urine, and pain in limbs.

These are not all the possible side effects of NATPARA. For more information, talk with your doctor. You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch or call 1-800-FDA-1088.

Enclosure: NATPARA Full Prescribing Information

©2022 Takeda Pharmaceuticals U.S.A., Inc. 300 Shire Way, Lexington, MA 02421. 1-877-TAKEDA-7. All rights reserved. TAKEDA and the TAKEDA logo are trademarks or registered trademarks of Takeda Pharmaceutical Company Limited. NATPARA® and the NATPARA logo are registered trademarks of Shire-NPS Pharmaceuticals, Inc., a Takeda company. OnePath® is a registered trademark of Shire Human Genetic Therapies, Inc.