PEDIATRIC USE ONLY

45 mg for 6-month administration





Do not remove from clamshell until ready to use. Recommended dosage: After mixing, a healthcare provider should immediately administer entire contents of syringe by intramuscular injection every 6 months. See prescribing information. **Only Activate Safety Device** 00300743575019

Post-Injection. Store at 20°C to 25°C (68°F to 77°F);

excursions 15-30°C (59-86°F)

Manufactured for: AbbVie Inc.

- North Chicago, IL 60064 by: Takeda Pharmaceutical
- Company Limited Osaka, Japan 540-8645

Product of Japan. Rx only



NDC 0074-3575-01 Single Dose Administration Kit with prefilled dual-chamber syringe.



Dispense the accompanying Medication Guide to each patient.

45 mg for 6-month administration

FOR INTRAMUSCULAR INJECTION

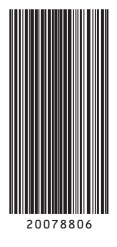
The front chamber contains: leuprolide acetate 45 mg • polylactic acid 169.9 mg • D-mannitol 39.7 mg • stearic acid 10.1 mg The second chamber contains: carboxymethylcellulose sodium 7.5 mg • D-mannitol 75.0 mg • polysorbate 80 1.5 mg • water for injection, USP, and glacial acetic acid. USP to control pH



PEDIATRIC USE ONLY

45 mg for 6-month administration

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