My ChoiceTM

Drug Facts

Active ingredient

Levonorgestrel, USP 1.5 mg

Purpose

Emergency contraceptive

Use

for women to reduce chance of pregnancy af ter unprotected sex (if a contraceptive failed or if you did not use bir th control)

Warnings

Allergy alert

Do not use if you have ever had an allergic reaction to levonorgestrel

Sexually transmitted diseases (STDs) alert

This product does **not** protect against HIV/AIDS or other STDs.

Do not use

- · if you are already pregnant (because it will not work)
- for regular bir th control

Ask a doctor or pharmacist before use if you are taking efavirenz (HIV medication) or rifampin (tuberculosis treatment) or medication for seizures (epilepsy). These medications may reduce the effectiveness of levonorgestrel.

When using this product you may have

- menstrual changes
- tiredness
- dizziness
- nausea
- · headache
- · breast pain
- · lower stomach (abdominal) pain
- vomiting

Keep out of reach of children. In case of overdose, get medical help or contact a Poison Control Center right away (1-800-222-1222).

Directions

- take as soon as possible within 72 hours (3 days) after unprotected sex. The sooner you take it, the better it will work.
- · if you vomit within 2 hours after taking the medication, call a healthcare professional to find out if you should repeat the dose

Other information

- · read the instructions, warnings, and enclosed product leaflet before use
- this product works mainly by preventing ovulation (egg release). It may also prevent fer tilization of a released egg (joining of sperm and egg) or at tachment of a fer tilized egg to the uterus (implantation).
- · do not use if carton is open or tear strip is removed or blister seal is broken or missing
- store at 20° to 25°C (68° to 77°F)

Inactive ingredients

colloidal silicon dioxide, corn starch, hypromellose, lactose monohydrate, magnesium stearate, talc

Questions or comments?

For more information, call toll free 1-800-818-4555 weekdays

Distributed by:

Ohm Laboratories Inc.

New Brunswick, NJ 08901

PRINCIPAL DISPLAY PANEL - 1.5 mg Tablet Blister Pack Carton

NDC 62756-720-60

[†]Compare To the active ingredient of Plan B One-Step®

See New Warning

My Choice™

Levonorgestrel Tablet 1.5 mg

Emergency Contraceptive

- Reduces the chance of pregnancy after unprotected sex
- Not for regular birth control
- · The sooner you take it, the more effective it will be
- Take as soon as possible within 72 hours (3 days) after unprotected sex
- · Will not harm an existing pregnancy

1 Tablet Levonorgestrel 1.5 mg

One Tablet. One Step.



MY CHOICE TM

levonorgestrel tablet

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Sou	irce)	NDC:627	756-720				
Route of Administration	ORAL								
Active Ingredient/Active	ve Moiety								
	Ingredient Name Bas				gth Strength				
LEVONORGESTREL (UNII: 5W7SIA7YZW) (LEVONORGESTREL - UNII:5W7SIA7YZW)			LEVO	1.5 mg					
Inactive Ingredients									
	Ingredient Name				Strength				
SILICON DIOXIDE (UNII: ET	J7Z6XBU4)								
STARCH, CORN (UNII: O8232NY3SJ)									
HYPROMELLOSE, UNSPECIFIED (UNII: 3NXW29V3WO)									
LACTOSE MONOHYDRATE (UNII: EWQ57Q815X)									
MAGNESIUM STEARATE (UNII: 70097M6I30)									
TALC (UNII: 7SEV7J4R1U)									
Product Characteristic	s								
Color	WHITE (white to off-white)	Score	Score		no score				
Shape	ROUND (circular)	Size	Size		8mm				
Flavor		Impri	Imprint Code		718				
Contains									
Packaging									
# Item Code	Package Description		Marketing Start Da	ite I	Marketing End Date				
1 NDC:62756-720-60	1 in 1 CARTON	04/01	/2018						
1	1 in 1 BLISTER PACK; Type 0: Not a Combination Product								
Marketing Information									
Marketing Category	Application Number or Monograph Citation		Marketing Start Date	. 1	Marketing End Date				
ANIDA	A NID A 202025	0.4/01/20	10						

Labeler - SUN PHARMACEUTICAL INDUSTRIES, INC. (146974886)

ANDA202635

ANDA

Establishment								
Name	Address	ID/FEI	Business Operations					
Sun Pharmaceutical Industries Limited		725959238	ANALYSIS(62756-720), MANUFACTURE(62756-720)					

04/01/2018

Revised: 4/2022 SUN PHARMACEUTICAL INDUSTRIES, INC.