

FEM CHOICE MORNING AFTER - levonorgestrel tablet
Aurohealth LLC

Fem Choice™ morning after
(levonorgestrel tablet 1.5 mg)

Drug Facts

Active ingredient (in each tablet)

Levonorgestrel USP 1.5 mg

Purpose

Emergency contraceptive

Use

for women to reduce chance of pregnancy after unprotected sex (if a contraceptive failed or if you did not use birth 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 birth 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
- nausea
- lower stomach (abdominal) pain
- tiredness

- headache
- dizziness
- breast pain
- vomiting

Keep out of reach of children.

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

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**
- **do not use if carton is open or tear strip is removed or the blister seal is broken or missing**
- store at 20° to 25°C (68° to 77°F)

Inactive ingredients

colloidal silicon dioxide, corn starch, lactose monohydrate, magnesium stearate, potato starch and talc.

Questions?

Call **1-855-274-4122**

Distributed by:

AUROHEALTH LLC

279 Princeton-Hightstown Road
East Windsor, NJ 08520

Made in India

Code: TS/DRUGS/22/2009

Patient Information

Fem Choice™ morning after (levonorgestrel tablet 1.5 mg)

Emergency Contraceptive

One Tablet. One Step.

What You Need to Know

What is Fem Choice™ morning after?

Fem Choice™ morning after is emergency contraception that helps prevent pregnancy after birth control failure or unprotected sex. It is a **backup** method of preventing pregnancy and should not be used as regular birth control.

What Fem Choice™ morning after is not.

Fem Choice™ morning after will not work if you are already pregnant and will not affect an existing pregnancy. Fem Choice™ morning after will not protect you from HIV infection (the virus that causes AIDS) and other sexually transmitted diseases (STDs).

When should I use Fem Choice™ morning after?

The sooner you take emergency contraception, the better it works. You should use Fem Choice™ morning after within 72 hours (3 days) **after you have had unprotected sex**.

Fem Choice™ morning after is a backup or emergency method of birth control you can use when:

- your regular birth control was used incorrectly or failed
- you did not use any birth control method

When not to use Fem Choice™ morning after.

Fem Choice™ morning after should not be used:

- as a regular birth control method, because it's not as effective as regular birth control.
- if you are already pregnant, because it will not work.
- if you are allergic to levonorgestrel or any other ingredients in Fem Choice™ morning after.

When should I talk to a doctor or pharmacist?

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 Fem Choice™ morning after and increase your chance of becoming pregnant. Your doctor may prescribe another form of

emergency contraception that may not be affected by these medications.

How does Fem Choice™ morning after work?

Fem Choice™ morning after works before release of an egg from the ovary. As a result, Fem Choice™ morning after usually stops or delays the release of an egg from the ovary. Fem Choice™ morning after is one tablet that contains a higher dose of levonorgestrel than birth control pills and works in a similar way to prevent pregnancy.

How can I get the best results from Fem Choice™ morning after?

You have 72 hours (3 days) to try to prevent pregnancy after birth control failure or unprotected sex. **The sooner you take Fem Choice™ morning after, the better it works.**

How effective is Fem Choice™ morning after?

If Fem Choice™ morning after is taken as directed, it can significantly decrease the chance that you will get pregnant. About 7 out of every 8 women who would have gotten pregnant will not become pregnant.

How will I know Fem Choice™ morning after worked?

You will know Fem Choice™ morning after has been effective when you get your next period, which should come at the expected time, or within a week of the expected time. If your period is delayed beyond 1 week, it is possible you may be pregnant. You should get a pregnancy test and follow up with your healthcare professional.

Will I experience any side effects?

- some women may have changes in their period, such as a period that is heavier or lighter or a period that is early or late. **If your period is more than a week late, you may be pregnant.**
- if you have severe abdominal pain, you may have an ectopic pregnancy, and should get immediate medical attention.
- when used as directed, Fem Choice™ morning after is safe and effective. Side effects may include changes in your period, nausea, lower stomach (abdominal) pain, tiredness, headache, dizziness, and breast tenderness.
- if you vomit within 2 hours of taking the medication, call a healthcare professional to find out if you should repeat the dose.

What if I still have questions about Fem Choice™ morning after?

If you have questions or need more information, call our toll-free number AUROHEALTH LLC at 1-855-274-4122.

Other Information

Keep out of reach of children:

In case of overdose, get medical help or contact a Poison Control Center right away at 1-800-222-1222.

Do not use if the blister seal is opened.

Store at room temperature 20° to 25°C (68° to 77°F).

Active ingredient: levonorgestrel 1.5 mg

Inactive ingredients: colloidal silicon dioxide, corn starch, lactose monohydrate, magnesium stearate, potato starch, and talc.

If you are sexually active, you should see a healthcare provider for routine checkups. Your healthcare provider will talk to you about and, if necessary, test you for sexually transmitted diseases, teach you about effective methods of routine birth control, and answer any other questions you may have.

Distributed by:

AUROHEALTH LLC

279 Princeton-Hightstown Road
East Windsor, NJ 08520

Manufactured by:

Aurobindo Pharma Limited

Hyderabad-500 032, India

Revised: 02/2023

PACKAGE LABEL-PRINCIPAL DISPLAY PANEL -1.5 mg (1 Tablet Carton Label)

†Compare to the active ingredient
of Plan B One-Step®

Fem

Choice™

morning after

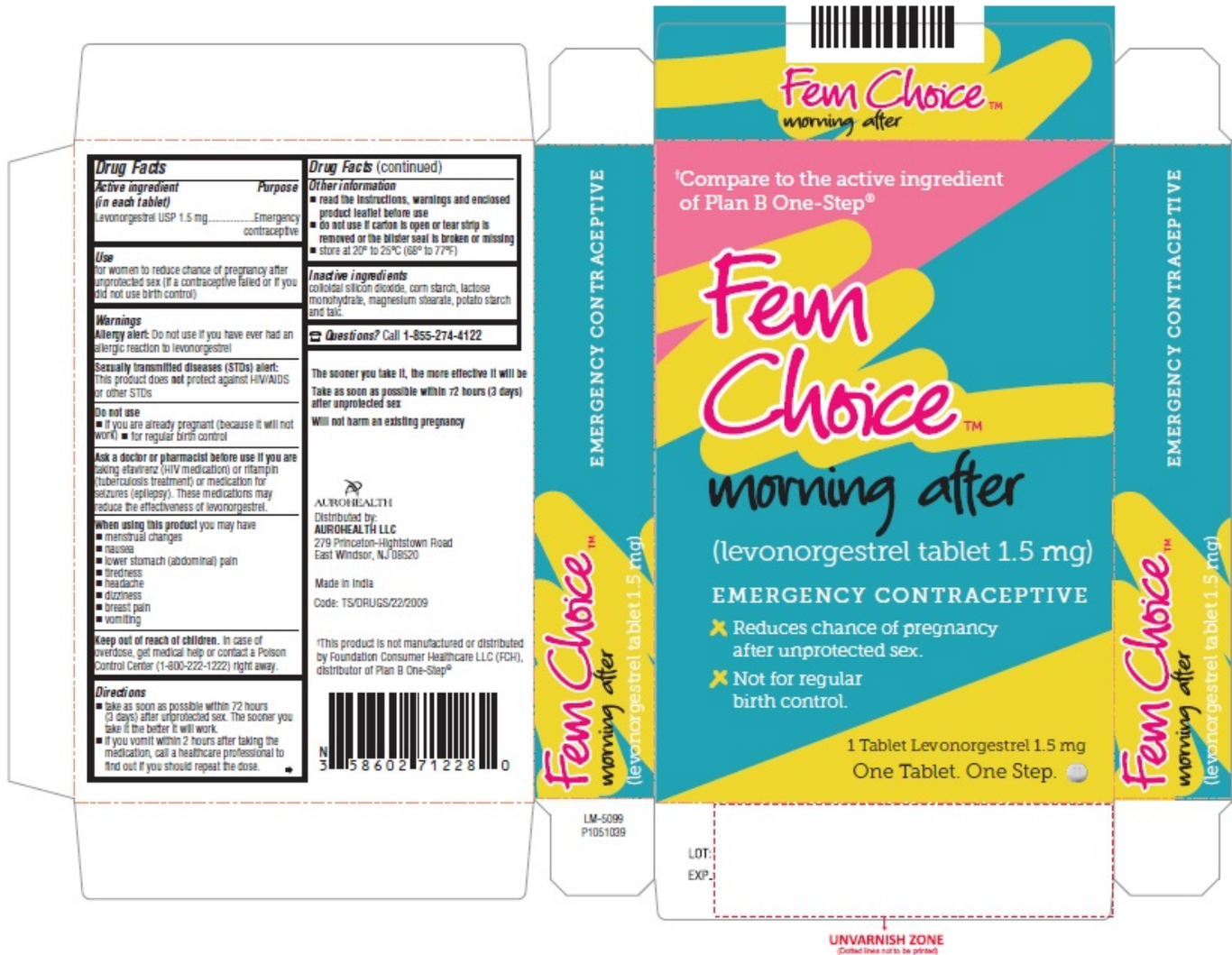
(levonorgestrel tablet 1.5 mg)

EMERGENCY CONTRACEPTIVE

x Reduces chance of pregnancy
after unprotected sex.

x Not for regular
birth control.

1 Tablet Levonorgestrel 1.5 mg
One Tablet. One Step.



FEM CHOICE MORNING AFTER

levonorgestrel tablet

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:58602-712
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
LEVONORGESTREL (UNII: 5W7SIA7YZW) (LEVONORGESTREL - UNII:5W7SIA7YZW)	LEVONORGESTREL	1.5 mg

Inactive Ingredients

Ingredient Name	Strength
-----------------	----------

SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
STARCH, CORN (UNII: O8232NY3SJ)	
LACTOSE MONOHYDRATE (UNII: EWQ57Q8I5X)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
STARCH, POTATO (UNII: 8I089SAH3T)	
TALC (UNII: 7SEV7J4R1U)	

Product Characteristics

Color	WHITE (White to off-white)	Score	no score
Shape	ROUND	Size	8mm
Flavor		Imprint Code	S;11
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:58602-712-28	1 in 1 CARTON	01/26/2022	
1		1 in 1 BLISTER PACK; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA206867	01/26/2022	

Labeler - Aurohealth LLC (078728447)

Establishment

Name	Address	ID/FEI	Business Operations
Aurobindo Pharma Limited		650381903	ANALYSIS(58602-712) , MANUFACTURE(58602-712)

Revised: 2/2023

Aurohealth LLC