# IBUPROFEN- ibuprofen tablet, film coated Pharbest Pharmaceuticals, Inc.

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## **Drug Facts**

#### Active ingredient (in each tablet)

Ibuprofen 200 mg (NSAID)\* \*nonsteroidal anti- inflammatory drug

## Purpose

Pain Reliever/ Fever Reducer

## Uses

- temporarily relieves minor aches and pain due to:
- headache
- toothache
- backache
- menstrual cramps
- the common cold
- muscular aches
- minor pain of arthritis
- temporarily reduces fever

## Warnings

**Allergy alert:** Ibuprofen may cause a severe allergic reaction, especially in people allergic to Aspirin. Symptoms may include

- hives
- facial swelling
- asthma (wheezing)
- shock
- skin reddening
- rash
- blisters

If an allergic reaction occurs, stop use and seek medical help right away.

**Stomach bleeding warning:** This product contains an NSAID, which may cause severe stomach bleeding. The chance is higher if you

- are age 60 or older
- have had stomach ulcers or bleeding problems
- take a blood-thinning (anticoagulant) or steroid drug
- take other drugs containing prescription or nonprescription NSAIDs (aspirin, ibuprofen, naproxen, or others)
- have 3 or more alcoholic drinks every day while using this product
- take more or for a longer time than directed

Do not use

- if you have ever had an allergic reaction to any other pain reliever/ fever reducer
- right before or after heart surgery

## Ask a doctor before use if

- stomach bleeding warning applies to you
- you have problems or serious side effects from taking pain relievers or fever reducers
- you have a history of stomach problems, such as heartburn
- you have high blood pressure, heart disease, liver cirrhosis, kidney disease, or asthma
- you are taking a diuretic

## Ask a doctor or pharmacist before use if you are

- under a doctor's care for any serious condition
- taking aspirin for heart attack or stroke, because ibuprofen may decrease this benefit of aspirin
- taking any other drug

## when using this product

- take with food or milk if stomach upset occurs
- the risk of heart attack or stroke may increase if you use more than directed or for longer than directed

## stop use and ask a doctor if

you experience any of the following signs of stomach bleeding:

- feel faint
- vomit blood
- have bloody or black stools
- have stomach pain that does not get better
- pain gets worse or lasts more than 10 days
- fever gets worse or lasts more than 3 days
- redness or swelling is present in the painful area
- any new symptoms appear

## If pregnant or breast-feeding,

ask a health professional before use. It is especially important not to use ibuprofen during the last 3 months of pregnancy unless definitely directed to do so by a doctor because it may cause problems in the unborn child or complications during delivery.

## 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

- Do not take more than directed
- The smallest effective dose should be used

adults and children 12 years of age Take 1 tablet every 4 to 6 hours while symptoms persist. If pain or fever does not respond to 1 tablet, 2 tablets may be used. Do not

| and over                   | exceed σ tablets 111 24 hours unless directed by a doctor |  |
|----------------------------|---|--|
| children under<br>12 years | consult a doctor  |  |

#### Other information

- Tamper Evident: do not use if safety seal under cap is broken or missing
- store at room temperature (20°- 25°C)
- avoid excessive heat above 40°C (104°F)

#### **Inactive ingredients**

carnuba wax, colloidal silicon dioxide, corn starch, hypromellose, lactose anhydrous, magnesium stearate, microcrystalline cellulose, polydextrose, polyethylene glycol, red iron oxide, stearic acid, sodium starch glycolate, titanium dioxide

#### **Questions or comments?**

(866) 562-2756 Mon-Fri: 8 AM to 4 PM

PHARBEST

NDC 16103-407-11

\*Compare to the active ingredient in ADVIL®

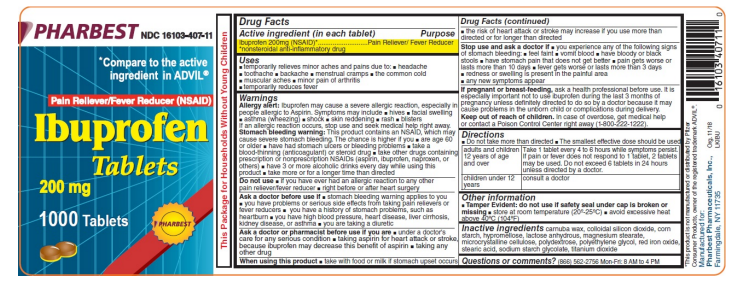
## Pain Reliever/Fever Reducer (NSAID)

Ibuprofen

Tablets

200 mg

1000 Tablets



## **IBUPROFEN**

ibuprofen tablet, film coated

#### **Product Information**

Product Type

HUMAN OTC DRUG

Item Code (Source)

NDC:16103-407

| Route of Administ   |   |  |  |   |                   |                      |                               |                  |                      |
|---|---|--|--|---|-------------------|----------------------|-------------------------------|------------------|----------------------|
|   | ration  | ORAL   |  |   |                   |                      |                               |                  |                      |
|   |   |  |  |   |                   |                      |                               |                  |                      |
| Active Ingredie   | nt/Active   | Moiety   |  |   |                   |                      |                               |                  |                      |
| Ingredient Name   |   |  |  |   |                   |                      | Basis of                      | Strength         | Strength             |
| BUPROFEN (UNII: \   | WK2XYI10QN  | M) (IBUPROFEI  | N - UNII:WK2X                                    | (YI10QM)  |                   |                      | IBUPRO FEN                    |                  | 200 mg               |
| ,   |   |  |  | . ,   |                   |                      |                               |                  |                      |
| nactive Ingred  | ients   |  |  |   |                   |                      |                               |                  |                      |
| Ingredient Name   |   |  |  |   |                   |                      |                               |                  | Strength             |
| CARNAUBA WAX (UNII: R12CBM0EIZ)   |   |  |  |   |                   |                      |                               |                  |                      |
| ILICON DIO XIDE   |   |  |  |   |                   |                      |                               |                  |                      |
| TARCH, CORN (UI   |   |  |  |   |                   |                      |                               |                  |                      |
| IYPRO MELLOSE,  | UNSPECIFIE  | E <b>D</b> (UNII: 3NXV   | W29V3WO)   |   |                   |                      |                               |                  |                      |
| NHYDRO US LACT  | <b>OSE</b> (UNII: 1   | 3SY5LH9PMK)  |  |   |                   |                      |                               |                  |                      |
| AGNESIUM STEA   | RATE (UNII:   | 70097M6I30)  |  |   |                   |                      |                               |                  |                      |
| CELLULOSE, MICR   | RO CRYSTAI  | LLINE (UNII: O   | P1R32D61U)                                       |   |                   |                      |                               |                  |                      |
| OLYDEXTROSE (   | UNII: VH2XO   | U12IE)   |  |   |                   |                      |                               |                  |                      |
| OLYETHYLENE G   | LYCOL, UN   | SPECIFIED (U   | JNII: 3WJQ0SI                                    | DW1A)   |                   |                      |                               |                  |                      |
| FERRIC OXIDE RED  | <b>)</b> (UNII: 1K09  | F3G675)  |  |   |                   |                      |                               |                  |                      |
| TEARIC ACID (UN   | II: 4ELV7Z65  | GAP)   |  |   |                   |                      |                               |                  |                      |
| SO DIUM STARCH O  | GLYCOLAT  | Ε ΤΥΡΕ Α ΡΟ  | TATO (UNII: 5                                    | 856J3G2   | 42)               |                      |                               |                  |                      |
| TTTANIUM DIO XID  | E (UNII: 15F12  | X9V2JP)  |  |   |                   |                      |                               |                  |                      |
|   |   | X9 V2JP)   |  |   |                   |                      |                               |                  |                      |
| Product Charac  | teristics   | X9 V2JP)<br>pro wn   | Sc   | ore   |                   |                      |                               | no score         |                      |
| Product Charac  | t <b>eristics</b>   | Dro wn   |  | ore   |                   |                      |                               | no score<br>10mm |                      |
| Product Charac<br>Color<br>Shape  | t <b>eristics</b>   |  | Siz  | ze  | de                |                      |                               | 10 mm            |                      |
| Product Charac<br>Color<br>Shape<br>Flavor  | t <b>eristics</b>   | Dro wn   | Siz  |   | de                |                      |                               |                  |                      |
| Product Charac<br>Color<br>Shape<br>Flavor  | t <b>eristics</b>   | Dro wn   | Siz  | ze  | de                |                      |                               | 10 mm            |                      |
| FITANIUM DIO XIDI<br>Product Charac<br>Color<br>Shape<br>Flavor<br>Contains<br>Packaging  | t <b>eristics</b>   | Dro wn   | Siz  | ze  | de                |                      |                               | 10 mm            |                      |
| Product Charac<br>Color<br>Shape<br>Plavor<br>Contains<br>Packaging   | t <b>eristics</b>   | prown<br>ROUND   | Siz  | ze<br>print Coo                                 | de                | Ma                   | rketing St<br>Date            | 10 mm<br>44;29 1 | arketing End<br>Date |
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10/01/2018

Labeler - Pharbest Pharmaceuticals, Inc. (557054835)

**Registrant** - Pharbest Pharmaceuticals, Inc. (557054835)

| Establishment                  |         |           |                                       |  |  |  |  |  |
|--------------------------------|---------|-----------|---------------------------------------|--|--|--|--|--|
| Name                           | Address | ID/FEI    | Business Operations                   |  |  |  |  |  |
| Pharbest Pharmaceuticals, Inc. |         | 557054835 | relabel(16103-407), repack(16103-407) |  |  |  |  |  |

Revised: 10/2018

Pharbest Pharmaceuticals, Inc.