

CENTER FOR DRUG EVALUATION AND RESEARCH

Approval Package for:

APPLICATION NUMBER:
ANDA 75-010 / S-003

Name: Ibuprofen Tablets USP, 200 mg (Brown)

Sponsor: LNK International, Inc.

Approval Date: August 15, 2002

CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER:
ANDA 75-010 / S-003

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APPLICATION NUMBER:
ANDA 75-010 / S-003

APPROVAL LETTER

cc:ANDA

Division File

Field Copy

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APPROVAL LETTER – MULTIPLE SUPPLEMENTS

Endorsements:

HFD-613/JBarlow

 7/26/22

**APPEARS THIS WAY
ON ORIGINAL**

CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER:
ANDA 75-010 / S-003

LABELING

FOLD

AUG 15 2002

Drug Facts

Active ingredient (in each brown tablet) **Purpose**
Pain Reliever/
Ibuprofen USP, 200 mg. Fever Reducer

Uses for the temporary relief of minor aches and pains associated with ■ the common cold ■ headache ■ toothache ■ muscular aches ■ backache ■ arthritis ■ menstrual cramps ■ For the reduction of fever

Ibuprofen Tablets USP, 200mg
CAPLETS
Pain Reliever/Fever Reducer

Drug Facts (continued)

Warnings Do not combine this product with any other (ibuprofen containing) product.

Do not use ■ if you have had a severe allergic reaction to aspirin, e.g., asthma, swelling, shock or hives, because even though this product contains no aspirin or salicylates, cross-reactions may occur in persons allergic to aspirin ■ for pain for more than 10 days ■ for fever for more than 3 days ■ with aspirin or acetaminophen

▼ LIFT HERE ▼

outside

FOLD

Drug Facts (continued)

Ask a doctor before use if ■ you are being treated for a serious condition ■ you have any condition that requires you to take prescription drugs ■ you have had any problems or serious side effects from taking any non-prescription pain reliever

Stop use and ask a doctor if ■ you experience any symptoms which are unusual or seem unrelated to the condition for which you took ibuprofen ■ pain or fever persists or gets worse ■ new symptoms occur ■ the painful area is red or swollen ■ more than mild heartburn, upset stomach or stomach pain occurs with use

If pregnant or breast-feeding, ask a health professional for advice before use. ESPECIALLY IMPORTANT: DO NOT USE IBUPROFEN DURING THE LAST 3 MONTHS OF PREGNANCY. IT MAY SPECIFICALLY DIRECTLY HARM THE FETUS. CAUSE PROBLEMS WITH THE BIRTH OF A CHILD OR COMPLICATE THE BIRTH DELIVERY.
Keep out of the reach of children. In case of overdose, get immediate medical attention or contact a Poison Control Center right away.

Alcohol Warning: If you drink alcohol or more alcoholic drinks than usual, ask your doctor whether you should take ibuprofen or other pain relievers/fever reducers. Ibuprofen may cause stomach bleeding.

inside: left

FOLD

Drug Facts (continued)

Directions. The smallest effective dose should be used. Take with food or milk if occasional or mild heartburn, upset stomach, or stomach pain occurs with use.

■ **adults:** 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 but do not take more than 6 tablets in 24 hours unless directed by a doctor.
■ **children:** do not give this product to children under 12 except under the advice and supervision of a doctor

Other information ■ **TAMPER EVIDENT: DO NOT USE IF IMPRINTED SAFETY SEAL UNDER CAP IS BROKEN OR MISSING.** ■ store at controlled room temperature ■ avoid excessive heat 40°C (104°F)

Inactive ingredients calcium waxes, cellulose, corn starch, fumaric silica gel, hydroxypropyl methylcellulose, lactose, magnesium stearate, polyacrylates, polyethylene glycol, red iron oxide, sodium starch glycolate, stearic acid, titanium dioxide

Made in USA
Distributed By
Reg. # 388000

AUG 15 2002

inside: right

N9146A

Ibuprofen Tablets USP, 200mg
CAPLETS
Pain Reliever/Fever Reducer
12 FILM COATED BROWN
CAPLETS*, 200 mg EACH
*Caplets (Capsule-Shaped Tablets)

Ibuprofen Tablets USP, 200mg
Pain Reliever/Fever Reducer
CAPLETS
12 FILM COATED BROWN
CAPLETS*, 200 mg EACH
*Caplets (Capsule-Shaped Tablets)
Compare to the active
ingredient in Adult Caplets.

Drug Facts (continued)
Directions The smallest effective dose should be used. Take with food or milk, if occasional or mild heartburn, upset stomach, or stomach pain occurs with use.
■ **adults:** 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 but do not take more than 6 tablets in 24 hours unless directed by a doctor.
■ **children:** do not give this product to children under 12 except under the advice and supervision of a doctor.
Other information ■ **TAMPER EVIDENT:** DO NOT USE IF IMPRINTED SHEET SEAL UNDER CAP IS BROKEN OR MISSING ■ Store at controlled room temperature ■ avoid excessive heat 40°C (104°F) ■ see end flap for expiration date and lot no.
Inactive ingredients carmelumina wax, cellulose, corn starch, tinned silica gel, hydroxypropyl methylcellulose, lactose, magnesium stearate, polyethylene glycol, polyethylene glycol, red iron oxide, sodium saccharin, glycolic stearic acid, titanium dioxide

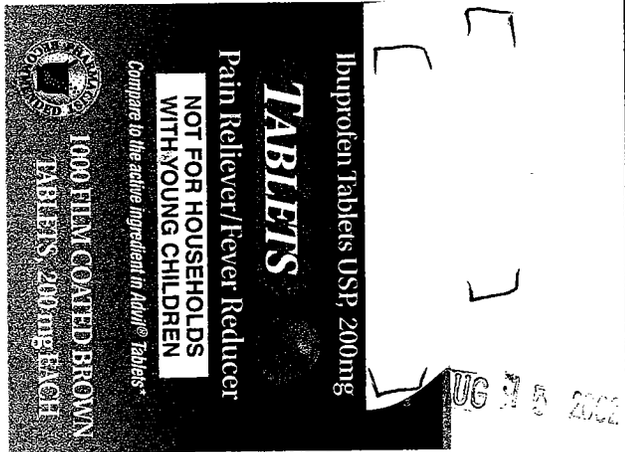
ITEM XXXXXX
UPC-FPO

Drug Facts
Active ingredient (in each brown tablet) Ibuprofen USP 200 mg
Purpose Pain Reliever/Fever Reducer

Uses for the temporary relief of minor aches and pains associated with:
■ the common cold ■ headache ■ toothache ■ muscular aches ■ backache ■ arthritis ■ menstrual cramps ■ for the reduction of fever
Warnings Do not combine this product with any other ibuprofen containing product.
Do not use ■ If you have had a severe allergic reaction to aspirin, e.g., asthma, swelling, hives, or rash, because even though this product contains no aspirin or salicylates, cross-reactions may occur in patients allergic to aspirin ■ for pain for more than 10 days ■ for fever for more than 3 days ■ with aspirin or acetaminophen
Ask a doctor before use if ■ you are being treated for a serious condition ■ you have any condition that requires you to take prescription drugs ■ you have had any

Drug Facts (continued)
problems or ~~fever~~ ~~fever~~ from taking any prescription or over-the-counter drugs.
Stop use and ask your doctor if you experience any symptoms that are unusual or seem unrelated to the condition for which you took ibuprofen ■ If you experience a painful area in red or swollen, or if you get worse ■ new symptoms such as dizziness, mild heartburn, upset stomach, or stomach pain occurs with use
If pregnant or breast-feeding, ask your health professional before use. It is especially important not to use ibuprofen during the last 3 months of pregnancy. Do not use unless specifically directed to do so by a doctor because it may cause complications during delivery.
Keep out of the reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.
Alcohol Warning: If you consume 3 or more alcoholic drinks every day, ask your doctor whether you should take ibuprofen or other pain relievers/fever reducers. Ibuprofen may cause stomach bleeding.
*This product is not manufactured or distributed by Wyeth-Robbins, owners of the registered trademark Advil® Caplets.

Drug Facts
Active ingredient (in each brown tablet) Ibuprofen USP 200 mg
Purpose Pain Reliever/Fever Reducer



APPROVED

Drug Facts

Active ingredient (in each brown tablet)	Purpose
Ibuprofen USP, 200 mg	Pain Reliever/Fever Reducer

Uses for the temporary relief of minor aches and pains associated with ■ the common cold ■ headache ■ toothache ■ muscular aches ■ backache ■ arthritis ■ menstrual cramps ■ For the reduction of fever

Warnings Do not combine this product with any other Ibuprofen containing product.

Do not use ■ if you have had a severe allergic reaction to aspirin, e.g., asthma, swelling, shock or hives, because even though this product contains no aspirin or salicylates, cross-reactions may occur in patients allergic to aspirin ■ for pain for more than 10 days ■ for fever for more than 3 days ■ with aspirin or acetaminophen

Ask a doctor before use if ■ you are being treated for a serious condition ■ you have any condition that requires you to take prescription drugs ■ you have had any problems or serious side effects from taking any non-prescription pain reliever

Stop use and ask a doctor if ■ you experience any symptoms which are unusual or seem unrelated to the condition for which you took Ibuprofen ■ pain or fever persists or gets worse ■ new symptoms occur ■ the painful area is red or swollen ■ more than mild heartburn, upset stomach or stomach pain occurs with use

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 SPECIFICALLY DIRECTED TO DO SO BY A DOCTOR BECAUSE IT MAY CAUSE PROBLEMS IN THE UNBORN CHILD OR COMPLICATIONS DURING DELIVERY.

Keep out of the reach of children. In case of overdose, get medical help or contact a Poison Control Center right away.

Alcohol Warning: If you consume 3 or more alcoholic drinks every day, ask your doctor whether you should take Ibuprofen or other pain relievers/fever reducers. Ibuprofen may cause stomach bleeding.

Directions The smallest effective dose should be used. Take with food or milk if occasional or mild heartburn, upset stomach, or stomach pain occurs with use.

- **adults:** 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 but do not take more than 6 tablets in 24 hours unless directed by a doctor
- **children:** do not give this product to children under 12 except under the advice and supervision of a doctor

Other information ■ TAMPER EVIDENT: DO NOT USE IF IMPRINTED SAFETY SEAL UNDER CAP IS BROKEN OR MISSING ■ store at controlled room temperature ■ avoid excessive heat 40°C (104°F)

Inactive ingredients carnauba wax, cellulose, corn starch, fumed silica gel, hydroxypropyl methylcellulose, lactose, magnesium stearate, polydextrose, polyethylene glycol, red iron oxide, sodium starch glycolate, stearic acid, titanium dioxide

*This product is not manufactured or distributed by Whitehall-Robins, owners of the registered trademark Advil® Tablets.
 Made in USA

Distributed by

100% Satisfaction Guaranteed

Rev. XXXXX
 ITEM XXXXX

UPC-FPO

CENTER FOR DRUG EVALUATION AND RESEARCH

APPLICATION NUMBER:
ANDA 75-010 / S-003

CORRESPONDENCE

LNK INTERNATIONAL, INC.

Over-The-Counter Pharmaceutical Manufacturer

60 Arkay Drive, Hauppauge, LI, NY 11788

ANDA 75-010 Ibuprofen USP 200mg Film Coated Brown Tablets
ANDA 75-139 Ibuprofen USP 200mg Film Coated White Tablets
Labeling Revision

February 11, 2002

Mr. J. Barlow
Food and Drug Administration
Office of Generic Drugs
CDER
8700 Standish Place
Metro Park North II
Rockville, MD 20855

NDA NO. 75-010 REF. NO. SL-603/AI
NDA SUPPL FOR Labeling Rev.

Supplement: Changes Being Effected
Labeling Change to Comply with OTC Requirements

ANDA 75-139; Ibuprofen USP, 200mg White Film Coated Tablets (RLD = Motrin)
ANDA ~~75-010~~ Ibuprofen USP, 200mg Brown Film Coated Tablets (RLD = Advil)

Dear Mr. Barlow:

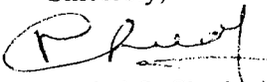
This is a Special Supplement - Changes Being Effected. The effective date is May 16, 2002 based on the **OVER-THE-COUNTER HUMAN DRUGS; LABELING REQUIREMENTS; PARTIAL EXTENSION OF COMPLIANCE DATES; FR FDA 06/20/2000 F 65 FR 38191**. Our understanding is that this date is a shipment date and has no affect on product previously shipped with the old label and labeling. LNK is responding to a fax request dated January 24, 2002. The fax had provided a copy of approved label and labeling for the Reference Listed Drug (RLD), Motrin IB NDA 19-012/S-024. We were instructed to make necessary revisions and submit as a CBE labeling supplement.

LNK had submitted a supplement for a second color (**Orange**) in our approved Abbreviated New Drug Application, ANDA 75-139, **Ibuprofen, USP, 200mg White Tablets**. This supplement included a model label and labeling based on the requirements in 21 CFR 201.66 and the approved label and labeling of Motrin IB. Attached are copies of smallest and largest size model label and labeling that will be used in all-label and labeling for the approved packaging sizes and in the third party product label and labeling that LNK manufactures and packages. LNK commits to submitting Final Printed Labels in our Annual Report.

LNK is certain that the information is sufficient for a comprehensive review of the supplemental request. If there is need for additional information, please call me at 631-543-3787.



Sincerely,


Pankaj S. Chudgar
Vice President

PSC/dju
Enc. Telephone: (631) 435-3500

Facsimile: (631) 435-3542

E-Mail: lnk01@ix.netcom.com