

MEDIQUE IPRIN- ibuprofen tablet, film coated
DOVER ADDAPRIN- ibuprofen tablet, film coated
MEDI-FIRST IBUPROFEN- ibuprofen tablet, film coated
MEDI-FIRST PLUS IBUPROFEN- ibuprofen tablet, film coated
OTIS CLAPP ULTRAPRIN- ibuprofen tablet, film coated
Unifirst First Aid Corporation

UniFirst First Aid Ibuprofen

Drug Facts

Active ingredient

Ibuprofen 200 mg (NSAID*)

*nonsteroidal anti-inflammatory drug

Purpose

Pain reliever/fever reducer

Uses

Temporarily relieves minor aches and pains associated with

■ headache ■ toothache ■ backache ■ menstrual cramps

■ common cold ■ muscular aches ■ minor arthritis pain

Temporarily reduces fever.

Warnings

Allergy Alert : Ibuprofen may cause a severe allergic reaction, especially in people allergic to aspirin. Symptoms may include:

■ hives ■ skin reddening ■ asthma (wheezing) ■ facial swelling ■ rash ■ shock ■ 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

Heart attack or stroke warning: NSAIDs, except aspirin, increase the risk of heart attack, heart failure, and stroke. These can be fatal. The risk is higher if you use more

than directed or for longer 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

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

Ask a doctor or pharmacist before use if you are

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

When using this product

- take with food or milk if stomach upset occurs

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
- you have symptoms of heart problems or stroke
 - chest pain
 - trouble breathing
 - weakness in one part or side of body
 - slurred speech
 - leg swelling
- pain gets worse or lasts for more than 10 days
- fever gets worse or lasts for more than 3 days
- redness or swelling is present in the painful area
- any new or unexpected symptoms occur

If pregnant or breast feeding, ask a health professional before use. It is especially important not to use ibuprofen at 20 weeks or later in 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 reach of Children. In case of overdose, get medical help or contact a Poison Control Center right away.

Directions

■ **do not use more than directed**

■ **the smallest effective dose should be used**

■ do not take longer than 10 days, unless directed by a doctor (see Warnings)

Adults and children:(12 years and older)

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 exceed 6 tablets in 24 hours, unless directed by a doctor.

Children under 12 years:

Ask a doctor

Other information

■ read all product information before using

■ store at 68-77°F (20-25°C)

■ avoid excessive heat 104°F (above 40°C)

■ tamper evident sealed packets

■ do not use any opened or torn packets

Inactive ingredients

carnauba wax*, corn starch, hypromellose*, iron oxide red, lactose*, magnesium stearate*, microcrystalline cellulose*, polydextrose*, polyethylene glycol, polyvinyl alcohol*, povidone K30*, silicon dioxide, sodium starch glycolate, stearic acid, talc*, titanium dioxide

****may contain***

Questions or comments? 1-800-634-7680

Medique Iprin Label

Medique® I-Prin

Ibuprofen 200 mg

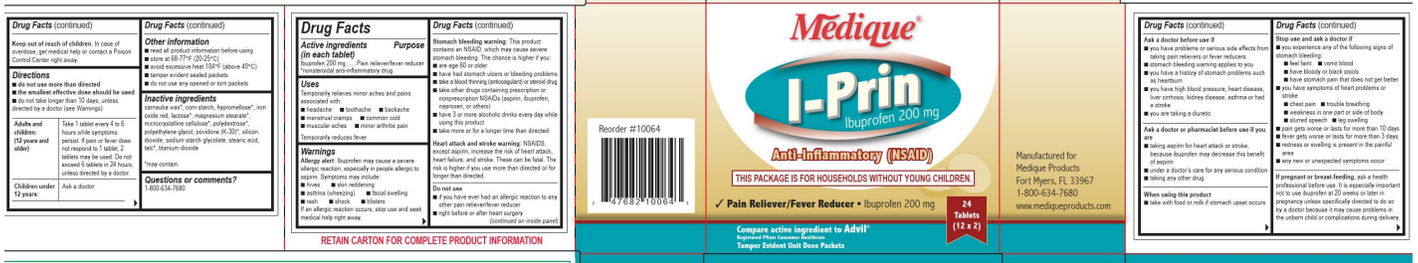
Anti-inflammatory (NSAID)

This Package is for Households without Young Children.

Pain Reliever/Fever Reducer • Ibuprofen 200 mg

24 Tablets (12 x 2)

Tamper Evident Unit Dose Packets



Medi-First Ibuprofen Label

Medi-First®

Ibuprofen 200 mg

100 tablets (50 x 2)

Pain Reliever/Fever Reducer

Aches, Fever • Ibuprofen (NSAID) 200mg

Pull to Open

Compare active ingredient to: Advil®

Registered Trademark of Pfizer Consumer Healthcare

This Package is for Households without Young Children.

Tamper Evident Unit Dose Packets



Medi-First Plus Ibuprofen Label

Medi-First® Plus

Ibuprofen

250 tablets (125 x 2's)

This Package is for Households without Young Children

Pull To Open

Ibuprofen 200 mg (NSAID)

Pain Reliever/Fever Reducer

Compare active ingredient to:

Advil®

Registered Trademark of Pfizer Consumer Healthcare

Tamper Evident Unit Dose Packets

Drug Facts
Active ingredient (in each tablet) Purpose
Ibuprofen 200 mg (NSAID) Pain reliever/fever reducer
*nonsteroidal anti-inflammatory drug

Uses
Temporarily relieves minor aches and pains associated with:
■ headache ■ backache ■ muscle aches ■ menstrual cramps
■ common cold ■ minor arthritis pain
Temporarily reduces fever.

Warnings
Allergy alert: Ibuprofen may cause a severe allergic reaction, especially in people allergic to aspirin. Symptoms may include:
■ hives ■ skin redness ■ asthma (wheezing)
■ facial swelling ■ rash ■ shock ■ 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)

Drug Facts (continued)
■ have 3 or more alcoholic drinks every day while using this product
■ take more or for a longer time than directed
Heart attack and stroke warning: NSAIDs, except aspirin, increase the risk of heart attack, heart failure, and stroke. These can be fatal. The risk is higher if you use more than directed or for longer 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
■ you have problems or serious side effects from taking pain relievers or fever reducers
■ you have high blood pressure, heart disease, liver cirrhosis, kidney disease, asthma or had a stroke
■ you have a history of stomach problems such as heartburn
■ you are taking a diuretic
Ask a doctor or pharmacist before use if you are
■ taking aspirin for heart attack or stroke, because ibuprofen may decrease the benefit of aspirin
■ under a doctor's care for any serious condition
■ taking any other drug (continued on opposite panel)

When using this product
■ take with food or milk if stomach upset occurs

Stop use and ask a doctor if
■ you experience any of the following signs of stomach bleeding:
■ red stool ■ vomit blood
■ have bloody or black stools
■ have stomach pain that does not get better
■ you have symptoms of heart problems or stroke
■ chest pain ■ trouble breathing
■ weakness in one part or side of body
■ slurred speech ■ leg swelling
■ skin gets worse or lasts for more than 10 days
■ fever gets worse or lasts for more than 3 days
■ redness or swelling is present in the painful area
■ any new or unexpected symptoms occur

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

Directions
■ do not use more than directed
■ the smallest effective dose should be used
■ do not take longer than 10 days, unless directed by a doctor (see Warnings)

Adults and children (12 years and older)
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 exceed 6 tablets in 24 hours, unless directed by a doctor.

Children under 12 years:
Ask a doctor

Other Information
■ read all product information before using
■ store at 68-77°F (20-25°C)
■ avoid excessive heat 104°F (above 40°C)
■ tamper evident sealed packets
■ do not use any opened or torn packets
(continued on opposite panel)

Drug Facts (continued)
Inactive ingredients
carnauba wax*, corn starch, hypromellose*, iron oxide red, lactose*, magnesium stearate*, microcrystalline cellulose*, polydextrose*, polyethylene glycol, povidone K-30†, silicon dioxide, sodium starch glycolate, stearic acid, talc*, titanium dioxide
*may contain
Questions or comments? 1-800-634-7686

Compare active ingredient to:
Advil®
Registered Trademark of Pfizer Consumer Healthcare

Tamper Evident Unit Dose Packets

RETAIN CARTON FOR COMPLETE PRODUCT INFORMATION

Barcode: 3 47682 90833

Model #0833

Manufactured by Med1FirstPlus
Fort Myers, FL 33947 USA
1-800-634-7686
www.med1products.com

Dover Addaprin Label

Dover Addaprin™

Pain Reliever-Fever Reducer

Ibuprofen 200 mg Tablets (NSAID)

This Package is for Households without Young Children.

Dover Pharmaceutical

Products of the highest quality and effectiveness

Tamper Evident

Sealed Packets

Unit Dose Packs

500 Tablets

(250 Packets of 2)

All Dover Pharmaceutical formulas conform to federal regulations

Antihistamine free

No danger of drowsiness

Sugar Free

for safer use by diabetics

Salt free

Minimizes high blood pressure

Caffeine Free

Avoides over stimulation

<p>All Dover Pharmaceutical formulas conform to federal regulations</p> <ul style="list-style-type: none"> Antihistamine Free No danger from the drowsiness side effects of antihistamines, on sodium restricted diets Sugar Free For safer use by those with hypertension (high blood pressure) or others on sodium restricted diets Sugar Free Contains no sucrose or dextrose for safer use by diabetics. Diabetes is the leading cause of blindness and third most frequent cause of death in the U.S. Caffeine Free Avoids possibility of excess stimulation and hypersensitivity. 	<p>All Dover Pharmaceutical formulas conform to federal regulations</p> <ul style="list-style-type: none"> Antihistamine Free No danger of drowsiness Sugar Free For safer use by diabetics Salt Free Minimizes high blood pressure Caffeine Free Avoids over-stimulation 	<p>All Dover Pharmaceutical formulas conform to federal regulations</p> <ul style="list-style-type: none"> Antihistamine Free No danger from the drowsiness side effects of antihistamines, on sodium restricted diets Salt Free For safer use by those with hypertension (high blood pressure) or others on sodium restricted diets Sugar Free Contains no sucrose or dextrose for safer use by diabetics. Diabetes is the leading cause of blindness and third most frequent cause of death in the U.S. Caffeine Free Avoids possibility of excess stimulation and hypersensitivity. 	<p>All Dover Pharmaceutical formulas conform to federal regulations</p> <ul style="list-style-type: none"> Antihistamine Free No danger of drowsiness Salt Free Minimizes high blood pressure Sugar Free For safer use by diabetics Caffeine Free Avoids over-stimulation
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Drug Facts	Purpose
Active ingredient (in each tablet) Ibuprofen 200 mg (NSAID)	Pain reliever/fever reducer
Uses Temporarily relieves minor aches and pains associated with: ● headache ● toothache ● menstrual cramps ● sore throat ● muscle aches ● minor arthritis pain	
Warnings Allergy alert: Ibuprofen may cause a severe allergic reaction, especially in people allergic to aspirin. Symptoms may include: ● hives ● skin redness ● asthma (wheezing) ● facial swelling ● rash ● shock 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 65 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 taken aspirin regularly every day while using this product ● take more or for a longer time than directed Heart attack or stroke warning: NSAIDs, except aspirin, increase the risk of heart attack, heart failure, and stroke. These can be fatal. The risk is higher if you use them from time to time or for longer than directed.	
Do not use ● if you have ever had an allergic reaction to any other pain reliever/fever reducer ● if you have had or after heart surgery	
Ask a doctor before use if ● you have problems or serious side effects from taking pain relievers or fever reducers ● stomach bleeding warning applies to you ● you have a history of stomach problems such as heartburn ● you have high blood pressure, heart disease, liver, kidney, or bladder problems ● you are taking a diuretic	
Ask a doctor or pharmacist before use if you are ● taking aspirin for heart attack or stroke, because ibuprofen may decrease this benefit ● aspirin ● under a doctor's care for any serious condition ● taking any other drug	
Questions or comments? 1-800-434-7880	

Dover **Addaprin™**
PAIN RELIEVER-FEVER REDUCER
Ibuprofen 200 mg Tablets (NSAID)

THIS PACKAGE IS FOR HOUSEHOLDS WITHOUT YOUNG CHILDREN.

Dover Pharmaceutical
Product of the highest quality and effectiveness

TAMPER EVIDENT Sealed Packets

UNIT DOSE PACKS
500 TABLETS
(250 PACKETS OF 2)

Dover **Addaprin™**
PAIN RELIEVER-FEVER REDUCER
Ibuprofen 200 mg Tablets (NSAID)

THIS PACKAGE IS FOR HOUSEHOLDS WITHOUT YOUNG CHILDREN.

Dover Pharmaceutical
Product of the highest quality and effectiveness

TAMPER EVIDENT Sealed Packets

UNIT DOSE PACKS
500 TABLETS
(250 PACKETS OF 2)

Dover **Addaprin™**
PAIN RELIEVER-FEVER REDUCER
Ibuprofen 200 mg Tablets (NSAID)

THIS PACKAGE IS FOR HOUSEHOLDS WITHOUT YOUNG CHILDREN.

Dover Pharmaceutical
Product of the highest quality and effectiveness

TAMPER EVIDENT Sealed Packets

UNIT DOSE PACKS
500 TABLETS
(250 PACKETS OF 2)



Reader No. 1625314
Manufactured for Medique Products, Fort Myers, FL 33967, 1-800-434-7880, www.mediqueproducts.com

Retain carton for complete product information

Otis Clapp Ultraprin Label
 OC Otis Clapp
 Quality & Integrity Since 1840
 Ultraprin™
 Pain Reliever-Fever Reducer (NSAID)
 Ibuprofen Tablets USP 200 mg
 For Deep Seated Pain
 See Warnings and Directions on Side Panel
 Tear Out Along Perforation To Dispense
 Professional Healthcare
 500 Tablets (250 Packets of 2)

Drug Facts	Purpose
Active ingredient (in each tablet) Ibuprofen 200 mg (NSAID)	Pain reliever/fever reducer
Uses Temporarily relieves minor aches and pains associated with: ● headache ● toothache ● menstrual cramps ● sore throat ● muscle aches ● minor arthritis pain	
Warnings Allergy alert: Ibuprofen may cause a severe allergic reaction, especially in people allergic to aspirin. Symptoms may include: ● hives ● skin redness ● asthma (wheezing) ● facial swelling ● rash ● shock 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 65 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 taken aspirin regularly every day while using this product ● take more or for a longer time than directed Heart attack or stroke warning: NSAIDs, except aspirin, increase the risk of heart attack, heart failure, and stroke. These can be fatal. The risk is higher if you use them from time to time or for longer than directed.	
Do not use ● if you have ever had an allergic reaction to any other pain reliever/fever reducer ● if you have had or after heart surgery	
Ask a doctor before use if ● you have problems or serious side effects from taking pain relievers or fever reducers ● stomach bleeding warning applies to you ● you have a history of stomach problems such as heartburn ● you have high blood pressure, heart disease, liver, kidney, or bladder problems ● you are taking a diuretic	
Ask a doctor or pharmacist before use if you are ● taking aspirin for heart attack or stroke, because ibuprofen may decrease this benefit ● aspirin ● under a doctor's care for any serious condition ● taking any other drug	
Questions or comments? 1-800-434-7880	

ULTRAPRIN™
Pain Reliever-Fever Reducer (NSAID)
Ibuprofen Tablets USP 200 mg
For Deep-Seated Pain

See Warning and Directions on Side Panel.

TEAR OUT ALONG PERFORATION TO DISPENSE

PROFESSIONAL HEALTHCARE

500 TABLETS (250 PACKETS OF 2)

ULTRAPRIN™
Ibuprofen Tablets USP 200 mg
For Deep-Seated Pain

Relieves headaches and pain
Relieves body aches
Reduces inflammation
Will not cause drowsiness

OTIS CLAPP
Quality & Integrity Since 1840
Mfg. for Medique Products, Fort Myers, FL 33967

MEDIQUE IPRIN
ibuprofen tablet, film coated

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:47682-600
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IBUPROFEN (UNII: WK2XY110QM) (IBUPROFEN - UNII:WK2XY110QM)	IBUPROFEN	200 mg

Inactive Ingredients

Ingredient Name	Strength
CARNAUBA WAX (UNII: R12CBM0EIZ)	
MAGNESIUM STEARATE (UNII: 70097M6130)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
STARCH, CORN (UNII: O8232NY3SJ)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
LACTOSE (UNII: J2B2A4N98G)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
POLYDEXTROSE (UNII: VH2XOU12IE)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	

Product Characteristics

Color	brown	Score	no score
Shape	ROUND	Size	10mm
Flavor		Imprint Code	44;291
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:47682-600-13	250 in 1 BOX	01/26/2017	
1	NDC:47682-600-99	2 in 1 PACKET; Type 0: Not a Combination Product		
2	NDC:47682-600-47	100 in 1 BOX	01/26/2017	
2		2 in 1 PACKET; Type 0: Not a Combination Product		
3	NDC:47682-600-64	12 in 1 BOX	01/26/2017	
3		2 in 1 PACKET; Type 0: Not a Combination Product		
4	NDC:47682-600-	2 in 1 BOX	01/26/2017	

4	69	2 in 1 PACKET; Type 0: Not a Combination Product	01/20/2017
4		2 in 1 PACKET; Type 0: Not a Combination Product	
5	NDC:47682-600-99	2 in 1 PACKET; Type 0: Not a Combination Product	02/01/2016

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA075010	02/01/2016	

DOVER ADDAPRIN

ibuprofen tablet, film coated

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:47682-614
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IBUPROFEN (UNII: WK2XYI10QM) (IBUPROFEN - UNII:WK2XYI10QM)	IBUPROFEN	200 mg

Inactive Ingredients

Ingredient Name	Strength
STARCH, CORN (UNII: O8232NY3SJ)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CARNAUBA WAX (UNII: R12CBM0EIZ)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
POLYDEXTROSE (UNII: VH2XOU12IE)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
LACTOSE (UNII: J2B2A4N98G)	

Product Characteristics

Color	brown	Score	no score
Shape	ROUND	Size	10mm
Flavor		Imprint Code	44;291
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:47682-614-13	250 in 1 BOX	01/26/2017	
1	NDC:47682-614-99	2 in 1 PACKET; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA075010	02/01/2016	

MEDI-FIRST IBUPROFEN

ibuprofen tablet, film coated

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:47682-608
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IBUPROFEN (UNII: WK2XYI10QM) (IBUPROFEN - UNII:WK2XYI10QM)	IBUPROFEN	200 mg

Inactive Ingredients

Ingredient Name	Strength
STARCH, CORN (UNII: O8232NY3SJ)	
CARNAUBA WAX (UNII: R12CBM0EIZ)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
LACTOSE (UNII: J2B2A4N98G)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
POLYDEXTROSE (UNII: VH2XOU12IE)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	

Product Characteristics

Color	brown	Score	no score
Shape	ROUND	Size	10mm
Flavor		Imprint Code	44;291
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:47682-608-30	4 in 1 BOX	01/26/2017	
1	NDC:47682-608-99	2 in 1 PACKET; Type 0: Not a Combination Product		
2	NDC:47682-608-33	50 in 1 BOX	01/26/2017	
2		2 in 1 PACKET; Type 0: Not a Combination Product		
3	NDC:47682-608-48	125 in 1 BOX	01/26/2017	
3		2 in 1 PACKET; Type 0: Not a Combination Product		
4	NDC:47682-608-13	250 in 1 BOX	01/26/2017	
4		2 in 1 PACKET; Type 0: Not a Combination Product		
5	NDC:47682-608-50	25 in 1 BOX	04/01/2019	
5		2 in 1 PACKET; Type 0: Not a Combination Product		
6	NDC:47682-608-99	2 in 1 PACKET; Type 0: Not a Combination Product	02/01/2016	

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA075010	02/01/2016	

MEDI-FIRST PLUS IBUPROFEN

ibuprofen tablet, film coated

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:47682-609
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IBUPROFEN (UNII: WK2XYI10QM) (IBUPROFEN - UNII:WK2XYI10QM)	IBUPROFEN	200 mg

Inactive Ingredients

Ingredient Name	Strength
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
CARNAUBA WAX (UNII: R12CBM0EIZ)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	
LACTOSE (UNII: J2B2A4N98G)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
HYPROMELLOSES (UNII: 3NXW29V3WO)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
POLYDEXTROSE (UNII: VH2XOU12IE)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ05DW1A)	
STARCH, CORN (UNII: O8232NY3SJ)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	

Product Characteristics

Color	brown	Score	no score
Shape	ROUND	Size	10mm
Flavor		Imprint Code	44;291
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:47682-609-13	250 in 1 BOX	01/26/2017	01/27/2017
1		2 in 1 PACKET; Type 0: Not a Combination Product		
2	NDC:47682-609-33	50 in 1 BOX	01/26/2017	
2		2 in 1 PACKET; Type 0: Not a Combination Product		
3	NDC:47682-609-48	125 in 1 BOX	01/26/2017	
3		2 in 1 PACKET; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA075010	02/01/2016	

OTIS CLAPP ULTRAPRIN

ibuprofen tablet, film coated

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:47682-602
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IBUPROFEN (UNII: WK2XYI10QM) (IBUPROFEN - UNII:WK2XYI10QM)	IBUPROFEN	200 mg

Inactive Ingredients

Ingredient Name	Strength
HYPROMELLOSES (UNII: 3NXW29V3WO)	
LACTOSE (UNII: J2B2A4N98G)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
POLYDEXTROSE (UNII: VH2XOU12IE)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
SODIUM STARCH GLYCOLATE TYPE A POTATO (UNII: 5856J3G2A2)	
STEARIC ACID (UNII: 4ELV7Z65AP)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	
CARNAUBA WAX (UNII: R12CBM0EIZ)	
STARCH, CORN (UNII: O8232NY3SJ)	
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
FERRIC OXIDE RED (UNII: 1K09F3G675)	

Product Characteristics

Color	brown	Score	no score
Shape	ROUND	Size	10mm
Flavor		Imprint Code	44;291
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:47682-602-13	250 in 1 BOX	02/01/2016	04/03/2017
1	NDC:47682-602-99	2 in 1 PACKET; Type 0: Not a Combination Product		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA075010	02/01/2016	04/03/2017

Labeler - Unifirst First Aid Corporation (832947092)

Revised: 9/2025

Unifirst First Aid Corporation