

IBUPROFEN- ibuprofen tablet, film coated
PD-Rx Pharmaceuticals, Inc.

IBUPROFEN 800 MG TABLETS

ibuprofen tablets 400 mg - 600 mg- 800 mg medguide

800 mg (white to off-white, capsule shaped, biconvex, film-coated tablets debossed with '123' on one side and plain on other side)

NDC 43063-858-06 Bottles of 6

800 mg label

TAKE _____ TABLET(S) _____ TIMES A DAY WITH FOOD.

PACKAGED BY PD-IDE PHARMACEUTICALS, ORLANDO, FL 32837

43063-858-06
IBUPROFEN
USP
800 MG
5 TABLETS
REORDER #222261
LOT F22E83 EXP 12/2023

43063-858-06
IBUPROFEN
USP
800 MG
5 TABLETS
REORDER #222261
LOT F22E83 EXP 12/2023

CALL YOUR DOCTOR FOR MEDICAL
ADVICE ABOUT SIDE EFFECTS

YOU MAY REPORT SIDE EFFECTS
TO THE FDA AT 1-800-FDA-1088

ORGANOLEPTIC MARKINGS:

WHITE 123 OBLONG



GTIN: 00343063858066

SNO: F22E83000044

EXP: 12/2023

LOT: F22E83

Rx Only

49483-6041
MARKSANS PHARMA LTD
VERNA GOA, 403 722 INDIA

NDC 43063-858-06

IBUPROFEN
USP

800 MG

6 TABLETS

EACH TABLET CONTAINS:
FILM COATED
IBUPROFEN USP 800 MG

WARNING:

KEEP OUT OF THE REACH OF CHILDREN

DOSAGE AND STORAGE:
SEE PACKAGE INSERT

IBUPROFEN

ibuprofen tablet, film coated

Product Information

Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:43063-858(NDC:49483- 604)
Route of Administration	ORAL		

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
IBUPROFEN (UNII: WK2XYI10QM) (IBUPROFEN - UNII:WK2XYI10QM)	IBUPROFEN	800 mg

Inactive Ingredients

Ingredient Name	Strength
SILICON DIOXIDE (UNII: ETJ7Z6XBU4)	
CROSCARMELOSE SODIUM (UNII: M28OL1HH48)	
MAGNESIUM STEARATE (UNII: 70097M6I30)	
CELLULOSE, MICROCRYSTALLINE (UNII: OP1R32D61U)	
POLYETHYLENE GLYCOL, UNSPECIFIED (UNII: 3WJQ0SDW1A)	
POLYVINYL ALCOHOL (UNII: 532B59J990)	
STARCH, PREGELATINIZED CORN (UNII: O8232NY3SJ)	
TALC (UNII: 7SEV7J4R1U)	
TITANIUM DIOXIDE (UNII: 15FIX9V2JP)	

Product Characteristics

Color	white	Score	no score
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Shape	CAPSULE	Size	19mm
Flavor		Imprint Code	123
Contains			

Packaging				
#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:43063-858-06	6 in 1 BOTTLE, PLASTIC; Type 0: Not a Combination Product	01/04/2019	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
ANDA	ANDA090796	12/30/2015	

Labeler - PD-Rx Pharmaceuticals, Inc. (156893695)

Registrant - PD-Rx Pharmaceuticals, Inc. (156893695)

Establishment			
Name	Address	ID/FEI	Business Operations
PD-Rx Pharmaceuticals, Inc.		156893695	repack(43063-858)

Revised: 3/2023

PD-Rx Pharmaceuticals, Inc.