

CIPROFLOXACIN HYDROCHLORIDE- ciprofloxacin hydrochloride powder

AX Pharmaceutical Corp

Ciprofloxacin Hydrochloride

Ciprofloxacin Hydrochloride

Caution: For pharmacy compounding use only. For veterinary use only. Use according to practitioner's prescription. Federal law prohibits dispensing without prescription. Keep tightly closed at controlled room temperature away from strong heat and light.



AX Pharmaceutical Corp

Ciprofloxacin HCl, USP

Retest Date: Dec 01, 2022

1kg

Original Reference: 105-181202-2

NDC: 73377-158-01

CAS: 86393-32-0

Relabelled by: AX Pharmaceutical Corp

Lot: E01-18L02SH

Original Manufacturer: Zhejiang Guobang Pharmaceutical CO., LTD
No.6 Wei Wu Road, Hangzhou Gulf Shangyu Industrial Zone, Zhejiang, China, 312369

100 TesmaWay, Unit 8, Concord, ON Canada L4K 0J9 Fax: 416 352 1618

Toll free: 1 866 305 0566

Harmful to aquatic life. May be harmful if swallowed. Causes serious eye irritation.
Avoid release to the environment. Wash thoroughly after handling.
Wear eye protection/face protection. IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses if present and easy to do - continue rinsing. IF eye irritation persists: Get medical advice/attention.



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AX Pharmaceutical Corp

Ciprofloxacin HCl, USP

Retest Date: Dec 01, 2022

50g

Original Reference: 105-181202-2

NDC: 73377-158-02

CAS: 86393-32-0

Relabelled by: AX Pharmaceutical Corp

Lot: E01-18L02SH

Original Manufacturer: Zhejiang Guobang Pharmaceutical CO., LTD
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Warning

CIPROFLOXACIN HYDROCHLORIDE

ciprofloxacin hydrochloride powder

Product Information

Product Type

Item Code (Source)

NDC:73377-158

Route of Administration

NOT APPLICABLE

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
CIPROFLOXACIN HYDROCHLORIDE (UNII: 4BA73M5E37) (CIPROFLOXACIN - UNII:5E8K9I0O4U)	CIPROFLOXACIN	1 g in 1 g

Product Characteristics

Color	yellow	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Marketing Start Date	Marketing End Date
1	NDC:73377-158-01	1000 g in 1 JAR		
2	NDC:73377-158-02	50 g in 1 JAR		

Marketing Information

Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
bulk ingredient for animal drug compounding		03/09/2022	

Labeler - AX Pharmaceutical Corp (204011316)

Establishment

Name	Address	ID/FEI	Business Operations
AX Pharmaceutical Corp		204011316	repack, relabel

Revised: 3/2022

AX Pharmaceutical Corp