

DEX-MOXI-KETOR- dexamethasone phosphate - moxifloxacin - ketorolac tromethamine injection, solution

Imprimis NJOF, LLC

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Store at 20° to 25° C (68° to 77° F)

Package Label

NDC 71384-513-01

Dexamethasone-Moxifloxacin-Ketorolac PF injection

(1/0.5/0.4) mg/ml

Volume: 1ml/vial

Quantity: 20

Lot: XXXXXX

Date Compounded:

DDMMYYYY

Expires on:

DDMMYYYY

Store at 20 to 25°C

Rev. 0

imprimis [®] **R_x**

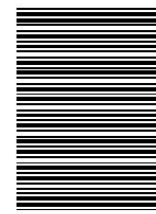
Imprimis NJOF, LLC.

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Ledgewood, NJ (844)446-6979

In case of adverse event contact:

www.fda.gov/medwatch or (800) FDA1088



This is a compounded drug.
NOT FOR RESALE
OFFICE USE ONLY

DEX-MOXI-KETOR

dexamethasone phosphate - moxifloxacin - ketorolac tromethamine injection, solution

Product Information

| | | | |
|--------------------------------|-------------------------|---------------------------|---------------|
| Product Type | HUMAN PRESCRIPTION DRUG | Item Code (Source) | NDC:71384-513 |
| Route of Administration | OPHTHALMIC | | |

Active Ingredient/Active Moiety

| Ingredient Name | Basis of Strength | Strength |
|---|--------------------------|-----------------|
| DEXAMETHASONE SODIUM PHOSPHATE (UNII: AI9376Y64P) (DEXAMETHASONE - UNII:7S5I7G3JQL) | DEXAMETHASONE PHOSPHATE | 1 mg in 1 mL |
| MOXIFLOXACIN HYDROCHLORIDE MONOHYDRATE (UNII: B8956S8609) (MOXIFLOXACIN - UNII:U188XYD42P) | MOXIFLOXACIN | 0.5 mg in 1 mL |
| KETOROLAC TROMETHAMINE (UNII: 4EVE5946BQ) (KETOROLAC - UNII:YZI5105V0L) | KETOROLAC | 0.4 mg in 1 mL |

Packaging

Marketing Start Marketing End

| # | Item Code | Package Description | Marketing Start Date | Marketing End Date |
|---|------------------|---|----------------------|--------------------|
| 1 | NDC:71384-513-01 | 20 in 1 BOX | 01/05/2018 | |
| 1 | | 1 mL in 1 VIAL, SINGLE-USE; Type 0: Not a Combination Product | | |

Marketing Information

| Marketing Category | Application Number or Monograph Citation | Marketing Start Date | Marketing End Date |
|-----------------------|--|----------------------|--------------------|
| unapproved drug other | | 01/05/2018 | |

Labeler - Imprimis NJOF, LLC (080431967)

Registrant - Imprimis NJOF, LLC (080431967)

Revised: 2/2020

Imprimis NJOF, LLC