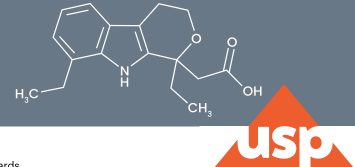
Etodolac Quality Solutions

Category: Analgesics



USP can support your development and manufacturing activities on Analgesics -based medicines with these existing and upcoming standards.

Authorized Distributor

MONOGRAPH 1

ETODOLAC - NEW Official as of 1-Dec-2024	
USP Etodolac Peak Identification Mixture RS	<u>USP-1268892</u>
USP Etodolac RS	<u>USP-1268706</u>

MONOGRAPH 2

TABLETS OFFICIAL AS OF 31-DEC-2012	
USP Etodolac RS	<u>USP-1268706</u>
USP Etodolac Related Compound A RS	<u>USP-1268728</u>

MONOGRAPH 3

ETODOLAC TABLETS Official as of 1-May-2024	
USP Etodolac RS	<u>USP-1268706</u>
USP Etodolac Related Compound A RS	<u>USP-1268728</u>
USP Etodolac Related Compound H RS	<u>USP-1268769</u>

MONOGRAPH 4

ETODOLAC CAPSULES Official as of 31-Dec-2012	
USP Etodolac RS	<u>USP-1268706</u>
USP Etodolac Related Compound A RS	<u>USP-1268728</u>
PHARMACEUTICAL ANALYTICAL IMPURITIES (PAI)*	
1-PROPYL ETODOLAC	<u>USP-1A07450</u>

<11> USP REFERENCE STANDARDS Official as of 01-Nov-2020

INCLUDED GENERAL CHAPTERS

<197> SPECTROSCOPIC IDENTIFICATION TESTS

Official as of 01-Sep-2021

<281> RESIDUE ON IGNITION

Official as of 31-Dec-2012

<541> TITRIMETRY

Officialas of O1-Aug-2024

<621> CHROMATOGRAPHY - NEW

Official as of O1-Dec-2024

<711> DISSOLUTION

Official as of 1-May-2023

USP Dissolution Performance Verification USP-1222818 Standard-Prednisone RS

<905> UNIFORMITY OF DOSAGE UNITS

Officialas of 01-Aug-2023

<921> WATER DETERMINATION

Official as of O1-May-2022

USP Sodium Tartrate Dihydrate RS USP-1614909

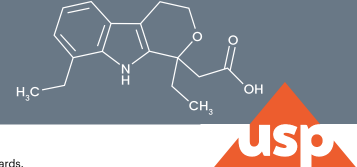
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^{*} Pharmaceutical Analytical Impurities (PAI products) are released using a process developed by USP's subject matter experts. The release process is based on internal policies, standard operating procedures, and requirements as defined by USP's Quality Management System. USP is an ISO 9001:2015 certified facility. PAI products are different from official USP Reference Standards. PAI products are not required for compendial compliance.

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INCLUDED EXCIPIENTS

LACTOSE MONOHYDRATE Official as of 1-Dec-2024	
USP Dextrose RS	<u>USP-1181302</u>
USP Fructose RS	<u>USP-1286504</u>
USP Lactose Monohydrate RS	<u>USP-1356701</u>
USP Sucrose RS	<u>USP-1623637</u>

TITANIUM DIOXIDEOfficial as of 1-Jun-2023

MAGNESIUM STEARATE Officialas of 1-Aug-2016	
USP Palmitic Acid RS	<u>USP-1492007</u>
USP Stearic Acid RS	USP-1621008

FERRIC OXIDE

MICROCRYSTALLINE CELLULOSE Official as of 1-Dec-2019	
USP Microcrystalline Cellulose RS	<u>USP-1098388</u>