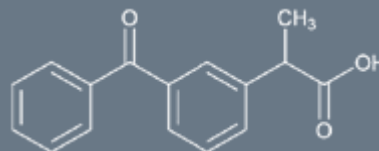


Ketoprofen

Quality Solutions

Category: Analgesic



Authorized Distributor

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

Official and Proposed Documentary Standards and Associated Physical Materials

MONOGRAPH 1

KETOPROFEN Official as of 01-May-2020	
USP Ketoprofen RS	USP-1356632
USP Ketoprofen Related Compound D RS	USP-1356563

MONOGRAPH 2

KETOPROFEN CAPSULES Official as of 01-May-2020	
USP Ketoprofen RS	USP-1356632
USP Ketoprofen Related Compound A RS	USP-1356643
USP Ketoprofen Related Compound C RS	USP-1356552
USP Ketoprofen Related Compound D RS	USP-1356563

MONOGRAPH 3

KETOPROFEN EXTENDED-RELEASE CAPSULES Official as of 01-May-2020	
USP Ketoprofen RS	USP-1356632
USP Ketoprofen Related Compound D RS	USP-1356563

RELATED EXCIPIENTS STANDARDS

SUCROSE Official as of 01-Jun-2023	
USP Sucrose RS	USP-1623637

RELATED EXCIPIENTS STANDARDS

TRIETHYL CITRATE Official as of 01-May-2023	
USP Triethyl Aconitate RS	USP-1683537
USP Triethyl Citrate RS	USP-1683606

RELATED EXCIPIENTS STANDARDS

MANNITOL Official as of 01-May-2020	
USP Mannitol RS	USP-1375105

RELATED EXCIPIENTS STANDARDS

GELATIN Official as of 01-Nov-2020	
USP Gelatin RS	USP-1288485

01-Oct-2023. For publications related to **Ketoprofen** in PF online after this date, please visit the integrated [USP-NF/PF online](#).

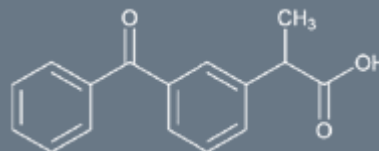
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Official and Proposed Documentary Standards and Associated Physical Materials

PHARMACEUTICAL ANALYTICAL IMPURITY(IES)*	
USP 2-(3-Carboxyphenyl)propionitrile PAI	USP-1A02750
USP 3-Carboxyphenylacetoneitrile PAI New	USP-1A02760
USP 2,4,5-Trimethylketoprofen PAI	USP-1A02780
USP Benzoylphenylacetoneitrile PAI New	USP-1A02910
USP Dimethylketoprofen PAI New	USP-1A02770

* 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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