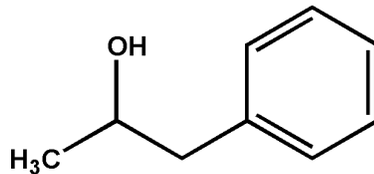




U.S. Pharmacopeia
The Standard of QualitySM

USP Certificate

Dextroamphetamine Related Compound A LOT F0H131



Molecular Formula

C₉H₁₂O

Molecular Weight

136.19

CAS Number

698-87-3

LABEL TEXT

USP[®] REFERENCE STANDARD

DEXTROAMPHETAMINE RELATED COMPOUND A 25 mg
(1- Phenyl - 2 - propanol)

Warning! Combustible liquid.

Do not dry. After opening the ampul, store in a tightly closed container. Protect from light.

Keep away from flames and hot surfaces-No smoking.
Wear protective gloves/eye protection/face protection.
In case of fire: Use appropriate media to extinguish.
Store in a well-ventilated place. Keep cool. Dispose of contents/container in accordance with local/regional/national/international regulations.

For use with specified USP compendial tests.
Not for use as a drug. See SDS prior to use at www.usp.org/sds.

KEEP AMPULE IN SECONDARY CONTAINER!
USP, 12601 Twinbrook Pkwy, Rockville, MD, +1-301-881-0666
CAT. NO. 1180015 Material mfd. in Belgium

LOT: F0H131



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Protect from light.

Intentionally over-labeled for GHS compliance

USP, 12601 Twinbrook Pkwy, Rockville, MD, +1-301-881-0666

CAT. NO. 1180015

Lot: F0H131

Material mfd in Belgium

USP certifies that the USP Reference Standards Committee, in accordance with their rules and procedures, determined that this USP Reference Standard lot is suitable to assess compliance with the monograph standards for which it is specified. The critical characteristics of this lot are usually determined independently in three or more laboratories, including USP, government, academic, and industrial collaborators.

Jeri L. Joth

QA Director

Calculation Value

Unless otherwise stated on the Reference Standard label, a value of 100.0% should be used in USP or NF compendial applications for which the use of this Reference Standard is intended. Please refer to the specific Reference Standard label for further information.

Expiration

Current lots are identified in the Official USP Reference Standards catalog. In some cases, the previous lot may still be considered official. If so, it is identified in the column marked "Previous Lot/Valid Use Date." Ordinarily, the previous lot is carried in official status for about one year after the current lot enters distribution.

It is the responsibility of each user to determine that this lot is current when used. To ensure up-to-date information, USP publishes the Official USP Reference Standards Catalog, which contains official lot designations. This information is also available on the USP web site, at www.usp.org, as well as in the bimonthly subscription publication, *Pharmacopeial Forum*.

Instructions for Use

Follow the instructions in the appropriate USP or NF Monographs and General Requirements for Tests and Assays of the current *USP–NF*. In the event that instructions on the label of this lot differ from those found in the current *USP–NF*, those on the label supersede any instructions listed in Chapter <11>.

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