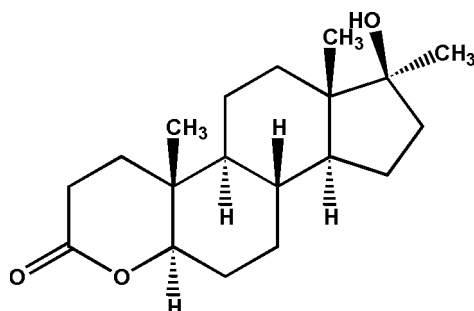




U.S. Pharmacopeia
The Standard of QualitySM

USP Certificate

Oxandrolone Related Compound B LOT F0F136



Molecular Formula

C₁₉H₃₀O₃

Molecular Weight

306.44

CAS Number

4424-45-7

LABEL TEXT



Lot No.: F0F136



REFERENCE STANDARD

OXANDROLONE RELATED COMPOUND B CIII 20 mg
(17beta-hydroxy-17alpha-methyl-4-oxa-5alpha-androstan-3-one)

NDC# 00216-4016-01



Danger! May cause cancer. May damage fertility or the unborn child.

Do not dry. Keep container tightly closed. Protect from light. Store in a refrigerator.

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

CAT No. 1482025

Material mfd in Hungary

Intentionally over-labeled for GHS compliance

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

Obtain special instructions before use. Do not handle until all safety precautions have been read and understood. Wear protective gloves/protective clothing/eye protection/face protection. If exposed or concerned: Get medical advice/attention. Store locked up. Dispose of contents/container in accordance with local/regional/national/international regulations.

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Jeri L. Toth

QA Director

Calculation Value

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