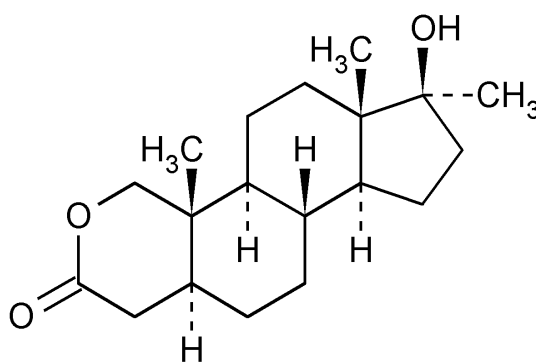




U.S. Pharmacopeia
The Standard of QualitySM

USP Certificate

Oxandrolone LOT H0E223



Molecular Formula

C₁₉H₃₀O₃

Molecular Weight

306.44

CAS Number

53-39-4

LABEL TEXT



Lot No.: H0E223



USP[®] REFERENCE STANDARD

OXANDROLONE CIII 50 mg
NDC # 00216-4005-01

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

Do not dry. For quantitative applications, use a value of 0.994 mg of oxandrolone per mg on the as-is basis. Keep container tightly closed. Protect from light.

USP, 12601 Twinbrook Pkwy, Rockville, MD, +1-301-881-0666
CAT No. 1482003 Material mfd in United States
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.

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, FDA, and academic or industrial collaborators.

Jeri L. Joth

QA Director

Calculation Value

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