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USP 31-NF 26 | USP-NF - USP-NF | USP-NF
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The electronic publications of Second Supplement to USP 31 NF 26 contain a version of <55> Biological Indicators Resistance Performance Tests that does not include the revised text with revision mark up, as it appeared in the First Supplement to USP 31 NF 26. This will be

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corrected in the online version of the Second Supplement to USP 31-NF 26 by July 25. It will be correct in the USP ...

Second Supplement to USP 31-NF 26 Online and CD includes ...

Revisions to USP 31-NF 26, Second Supplement (published May 2008) Published April 2008 General Chapters
Monographs: A-C D-N O-S T-Z Monograph Title Section
Head Scientific Liaison DANTROLENE SODIUM CAPSULES
PF 33(4) Pg. 645 Dissolution <711> DEHYDROACETIC
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Transfer a 100-mL specimen to a tared, pear-shaped, 100-mL centrifuge tube containing a few boiling stones, and weigh.

Suspend a thermometer in the liquid, and place the tube in a medium maintained at a temperature of 32 above the

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expected boiling temperature. When the thermometer reading becomes constant, record as the boiling temperature the thermometer reading after at least 5% of the ...

usp31nf26s1_c601, General Chapters: <601> AEROSOLS, NASAL ...

A complete list of Packings (L), Phases (G), and Supports (S) used in USP-NF tests and assays is located under Chromatographic Reagents in the Reagents, Indicators, and Solutions section. This list is intended to be a convenient reference for the chromatographer to identify the pertinent chromatographic reagent specified in the individual monograph. [note- Particle sizes given in the ...

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usp31nf26s1_c621, General Chapters: <621>

CHROMATOGRAPHY

USP-NF is a combination of two compendia, the United States Pharmacopeia (USP) and the National Formulary (NF). Monographs for drug substances, dosage forms, and compounded preparations are featured in the USP.

Monographs for dietary supplements and ingredients appear in a separate section of the USP. Excipient monographs are in the

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Fatty Products Dissolve in isopropyl myristate sterilized by filtration, or mix the product to be examined with the minimum necessary quantity of sterile polysorbate 80 or another

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noninhibitory sterile surface-active reagent heated, if necessary, to not more than 40 or, in exceptional cases, to not more than 45. Mix carefully and if necessary maintain the temperature in a water bath.

usp31nf26s1_c61h, General Chapters: <61>

MICROBIOLOGICAL ...

USP-NF. Three New Revision Bulletins (posted 25-Sep-2020) Two New Interim Revision Announcements (posted 25-Sep-2020) One New General Chapter Prospectus (posted 25-Sep-2020) One new Revision Bulletin (posted 25-Sep-2020) Cumulative List Updated (posted 26-June-2020) Reference Standards. Updates on USP Reference Standards in response to COVID-19. Download

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full list as: EXCEL ...

U.S. Pharmacopeia

If the substance is a solid □ Transfer the weighed quantity of the test substance to a clean, dry, 100-mL Kjeldahl flask. [note □ A 300-mL flask may be used if the reaction foams excessively.] Clamp the flask at an angle of 45, and add a sufficient quantity of a mixture of 8 mL of sulfuric acid and 10 mL of nitric acid to moisten the substance thoroughly.

usp31nf26s1_c231, General Chapters: <231> HEAVY METALS

Mix the L-cystine, sodium chloride, dextrose, yeast extract, and pancreatic digest of casein with the purified water, and

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heat until solution is effected. Dissolve the sodium thioglycollate or thioglycolic acid in the solution and, if necessary, add 1 N sodium hydroxide so that, after sterilization, the solution will have a pH of 7.1 ± 0.2 .

usp31nf26s1_c71, General Chapters: <71> STERILITY TESTS

This proposed revision suggested a May 14, 2014 date linking General Chapters <232> Elemental Impurities—Limits and <233> Elemental Impurities—Procedures to drug product monographs in the United States Pharmacopeia (USP). As such, section 5.60.30 will not be included in the General Notices that will be published in USP 37—NF 32, and therefore there is no requirement for any drug product ...

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Elemental Impurities Updates | USP

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The United States Pharmacopeia (USP) recently announce that its official standards publication, the United States Pharmacopeia 27 and National Formulary 22 (USP 27-NF 22) became official on Jan. 1, 2004. Two Supplements will follow in February and June-becoming official in April and August 2004, respectively. This edition of the USP-NF contains several new features and

USP 27- NF 22 Becomes Official - Third Annual Edition ...
omitted from the USP-NF in the indicated Book or Supplement. The requirements stated in the . General Notices and Requirements section of the USP-NF apply to all articles recognized in the USP -NF and to all general

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chapters unless specifically stated otherwise. Although this revision (USP 38 NF 33) is generally official beginning May 1, 2015, particular provisions may indicate another ...

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