

[x](#)

How to Use

- **Searching:** Type keyword in search field at top of page. Search by all or part of a monograph title. For searches using multiple criteria, you will find items that match each of the specified criteria unless quotation marks are used.
 - For example, a search on Aminosalicyclic Acid Tablets will result in anything that contains “Aminosalicyclic” OR “Acid” OR “Tablets”
 - A search for “Aminosalicyclic Acid Tablets” will result in anything that specifically contains “Aminosalicyclic Acid Tablets”
- **Sorting:** Click on any column header title to sort alphabetically or chronologically in ascending or descending order. Note: the page load column is sorted alphabetically so that a number is ordered by first digit vs. by the actual number; thus, numbers will not always be in order.
 - For example, page 2178 will come before page 74 on a page sort.
- **Downloading:** You can download the Errata table in Comma-separated Value (.csv). The download will include the Errata that you have filtered on.
- **Importing:** You will need to import the file into Excel or Open Office with UTF-8 encoding, as opposed to simply opening it. To import, open Excel or Open Office and select import from the File drop-down. Depending on the version you are using, you should be presented with import formatting options to include UTF-8 as one of the first steps. Importing via UTF-8 should eliminate odd character conversions.

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
POTASSIUM M SPECIFIC	USP36–NF31	2172	27-Sep-2013	1-Oct-2013	USP38–NF33	First	Line 1: Change

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Errata Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
ETAPHOSPHA TE	TESTS/ <i>Viscosity—Capillary Viscometer Methods <911> and Rotational Rheometer Methods <912></i>							<i>Supplement to USP37–NF32</i>	and to: or
ISOSORBIDE MONONITRATE EXTENDED- RELEASE TABLETS	IMPURITIES/ <i>Organic Impurities, Procedure 2</i>	<i>First Supplement to USP36–NF31</i>	5996	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 3: Change the section head <i>Isosorbide mononitrate related compound A stock solution:</i> to: <i>Isosorbide mononitrate related compound A standard stock solution:</i> AND Line 7: Change the section head <i>Isosorbide dinitrate stock solution:</i> to: <i>Isosorbide</i>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<i>dinitrate standard stock solution:</i> AND Line 3 of <i>Standard stock solution:</i> Change <i>Isosorbide mononitrate related compound A stock solution and Isosorbide dinitrate stock solution,</i> to: <i>Isosorbide mononitrate related compound A standard stock solution and Isosorbide dinitrate standard stock solution,</i> Line 1 of Sample: Change 1 g of the
SORBITAN SE IDENTIFICATIO SQUIOLEATE N/A.	USP36–NF31	2215	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
PANCURONIUM BROMIDE INJECTION	IM PUR ITIES/ <i>Organic Impurities</i>	<i>Second Supplement to USP36–NF31</i>	6677	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	residue obtained in the <i>Assay for Fatty Acids</i> to: Residue obtained in the <i>Assay for Fatty Acids</i> AND Line 2 of <i>Acceptance criteria</i> : Change 192–204 to: 192–204 on 1-g sample Footnote a of <i>Table 1</i> : Change Piperidinium, 1-[(2,3,5,16,17)-17-acetyloxy-3-hydroxy-2-(1-piperidinyl)androstan-16-yl]-1-methyl bromide. to: Piperidinium, 1-[(2,3,5,16,17)-17-acetyloxy-3-h

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
CAFFEINE CITRATE INJECTION	Assay	USP36–NF31	2732	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	hydroxy-2-(1-piperidinylandrost an-16-yl]-1-methyl. Line 3 of <i>Chromatographic system</i> : Change 150-cm column to: 15-cm column
BEHENOYL POIM LYOXYLGLYC ERIDES	PURITIES/ <i>Limit of Free Glycerol /Titrimetric system</i>	USP36–NF31	1892	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of <i>Mode</i> : Change Direct titration to: Residual titration
ESOMEPRAZO LE MAGNESIUM	SPECIFIC TESTS/ <i>Color of Solution</i>	USP36–NF31	3464	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of <i>Sample solution</i> : Change 20 mg/mL of Esomeprazole Magnesium in methanol to: 20 mg/mL of Esomeprazole Magnesium in methanol, filtered

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Errata Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
EGG PHOSPH OLIPIDS	ASSAY/ <i>Content of Phospholipids</i>	USP36–NF31	2000	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 2 of <i>Solution A</i> : Change acetic acid to: glacial acetic acid AND Line 2 of <i>Solution B</i> : Change acetic acid to: glacial acetic acid
OCTOCRYLEN E	<i>Identification, Ultraviolet Absorption <197U></i>	USP36–NF31	4557	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 4: Change Absorptivities, calculated on the as-is basis, to: Absorptivity at 303 nm, calculated on the as-is basis,
LAUROYL POL IM YOXYLGLYCE RIDES	PURITIES/ <i>Limit of Free Glycerol</i>	USP36–NF31	2064	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Mode in Titrimetric system</i> : Change Direct titration to: Residual titration

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>AND</p> <p>Line 10 of <i>Analysis</i>: Change $(V_S ? V_B)$ to: $(V_B ? V_S)$</p> <p>AND</p> <p>Line 11 of <i>Analysis</i>: Change $V_S = \textit{Titrant}$ volume consumed by the <i>Sample</i> (mL) $V_B = \textit{Titrant}$ volume consumed by the <i>Blank</i> (mL) to: $V_B = \textit{Titrant}$ volume consumed by the <i>Blank</i> (mL) $V_S = \textit{Titrant}$ volume consumed by the <i>Sample</i> (mL)</p>

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Errata Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
RIMEXOLONE OPTHALMIC SUSPENSION	<i>Viscosity—Capillary Viscometer Methods <911> and Rotational Rheometer Methods <912></i>	USP36–NF31	5053	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1: Change and to: or
POLYOXYL 20 CETOSTEARYL ETHER	SPECIFIC TESTS/Average Polymer Length	USP36–NF31	2155	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 16 of Analysis: Change Result = $[32 \times A_2 / (A_1 \times 3)] / 4$ to: Result = $[(32 \times A_2 / A_1) \times 3] / 4$
URSODIOL TABLETS	IMPURITIES/Organic Impurities/Procedure	USP36–NF31	5520	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 2 of Sample solution: Change Tablets, equivalent to about 25 mg of ursodiol, to: Tablets, equivalent to about 250 mg of ursodiol,
SORBITAN MONOPALMITATE	IDENTIFICATION	USP36–NF31	2213	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of Sample: Change 1 g of the

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
									residue obtained in the <i>Assay for Fatty Acids</i> to: Residue obtained in the <i>Assay for Fatty Acids</i> AND Line 2 of <i>Acceptance criteria</i> : Change 210–225 to: 210–225 on 1-g sample
BETHANECHOL CHLORIDE	IMPURITIES/Heavy Metals, Method 1 <231>	<i>Second Supplement to USP36–NF31</i>	6568	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Test preparation</i> : Change Bethacholine Chloride to: Bethanechol Chloride
STEARYL ALCOHOL	SPECIFIC TESTS/ <i>Fats and Fixed Oils, Hydroxyl Value</i> <401>	<i>USP36–NF31</i>	2252	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 14 of <i>Analysis</i> : Change $[(V_S - V_B) \times N \times F]/W$ to:

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p> $[(V_B - V_S) \times N \times M_r] / W$ AND Line 15 of <i>Analysis:</i> Change V_S = volume of 1 N sodium hydroxide consumed by the sample (mL) V_B = volume of 1 N sodium hydroxide consumed by the blank test (mL) to: V_B = volume of 1 N sodium hydroxide consumed by the blank test (mL) V_S = volume of 1 N sodium hydroxide consumed by the sample (mL) AND Line 21 of </p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
ACETYLTRIBUTYL CITRATE IDENTIFICATION N/B.	USP36–NF31	1869	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	<p><i>Analysis:</i> Change F = molecular weight of potassium hydroxide, 56.11 to: M_r = molecular weight of potassium hydroxide, 56.11</p> <p>Line 2: Change USP Acetyltriethyl Citrate RS to: USP Acetyltributyl Citrate RS</p>
CLAVULANATE IM POTASSIUM PURITIES/Organic Impurities/Procedure 3: Limit of Aliphatic Amines	USP36–NF31	3022	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	<p>In the definition list in <i>Analysis:</i> Change C_U = nominal concentration of Clavulanate Potassium in the <i>Standard solution</i> to:</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
CETOSTEARY SPECIFIC L ALCOHOL TESTS/ <i>Fats and Fixed Oils, Hydroxyl Value <401></i>	USP36–NF31	1954	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	<p>C_U = nominal concentration of Clavulanate Potassium in the <i>Sample solution</i></p> <p>Line 1 of <i>Mode in Titrimetric system</i>: Change Direct titration to: Residual titration AND Line 16 of <i>Analysis</i>: Change Result = $[(V_S ? V_B) \times F]/W$ to: Result = $[(V_B ? V_S) \times N \times M_r]/W$ AND Line 17 of <i>Analysis</i>: Change $V_S = \text{Titration volume consumed by the Sample (mL)}$</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
MERCAPTOPYRIMIDINE TABLETS PUR ITIES/Organic Impurities/Procedure	USP36–NF31	4249	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	V_B = Titrant volume consumed by the <i>Blank</i> (mL) F = calculation factor, 56.1 to: V_B = Titrant volume consumed by the <i>Blank</i> (mL) V_S = Titrant volume consumed by the <i>Sample</i> (mL) N = actual normality of 1 N sodium hydroxide (mEq/mL) M_r = molecular weight of potassium hydroxide, 56.11 Line 6 of <i>Sample stock</i> <i>solution:</i> Change Dilute with

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
GLYCERYL MOASSAY/ NOLINOLEATE	<i>Procedure</i>	USP36–NF31	2030	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	water to volume. to: Dilute with <i>Solution A</i> to volume. Line 13 of <i>Chromatographic system</i> : Change <i>Column temperature</i> : 40° to: <i>Temperatures Detector</i> . 40° <i>Column</i> : 40°
QUININE SULFATE	IM PUR ITIES/ <i>Dihydroquinine Sulfate</i>	USP36–NF31	4995	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 5 of <i>Analysis</i> : Change Result = $(r_U/r_S) \times 100$ to: Result = $r_U/(r_U + r_S) \times 100$
POLYISOBUTYLENE	IM PURITIES/ <i>Lead <251></i>	USP36–NF31	2149	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Acceptance criteria</i> : Change NMT 3 mg/g to: NMT 3 ?g/g

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Errata Post Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
TERAZOSIN CAPSULES	PERFORMANC E TESTS/ Dissolution <711>/Test 1/Spectrometric conditions	USP36–NF31	5308	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of Analytical wavelength: Change UV 245 nm to: UV 246 nm AND Line 3 of Cell length: Change 0.2 cm for Capsules labeled to contain 10 mg to: 0.5 cm for Capsules labeled to contain 10 mg
SORBITAN MO IDENTIFICATIO NOLAURATE	N/A.	USP36–NF31	2212	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of Sample: Change 1 g of the residue obtained in the Assay for Fatty Acids to: Residue obtained in the Assay for Fatty

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
									Acids AND Line 2 of <i>Acceptance criteria</i> : Change 260–280 to: 260–280 on 1-g sample
LOPINAVIR AND RITONAVIR TABLETS	PERFORMANC E TESTS/ <i>Dissolution</i> <711>	<i>First Supplement to USP36–NF31</i>	6005	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Tolerances</i> : Change 80.0% to: 80%
SORBITAN TRIOLEATE	IDENTIFICATIO N/A.	<i>USP36–NF31</i>	2216	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Sample</i> : Change 1 g of the residue of oleic acid obtained in the <i>Assay for Fatty Acids</i> to: Residue obtained in the <i>Assay for Fatty Acids</i> AND Line 2 of <i>Acceptance</i>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
POLYMYXIN B IM SULFATE PUR ITIES/Organic Impurities	<i>Second Supplement to USP36–NF31</i>	6686	27-Sep-2013	1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	<i>criteria: Change 192–204 to: 192–204 on 1-g sample</i> Line 2: Change <i>Buffer, Mobile phase, Diluent, Sensitivity solution,</i> to: <i>Buffer, Mobile phase, Diluent, Standard solution, Sensitivity solution, AND</i> Line 1 of <i>Samples in Analysis: Change Standard solution, Sample solution, and Sensitivity solution</i> to: <i>Sample solution</i> Line 17 of
ELEMENTAL I DRUG	<i>USP36–NF31</i>	151	27-Sep-2013	1-Oct-2013	<i>USP38–NF33</i>	<i>First</i>	

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
MPURITIES—LIP MITS	RO DUCTS/ <i>Options for Demonstrating Compliance</i>							<i>Supplement to USP37–NF32</i>	<i>Summation Option: Change the manufacturer must validate to: the manufacturer must ensure</i>
CAFFEINE CITRATE ORAL SOLUTION	<i>Assay</i>	<i>USP36–NF31</i>	2733	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 3 of <i>Chromatographic system: Change 150-cm column to: 15-cm column</i>
BETADEX	IM PURITIES/ <i>Limit of Reducing Sugars</i>	<i>USP36–NF31</i>	1905	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 3 of <i>Tartrate solution: Change 20 mg/mL of anhydrous sodium sulfate to: 200 mg/mL of anhydrous sodium sulfate</i>
ETHIODIZED OIL INJECTION	<i>Viscosity—Capillary Viscometer Methods <911> and Rotational</i>	<i>USP36–NF31</i>	3505	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1: <i>Change and to: or</i>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
DIPHENHYDR AMINE HYDRO CHLORIDE INJECTION	<i>Rheometer</i> <i>Methods <912></i> Assay USP36–NF31	3276	26-Jul-2013	1-Aug-2013	USP38–NF33	<i>First</i> <i>Supplement to</i> <i>USP37–NF32</i>	Line 2: Change <i>Mobile phase,</i> <i>Standard</i> <i>preparation,</i> <i>System</i> <i>suitability</i> <i>solution,</i> and <i>Chromatographi</i> <i>c</i> <i>system</i> —Prepare as directed in the Assay under <i>Diphenhydrami</i> <i>ne</i> <i>Hydrochloride.</i> to: <i>Mobile</i> <i>phase</i> —Prepare a solution of acetonitrile, water, and triethylamine (50: 50: 0.5), adjust with glacial acetic acid to a pH of 6.5, filter, and degas. Make adjustments if

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>necessary (see <i>System Suitability</i> under <i>Chromatography</i> <621>). <i>Standard preparation</i></p> <p>—Dissolve an accurately weighed quantity of USP Diphenhydramine Hydrochloride RS in water to obtain a solution having a known concentration of about 0.5 mg per mL.</p> <p>AND</p> <p>After the <i>Assay preparation</i> subsection: Add <i>System suitability solution</i></p> <p>—Dissolve about 5 mg of</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>benzophenone in 5 mL of acetonitrile, dilute with water to 100 mL, and mix. Transfer 1.0 mL of this solution and 5 mg of diphenhydramine hydrochloride to a 10-mL volumetric flask, dilute with water to volume, and mix.</p> <p><i>Chromatographic system (see Chromatography <621>)</i>—The liquid chromatograph is equipped with a 254-nm detector and a 4.6-mm x 25-cm column that contains packing L10. The flow rate is about 1 mL per</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>minute.</p> <p>Chromatograph the <i>System suitability solution</i>, and record the peak responses as directed for <i>Procedure</i>; the resolution, <i>R</i>, between the benzophenone and diphenhydramine peaks is not less than 2.0.</p> <p>Chromatograph replicate injections of the <i>Standard preparation</i>, and record the peak responses as directed for <i>Procedure</i>; the relative standard deviation is not more than 2.0%, and the tailing factor for</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>the diphenhydramine hydrochloride peak is not more than 2.0. AND</p> <p>Line 1 of <i>Procedure:</i> Change Proceed as directed for <i>Procedure</i> in the Assay under <i>Diphenhydramine Hydrochloride</i>. to:</p> <p>Separately inject equal volumes, about 10 µL, of the <i>Standard preparation</i> and the <i>Assay preparation</i> into the chromatograph, record the chromatograms, and measure the responses</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
FEXOFENADIN ASSAY/ E HYDROCHL ORIDE TABLETS	<i>USP36–NF31 Procedure</i>	3576	26-Jul-2013	1-Aug-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	for the major peaks. Line 6 of <i>Sample stock solution:</i> Change (equivalent to 80% of the total flask volume) to: (sufficient to fill the flask to 80% of its volume)
NIFEDIPINE EXPERFORMANC TENDED- RELEASE TABLETS	<i>USP36–NF31 E TESTS/ Dissolution <711>/Test 4/Instrumental conditions</i>	4509	26-Jul-2013	1-Aug-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Cell:</i> Change 0.5 cm to: 1 cm
GYMNEMA	<i>IDENTIFICATIO First N/B. Thin-Layer Supplement to Chromatograph USP36–NF31 y/ Chromatographi c system</i>	5880	26-Jul-2013	1-Aug-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 2 of <i>Adsorbent:</i> Change 5m to: 5 µm
POWDERED STINGING NETTLE EXTRACT	<i>COMPOSITION /Content of Total Amino Acids</i>	1608	26-Jul-2013	1-Aug-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Reagent solution:</i> Change Solution

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
OXCARBAZEPIIM NE	PUR ITIES/ <i>Organic Impurities, Procedure 1</i>	6035	26-Jul-2013	1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	containing 1.00 g of ninhydrin, 1.50 g of hydrindantin, to: Solution containing 1.00 g of ninhydrin, 150 mg of hydrindantin, Row 4 of Column 1 of Table 1: Change Dibenzazepinone ^b to: Oxcarbazepine related compound E AND Delete footnote b AND Reletter the following footnotes in both the table and footnote definitions: c to b

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
CALCIUM CARBONATE	IMPURITIES	USP36–NF31	2747	26-Jul-2013		1-Aug-2013	USP38–NF33	First Supplement to USP37–NF32	d to c e to d Line 1 of Acceptance criteria in <i>Limit of Fluoride</i> : Change 50 ppm to: NMT 50 ppm AND Line 1 of Acceptance criteria in <i>Mercury, Method IIa</i> <261>: Change 0.5 ppm to: NMT 0.5 ppm
CRYOPRESERVED HUMAN FIBROBLAST-DERIVED DERMAL SUBSTITUTE	<i>Total collagen content</i>	USP36–NF31	3155	26-Jul-2013		1-Aug-2013	USP38–NF33	First Supplement to USP37–NF32	Line 7 of <i>Collagen calibration standards</i> : Change by adding 25 mL, 50 mL, 100 mL, and 200 mL, to: by adding 25

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
EDETATE DISODIUM	ASSAY/ <i>Procedure</i>	USP36–NF31	3370	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	?L, 50 ?L, 100 ?L, and 200 ?L, Line 16 of <i>Analysis:</i> Change Calculate the percentage of edetate disodium to: Calculate the weight of edetate disodium AND Line 19 of <i>Analysis:</i> Change Result = $(V_T/V_U) \times W \times (M_{r1}/M_{r2}) \times 100$ to: Result = $(V_T/V_U) \times W \times (M_{r1}/M_{r2})$
LEVOFLOXACIN	ADDITIONAL REQUIREMENT S/USP <i>Reference Standards <11></i>	USP36–NF31	4099	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 2 of USP Levofloxacin Related Compound A RS: Change (S

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>)-9-Fluoro-3-methyl-10-(piperazin-1-yl)-7-oxo-2,3-dihydro-7H-pyrido[1,2,3-de][1,4-benzoxazine-6-carboxylic acid.</p> <p>to:</p> <p>(S)-9-Fluoro-3-methyl-10-(piperazin-1-yl)-7-oxo-2,3-dihydro-7H-pyrido[1,2,3-de][1,4]benzoxazine-6-carboxylic acid.</p> <p>AND</p> <p>Line 2 of USP Levofloxacin Related Compound B RS: Change (S</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
)-9,10-Difluoro-3-methyl-7-oxo-2,3-di hydro-7H -p yrido [1,2,3- <i>de</i>] [1,4-benzoxazine-6-carboxylic acid. to: (S)-9,10-Difluoro-3-methyl-7-oxo-2,3-di hydro-7H -p yrido [1,2,3- <i>de</i>] [1,4]benzoxazine-6-carboxylic acid.
VENLAFAXINE ADDITIONAL R HYDROCHLOR EQUIREMENT IDE	USP36–NF31 S/USP Reference Standards <11>	5551	26-Jul-2013	1-Aug-2013	USP38–NF33	First Supplement to USP37–NF32	Line 2 of Venlafaxine Related Compound A RS: Change 1-(1-(4-methoxy phenyl)-2-(meth

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
									ylamino)ethyl)cyclohexanol. C ₁₆ H ₂₅ NO ₂ 263.38 to: 1-(1-(4-Methoxyphenyl)-2-(methylamino)ethyl)cyclohexanol hydrochloride. C ₁₆ H ₂₅ NO ₂ · HCl 299.84
STINGING NETTLE	COMPOSITION <i>/Content of Total Amino Acids</i>	USP36–NF31	1604	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Reagent solution</i> : Change Solution containing 1.00 g of ninhydrin, 1.50 g of hydrindantin, to: Solution containing 1.00 g of ninhydrin, 150 mg of hydrindantin,
GENTAMICIN SULFATE	IM PURITIES/ <i>Limit of Methanol</i>	<i>First Supplement to USP36–NF31</i>	5990	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 11 of <i>Analysis</i> : Change

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>Result = $(R_U/R_S) \times (C_S/C_U) \times D \times F \times 100$ to: Result = $(R_U/R_S) \times (C_S/C_U) \times D \times F$ AND Line 18 of <i>Analysis:</i> Change $C_S =$ percentage of methanol in the <i>Standard solution</i> (% v/v) to: $C_S =$ percentage of methanol in the <i>Standard solution</i> AND Line 23 of <i>Analysis:</i> Change $F =$ conversion factor, 0.001 g/mg to:</p>

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
AMOXICILLIN	IM PUR ITIES/ <i>Organic Impurities/Procedure</i>	USP36–NF31	2477	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	<i>F = conversion factor, 1000 mg/g</i> Line 2 of <i>Acceptance criteria</i> : Change [Note—The reporting limit is 0.03% of the amoxicillin peak from the <i>Standard solution</i> .] to: [Note—The reporting limit is 0.03 times the amoxicillin peak from the <i>Standard solution</i> .]
CLARITHROMYCIN EXTENDED-RELEASE TABLETS	PERFORMANCE TESTS/ <i>Dissolution <711>/Test 4</i>	USP36–NF31	3019	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 2 of <i>Standard solution</i> : Change and <i>Medium</i> (96:4). to: and <i>Medium</i> (4:96).
DIPHENHYDR	<i>Assay</i>	USP36–NF31	3277	26-Jul-2013		1-Aug-2013	USP38–NF33	<i>First</i>	Line 2: Change

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
AMINE HYDRO CHLORIDE ORAL SOLUTION						<i>Supplement to USP37–NF32</i>	<i>Mobile phase, Standard preparation, System suitability solution, and Chromatographic system—Prepare as directed in the Assay under Diphenhydramine Hydrochloride. to: Mobile phase—Prepare a solution of acetonitrile, water, and triethylamine (50: 50: 0.5), adjust with glacial acetic acid to a pH of 6.5, filter, and degas. Make adjustments if necessary (see System Suitability under</i>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p><i>Chromatography</i> <621>). <i>Standard preparation</i> —Dissolve an accurately weighed quantity of USP Diphenhydramine Hydrochloride RS in water to obtain a solution having a known concentration of about 0.5 mg per mL.</p> <p>AND</p> <p><i>After the Assay preparation subsection: Add System suitability solution</i> —Dissolve about 5 mg of benzophenone in 5 mL of acetonitrile,</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p>dilute with water to 100 mL, and mix. Transfer 1.0 mL of this solution and 5 mg of diphenhydramine hydrochloride to a 10-mL volumetric flask, dilute with water to volume, and mix.</p> <p><i>Chromatographic system (see Chromatography <621>)</i>—The liquid chromatograph is equipped with a 254-nm detector and a 4.6-mm x 25-cm column that contains packing L10. The flow rate is about 1 mL per minute.</p> <p><i>Chromatograph the System</i></p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
							<p><i>suitability solution</i>, and record the peak responses as directed for <i>Procedure</i>; the resolution, <i>R</i>, between the benzophenone and diphenhydramine peaks is not less than 2.0. Chromatograph replicate injections of the <i>Standard preparation</i>, and record the peak responses as directed for <i>Procedure</i>; the relative standard deviation is not more than 2.0%, and the tailing factor for the diphenhydramine hydrochloride</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
FLUPHENAZIN Assay	USP36–NF31	3639	26-Jul-2013	1-Aug-2013	USP38–NF33	First	<p>peak is not more than 2.0. AND Line 1 of <i>Procedure:</i> Change Proceed as directed for <i>Procedure</i> in the Assay under <i>Diphenhydramine Hydrochloride.</i> to: Separately inject equal volumes, about 10 µL, of the <i>Standard preparation</i> and the Assay <i>preparation</i> into the chromatograph, record the chromatograms, and measure the responses for the major peaks. Line 11 of</p>

Monograph Title Section	Source Publication	Page Number	Errata Post Date Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
E DECANOATE INJECTION						<i>Supplement to USP37–NF32</i>	<i>Standard preparation: Delete (1:5) AND Line 8 of Assay preparation: Delete (1:5)</i>
OLMESARTAN IM MEDOXOMIL PURITIES/ <i>Organic Impurities/Impurity Table</i>	<i>USP36–NF31</i>	4570	26-Jul-2013	1-Aug-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Footnote d: Change ((5-Methyl-2-oxo-1,3-dioxol-4-yl)methyl 4-(2-hydroxypropan-2-yl)-2-propyl-1-((2'-(1-trityl-1H-tetrazol-5-yl)biphenyl-4-yl)methyl)-1H-imidazole-5-carboxylate. to: (5-Methyl-2-oxo-1,3-dioxol-4-yl)methyl 4-(2-hydroxypropan-2-yl)-2-propyl-1-((2'-(2-trityl-1H

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Sort ascending	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
									-tetrazol-5-yl)biphenyl-4-yl)methyl)-1 <i>H</i> -imidazole-5-carboxylate.

Pagination

- [First page « First](#)
- [Previous page ‹ Previous](#)
- ...
- [Page 32](#)
- [Page 33](#)
- [Page 34](#)
- [Page 35](#)
- [Page 36](#)
- [Page 37](#)
- [Page 38](#)
- [Page 39](#)
- [Page 40](#)
- [Next page Next ›](#)
- [Last page Last »](#)