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Monograph Title	Section	Source	Page Number	Errata Post	Errata Official	Target Errata	Target Online	Description
		Publication		Date	Date	Print Publication	Fix Publication	
THIMEROSAL	SPECIFIC	USP36–NF31	5369	31-May-2013	1-Jun-2013	USP37–NF32	USP37–NF32	Line 4 of

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TOPICAL AEROSOL	TESTS/ <i>Alcohol Determination, Method II</i>								<p><i>Analysis:</i> Change Determine the alcohol content of the sample thus prepared by the <i>Gas-Liquid Chromatographic Method (see Method II in Alcohol Determination <611>, using methyl ethyl ketone as the internal standard in place of acetone.</i></p> <p>to: Determine the alcohol content of the sample thus prepared by the <i>Gas Chromatographic Method (see Method II in Alcohol D</i></p>

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CARAWAY OIL	DEFINITION	USP36–NF31	1924	31-May-2013		1-Jun-2013	USP37–NF32	USP37–NF32	<p><i>ete rminat ion <611></i>), using methyl ethyl ketone as the internal standard in place of acetonitrile.</p> <p>Line 3: Change It contains NMT 50.0% of <i>d</i>-carvone (C₁₀H₁₄O). to: It contains NLT 50.0% of <i>d</i>-carvone (C₁₀H₁₄O).</p>
NORTRIPTYLIN NE HYDROCHLORIDE	IMPURITIES/Organic Impurities	<i>First Supplement to USP36–NF31</i>	6027	31-May-2013		1-Jun-2013	USP37–NF32	USP37–NF32	<p>Line 3 of <i>Acceptance criteria</i>: Change <i>Standard solution</i> to: <i>Sample solution</i></p> <p>Line 6: Change C₁₄H₁₅NO₂ to: C₁₄H₁₃NO₂</p>
FOSPHENYTOIN SODIUM INJECTION	USP Reference standards <11>	USP36–NF31	3680	31-May-2013		1-Jun-2013	USP37–NF32	USP37–NF32	<p>Line 6: Change C₁₄H₁₅NO₂ to: C₁₄H₁₃NO₂</p>
MINOCYCLINE	Assay	USP36–NF31	4378	31-May-2013		1-Jun-2013	USP37–NF32	USP37–NF32	<p>Line 2: Change</p>

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HYDROCHLORIDE TABLETS							<p><i>Mobile phase, Standard preparation, Resolution solution, and Chromatographic system</i></p> <p>—Proceed as directed in the Assay under <i>Minocycline Hydrochloride</i>.</p> <p>to:</p> <p><i>Mobile phase</i>—Prepare a mixture of 0.2 M ammonium oxalate, 0.01 M edetate disodium, dimethylformamide, and tetrahydrofuran (600:180:120:80). Adjust with ammonium hydroxide to a pH of 7.2, and pass through a filter of 0.5-μm</p>

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							<p>or finer pore size. Make adjustments if necessary (see <i>System Suitability</i> under <i>Chromatography</i> <621>). <i>Standard preparation</i></p> <p>—Dissolve an accurately weighed quantity of USP Minocycline Hydrochloride RS in water to obtain a solution having a known concentration of about 500 µg of minocycline (C₂₃H₂₇N₃O₇) per mL. Use this solution within 3 hours. <i>Resolution solution</i></p> <p>—Transfer 10</p>

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							<p>mg of USP Minocycline Hydrochloride RS to a 25-mL volumetric flask, add 20 mL of 0.2 M ammonium oxalate, and swirl to dissolve. Heat on a water bath at 60° for 180 minutes, and allow to cool. Dilute with water to volume, and mix.</p> <p><i>Chromatographic system (see Chromatography <621>)</i>—The liquid chromatograph is equipped with a 280-nm detector and a 4.6-mm x 25-cm column that contains</p>

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							<p>5-μm packing L1, and is maintained at a constant temperature of about 40°. The flow rate is about 1.5 mL per minute. Chromatograph the <i>Standard preparation</i>, and record the peak responses as directed for <i>Procedure</i>: the capacity factor, k', is not less than 5.0 and not more than 11.5; the tailing factor for the analyte peak is not less than 0.9 and not more than 2.0; and the relative standard deviation for replicate injections is not</p>

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							<p>more than 2.0%. Chromatograph the <i>Resolution solution</i>, and record the peak responses as directed for <i>Procedure</i>: the relative retention times are about 0.7 for epiminocycline and 1.0 for minocycline; and the resolution, <i>R</i>, between epiminocycline and minocycline is not less than 4.6.</p> <p>AND</p> <p>Line 1 of <i>Procedure</i>: Change Proceed as directed for <i>Procedure</i> in the Assay under</p>

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							<p><i>Minocycline Hydrochloride.</i></p> <p>to:</p> <p>Separately inject equal volumes (about 20 µL) of the <i>Standard preparation</i> and the <i>Assay preparation</i> into the chromatograph, record the chromatograms, and measure the responses for the major peaks.</p> <p>Line 3 of <i>System suitability.</i> Change [Note—The retention time for squalene is about 18 min; the relative retention times for tetradecane, hexadecane,</p>
HYDROGENAT ASSAY/ ED POLYDECENE	<i>Content of Decene Oligomer</i> USP36–NF31	2133	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	

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CLOTRIMAZOL ASSAY/ E AND BETAM ETHASONE DI PROPIONATE CREAM	USP36–NF31	3075	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	and squalene are about 0.5, 0.6, and 1.0, respectively.] to: [Note—The retention time for squalene is about 18 min; the relative retention times for tetradecane, hexadecane, and squalene are about 0.5, 0.6, and 1.0, respectively.] Line 3 of <i>Betamethasone dipropionate stock solution</i> : Change <i>J</i> being the ratio (in mg/g) of betamethasone to clotrimazole in the Cream to: <i>J</i> being the ratio of the labeled amount of

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CALCIUM SULFATE	SPECIFIC TESTS/ <i>Loss on Drying</i> <731>	USP35–NF30	1724	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	betamethasone (in mg/g) to the labeled amount of clotrimazole (in mg/g) in the Cream Line 1 of <i>Acceptance criteria</i> : Change NMT 1.5% for the anhydrous form and NMT 19.0%–23.0% for the dihydrate to: NMT 1.5% for the anhydrous form and 19.0%–23.0% for the dihydrate
TAPIOCA STARCH	<i>Limit of oxidizing substances</i>	USP35–NF30	1987	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 8: Change Add 1 mL of starch TS, and titrate with 0.002 N sodium thiosulfate VS to the disappearance of the starch–iodide color.

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LEVETIRACETAM	ADDITIONAL REQUIREMENTS	USP35–NF30	3659	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	to: Add 1 mL of starch TS, and titrate with 0.002 N sodium thiosulfate VS to the disappearance of the starch–iodine color. Line 9 of <i>USP Reference Standards</i> <11>: Change C ₈ H ₁₄ ClNO ₃ 207.65 to: C ₈ H ₁₅ ClN ₂ O ₂ 206.67
DULOXETINE DELAYED-RELEASE CAPSULES	PERFORMANCE TESTS/ <i>Dissolution <711>/ Chromatographic system</i>	<i>Second Supplement to USP35–NF30</i>	5940	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 1 of <i>Column</i> : Change 4.6-mm x 7.5-cm; 3-µm packing L7 to: 4.6-mm x 7.5-cm; 3- or 3.5-µm packing L7

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HYMETELLOS E	IM PUR ITIES/ <i>Chloride and Sulfate, Chloride <221></i>	USP36–NF31	2044	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Change the subsection title <i>Standard solution</i> to: <i>Control solution</i> AND Line 4 of <i>Analysis: Change Standard solution</i> to: <i>Control solution</i> AND Line 2 of <i>Acceptance criteria: Change Standard solution</i> to: <i>Control solution</i>
BENZTROPINE MESYLATE	CHEMICAL INFORMATION	USP36–NF31	2628	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 2: Change 8-Azabicyclo[3.2.1]octane, 3-(di phenylmethoxy) -, <i>endo</i> -, methanesulfonate; to: 8-Azabicyclo[3.2.1]octane, 3-(di

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LORAZEPAM TABLETS	IMPURITIES/Organic Impurities/System suitability/Suitability requirements	USP36–NF31	4153	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	phenylmethoxy)-N-methyl-, endo-, methane sulfonate; Line 1 of <i>Tailing factor</i> . Change 2.0, <i>Standard solution</i> to: NMT 2.0, <i>Standard solution</i>
BRINZOLAMIDE	Related compounds/Test 2	USP35–NF30	2385	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 15 of <i>Procedure</i> : Change relative retention time greater than 6. to: relative retention greater than 6.
VINORELBINE INJECTION	Assay	USP35–NF30	5028	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 1: Change <i>Phosphate buffer</i> , <i>Mobile phase</i> , and <i>System suitability solution</i> —Proceed as directed in the

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							<p>Assay under <i>Vinorelbine Tartrate</i>. to: <i>Phosphate buffer</i>—Dissolve 6.9 g of monobasic sodium phosphate in 900 mL of water. Adjust with phosphoric acid to a pH of 4.2, dilute with water to 1000 mL, and mix.</p> <p><i>Mobile phase</i>—Dissolve 1.22 g of sodium 1-decanesulfonate in 620 mL of methanol. Add 380 mL of <i>Phosphate buffer</i>, mix, filter, and degas. Make adjustments if necessary (see</p>

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							<p><i>System Suitability</i> under <i>Chromatography</i> <621>).</p> <p><i>System suitability solution</i></p> <p>—Dissolve accurately weighed quantities of USP Vinorelbine Tartrate RS and USP Vinorelbine Related Compound A RS in water, and dilute quantitatively, and stepwise if necessary, with water to obtain a solution having known concentrations of about 1.4 mg per mL and 0.01 mg per mL,</p>

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POWDERED BLACK PEPPER EXTRACT	DEFINITION	USP36–NF31	1365	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	<p>respectively. Expose a portion of this solution in a suitable xenon lamp apparatus capable of supplying a dose of 1600 KJ/m² between 310 and 800 nm at a power of 500 W/m² for about 1 h, in order to generate an additional degradation product 3,6-epoxy vinorelbine having a relative retention time of about 0.8.</p> <p>Line 5: Change It contains NLT 90.0% and NMT 110.0% of the labeled amount of piperine.</p>

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POLYVINYL ACETATE PHTHALATE	IM PURITIES/ <i>Free Phthalic Acid</i>	USP36–NF31	2168	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	to: It contains NLT 90.0% and NMT 110.0% of the labeled amount of piperine, calculated on the dried basis. Line 1 of <i>Sample solution</i> : Change 6 mg/mL of polyvinyl acetate to: 6 mg/mL of polyvinyl acetate phthalate
DILTIAZEM HYDROCHLORIDE ORAL SUSPENSION	ASSAY/ <i>Procedure</i>	USP36–NF31	3263	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 6 of <i>Sample solution</i> : Change Pipet 1.0 mL of the sample solution to: Pipet 1.0 mL of the sample
IFOSFAMIDE	<i>Chloroform-</i>	USP35–NF30	3477	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 18 of <i>Test</i>

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									<i>preparation:</i> Change ammonium hydroxide solution. to: ammonium hydroxide.
ATROPINE SULFATE TABLETS	Assay	USP35–NF30	2272	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 9 of <i>Procedure:</i> Change R_U and R_S are as defined therein. to: R_U and R_S are the peak area ratios of atropine to homatropine.
VANCOMYCIN INJECTION	SPECIFIC TESTS/ <i>Composition of Vancomycin</i>	USP35–NF30	5003	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 16 of <i>Analysis:</i> Change D = dilution factor, <i>Sample stock solution</i> to <i>Sample solution</i> , 25 to: D = dilution factor, <i>Sample</i>

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									<p><i>stock solution to Sample solution AND</i></p> <p>Line 29 of <i>Analysis:</i> Change <i>D</i> = dilution factor, <i>Sample stock solution to Sample solution</i>, 25 to: <i>D</i> = dilution factor, <i>Sample stock solution to Sample solution</i></p>
SUMATRIPTAN INJECTION	SPECIFIC TESTS/ <i>Osmolality and Osmolarity <785></i>	<i>Second Supplement to USP35–NF30</i>	5996	29-Mar-2013		1-Apr-2013	<i>USP37–NF32</i>	<i>USP37–NF32</i>	<p>Line 1: Change 270–330 mOsmol to: 270–330 mOsmol/kg</p>
INOSITOL	SPECIFIC TESTS/ <i>Conductivity</i>	<i>USP36–NF31</i>	2049	29-Mar-2013		1-Apr-2013	<i>USP37–NF32</i>	<i>USP37–NF32</i>	<p>Line 1 of <i>Sample solution:</i> Change Transfer 10.0 g of Inositol, weighed and calculated on the dried basis,</p>

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CLONAZEPAM ASSAY/ ORAL <i>Procedure</i> SUSPENSION	USP36–NF31	3053	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	to a 50-mL volumetric flask, and dissolve in and dilute with water (previously boiled and cooled to room temperature) to volume. to: 0.2 g/mL of Inositol in water (previously boiled and cooled to room temperature). Line 6 of <i>Sample solution</i> : Change Pipet 2.5 mL of the <i>Sample solution</i> to: Pipet 2.5 mL of the sample
METHENAMIN ASSAY/ E ORAL <i>Procedure</i> SOLUTION	USP36–NF31	4288	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	Line 10 of <i>Analysis</i> : Change <i>B</i>

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AZITHROMYCI N	IM PUR ITIES/ <i>Organic I mpurities/Proce dure 2</i>	USP35–NF30 2279	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	<p>$S =$ absorbance of the <i>Sample blank</i> to: $B_S =$ absorbance of the <i>Standard blank</i></p> <p>Line 15 of <i>Analysis:</i> Change $C_S =$ concentration of USP Azithromycin RS in the <i>Standard solution</i> ($\mu\text{g/mL}$) to: $C_S =$ concentration of USP Azithromycin RS in the <i>Standard solution</i> (mg/mL) AND Add after C_U: $P =$ potency of USP</p>

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ATRACURIUM BESYLATE INJECTION	IM PUR ITIES/ <i>Organic Impurities/Acceptance criteria/</i> Table 2	<i>Second Supplement to USP35–NF30</i>	5909	29-Mar-2013		1-Apr-2013	<i>USP37–NF32</i>	<i>USP37–NF32</i>	Azithromycin RS (µg/mg of azithromycin) Footnote b: Change <i>cis</i> isomer of the hydroxy compound. to: <i>trans</i> isomer of the hydroxy compound. AND Footnote c: Change <i>trans</i> isomer of the hydroxy compound. to: <i>cis</i> isomer of the hydroxy compound. AND Footnote d: Change <i>cis</i> isomer of the monoacrylate. to: <i>trans</i> isomer of the

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ZINC SULFATE TABLETS	Identification/B. <i>Zinc</i> <i>USP35–NF30</i>	5077	29-Mar-2013	1-Apr-2013	<i>USP37–NF32</i>	<i>USP37–NF32</i>	monoacrylate. AND Footnote e: Change <i>trans</i> isomer of the monoacrylate. to: <i>cis</i> isomer of the monoacrylate. Line 1 of <i>Sodium hydroxide solution:</i> Change 42 mg/mL of sodium hydroxide to: 420 mg/mL of sodium hydroxide
DIETHYL SEBACATE	DEFINITION <i>USP36–NF31</i>	1994	29-Mar-2013	1-Apr-2013	<i>USP37–NF32</i>	<i>USP37–NF32</i>	Line 2: Change Diethyl Sebacate consists of the diester of alcohol and sebacic acid. to:

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PROPYLENE GLYCOL MONOLAURATE	IM PURITIES/ <i>Limit of Propylene Glycol</i>	USP36–NF31	2180	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Diethyl Sebacate consists of the diester of alcohol (ethanol) and sebacic acid. Line 8 of <i>Analysis</i> : Change Calculate the percentage of free propylene glycol in the portion of Propylene Glycol Monocaprylate taken: to: Calculate the percentage of free propylene glycol in the portion of Propylene Glycol Monolaurate taken: Line 5 of <i>Analysis</i> :
GLUCONOLAC TONE	IDENTIFICATIO N/A.	USP36–NF31	3742	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	

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METHOTREXATE IM PUR ITIES/ <i>Organic Impurities/Procedure 1: Related Compounds</i>	USP35–NF30	3855	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	Change crystals of the phenylhydrazine of gluconic acid to: crystals of the phenylhydrazide of gluconic acid Footnote b of <i>Impurity Table 1</i> : Change (S)-2-{4-[(2-Amino-4-oxo-1,4-dihydropteridin-6-yl)methylamino]-N-methylbenzamide}pentanedioic acid. to: (S)-2-(4-[(2-Amino-4-oxo-1,4-dihydropteridin-6-yl)methylamino]benzamide)pentanedioic acid.

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BETAMETHAS ONE ORAL SOLUTION	<i>Identification/A:</i> USP35–NF30	2336	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	acid. Line 1: Change A: to: <i>A: Thin-Layer Chromatographic Identification Test <201></i> —
VINORELBINE INJECTION	<i>Related compounds</i> USP35–NF30	5028	29-Mar-2013	1-Apr-2013	USP37–NF32	USP37–NF32	Delete the subsection <i>Standard solution</i> and <i>Diluted standard solution</i> . Replace with: <i>Standard solution</i> —Dissolve an accurately weighed quantity of USP Vinorelbine Tartrate RS in <i>Mobile phase</i> to obtain a solution having a known concentration of about 1.4 mg per mL.

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							<p><i>Diluted standard solution</i> —Transfer 1.0 mL of the <i>Standard solution</i> to a 50-mL volumetric flask, and dilute with <i>Mobile phase</i> to volume. Pipet 1.0 mL of this solution into a 100-mL volumetric flask, and dilute with <i>Mobile phase</i> to volume.</p> <p>AND</p> <p>Line 1 of <i>Procedure</i>: Change Proceed as directed for <i>Procedure</i> in the test for <i>Related compounds</i> under <i>Vinorelbine</i></p>


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							<p><i>Tartrate.</i> to: Separately inject equal volumes (about 20 µL) of the <i>Test solution</i> and the <i>Diluted standard solution</i> into the chromatograph, record the chromatograms, and measure the areas for the major peaks. Record the chromatograms for three times the retention time of the vinorelbine peak. Disregard any peaks with an area less than or equal to one-half of the area of the peak obtained for vinorelbine in</p>

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									<p>the <i>Diluted standard solution</i>. Calculate the percentage of each impurity in the portion of Injection taken by the formula: $100(r_i/r_s)$ in which r_i is the peak response for each impurity obtained from the <i>Test solution</i>; and r_s is the sum of the responses of all the peaks.</p>
GOOD STORAGE AND DISTRIBUTION PRACTICES FOR DRUG PRODUCTS	QUALITY MANAGEMENT SYSTEM/Storage Management System/Receiving and Transferring Drug Products	USP36–NF31	693	29-Mar-2013		1-Apr-2013	USP37–NF32	USP37–NF32	Line 1 of footnote 1: Change JP Edmond, to: JP Emond,
ACESULFAME POTASSIUM	IMPURITIES/Limit	USP35–NF30	1680	31-Jan-2013		1-Feb-2013	USP37–NF32	Second Supplement to	Line 25 of Analysis:

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								USP36–NF31	Change C = concentration of fluoride in the <i>Sample solution</i> , from the standard curve (mg/mL) to: C = concentration of fluoride in the <i>Sample solution</i> , from the standard curve (µg/mL)
AMANTADINE HYDROCHLORIDE CAPSULES	PERFORMANCE TESTS/ <i>Dissolution</i> <711>/Test 2	USP35–NF30	2153	31-Jan-2013		1-Feb-2013	USP37–NF32	Second Supplement to USP36–NF31	Line 5 of <i>Chromatographic system</i> : Change Column: 0.32-mm x 30-cm, 0.25-µm film, phase G1 to: Column: 0.32-mm x 30-m, 0.25-µm film, phase G1
OXYBUTYNIN CHLORIDE EX E	PERFORMANCE	USP35–NF30	4167	31-Jan-2013		1-Feb-2013	USP37–NF32	Second Supplement to	Line 3 of <i>Working</i>

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TENDED-RELEASE TABLETS	TESTS/ <i>Dissolution</i> <711>/Test 2							USP36–NF31	<i>standard solution:</i> Change or transfer 10 mL for Tablets labeled to contain 10 mg, to a 100-mL volumetric flask. to: transfer 10 mL for Tablets labeled to contain 10 mg, or transfer 15 mL for Tablets labeled to contain 15 mg to a 100-mL volumetric flask.
ALLANTOIN	IDENTIFICATION	<i>First Supplement to USP35–NF30</i>	5429	31-Jan-2013		1-Feb-2013	USP37–NF32	<i>Second Supplement to USP36–NF31</i>	Line 1 of <i>B. Thin-Layer Chromatographic Identification Test <201>:</i> Change The R_F value of the principal spot from <i>Sample solution A</i> corresponds

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CAPTOPRIL ORAL SUSPENSION	Assay <i>USP35–NF30</i>	2477	31-Jan-2013	1-Feb-2013	<i>USP37–NF32</i>	<i>Second Supplement to USP36–NF31</i>	<p>to that from <i>Standard solution A</i>, as described in the test for <i>Organic Impurities</i>.</p> <p>to:</p> <p>The R_F value of the principal spot from <i>Sample solution B</i> corresponds to that from <i>Standard solution A</i>, as described in the test for <i>Organic Impurities</i>.</p> <p>Line 1 of <i>Mobile phase</i>: Change Prepare a filtered and degassed mixture of methanol and water (11:9) containing 0.5 mL of phosphoric acid.</p> <p>to:</p>

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CHROMATOGRAPHY SYSTEMS UIT ABILITY/ <i>Stationary Phase</i>	USP35–NF30	258	31-Jan-2013	1-Feb-2013	USP37–NF32	Second Supplement to USP36–NF31	<p>Methanol and water (55:45) containing 0.5 mL/L of phosphoric acid. Filter, and degas.</p> <p>Line 3 of <i>Flow Rate (HPLC)</i>: Change </p> <p>in which F_1 is the flow rate indicated in the monograph, in mL/min; F_2 is the adjusted flow rate, in mL/min; l_1 is the length of the column indicated in the monograph; l_2 is the length of the column used; d_1 is the column inner diameter indicated in the monograph; and d_2 is the</p>

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LOW-SUBSTIT Assay	USP35–NF30	1822	31-Jan-2013	1-Feb-2013	USP37–NF32	Second	<p>internal diameter of the column used. Additionally, the flow rate can be adjusted by $\pm 50\%$.</p> <p>to:</p> <p>⊗</p> <p>in which F_1 is the flow rate indicated in the monograph, in mL/min; F_2 is the adjusted flow rate, in mL/min; d_1 is the column inner diameter indicated in the monograph; and d_2 is the internal diameter of the column used. Additionally, the flow rate can be adjusted by $\pm 50\%$.</p> <p>Line 2: Change</p>

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UTED HYDROXYPROPYL CELLULOSE								<i>Supplement to USP36–NF31</i>	<i>Hypromellose 2906, except to substitute Low-Substituted Hydroxypropyl Cellulose for Hypromellose 2906 throughout. to: Hypromellose, except to substitute Low-Substituted Hydroxypropyl Cellulose for Hypromellose throughout.</i>
NORGESTIMATE	<i>Limit of residual solvents</i>	<i>USP35–NF30</i>	4083	31-Jan-2013		1-Feb-2013	<i>USP37–NF32</i>	<i>Second Supplement to USP36–NF31</i>	Line 1 of <i>Limit of residual solvents</i> : Change to: <i>Limit of residual solvents <467></i>
VINCRISTINE SULFATE	IMPURITIES/ <i>Organic Impurities</i>	<i>USP35–NF30</i>	5022	31-Jan-2013		1-Feb-2013	<i>USP37–NF32</i>	<i>Second Supplement to USP36–NF31</i>	Line 5 of <i>Analysis</i> : Change Result = $[r_{UA} / (r_{UA} + 25r$

Monograph Title	Section	Source Publication	Page Number	Errata Post Date	Errata Official Date	Target Errata Print Publication	Target Online Fix Publication	Description
								$\frac{r_{UB}}{r_{UA} + 30r_{UB}} \times 100$ to: Result = $\frac{r_{UA}}{r_{UA} + 30r_{UB}} \times 100$ AND Change line 12 of <i>Analysis</i> : Result = $\frac{r_{UA}}{r_{UA} + 25r_{UB}} \times 100$ to: Result = $\frac{r_{UA}}{r_{UA} + 30r_{UB}} \times 100$
ADAPALENE	IM PUR ITIES/ <i>Residual Solvent: Limit of 2012)</i> <i>Triethylamine</i>	<i>Revision Bulletin (Official December 01,</i>	Online	31-Jan-2013	1-Feb-2013	<i>USP37–NF32</i>	<i>Second Supplement to USP36–NF31</i>	Line 1 of <i>Diluent</i> : Change Dimethyl sulfoxide and 1 N sodium hydroxide solution (4:1) to: Dimethyl sulfoxide AND Line 1 of <i>Standard solution</i> : Change 3.2 ?g/mL of

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							<p>USP Triethylamine RS in <i>Diluent</i> to: 4.0 ?g/mL of USP Triethylamine RS in <i>Diluent</i>. Transfer 4.0 mL of this solution to a 20-mL headspace vial, and add 1.0 mL of 1 N NaOH solution. AND Line 1 of <i>Sample solution</i>: Change 40 mg/mL of Adapalene in <i>Diluent</i> to: 50 mg/mL of Adapalene in <i>Diluent</i>. Transfer 4.0 mL of this solution to a 20-mL headspace vial,</p>

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GLYCERYL BEHENATE	IMPURITIES	USP35–NF30	1811	31-Jan-2013		1-Feb-2013	USP37–NF32	Second Supplement to USP36–NF31	<p>and add 1.0 mL of 1 N NaOH solution.</p> <p>Line 34 of <i>Content of 1-Monoglycerides/Analysis:</i> Change <i>F</i> = equivalency factor of glyceryl monobehenate, 207.3 mg/mEq to: <i>F</i> = equivalency factor of glyceryl monobehenate, 0.2073 g/mEq AND Line 19 of <i>Limit of Free Glycerin/Analysis:</i> Change <i>F</i> = equivalency factor of glycerin, 23.0 mg/mEq to: <i>F</i> = equivalency</p>

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CAPTOPRIL ORAL SOLUTION	Assay	USP35–NF30	2477	31-Jan-2013		1-Feb-2013	USP37–NF32	Second Supplement to USP36–NF31	factor of glycerin, 0.023 g/mEq Line 1 of <i>Mobile phase</i> : Change Prepare a filtered and degassed mixture of methanol and water (11:9) containing 0.5 mL of phosphoric acid. to: Methanol and water (55:45) containing 0.5 mL/L of phosphoric acid. Filter, and degas.
PHENYLALANINE IMPURITIES		USP35–NF30	4296	31-Jan-2013		1-Feb-2013	USP37–NF32	Second Supplement to USP36–NF31	Line 1 of <i>Heavy Metals, Method I</i> <231>: Change <i>Method I</i> to: <i>Method II</i>
ELEMENTAL I	Drug	Second	5633	31-Jan-2013		1-Feb-2013	USP37–NF32	Second	Row 16 of

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MPURITIES--LI MITS	<i>Products/Large Volume Parenterals</i>	<i>Supplement to USP35–NF30</i>						<i>Supplement to USP36–NF31</i>	Column 4 of <i>Table 1:</i> Change 70 to: 100 AND Row 16 of Column 5 of <i>Table 1:</i> Change 25 to: 10

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