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CAPSICUM	SPECIFIC	<i>Second</i>	6577	22-Nov-2013	1-Dec-2013	<i>USP38–NF33</i>	<i>Second</i>	Line 1 of

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OLEORESIN	TESTS/ <i>Limit of Nonivamide</i>	<i>Supplement to USP36–NF31</i>						<i>Supplement to USP37–NF32</i>	<i>Acceptance criteria: Change on the dried basis to: on the anhydrous basis</i>
RISPERIDONE ORAL SOLUTION	ADDITIONAL REQUIREMENT S/USP Reference Standards <11>	<i>Second Supplement to USP36–NF31</i>	6690	22-Nov-2013		1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	Line 2 of USP Risperidone Related Compounds Mixture RS: Change Contains a 98.9: 0.5: 0.30: 0.3 (area %) mixture of four compounds: 98.9% of <i>Risperidone</i> . 0.5% of <i>Risperidone cis-N-oxide: cis -3-[2-[4-(6-Fluoro-1,2-benzisoxazol-3-yl)-1-piperidinyl]ethyl]-6,7, 8,9-tetrahydro-2-methyl-4H</i>

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							<p>-pyri do[1,2-a]pyrimidin-4-one]. 0.3% of <i>Bicyclorisperido ne</i>: 3-(4-Fluoro- 2-hydroxypheny l)-1-[2-(6,7,8,9-t etrahydro-2-met hyl-4 -oxo-4<i>H</i> -pyr ido-[1,2 -a]pyrimidin-3-yl)e thyl]-1-aza-2-az oniabicyclo[2.2. 2]oct-2-ene iodide. 0.3% of <i>Z</i>- <i>Oxime</i>: (<i>Z</i>)-3-[2-[4-(2,4-Dif luorophenyl) (hy droxyimino)met hyl]-1-piperidiny l]ethyl]-6,7,8,9-t etrahydro-2-met hyl-4<i>H</i> -pyri</p>

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							<p>do[1,2-a]pyrimidin-4-one</p> <p>.</p> <p>to:</p> <p>Contains a mixture of the following four compounds:</p> <p>98.9% of <i>Risperidone</i>.</p> <p>0.5% of <i>Risperidone cis-N-oxide: cis</i>-3-[2-[4-(6-Fluoro-1,2-benzisoxazol-3-yl)-1-piperidinyl]ethyl]-6,7,8,9-tetrahydro-2-methyl-4H-pyrido[1,2-a]pyrimidin-4-one-N-oxide.</p> <p>0.3% of <i>Bicyclorisperidone: 3-(4-Fluoro-2-hydroxyphenyl)-1-[2-(6,7,8,9-tetrahydro-2-me</i></p>

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									<p>thyl- 4-oxo-4H -pyr ido-[1,2 -a]pyrimidin-3-yl)e thyl]-1-aza-2-az oniabicyclo[2.2. 2]oct-2-ene iodide. 0.3% of Z- Oxime: (Z)-3-[2-[4-(2,4-Dif luorophenyl) (hy droxyimino)met hyl]-1-piperidiny l]ethyl]-6,7,8,9-t etrahydro-2-met hyl-4H -pyri do[1,2-a]pyrimidin-4-one .</p>
BOSWELLIA SERRATA	COMPOSITION /Content of Keto- Derivatives of ?-Boswellic Acids/System s uitability/Suitabil ity requirements	USP36–NF31	1366	22-Nov-2013		1-Dec-2013	USP38–NF33	Second Supplement to USP37–NF32	Line 1 of <i>Tailing factor</i> . Change 11-keto-?-boswellic acid peak to: 3-acetyl-11-ket

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APROTININ	<i>Limit of des-Ala-aprotinin and des-Ala-des-Gly-aprotinin</i>	USP36–NF31	2522	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	o-?-boswellic acid peak Change the subsection head <i>Capillary zone electrophoresis system (see Capillary Electrophoresis under Biotechnology-Derived Articles—Test <1047>)</i> — to: <i>Capillary zone electrophoresis system—</i>
KETOROLAC T IM ROMETHAMIN PUR E TABLETS	<i>ITIES/Organic Impurities</i>	USP36–NF31	4042	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	Line 1 of <i>Sample solution:</i> Change Proceed as directed for the <i>Sample stock solution</i> in the Assay. to: Proceed as directed for the

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RISPERIDONE <i>USP Reference standards <11></i>	<i>USP36–NF31</i>	5063	22-Nov-2013	1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	<p><i>Sample solution in the Assay.</i></p> <p>Line 7 of USP Risperidone System Suitability Mixture RS: Change 9-Hydroxyrisperidone-(6RS)-3-[2-[4-(6-fluoro-1,2-benzisoxazol-3-yl)piperidin-1-yl]ethyl]-2,6-dimethyl-6,7,8,9-tetrahydro-4H-pyrido[1,2-a]pyrimidin-4-one .</p> <p>to:</p> <p>9-Hydroxyrisperidone: (9RS)-3-{2-[4-(6-fluoro-1,2-benzisoxazol-3-yl)piperidin-1-yl]ethyl}-9-h</p>

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CHINESE SALVIA	COMPOSITION	<i>Second Supplement to Salviolic Acid USP36–NF31 B</i>	6331	22-Nov-2013		1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	ydroxy-2-methyl-6,7,8,9-tetrahydro-4H-pyrido[1,2-a]pyrimidin-4-one . Line 16 of Analysis: Change <i>W</i> = weight of Chinese Salvia used to prepare the <i>Sample solution</i> (mg) to: <i>W</i> = weight of Chinese Salvia used to prepare the <i>Sample stock solution</i> (mg)
LORATADINE	IMPURITIES/Organic Impurities, Procedure 1	<i>Second Supplement to USP36–NF31</i>	6650	22-Nov-2013		1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	Line 3 of Note: Change 4,8-dichloro-6,11-dihydro-5H-benzo[5,6]cyclohepta[1,2-b]pyridin-11-one

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POLYETHYLENE GLYCOL MONOMETHYL ETHER	IM PURITIES/ <i>Limit of 2-Methoxyethanol</i>	USP36–NF31	2142	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	to: 4,8-dichloro-5,6-dihydro-11H-benzo[5,6]cyclohepta[1,2-b]pyridin-11-one Line 11 of <i>Calibration</i> : Change On the two <i>Calibration</i> plots, to: On the <i>Calibration</i> plot, Line 2: Change gadoteridol. to: Gadoteridol Injection. Line 3 of <i>Chromatographic system</i> : Change a 0.32-mm × 25-m column that contains coating G1, to:
GADOTERIDOL INJECTION	<i>Bacterial endotoxins <85></i>	USP36–NF31	3701	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	to: Gadoteridol Injection. Line 3 of <i>Chromatographic system</i> : Change a 0.32-mm × 25-m column that contains coating G1, to:
OCTINOXATE	<i>Assay</i>	USP36–NF31	4556	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	to: Gadoteridol Injection. Line 3 of <i>Chromatographic system</i> : Change a 0.32-mm × 25-m column that contains coating G1, to:

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ISOTRETINOIN CAPSULES	PERFORMANCE TESTS/ Dissolution <711>/Test 4	<i>First Supplement to USP36–NF31</i>	6000	22-Nov-2013		1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	a 0.32-mm x 25-m column with 0.25-µm thickness of phase G1 coating, Line 13 of <i>System suitability</i> . Add section heads before "Calculate the percentage....": <i>Analysis Samples: Standard solution and Sample solution</i>
CYCLOBENZAPRINE HYDROCHLORIDE	ADDITIONAL REQUIREMENT S/USP Reference Standards <11>	<i>Second Supplement to USP36–NF31</i>	6585	22-Nov-2013		1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	Line 2 of USP C yclobenzaprine Related Compound A RS: Change 5-[3-(Dimethyl amino)propyl]-10,11-dihydro-5H-benz[<i>a,d</i>

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TRAMADOL HYASSAY/ DROCHLORID Procedure E	<i>Second Supplement to USP36–NF31</i>	6715	22-Nov-2013	1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>]cyclohepten-5-ol. to: 5-[3-(Dimethylamino)propyl]-5H-dibenzo[a,d]cyclohepten-5-ol. Line 2 of <i>System suitability solution</i> : Change USP Tramadol Hydrochloride Related Compound A RS to: USP Tramadol Related Compound A RS
BOSWELLIA SERRATA EXTRACT	COMPOSITION <i>USP36–NF31</i> <i>/Content of Keto-Derivatives of ?-Boswellic</i>	1367	22-Nov-2013	1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	Line 1 of <i>Tailing factor</i> . Change 11-keto-?-boswellic acid

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									peak to: 3-acetyl-11-keto- β -boswellic acid peak
CARVEDILOL	IM PURITIES/Organic Impurities, Procedure 3: Carvedilol Related Compound F	USP36–NF31	2822	22-Nov-2013		1-Dec-2013	USP38–NF33	Second Supplement to USP37–NF32	Line 2 of Sample solution: Change Use about 1.9 mL of Diluent per mg of the Carvedilol, and sonicate briefly to facilitate dissolution. to: Initially add Diluent to fill about 80% of the total volume. Sonicate briefly to facilitate dissolution. Cool, and dilute with Diluent to volume.
LEVALBUTEROL INHALATION	ADDITIONAL REQUIREMENT S/USP	USP36–NF31	4080	22-Nov-2013		1-Dec-2013	USP38–NF33	Second Supplement to USP37–NF32	Line 35: Delete USP Levalbuterol

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SOLUTION		<i>Reference Standards <11></i>							Related Compound H RS 4-[2-(<i>tert</i> -Butylamino)-1-methoxyethyl]-2-(hydroxymethyl)phenol. C ₁₄ H ₂₃ NO ₃ 253.34
RISPERIDONE ADDITIONAL R ORALLY DISINTEGRATING TABLETS		<i>USP36–NF31 S/USP Reference Standards <11></i>	5067	22-Nov-2013		1-Dec-2013	<i>USP38–NF33</i>	<i>Second Supplement to USP37–NF32</i>	Line 2 of USP Risperidone Related Compounds Mixture RS: Change Contains a 98.9/0.5/0.3/0.3 (area %) mixture of the following four compounds: Risperidone (98.9%) Risperidone <i>cis</i> - <i>N</i> -oxide (0.5%): <i>cis</i> -3-[2-[4-(6-fluoro-1,2-benzisoxazol-3-yl)-1-piperidinyl]ethyl]-6,7,

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							<p>8,9-tetrahydro-2-methyl-4H-pyrido[1,2-a]pyrimidin-4-one, N-oxide monohydrate. $C_{23}H_{29}FN_4O_4$ 444.50 Bicyclorisperidone (0.3%): 3-(4-fluoro-2-hydroxyphenyl)-1-[2-(6,7,8,9-tetrahydro-2-methyl-4-oxo-4H-pyrido[1,2-a]pyrimidin-3-yl)ethyl]-2-aza-1-azoniabicyclo[2.2.2]oct-2-ene iodide. $C_{23}H_{28}FIN_4O_4$ 538.40 Z-oxime (0.3%): (Z)-3-[2-[4-(2,4-Difluorophenyl)(hy</p>

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							<p>droxyimino)met hyl]-1-piperidiny l]ethyl]-6,7,8,9-t etrahydro-2-met hyl-4<i>H</i> -pyri do[1,2-<i>a</i>]pyrimidin-4-one . C₂₃H₂₈F₂N₄O₂ 430.29 to: Contains a mixture of four compounds: 98.9% of <i>Risperidone</i>. 0.5% of <i>Ris peridon ecis-N-oxide: cis</i> -3-[2-[4-(6-Fluor o-1,2-benzisoxa zol-3-yl)-1-piperi diny]ethyl]-6,7, 8,9-tetrahydro-2 -methy l-4<i>H</i> -pyri do[1,2-<i>a</i></p>

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							<p>]pyrimidin-4-one -<i>N</i>-oxide. 0.3% of <i>Bicyclorisperido</i> <i>ne</i>: 3-(4-Fluoro- 2-hydroxypheny l)-1-[2-(6,7,8,9-t etrahydro-2-met hyl-4 -oxo-4<i>H</i> -pyr ido-[1,2 -a]pyrimidin-3-yl)e thyl]-1-aza-2-az oniabicyclo[2.2. 2]oct-2-ene iodide. 0.3% of <i>Z</i>- <i>Oxime</i>: (<i>Z</i>)-3-[2-[4-(2,4-Dif luorophenyl)(hy droxyimino)met hyl]-1-piperidiny l]ethyl-6,7,8,9-te trahydro-2-meth yl-4<i>H</i> -pyri do[1,2-<i>a</i>]pyrimidin-4-one</p>

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VALERIAN TINCTURE	ST REN	<i>Second Supplement to GTH/Content of Valerenic Acids</i>	6352	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	Line 1 of <i>Acceptance criteria</i> : Change 90.0%–120.0% to: NLT 0.015% of valerenic acids, calculated as the sum of hydroxyvalerenic acid, acetoxyvalerenic acid, and valerenic acid
PARICALCITOLIM INJECTION	PUR ITIES/ <i>Organic Impurities/Chromatographic system/Columns</i>	<i>Second Supplement to USP36–NF31</i>	6678	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	Line 1 of <i>Guard</i> : Change 4.6-mm x 7.5-cm to: 4.6-mm x 7.5-mm
FERRIC AMMONIUM CITRATE	Mercury	USP36–NF31	2469	22-Nov-2013		1-Dec-2013	USP38–NF33	<i>Second Supplement to USP37–NF32</i>	Line 5 of <i>Standard solutions</i> : Change 2.5, 5.0, 10.0, and 35.0 µg to: 2.5, 5.0, 10.0,

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GRANISETRO N HYDROCHL ORIDE INJECTION	Assay	USP36–NF31	3772	22-Nov-2013		1-Dec-2013	USP38–NF33	Second Supplement to USP37–NF32	and 35.0 ng Line 10 of <i>Procedure:</i> Change 100(312.41 / 348.87)(<i>C</i> / <i>L</i>)(<i>r</i> _U / <i>r</i> _S) to: 100(312.41 / 348.87) (<i>C</i> / <i>C</i> _U)(<i>r</i> _U / <i>r</i> _S) AND Line of 14 of <i>Procedure:</i> Change <i>L</i> is as defined above; to: <i>C</i> _U is the nominal concentration, in mg per mL, of granisetron in the Assay <i>preparation</i> ; Change: and to: or
ETHYL OLEATE	SPECIFIC TESTS/ <i>Viscosity—Capillary Viscometer Methods <911> and Rotational</i>	USP36–NF31	2006	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Change: and to: or

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<p style="text-align: center;"><i>Rheometer Methods <912></i></p> <p>OXYMETAZOLINE HYDROCHLORIDE OPTHALMIC SOLUTION</p>	USP36–NF31	4652	27-Sep-2013	1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	<p>Line 1: Change A volume of Ophthalmic Solution, equivalent to about 2.5 mg of oxymetazoline hydrochloride, responds to the <i>Id</i> <i>entification</i> test under <i>Oxymetazoline Hydrochloride Nasal Solution</i>. to:</p> <p>Place a volume of Ophthalmic Solution, equivalent to about 2.5 mg of oxymetazoline hydrochloride, in a 60-mL separator, and add water to make about 10 mL. Add 2 mL of sodium</p>

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							carbonate solution (1 in 10), extract with 10 mL of chloroform, and transfer the chloroform extract to a second 60-mL separator. Extract the chloroform solution with 10 mL of 0.1 N hydrochloric acid, allow to separate, and discard the chloroform layer. Transfer 8 mL of the acidic aqueous layer to a test tube, neutralize by the dropwise addition of 1 N sodium hydroxide, add 1 drop of 1 N sodium hydroxide in

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LINOLEOYL P IM OLYOXYLGLY PURITIES/ <i>Limit</i> CERIDES <i>of Free Glycerol</i>	USP36–NF31	2068	27-Sep-2013	1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	excess, and mix. Add a few drops of sodium nitroferricyanide TS and 2 drops of sodium hydroxide solution (15 in 100), mix, and allow to stand for 10 minutes. Add 0.1 N hydrochloric acid dropwise until the pH is between 8 and 9, and allow to stand for 10 minutes. A violet color develops. Line 1 of <i>Mode in Titrimetric system</i> : Change Direct titration to: Residual titration AND Line 10 of <i>Analysis</i> :

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							Change (V_S ? V_B) to: (V_B ? V_S) AND 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)
RIVASTIGMINE CHEMICAL TARTRATE INFORMATION	USP36–NF31	5073	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1: Change 398.41 to:

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POLYSORBAT E 80	SPECIFIC TESTS/ <i>Viscosity—Capillary Viscometer Methods <911> and Rotational Rheometer Methods <912></i>	<i>USP36–NF31</i>	2163	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	400.42 Line 1: Change and to: or
BACLOFEN	ASSAY/ <i>Procedure/Chromatographic system</i>	<i>First Supplement to USP36–NF31</i>	5951	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of Column: Change 250-cm to: 25.0-cm
SORBITAN MONOSTEARATE	IDENTIFICATION N/A.	<i>USP36–NF31</i>	2214	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	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 Acids AND Line 2 of Acceptance

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FILGRASTIM	ASSAY/Potency	<i>Second Supplement to USP36–NF31</i>	6606	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	<i>criteria: Change 200–215 to: 200–215 on 1-g sample</i> Line 7 of <i>Preparation of cells for analysis:</i> Change Column 1 is filled with 50 L of <i>Medium B.</i> to: Column 1 is filled with 50 µL of <i>Medium B.</i>
BUTABARBITA L SODIUM TABLETS	<i>Identification, Infrared Absorption <197K></i>	USP36–NF31	2716	27-Sep-2013		1-Oct-2013	USP38–NF33	<i>First Supplement to USP37–NF32</i>	Line 5 of <i>Test specimen:</i> Change Proceed as directed for <i>Column Partition Chromatography</i> under <i>Chromatography <621></i> , packing the chromatographic tube as follows.

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DILUTED ALCOHOL	ADDITIONAL REQUIREMENTS	USP36–NF31	1874	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	to: Pack a chromatographic tube as follows. Delete <i>USP Reference Standards <11></i> section
DROSPIRENONE	IMPURITIES/ <i>Organic Impurities/Procedure 2</i>	USP36–NF31	3349	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	In footnote b of <i>Table 4</i> : Change 5?,17?-Dihydroxy-6?,7?:15?,16?-dimethylene-17?-pregnan-21-carboxylic acid, ?-lactone. to: 5?,17-Dihydroxy-6?,7?:15?,16?-dimethylene-3-oxo-17?-pregnan-21-carboxylic acid, ?-lactone.
CETYL ALCOHOL	SPECIFIC TESTS/ <i>Fats and Fixed Oils, Hydroxyl Value <401></i>	USP36–NF31	1956	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of <i>Mode in Titrimetric system</i> : Change Direct titration to: Residual titration

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							<p>AND</p> <p>Line 9 of <i>Analysis:</i> Change Result = $[(V_S ? V_B) \times F]/W$ to: Result = $[(V_B ? V_S) \times N \times M_r]/W$</p> <p>AND</p> <p>Line 10 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) $F = \textit{factor}$, 56.1 to: $V_B = \textit{Titrant}$ volume consumed by the <i>Blank</i> (mL) $V_S = \textit{Titrant}$ volume consumed by</p>

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METHYLPHENIDATE HYDROCHLORIDE EXTENDED-RELEASE TABLETS	USP36–NF31	4327	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	the <i>Sample</i> (mL) <i>N</i> = actual normality of 1 N sodium hydroxide (mEq/mL) <i>M_r</i> = molecular weight of potassium hydroxide, 56.11 Line 1 of <i>Capacity factor</i> . Change NMT 2 to: NLT 2
HYDROXYETHYL CELLULOSE SPECIFIC TESTS/ <i>Viscosity—Capillary Viscometer Methods <911> and Rotational Rheometer Methods <912></i>	USP36–NF31	2038	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1: Change and to: or
RIFAMPIN, ISONIAZID, PYRAZINAMIDE, AND	USP36–NF31	5047	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 12 of <i>Procedure</i> : Change the <i>Standard</i>

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ETHAMBUTOL HYDROCHLOR IDE TABLETS							<i>preparation and the Assay preparation, respectively. to: the Assay preparation and the Standard preparation, respectively.</i>
POLYOXYL 10 SPECIFIC OLEYL ETHER TESTS/ <i>Average Polymer Length</i>	USP36–NF31	2150	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 9 of Analysis: Change Result = [(31 A_2/A_1 ? 3)]/4 to: Result = [(31 × A_2/A_1) ? 3]/4
COMPOUND Assay for zinc UNDECYLENIC <i>undecylenate</i> ACID OINTMENT	USP36–NF31	5516	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 15 of Procedure: Change 431.94 is the molecular weight of zinc undecylenate; to: M_r is the molecular weight of zinc undecylenate, 431.94;

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SORBITAN IDENTIFICATIO MONOOLEATE N/A.	USP36–NF31	2213	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	<p>AND Line 16 of <i>Procedure:</i> Change 65.39 is the atomic weight of zinc; to: A_r is the atomic weight of zinc, 65.39; AND Line 19 of <i>Procedure:</i> Change C_H and C_L are the concentrations, in μg per mL, to: C_{s1} and C_{s2} are the concentrations, in μg per mL, Line 1 of <i>Sample:</i> Change 1 g of the residue obtained in the <i>Assay for Fatty</i></p>

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									<p><i>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</p>
QUININE SULFATE TABLETS	ASSAY/ <i>Procedure</i>	<i>First Supplement to USP36–NF31</i>	6046	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	<p>Line 4 of <i>Analysis</i>: Change dihydroquinone sulfate to: dihydroquinine sulfate</p>
STEAROYL PO IM LYOXYLGLYC ERIDES	PURITIES/ <i>Limit of Free Glycerol</i>	<i>USP36–NF31</i>	2250	27-Sep-2013		1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	<p>Line 11 of <i>Analysis</i>: Change (V_S ? V_B) to: (V_B ? V_S) AND Line 12 of <i>Analysis</i>:</p>

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							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)
ELEMENTAL I ANALYTICAL MPURITIES—LITESTING MITS	USP36–NF31	151	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 2: Change If, by validated processes and supply-chain control, to: If, by process monitoring and supply-chain control,
CARISOPROD	USP Reference USP36–NF31	2813	27-Sep-2013	1-Oct-2013	USP38–NF33	First	Line 2 of USP

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OL, ASPIRIN, standards <11> AND CODEINE PHOSPHATE TABLETS						<i>Supplement to USP37–NF32</i>	Codeine N -Oxide RS: Change $C_{18}H_{21}O_4$ to: $C_{18}H_{21}NO_4$
CAPRYLOCAP IM ROYL POLYOX PURITIES/ <i>Limit</i> YLGLYCERIDE of <i>Free Glycerol</i> S	<i>USP36–NF31</i>	1922	27-Sep-2013	1-Oct-2013	<i>USP38–NF33</i>	<i>First Supplement to USP37–NF32</i>	Line 1 of <i>Mode</i> in <i>Titrimetric</i> system: Change Direct titration to: Residual titration AND Line 10 of <i>Analysis</i> : Change ($V_T ? V_B$) to: ($V_B ? V_S$) AND Line 11 of <i>Analysis</i> : Change $V_T = \textit{Titrant}$ volume consumed by the <i>Sample</i> (mL) $V_B = \textit{Titrant}$ volume

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FERUMOXASIL ORAL SUSPENSION	Viscosity—Capillary Viscometer Methods <911> and Rotational Rheometer Methods <912>	USP36–NF31	3572	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	consumed by the <i>Blank</i> (mL) to: $V_B = \text{Titrant volume consumed by the } Blank \text{ (mL)}$ $V_S = \text{Titrant volume consumed by the } Sample \text{ (mL)}$ Line 1: Change and to: or
FERROSOFERRIC OXIDE	IMPURITIES	USP36–NF31	2018	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of <i>Sample solution C</i> in <i>Limit of Lead (Pb)</i> : Change <i>Sample solution</i> to: <i>Sample stock solution</i> AND Line 1 of <i>Sample solution D</i> in <i>Limit of</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>Lead (Pb):</i> Change <i>Sample solution</i> to: <i>Sample stock solution</i> AND Line 1 of <i>Sample solution A in Limit of Mercury (Hg) and Nickel (Ni):</i> Change <i>Sample solution</i> to: <i>Sample stock solution</i> AND Line 1 of <i>Sample solution B in Limit of Mercury (Hg) and Nickel (Ni):</i> Change <i>Sample solution</i> to: <i>Sample stock solution</i> AND Line 1 of <i>Sample solution</i></p>

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POLYVINYL ALCOHOL	SPECIFIC TESTS/ <i>Viscosity—Capillary Viscometer Methods <911>, Rotational Rheometer Methods <912>, and Rolling Ball Viscometer Method <913></i>	USP36–NF31	4830	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	<i>C in Limit of Mercury (Hg) and Nickel (Ni):</i> Change <i>Sample solution</i> to: <i>Sample stock solution</i> Line 2: Change and to: or
OLEOYL POLY OXYLGLYCERIDES	IMPURITIES/ <i>Limit of Free Glycerol</i>	USP36–NF31	2112	27-Sep-2013		1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 1 of <i>Mode</i> in <i>Titrimetric system</i> : Change Direct titration to: Residual titration AND Line 10 of <i>Analysis</i> : Change (V

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>s? V_B) to: (V_B ? V_S) AND 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>
SALICYLIC ACID	USP Reference standards <11>	USP36–NF31 5098	27-Sep-2013	1-Oct-2013	USP38–NF33	First Supplement to USP37–NF32	Line 3 of USP Salicylic Acid Related Compound A

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							RS: Change [CAS-99-96-7]. to: [99-96-7]. AND Line 3 of USP Salicylic Acid Related Compound B RS: Change C ₈ H ₆ O ₄ to: C ₈ H ₆ O ₅

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