Add the following:

Cetirizine Hydrochloride Tablets

DEFINITION

Cetirizine Hydrochloride Tablets contain NLT 90.0% and NMT 110.0% of cetirizine hydrochloride (C₂₁H₂₅ClN₂O₃ · 2HCl).

IDENTIFICATION

• A. The retention time of the major peak of the Sample solution corresponds to that of the Standard solution, as obtained in the Assay.

ASSAY

PROCEDURE

- Solution A: 2 N sulfuric acid and water (2:33)
- Buffer: 2.9 mL/L of phosphoric acid in water
- Mobile phase: Acetonitrile and Buffer (3:7)
- **Diluent:** Acetonitrile, *Solution A*, and water (100:1:100) **Standard solution:** 0.2 mg/mL of USP Cetirizine Hydro-
- chloride RS in *Diluent* Sample solution: 0.2 mg/mL of cetirizine hydrochloride in *Diluent* from NLT 20 powdered Tablets. [NOTE—Soni-
- cate, if necessary.] Chromatographic system

(See Chromatography (621), System Suitability.)

- Mode: LC

Detector: UV 230 nm **Column:** 4.6-mm × 25-cm; 5-μm packing L1

- Flow rate: 1.5 mL/min

Injection volume: 10 μ L Run time: 1.3 times the retention time of cetirizine System suitability

Sample: Standard solution

- Suitability requirements
- Tailing factor: NMT 2.0
- Relative standard deviation: NMT 2.0% Analysis
- Samples: Standard solution and Sample solution Calculate the percentage of the labeled amount of cetirizine hydrochloride (C21H25CIN2O3 · 2HCl) in the portion of Tablets taken:

Result =
$$(r_U/r_S) \times (C_S/C_U) \times 100$$

- = peak response from the Sample solution r_U
- = peak response from the Standard solution rs
- = concentration of USP Cetirizine Hydrochloride Cs RS in the Standard solution (mg/mL)
- C_U = nominal concentration of cetirizine hydrochloride in the Sample solution (mg/mL)

Acceptance criteria: 90.0%–110.0%

PERFORMANCE TESTS

Change to read:

• **DISSOLUTION** $\langle 711 \rangle$ Test 1. (RB 1-Aug-2012) Medium: Water; 900 mL, degassed Apparatus 2: 50 rpm **Time:** 30 min **Buffer:** 2.9 mL/L of phosphoric acid in water Mobile phase: Acetonitrile and Buffer (2:3) **Standard solution:** 11 μg/mL of USP Cetirizine Hydrochloride RS in water. This solution can be stored for 48 h at room temperature. Sample solution: Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size.

Chromatographic system (See Chromatography (621), System Suitability.) Mode: LC **Detector:** UV 230 nm **Column:** 4.6-mm × 25-cm; 5-μm packing L1 Flow rate: 1 mL/min Injection volume: 50 µL Run time: 1.3 times the retention time of cetirizine System suitability Sample: Standard solution Suitability requirements Tailing factor: NMT 2.0 Relative standard deviation: NMT 2.0% Analysis Samples: Standard solution and Sample solution Calculate the percentage of the labeled amount of cetirizine hydrochloride ($C_{21}H_{25}CIN_2O_3 \cdot 2HCI$) dissolved: Result = $(r_U/r_S) \times (C_S/L) \times V \times 100$ = peak response from the \circ Sample solution \circ (RB 1r = peak response from the \bullet Standard solution (RB rs = concentration of the Standard solution Cs (mg/mL) = label claim (mg/Tablet) L V = volume of *Medium*, 900 mL Tolerances: NLT 80% (Q) of the labeled amount of cetirizine hydrochloride (C21H25CIN2O3 · 2HCI) is dissolved. **Test 2:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 2*. Medium: Water; 900 mL Apparatus 2: 75 rpm **Time:** 30 min **Buffer:** 0.4 g/L of 1-heptane sulfonic acid sodium salt **Mobile phase:** Acetonitrile and *Buffer* (50:50). Adjust with 0.1 N sulfuric acid to a pH of 3.5. **Standard solution:** $11 \ \mu$ g/mL of USP Cetirizine Hydrochloride RS in Medium Sample solution: Pass a 20-mL portion of the solution under test through a nylon filter of 0.45-µm pore size. Discard the first 10 mL of filtrate. Chromatographic system (See Chromatography (621), System Suitability.) Mode: LC Detector: UV 210 nm **Column:** 3.9-mm × 30-cm; 10-μm packing L1 Flow rate: 1.5 mL/min Injection volume: 50 µL Run time: 1.6 times the retention time of cetirizine System suitability Sample: Standard solution Suitability requirements
 Tailing factor:
 NMT 2.0

 Relative standard deviation:
 NMT 2.0%
Analysis Samples: Standard solution and Sample solution Calculate the percentage of the labeled amount of cetirizine hydrochloride (C₂₁H₂₅ClN₂O₃ · 2HCl) dissolved: Result = $(r_U/r_s) \times (C_s/L) \times V \times 100$ = peak response from the Sample solution rυ

- = peak response from the Standard solution
- rs Cs = concentration of USP Cetirizine Hydrochloride
- RS in the Standard solution (mg/mL) L
 - = label claim (mg/Tablet)
- V = volume of *Medium*, 900 mL

Cetirizine 1

Tolerances: NLT 80% (Q) of the labeled amount of cetirizine hydrochloride $(C_{21}H_{25}CIN_2O_3 \cdot 2HCI)$ is dissolved. **Test 3:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 3*.

Medium: Water; 900 mL

Apparatus 2: 50 rpm

Time: 30 min

Standard solution: (L/900) mg/mL of USP Cetirizine Hydrochloride RS in water, where L is the label claim of cetirizine hydrochloride, in mg/Tablet Sample solution: Centrifuge a portion of the solution under test for NLT 15 min at 3000 rpm.

Instrumental conditions

(See Spectrophotometry and Light-Scattering (851).) Mode: UV Analytical wavelength: UV 231 nm

Blank: Medium

Path length: 1 cm

Analysis

Samples: Standard solution and Sample solution Calculate the percentage of the labeled amount of cetirizine hydrochloride (C21H25CIN2O3 · 2HCI) dissolved:

Result = $(A_U/A_S) \times (C_S/L) \times V \times 100$

- = absorbance of the Sample solution Aυ
- = absorbance of the Standard solution
- = concentration of USP Cetirizine Hydrochloride RS in the *Standard solution* (mg/mL) C_{s}
- = label claim (mg/Tablet)
- V = volume of *Medium*, 900 mL **Tolerances:** NLT 80% (Q) of the labeled amount of cetirizine hydrochloride (C21H25CIN2O3 · 2HCI) is
- dissolved. (RB 1-Aug-2012) UNIFORMITY OF DOSAGE UNITS (905): Meet the requirements

IMPURITIES

Change to read:

ORGANIC IMPURITIES

Solution A: 2 N sulfuric acid and water (2:33) Buffer: 3.4 g/L of tetrabutyl ammonium hydrogen sulfate in water

Diluent: Acetonitrile, Solution A, and water (910:27:63) Mobile phase: Acetonitrile, Solution A, and Buffer (93:5:2

- Standard solution: 1.5 µg/mL of USP Cetirizine Hydrochloride RS in Diluent
- Sample solution: 0.5 mg/mL of cetirizine hydrochloride in *Diluent* from NLT 20 powdered Tablets. [NOTE— Sonicate, if necessary.]
- Chromatographic system

Mode: LC Detector: UV 230 nm Column: 4.0-mm × 25-cm; 5-µm packing L3 Flow rate: 0.8 mL/min Injection volume: 20 µL Run time: 2.5 times the retention time of cetirizine System suitability Sample: Standard solution Suitability requirements Tailing factor: NMT 2.0 Relative standard deviation: NMT 10.0% Analysis Samples: Standard solution and Sample solution Calculate the percentage of each impurity in the portion of Tablets taken:

Result =
$$(r_U/r_S) \times (C_S/C_U) \times (1/F) \times 100$$

- = peak response of each impurity from the r_U Sample solution
- = peak response of cetirizine from the Standard rs solution
- Cs = concentration of USP Cetirizine Hydrochloride RS in the Standard solution (mg/mL)
- Cu = nominal concentration of cetirizine hydrochloride in the Sample solution (mɡ/mL)
- = relative response factor (see Table 1)

Acceptance criteria: See Table 1.

Table 1

Name	Relative Retention Time	Relative Response Factor	Acceptance Criteria, NMT (%)
Cetirizine lactose ester ^a	0.56	1.0	0.40
Cetirizine	1.0	_	_
Cetirizine ethanol ^b	1.67	1.2	•0.2• (RB 1- Aug-2012)
Any unspecified degradation product	_	_	0.2
Total impurities	_	_	0.8

^a 6-O-[2-(2-{4-[(4-Chlorophenyl)(phenyl)methyl]piperazin-1yl}ethoxy)acetyl]- β -D-galactopyranosyl- $(1 \rightarrow 4)\beta$ -D-glucopyranose. ^b 2-[4-[(4-Chlorophenyl)phenylmethyl]piperazin-1-yl]ethanol.

ADDITIONAL REQUIREMENTS

PACKAGING AND STORAGE: Preserve in well-closed containers, and store below 30°.

Add the following:

• **LABELING:** When more than one *Dissolution* test is given, the labeling states the Dissolution test used only if *Test 1* is not used. (RB 1-Aug-2012) USP REFERENCE STANDARDS (11)

USP Cetirizine Hydrochloride RS [USP35]

⁽See Chromatography (621), System Suitability.)