

## **Metformin Hydrochloride Extended-Release Tablets**

Type of PostingRevision BulletinPosting Date26-Feb-2021Official Date1-Mar-2021

**Expert Committee** Small Molecules 3

In accordance with the Rules and Procedures of the Council of Experts, the Small Molecules 3 Expert Committee has revised the Metformin Hydrochloride Extended-Release Tablets monograph. The purpose for the revision is to add *Dissolution Test 24* to accommodate FDA-approved drug products with different dissolution conditions and/or tolerances than the existing dissolution tests.

The Metformin Hydrochloride Extended-Release Tablets Revision Bulletin supersedes the currently official monograph.

Should you have any questions, please contact Andrea F. Carney, Scientific Liaison (301-816-8155 or <a href="mailto:afc@usp.org">afc@usp.org</a>).

# **Metformin Hydrochloride Extended-Release Tablets**

#### **DEFINITION**

Metformin Hydrochloride Extended-Release Tablets contain NLT 90.0% and NMT 110.0% of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ).

#### **IDENTIFICATION**

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

### **ASSAY**

#### PROCEDURE

**Buffer solution:** 0.5 g/L of <u>sodium 1-heptanesulfonate</u> and 0.5 g/L of <u>sodium chloride</u> in water. Before final dilution, adjust with 0.06 M <u>phosphoric acid</u> to a pH of 3.85.

**Mobile phase:** Acetonitrile and Buffer solution (1:9). [Note—To improve the separation, the composition of acetonitrile and Buffer solution may be changed to 1:19, if necessary.]

Diluent: 1.25% solution of acetonitrile in water

**Standard solution:** (*L*/4000) mg/mL of <u>USP Metformin Hydrochloride RS</u> in *Diluent*, where *L* is the labeled quantity, in mg, of metformin hydrochloride in each Tablet

**System suitability stock solution:** 12.5 μg/mL each of <u>USP Metformin Related Compound B RS</u> and <u>USP Metformin Related Compound C RS</u> in *Diluent* 

System suitability solution: Dilute 0.5 mL of the System suitability stock solution with the Standard solution to 50 mL.

Sample stock solution: Finely powder NLT 10 Tablets. Transfer powder, equivalent to the average Tablet weight, to a homogenization vessel, and add 500 mL of a 10% acetonitrile solution. Alternately, homogenize and allow to soak until the sample is fully homogenized. [Note—A suggested homogenization sequence is as follows. Homogenize the sample using five pulses, each of 5 s, at about 20,000 rpm, and allow to soak for 2 min. Repeat these steps two additional times.]

**Sample solution:** Pass a portion of the *Sample stock solution* through a suitable filter of 0.45-µm pore size, discarding the first 3 mL of filtrate. Transfer 25 mL of the filtrate to a 200-mL volumetric flask, and dilute with water to volume.

### **Chromatographic system**

(See Chromatography (621), System Suitability.)

Mode: LC

Detector: UV 218 nm

Column: 3.9-mm  $\times$  30-cm; 10- $\mu$ m packing L1

Column temperature: 30° Flow rate: 1 mL/min Injection volume: 10 µL

Run time: Until after the elution locus of metformin related compound C

System suitability

Sample: System suitability solution

[Note—The relative retention times for metformin related compound B, metformin, and metformin related compound C are 0.86, 1.0, and 2.1–2.3, respectively. Metformin related compound C can have a variable retention time. The composition of the *Mobile phase* may be changed to 1:19, if it elutes at a relative retention time of less than 2.1.]

### Suitability requirements

Resolution: NLT 1.5 between the peaks due to metformin related compound B and metformin

**Tailing factor:** NLT 0.8 and NMT 2.0 for the metformin peak

**Relative standard deviation:** NMT 1.5% for the metformin peak and NMT 10% for each of the peaks due to metformin related compound B and metformin related compound C

#### **Analysis**

Samples: Standard solution and Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) in the portion of Tablets taken:

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

 $r_{II}$  = peak response from the Sample solution

 $r_S$  = peak response from the *Standard solution* 

 $C_S$  = concentration of <u>USP Metformin Hydrochloride RS</u> in the *Standard solution* (mg/mL)

 $C_{II}$  = nominal concentration of metformin hydrochloride in the Sample solution

Acceptance criteria: 90.0%-110.0%

### **PERFORMANCE TESTS**

### Change to read:

• **Dissolution** (711)

Test 1

Medium: pH 6.8 phosphate buffer solution; 1000 mL

**Apparatus 1:** 100 rpm for Tablets labeled to contain 750 mg **Apparatus 2:** 100 rpm for Tablets labeled to contain 500 mg

Times: 1, 3, and 10 h
Detector: UV 232 nm

Standard solution: USP Metformin Hydrochloride RS in Medium

 $\textbf{Sample solution:} \ \text{Pass a portion of the solution under test through a suitable hydrophilic polyethylene filter of } 0.45-\mu\text{m}$ 

pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*. **Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) released at each

time point:

$$\mathsf{Result} = [(A_U/A_S) \times C_S \times (V - V_S) + (C_{60} \times V_S) + (C_{180} \times V_S)] \times (100/L)$$

 $A_U$  = absorbance of the Sample solution

 $A_S$  = absorbance of the Standard solution

 $C_S$  = concentration of the *Standard solution* (mg/mL)

V = initial volume of *Medium* in the vessel (mL)

 $V_S$  = volume withdrawn from the vessel for previous samplings (mL)

 $C_{60}$  = concentration of metformin hydrochloride in *Medium* determined at 1 h (mg/mL)

 $C_{180}$  = concentration of metformin hydrochloride in *Medium* determined at 3 h (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See <u>Table 1</u>.

Table 1

Time (h)	Amount Dissolved, 500-mg Tablet (%)	Amount Dissolved, 750-mg Tablet (%)
1	20-40	22-42
3	45-65	49-69
10	NLT 85	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

**Test 2:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 2*.

Medium: Prepare as directed for Test 1; 1000 mL.

**Apparatus 2:** 100 rpm **Times:** 1, 2, 6, and 10 h

Detector: UV 232 nm

Standard solution: USP Metformin Hydrochloride RS in Medium

 $\textbf{Sample solution:} \ \text{Pass a portion of the solution under test through a suitable polyethylene filter of 0.45-$\mu m$ pore size.}$ 

Dilute, if necessary, with *Medium* to a concentration that is similar to that of the *Standard solution*.

**Analysis:** Calculate, in mg/mL, the content of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$   $(C_t)$ , in *Medium* at each time point (t):

Result = 
$$(A_{IJ} \times C_S \times D_{IJ})/A_S$$

 $A_{II}$  = absorbance of the Sample solution

 $C_S$  = concentration of metformin hydrochloride in the Standard solution (mg/mL)

 $D_{II}$  = dilution factor of the solution under test

 $A_{S}$  = absorbance of the Standard solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at each time point by the following formulas.

Percentage dissolved at the first time point (1 h):

Result = 
$$(C_1 \times V \times 100)/L$$

 $C_1$  = content of metformin hydrochloride in *Medium* at the first time interval (mg/mL)

V = volume of Medium, 1000 mL

L = label claim (mg/Tablet)

Percentage dissolved at the second time point (2 h):

Result = 
$$[C_2 \times (V - SV_1) + C_1 \times SV_1] \times (100/L)$$

 $C_2$  = content of metformin hydrochloride in *Medium* at the second time interval (mg/mL)

V = volume of *Medium*, 1000 mL

 $SV_1$  = volume of the sample withdrawn at 1 h (mL)

 $C_1$  = content of metformin hydrochloride in *Medium* at 1 h (mg/mL)

L = label claim (mg/Tablet)

Percentage dissolved at the *n*th time point:

Result = 
$$\{C_n \times [V - (n-1)V_S] + (C_1 + C_2 + ... + C_{n-1}) \times V_S\} \times (100/L)$$

 $C_n$  = content of metformin hydrochloride in *Medium* at the *n*th time interval (mg/mL)

V = volume of Medium, 1000 mL

n = time interval of interest

 $V_{\rm S}$  = volume of sample withdrawn at each time interval (mL)

 $C = as C_1, C_2, C_3, ... C_{n-1}$ , the content of metformin hydrochloride in *Medium* at each time interval (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See Table 2.

Table 2

Time	Amount Dissolved
(h)	(%)
1	20–40

Time (h)	Amount Dissolved (%)
2	35-55
6	65-85
10	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

Test 3: If the product complies with this test, the labeling indicates that it meets USP Dissolution Test 3.

Medium, Apparatus 1, and Apparatus 2: Proceed as directed in Test 1.

Times: 1, 2, 5, and 12 h for Tablets labeled to contain 500 mg; and 1, 3, and 10 h for Tablets labeled to contain 750 mg

Detector: UV 232 nm

Standard solution: USP Metformin Hydrochloride RS in Medium

**Sample solution:** Pass a portion of the solution under test through a suitable hydrophilic polyethylene filter of 0.45-µm pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*.

**Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) released at each time point:

$$\mathsf{Result} = \{ [(A_U/A_S) \times C_S \times (V - V_S) + (C_{60} \times V_S) + (C_{120} \times V_S) + (C_{300} \times V_S) + (C_{720} \times V_S)] \times 100 \} / L$$

 $A_{II}$  = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_S$  = concentration of the *Standard solution* (mg/mL)

V = initial volume of *Medium* in the vessel (mL)

 $V_{\rm S}$  = volume withdrawn from the vessel for previous samplings (mL)

 $C_{60}$  = concentration of metformin hydrochloride in *Medium* determined at 1 h (mg/mL)

 $C_{120}$  = concentration of metformin hydrochloride in *Medium* determined at 2 h (mg/mL)

 $C_{300}$  = concentration of metformin hydrochloride in *Medium* determined at 5 h (mg/mL)

 $C_{720}$  = concentration of metformin hydrochloride in *Medium* determined at 12 h (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See <u>Tables 3</u> and <u>4</u>.

Table 3. For Tablets Labeled to Contain 500 mg

Time (h)	Amount Dissolved (%)
1	20-40
2	35–55
5	60-80
12	NLT 85

Table 4. For Tablets Labeled to Contain 750 mg

Time (h)	Amount Dissolved (%)
1	22–42
3	49-69
10	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

**Test 4:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 4*.

Medium: Prepare as directed for Test 1; 1000 mL.

**Apparatus 2:** 100 rpm **Times:** 1, 3, 6, and 10 h

Detector: UV 250 nm (shoulder)

Standard solution: USP Metformin Hydrochloride RS in Medium

**Sample solution:** Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*.

**Analysis:** Calculate, in mg/mL, the content of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$   $(C_t)$ , in *Medium* at each time point (t), by the formulas specified in *Test 2*.

Tolerances: See <u>Table 5</u>.

Table 5

Time (h)	Amount Dissolved (%)
1	20-40
3	45–65
6	65–85
10	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

**Test 5:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 5*.

Medium: pH 6.8 phosphate buffer solution; 900 mL, deaerated

**Apparatus 1:** 100 rpm, with the vertical holder described in *Figure 1* and *Figure 2* 

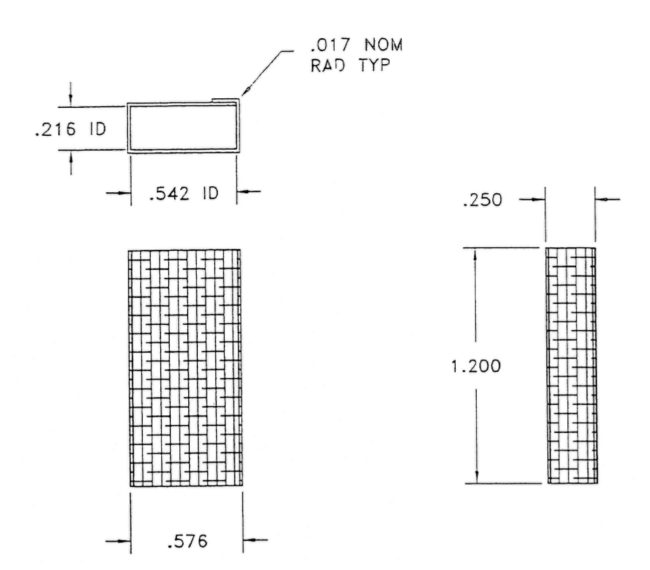
Times: 2, 8, and 16 h Detector: UV 250 nm

Standard solution: <u>USP Metformin Hydrochloride RS</u> in *Medium* 

**Sample solution:** Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*.

**Analysis:** Place a vertical sample holder into each basket (see <u>Figures 1</u> and  $\underline{2}$ ). Place 1 Tablet inside the sample holder, making sure that the Tablets are vertical at the bottom of the baskets.

Calculate, in mg/mL, the content of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) ( $C_t$ ), in *Medium* at each time point (t), by the formulas specified in *Test 2*.

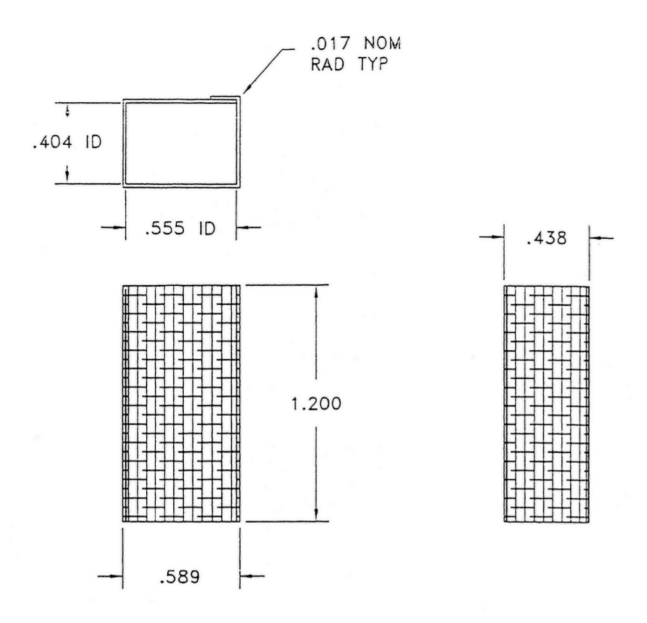


# NOTES:

- 1. MATERIAL: 316SS OR EQUIVALENT .017 WIRE VERTICAL MEAS SQUARE WEAVE WITH .039 SQUARE OPENINGS.
- 2. ALL DIMENSIONS ARE IN INCHES. TOLERANCES TO BE +/-.010

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Figure 1



# NOTES:

- 1. MATERIAL: 316SS OR EQUIVALENT .017 WIRE VERTICAL MEAS SQUARE WEAVE WITH .039 SQUARE OPENINGS.
- 2. ALL DIMENSIONS ARE IN INCHES. TOLERANCES TO BE +/-.010

Click image to enlarge

Figure 2

Tolerances: See <u>Table 6</u>.

Table 6

Time (h)	Amount Dissolved, 500-mg Tablet (%)	Amount Dissolved, 1000-mg Tablet (%)
2	NMT 30	NMT 30
8	60-85	65-90

Time (h)	Amount Dissolved, 500-mg Tablet (%)	Amount Dissolved, 1000-mg Tablet (%)
16	NLT 90	NLT 90

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 6:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 6*.

Medium: pH 6.8 phosphate buffer solution; 1000 mL, deaerated

Apparatus 2: 100 rpm, with USP sinker, if necessary

Detector: UV 233 nm

Standard solution: USP Metformin Hydrochloride RS in Medium

**Sample solution:** Pass a portion of the solution under test through a suitable hydrophilic polyethylene filter of 0.45-µm pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*.

**Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) released at each time point:

$$\text{Result} = \{ [(A_U/A_S) \times C_S \times (V - V_S) + (C_{60} \times V_S) + (C_{180} \times V_S) + (C_{600} \times V_S)] \times 100 \} / L$$

 $A_{II}$  = absorbance of the Sample solution

 $A_S$  = absorbance of the Standard solution

 $C_S$  = concentration of the Standard solution (mg/mL)

V = initial volume of *Medium* in the vessel (mL)

 $V_{\rm S}$  = volume withdrawn from the vessel for previous samplings (mL)

 $C_{60}$  = concentration of metformin hydrochloride in *Medium* determined at 1 h (mg/mL)

 $C_{180}$  = concentration of metformin hydrochloride in *Medium* determined at 3 h (mg/mL)

 $C_{600}$  = concentration of metformin hydrochloride in *Medium* determined at 10 h (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See <u>Table 7</u>.

Table 7

Time (h)	Amount Dissolved, 500-mg Tablet (%)	Amount Dissolved, 750-mg Tablet (%)
1	20-40	20-40
3	45–65	45–65
10	NLT 85	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 7:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 7*.

**Medium:** Prepare as directed in *Test 1*; 1000 mL.

Apparatus 1: 100 rpm for Tablets labeled to contain 750 mg

Apparatus 2: 50 rpm, with USP sinker, for Tablets labeled to contain 500 mg

Times: 1, 3, and 10 h
Detector: UV 232 nm

Standard solution: USP Metformin Hydrochloride RS in Medium

**Sample solution:** Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*.

**Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) released at each time point:

Result = 
$$\{[(A_U/A_S) \times C_S \times (V - V_S) + (C_{60} \times V_S) + (C_{180} \times V_S) + (C_{600} \times V_S)] \times 100\}/L$$

 $A_{II}$  = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_S$  = concentration of the Standard solution (mg/mL)

V = initial volume of *Medium* in the vessel (mL)

 $V_{\rm S}$  = volume withdrawn from the vessel for previous samplings (mL)

 $C_{60}$  = concentration of metformin hydrochloride in *Medium* determined at 1 h (mg/mL)

 $C_{1RO}$  = concentration of metformin hydrochloride in *Medium* determined at 3 h (mg/mL)

 $C_{600}$  = concentration of metformin hydrochloride in *Medium* determined at 10 h (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See Table 8.

Table 8

Time (h)	Amount Dissolved, 500-mg Tablet (%)	Amount Dissolved, 750-mg Tablet (%)
1	20-40	20-40
3	45-65	40-60
10	NLT 85	NLT 80

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 8:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 8*.

Medium: Prepare as directed in Test 1; 1000 mL.

Apparatus 1: 100 rpm for Tablets labeled to contain 750 mg

Apparatus 2: 100 rpm, with sinker, for Tablets labeled to contain 500 mg

**Times:** 1, 2, 6, and 10 h **Detector:** UV 232 nm

Standard solution: <u>USP Metformin Hydrochloride RS</u> in *Medium* 

**Sample solution:** Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size. Dilute, if necessary, with *Medium* to a concentration similar to that of the *Standard solution*.

**Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) released at each time point:

Result = {[
$$(A_U/A_S) \times C_S \times (V - V_S) + (C_{60} \times V_S) + (C_{120} \times V_S) + (C_{360} \times V_S) + (C_{600} \times V_S)] \times 100$$
}/L

 $A_U$  = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_c$  = concentration of the *Standard solution* (mg/mL)

V = initial volume of *Medium* in the vessel (mL)

 $V_S$  = volume withdrawn from the vessel for previous samplings (mL)

 $C_{60}$  = concentration of metformin hydrochloride in *Medium* determined at 1 h (mg/mL)

 $C_{120}$  = concentration of metformin hydrochloride in *Medium* determined at 2 h (mg/mL)

 $C_{360}$  = concentration of metformin hydrochloride in *Medium* determined at 6 h (mg/mL)

 $C_{600}$  = concentration of metformin hydrochloride in *Medium* determined at 10 h (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See <u>Table 9</u>.

Table 9

Time (h)	Amount Dissolved, 500-mg Tablet (%)	Amount Dissolved, 750-mg Tablet (%)
1	20-40	20-40
2	30-50	35-55
6	65-85	75-95
10	NLT 85	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 9:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 9*.

Medium: 0.05 M phosphate buffer, pH 6.8; 1000 mL

**Apparatus 1:** 100 rpm, for Tablets labeled to contain 750 mg **Apparatus 2:** 100 rpm, for Tablets labeled to contain 500 mg

Times: 1, 5, 12, and 20 h for Tablets labeled to contain 500 mg; and 1, 4, 10, and 24 h for Tablets labeled to contain 750

Standard solution: 0.5 mg/mL of USP Metformin Hydrochloride RS in Medium

Sample solution: Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size.

Detector: UV 232 nm

Path length: 0.01 cm, flow cell

Blank: Medium

**Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) released at each time point:

$$\text{Result} = \{ [(A_U/A_S) \times C_S \times (V - V_S) + (C_1 \times V_S) + (C_2 \times V_S) + (C_3 \times V_S) + (C_4 \times V_S)] \times 100 \} / L$$

 $A_{II}$  = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_{S}$  = concentration of the Standard solution (mg/mL)

V = initial volume of Medium in the vessel (mL)

 $V_{\rm S}$  = volume withdrawn from the vessel for previous samplings (mL)

 $C_{\star}$  = concentration of metformin hydrochloride in *Medium* determined at the first time point (mg/mL)

 $C_2$  = concentration of metformin hydrochloride in *Medium* determined at the second time point (mg/mL)

 $C_3$  = concentration of metformin hydrochloride in *Medium* determined at the third time point (mg/mL)

 $C_4$  = concentration of metformin hydrochloride in *Medium* determined at the fourth time point (mg/mL)

L = label claim (mg/Tablet)

Tolerances: See <u>Tables 10</u> and <u>11</u>.

Table 10. For Tablets Labeled to Contain 500 mg

Time (h)	Amount Dissolved (%)
1	20–40
5	45–65
12	70–90
20	NLT 85

Table 11. For Tablets Labeled to Contain 750 mg

Time (h)	Amount Dissolved (%)
1	20-45
4	45-70
10	70-95
24	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

**Test 10:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 10*.

**Medium:** 0.05 M phosphate buffer (prepared by dissolving 6.8 g of <u>monobasic potassium phosphate</u> in 250 mL of water, adding 77 mL of <u>0.2 N sodium hydroxide</u> and 500 mL of water, adjusting with <u>2 N sodium hydroxide</u> or 2 N hydrochloric acid to pH 6.8, and diluting with water to 1000 mL)

**Apparatus 1:** 100 rpm for Tablets labeled to contain 750 mg **Apparatus 2:** 100 rpm for Tablets labeled to contain 500 mg

Times: 1, 3, and 10 h

**Standard solution:** (L/100,000) mg/mL of <u>USP Metformin Hydrochloride RS</u> in *Medium*, where L is the label claim, in mg/Tablet. This solution is stable for 72 h at room temperature.

**Sample solution:** At the times specified, withdraw 10 mL of the solution under test and replace with 10 mL of *Medium* previously equilibrated at  $37.0 \pm 0.5^{\circ}$ . Centrifuge at 2500 rpm for 10 min. Dilute a portion of the supernatant with *Medium* to obtain a theoretical concentration of (L/100,000) mg/mL, where L is the label claim, in mg/Tablet.

**Detector:** UV 233 nm **Path length:** 1 cm **Blank:** *Medium* 

**Analysis:** Calculate the concentration, in mg/mL, of metformin hydrochloride  $(C_i)$  at each time point:

$$C_i = (A_{IJ}/A_S) \times C_S$$

 $A_{II}$  = absorbance of the Sample solution

 $A_{\varsigma}$  = absorbance of the *Standard solution* 

 $C_S$  = concentration of the Standard solution (mg/mL)

Calculate the cumulative percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved ( $Q_i$ ) at each time point (i):

At i = 1:

$$Q_1 = (C_1 \times V/L) \times 100$$

At i = 3:

 $Q_3 = [C_3(V - V_S) + (C_1 \times V_S)] \times 100/L$ 

At i = 10:

 $Q_{10} = [C_{10}(V - 2V_S) + (C_1 + C_3)V_S] \times 100/L$ 

V = initial volume of Medium, 1000 mL

 $V_{S}$  = sampling volume, 10 mL

L = label claim (mg/Tablet)

Tolerances: See <u>Table 12</u>.

Table 12

Time (h)	Amount Dissolved (%)
1	25-45
3	50-70
10	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 11:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 11*.

Medium: pH 6.8 phosphate buffer solution; 1000 mL

**Apparatus 1:** 100 rpm for Tablets labeled to contain 750 mg **Apparatus 2:** 100 rpm for Tablets labeled to contain 500 mg

Times: 1, 3, and 10 h

Standard solution: 7.5 µg/mL of USP Metformin Hydrochloride RS in Medium

**Sample solution:** At the times specified, withdraw 10 mL of the solution under test, and pass it through a suitable filter of 0.45-µm pore size, discarding the first 3 mL of filtrate. Dilute 3.0 mL of the filtrate with *Medium* to 200 mL. For Tablets labeled to contain 750 mg, dilute 2.0 mL of the filtrate with *Medium* to 200 mL. Replace the volume of *Medium* taken with the same volume of *Medium* preheated at 37.0  $\pm$  0.5°.

**Detector:** UV 232 nm **Path length:** 1 cm **Blank:** *Medium* 

**Analysis:** Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at each time point:

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

At 1 h:

Result =  $Q_1$ 

At 3 h:

Result =  $Q_3 + [(Q_1 \times 10)/V]$ 

At 10 h:

Result =  $Q_{10}$  + {[ $(Q_1 \times 10)/V$ ] + [ $(Q_3 \times 10)/V$ ]}

 $A_{II}$  = absorbance of the Sample solution

 $A_{S}$  = absorbance of the *Standard solution* 

 $C_{S}$  = concentration of the Standard solution (mg/mL)

L = label claim (mg/Tablet)

V = volume of Medium, 1000 mL

D = dilution factor of the Sample solution

Tolerances: See Table 13.

Table 13

Time (h)	Amount Dissolved (%)
1	25-45
3	50-70
10	NLT 80

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 12:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 12*.

Medium: pH 6.8 phosphate buffer solution; 1000 mL

**Apparatus 1:** 100 rpm **Times:** 1, 4, and 12 h

Standard stock solution: 0.2 mg/mL of USP Metformin Hydrochloride RS in Medium

**Standard solution:** 0.01 mg/mL of <u>USP Metformin Hydrochloride RS</u> in water, from the *Standard stock solution* 

**Sample solution:** At the times specified, withdraw 10 mL of the solution under test, and replace with 10 mL of *Medium* previously equilibrated at 37.0  $\pm$  0.5°. Pass it through a suitable filter, discarding the first few mL of the filtrate.

For Tablets labeled to contain 500 mg: Dilute 2.0 mL of the filtrate with water to 100 mL. For Tablets labeled to contain 1000 mg: Dilute 1.0 mL of the filtrate with water to 100 mL.

Detector: UV 232 nm

Blank: Dilute 1 mL of Medium with water to 100 mL.

**Analysis:** Calculate the concentration  $(C_i)$ , in mg/mL, of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$  in the sample withdrawn at each time point (i):

$$Result_i = (A_I/A_S) \times C_S \times D$$

A,, = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_S$  = concentration of the Standard solution (mg/mL)

D = dilution factor of the Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved ( $Q_i$ ) at each time point (i):

$$\mathsf{Result}_1 = C_1 \times V \times (1/L) \times 100$$

$$\mathsf{Result}_2 = \{ [C_2 \times V] + [C_1 \times V_S] \} \times (1/L) \times 100$$

Result<sub>3</sub> = {
$$[C_3 \times V] + [(C_2 + C_1) \times V_S]$$
} × (1/L) × 100

 $C_i$  = concentration of metformin hydrochloride in the portion of sample withdrawn at time point i (mg/mL)

V = initial volume of *Medium*, 1000 mL

L = label claim (mg/Tablet)

Tolerances: See Table 14.

Table 14

Time Point (i)	Time (h)	Amount Dissolved (%)
1	1	NMT 15
2	4	35-65
3	12	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

**Test 13:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 13*.

Medium: pH 6.8 phosphate buffer solution; 1000 mL

**Apparatus 1:** 100 rpm **Times:** 1, 4, 6, and 14 h

**Standard stock solution:** 0.2 mg/mL of <u>USP Metformin Hydrochloride RS</u> prepared as follows. Transfer a suitable amount of <u>USP Metformin Hydrochloride RS</u> into an appropriate volumetric flask. Dissolve by adding *Medium* to fill 50% of the flask volume and dilute with *Medium* to volume.

Standard solution: 0.01 mg/mL of USP Metformin Hydrochloride RS from Standard stock solution in water

**Sample stock solution:** At the times specified, withdraw 10 mL of the solution under test, and replace with the same volume of *Medium* preheated at 37.0  $\pm$  0.5°. Pass a portion of the solution under test through a suitable filter of 0.45- $\mu$ m pore size, discard the first few mL, and use the filtrate.

#### Sample solution

For Tablets labeled to contain 500 mg: Dilute 2 mL of Sample stock solution with water to 100 mL. For Tablets labeled to contain 1000 mg: Dilute 1 mL of Sample stock solution with water to 100 mL.

## **Instrumental conditions**

(See <u>Ultraviolet-Visible Spectroscopy (857)</u>.)

Mode: UV

Analytical wavelength: 232 nm

Blank

For Tablets labeled to contain 500 mg: Dilute 2 mL of *Medium* with water to 100 mL. For Tablets labeled to contain 1000 mg: Dilute 1 mL of *Medium* with water to 100 mL.

System suitability

**Sample:** Standard solution **Suitability requirements** 

Relative standard deviation: NMT 2.0%

**Analysis** 

Samples: Standard solution, Sample solution, and Blank

Calculate the concentration  $(C_i)$ , in mg/mL, of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$  in the sample withdrawn from the vessel at each time point (i):

$$Result_i = (A_U/A_S) \times C_S \times D$$

 $A_{II}$  = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_S$  = concentration of the *Standard solution* (mg/mL)

D = dilution factor of the Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at each time point (*i*):

$$Result_1 = C_1 \times V \times (1/L) \times 100$$

Result<sub>2</sub> = 
$$[(C_2 \times V) + (C_1 \times V_S)] \times (1/L) \times 100$$

Result<sub>3</sub> = {
$$[C_3 \times V] + [(C_2 + C_1) \times V_S]$$
} × (1/L) × 100

Result<sub>4</sub> = {
$$[C_4 \times V] + [(C_3 + C_2 + C_1) \times V_5]$$
} × (1/L) × 100

 $C_i$  = concentration of metformin hydrochloride in the portion of sample withdrawn at the specified time point (mg/mL)

V = volume of Medium, 1000 mL

L = label claim (mg/Tablet)

 $V_{\rm S}$  = volume of the Sample solution withdrawn at each time point and replaced with Medium (mL)

Tolerances: See <u>Table 15</u>.

Table 15

Time Point (i)	Time (h)	Amount Dissolved (%)
1	1	NMT 20
2	4	45-65
3	6	65-85
4	14	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

Test 14: If the product complies with this test, the labeling indicates that it meets USP Dissolution Test 14.

Medium: pH 6.8 phosphate buffer solution; 1000 mL

**Apparatus 1:** 100 rpm for Tablets labeled to contain 750 mg **Apparatus 2:** 100 rpm for Tablets labeled to contain 500 mg

Times: 1, 3, and 10 h

Standard solution: 7.5 µg/mL of USP Metformin Hydrochloride RS in Medium

**Sample solution:** At the times specified, withdraw 10 mL of the solution under test and replace with the same volume of *Medium*. Pass the solution under test through a suitable filter of  $10-\mu m$  pore size. Pass a portion of the filtered solution through a suitable filter of  $0.45-\mu m$  pore size, discarding the first few milliliters. Dilute with *Medium* to a concentration similar to that of the *Standard solution*.

#### **Instrumental conditions**

Mode: UV

Analytical wavelength: 232 nm

Blank: Medium

**Analysis** 

Samples: Standard solution, Sample solution, and Blank

Calculate the concentration  $(C_i)$ , in mg/mL, of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$  in the sample withdrawn from the vessel at each time point (i):

$$Result_i = (A_U/A_S) \times C_S \times D$$

A,, = absorbance of the Sample solution

 $A_c$  = absorbance of the Standard solution

 $C_S$  = concentration of the Standard solution ( $\mu$ g/mL)

D = dilution factor of the Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at each time point (*i*):

$$Result_1 = C_1 \times V \times (1/L) \times 100$$

Result<sub>2</sub> = 
$$[(C_2 \times V) + (C_1 \times V_S)] \times (1/L) \times 100$$

Result<sub>3</sub> = 
$$\{(C_3 \times V) + [(C_2 + C_1) \times V_S]\} \times (1/L) \times 100$$

Result<sub>4</sub> = 
$$\{(C_4 \times V) + [(C_3 + C_2 + C_1) \times V_5]\} \times (1/L) \times 100$$

 $C_i$  = concentration of metformin hydrochloride in the portion of sample withdrawn at the specified time point (mg/mL)

V = volume of Medium, 1000 mL

L = label claim (mg/Tablet)

 $V_c$  = volume of the Sample solution withdrawn at each time point and replaced with Medium (mL)

Tolerances: See Table 16.

Table 16

Time Point	Time	Amount Dissolved (%)	
(i)	(h)	500 mg Tablets	750 mg Tablets
1	1	30-50	25-45
2	3	55-75	50-70
3	10	NLT 85	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 16:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 16*.

Medium: pH 6.8 phosphate buffer solution; 1000 mL

Apparatus 1: 100 rpm Times: 1, 3, and 10 h

Standard solution: 0.015 mg/mL of USP Metformin Hydrochloride RS in Medium. Sonicate as needed.

**Sample stock solution:** At the times specified, withdraw 10 mL of the solution under test and pass it through a suitable filter.

### Sample solution

For Tablets labeled to contain 500 mg: Dilute 3 mL of the Sample stock solution with Medium to 100 mL. For Tablets labeled to contain 750 mg: Dilute 2 mL of the Sample stock solution with Medium to 100 mL.

# Instrumental conditions

(See <u>Ultraviolet-Visible Spectroscopy (857)</u>.)

Mode: UV-Vis

Analytical wavelength: UV 232 nm

Path length: 1 cm Blank: Medium System suitability

**Sample:** Standard solution **Suitability requirements** 

Relative standard deviation: NMT 2.0%

### **Analysis**

Samples: Standard solution and Sample solution

Calculate the concentration  $(C_i)$  of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$  in the sample withdrawn from the vessel at each time point (i):

Result = 
$$(A_U/A_S) \times C_S \times D$$

 $A_{II}$  = absorbance of the Sample solution

 $A_{S}$  = absorbance of the Standard solution

 $C_{S}$  = concentration of the Standard solution (mg/mL)

D = dilution factor for the Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the specified time point:

$$\mathsf{Result}_1 = C_1 \times V \times (1/L) \times 100$$

$$\mathsf{Result}_2 = \{ [C_2 \times (V - V_S)] + (C_1 \times V_S) \} \times (1/L) \times 100$$

$$\mathsf{Result}_3 = (\{C_3 \times [V - (2 \times V_S)]\} + [(C_2 + C_1) \times V_S]) \times (1/L) \times 100$$

 $C_i$  = concentration of metformin hydrochloride in the portion of sample withdrawn at the specified time point (mg/mL)

V = volume of Medium, 1000 mL

L = label claim (mg/Tablet)

 $V_{\rm S}$  = volume of the Sample solution withdrawn at each time point (mL)

Tolerances: See <u>Table 17</u>.

Table 17

Time Point (i)	Time (h)	Amount Dissolved (%)
1	1	25-45
2	3	50-70
3	10	NLT 80

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to *Dissolution* (711), *Acceptance Table 2*.

**Test 18:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 18*.

**Medium:** pH 6.8 phosphate buffer solution; 900 mL, deaerated **Apparatus 1:** 100 rpm, with vertical holder described in *Figure 1* 

Times: 1, 4, and 10 h

Standard solution: 0.044 mg/mL of <u>USP Metformin Hydrochloride RS</u> in *Medium*. Sonicate as needed.

**Sample stock solution:** At the times specified, withdraw a suitable amount of solution under test and replace with a suitable amount of *Medium*. Pass the solution under test through a suitable filter and discard the first few milliliters.

### Sample solution

For Tablets labeled to contain 500 mg: Dilute 2 mL of the Sample stock solution with Medium to 25 mL. For Tablets labeled to contain 1000 mg: Dilute 2 mL of the Sample stock solution with Medium to 50 mL.

### **Instrumental conditions**

(See <u>Ultraviolet-Visible Spectroscopy (857)</u>.)

Mode: UV-Vis

Analytical wavelength: UV 250 nm

Path length: 1 cm Blank: Medium System suitability

**Sample:** Standard solution **Suitability requirements** 

### Relative standard deviation: NMT 2.0%

### **Analysis**

Samples: Standard solution and Sample solution

Place a vertical sample holder into each basket (see <u>Figure 1</u>). Place 1 Tablet inside the sample holder, making sure that the Tablets are vertical at the bottom of the baskets.

Calculate the concentration ( $C_i$ ) of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) in the sample withdrawn from the vessel at each time point (i):

Result = 
$$(A_{IJ}/A_S) \times C_S \times D$$

 $A_{II}$  = absorbance of the Sample solution

 $A_S$  = absorbance of the *Standard solution* 

 $C_S$  = concentration of the Standard solution (mg/mL)

D = dilution factor of the Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at each time point (i):

$$Result_1 = C_1 \times V \times (1/L) \times 100$$

Result<sub>2</sub> = 
$$[(C_2 \times V) + (C_1 \times V_S)] \times (1/L) \times 100$$

Result<sub>3</sub> = 
$$\{(C_3 \times V) + [(C_2 + C_1) \times V_S]\} \times (1/L) \times 100$$

 $C_i$  = concentration of metformin hydrochloride in the portion of sample withdrawn at the specified time point (mg/mL)

V = volume of Medium, 900 mL

L = label claim (mg/Tablet)

 $V_{\rm S}$  = volume of the Sample solution withdrawn at each time point and replaced with Medium (mL)

Tolerances: See Table 18.

Table 18

Time Point (i)	Time (h)	Amount Dissolved (%)
1	1	NMT 30
2	4	45-70
3	10	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

**Test 23:** If the product complies with this test, the labeling indicates that it meets USP *Dissolution Test 23*.

Medium: pH 6.8 phosphate buffer; 900 mL, deaerated

Apparatus 1: 100 rpm Times: 1, 4, and 10 h

**Buffer:** 2.8 g/L of <u>sodium phosphate, monobasic</u>, 2.0 g/L of <u>sodium 1-heptanesulfonate</u>, and 2 mL/L of <u>triethylamine</u> prepared as follows. Dissolve 2.8 g of <u>sodium phosphate, monobasic</u> and 2.0 g of <u>sodium 1-heptanesulfonate</u> in 800 mL of <u>water</u>. Add 2 mL of <u>triethylamine</u>, and dilute with <u>water</u> to volume. Adjust with <u>phosphoric acid</u> to a pH of 3.5.

Mobile phase: Methanol and Buffer (40:60)

**Standard solution:** (L/900) mg/mL of <u>USP Metformin Hydrochloride RS</u> in *Medium*, where L is the label claim in mg/Tablet. Sonicate to dissolve, if necessary.

**Sample solution:** At the times specified, withdraw a suitable amount of solution under test. Pass the solution under test through a suitable filter, and discard the first 3 mL.

## **Chromatographic system**

(See Chromatography (621), System Suitability.)

Mode: LC

Detector: UV 260 nm

**Column:** 4.6-mm  $\times$  15-cm; 5- $\mu$ m packing  $\boxed{1}$ 

Column temperature: 35° Flow rate: 1 mL/min Injection volume: 10  $\mu$ L

Run time: NLT 2 times the retention time of metformin

System suitability

Sample: Standard solution
Suitability requirements
Tailing factor: NMT 2.5

Relative standard deviation: NMT 3.0%

### **Analysis**

Samples: Standard solution and Sample solution

Calculate the concentration ( $C_i$ ) of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) in the sample withdrawn from the vessel at each time point (i):

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

 $r_U$  = peak response of metformin from the Sample solution

 $r_S$  = peak response of metformin from the *Standard solution* 

 $C_{\rm S}$  = concentration of <u>USP Metformin Hydrochloride RS</u> in the *Standard solution* (mg/mL)

Calculate the percentage of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the specified time point (i):

$$\begin{aligned} \text{Result}_1 &= C_1 \times V \times (1/L) \times 100 \\ \text{Result}_2 &= \{ [C_2 \times (V - V_S)] + (C_1 \times V_S) \} \times (1/L) \times 100 \\ \\ \text{Result}_3 &= (\{C_3 \times [V - (2 \times V_S)]\} + [(C_2 + C_1) \times V_S]) \times (1/L) \times 100 \end{aligned}$$

 $C_i$  = concentration of metformin hydrochloride in the portion of sample withdrawn at the specified time point (mg/mL)

V = volume of the Medium, 900 mL

L = label claim (mg/Tablet)

 $V_S$  = volume of the Sample solution withdrawn at each time point (mL)

Tolerances: See Table 19.

Table 19

Time Point (i)	Time (h)	Amount Dissolved (%)
1	1	10-30
2	4	50-70
3	10	NLT 85

The percentages of the labeled amount of metformin hydrochloride ( $C_4H_{11}N_5 \cdot HCI$ ) dissolved at the times specified conform to <u>Dissolution (711)</u>, <u>Acceptance Table 2</u>.

▲Test 24: If the product complies with this test, the labeling indicates that it meets USP Dissolution Test 24.

Medium: pH 6.8 phosphate buffer solution; 1000 mL

**Apparatus 1:** 100 rpm for Tablets labeled to contain 750 mg **Apparatus 2:** 100 rpm for Tablets labeled to contain 500 mg

Times: 1, 3, and 10 h

Standard solution: 7.5 µg/mL of USP Metformin Hydrochloride RS in Medium

**Sample stock solution:** At the times specified, withdraw 10 mL of the solution under test and replace with the same amount of *Medium*. Pass a portion of the solution under test through a suitable filter of 0.45-μm pore size, discarding the first 3 mL.

### Sample solution

For Tablets labeled to contain 500 mg: Dilute 3 mL of the Sample stock solution with Medium to 200 mL. For Tablets labeled to contain 750 mg: Dilute 2 mL of the Sample stock solution with Medium to 200 mL.

#### **Instrumental conditions**

(See <u>Ultraviolet-Visible Spectroscopy (857)</u>.)

Mode: UV-Vis

Analytical wavelength: UV 232 nm

Blank: Medium

**Analysis** 

Samples: Standard solution and Sample solution

Calculate the concentration  $(C_i)$  of metformin hydrochloride  $(C_4H_{11}N_5 \cdot HCI)$  in the sample withdrawn from the vessel at each time point (i):

Result = 
$$(A_U/A_S) \times C_S \times D$$

 $A_{II}$  = absorbance of the Sample solution

A<sub>S</sub> = absorbance of the *Standard solution* 

 $C_S$  = concentration of <u>USP Metformin Hydrochloride RS</u> in the *Standard solution* (mg/mL)

D = dilution factor for the Sample solution

Calculate the percentage of the labeled amount of metformin hydrochloride (C<sub>4</sub>H<sub>11</sub>N<sub>5</sub>·HCl) dissolved at the specified time point:

$$Result_1 = C_1 \times V \times (1/L) \times 100$$

$$Result_2 = [(C_2 \times V) + (C_1 \times V_S)] \times (1/L) \times 100$$

Result<sub>3</sub> = {
$$[C_3 \times V] + [(C_2 + C_1) \times V_S]$$
} × (1/L) × 100

C<sub>i</sub> = concentration of metformin hydrochloride in the portion of sample withdrawn at the specified time point (mg/mL)

V = volume of Medium, 1000 mL

L = label claim (mg/Tablet)

 $V_{\rm c}$  = volume of the Sample solution withdrawn at each time point (mL)

Tolerances: See Table 20.

Table 20

Time Point (i)	Time (h)	Amount Dissolved (%)
1	1	25-45
2	3	45-65
3	10	NLT 80

The percentages of the labeled amount of metformin hydrochloride (C<sub>4</sub>H<sub>11</sub>N<sub>5</sub>·HCl) dissolved at the times specified conform to *Dissolution* (711), Acceptance Table 2. ▲ (RB 1-Mar-2021)

• **Uniformity of Dosage Units** (905): Meet the requirements

#### **IMPURITIES**

#### • ORGANIC IMPURITIES

Mobile phase, Sample solution, and Chromatographic system: Proceed as directed in the Assay.

Analysis: From the chromatogram of the Sample solution obtained in the Assay, calculate the percentage of each impurity in the portion of Tablets taken:

Result = 
$$(r_U/r_T) \times 100$$

= peak response for each impurity  $r_{IJ}$ 

= sum of all the peak responses  $r_T$ 

### Acceptance criteria

Individual impurities: NMT 0.1% Total impurities: NMT 0.6%

[Note—Disregard any peak less than 0.05%, and disregard any peak observed in the blank.]

### **ADDITIONAL REQUIREMENTS**

- Packaging and Storage: Preserve in well-closed, light-resistant containers, and store at controlled room temperature.
- LABELING: When more than one dissolution test is given, the labeling states the Dissolution test used only if Test 1 is not used.
- USP REFERENCE STANDARDS (11)

USP Metformin Hydrochloride RS

USP Metformin Related Compound B RS

1-Methylbiguanide hydrochloride.

C<sub>3</sub>H<sub>0</sub>N<sub>5</sub>HCl

151.60 USP Metformin Related Compound C RS

*N,N*-Dimethyl-[1,3,5]triazine-2,4,6-triamine.

154.17  $C_5H_{10}N_6$ 

### **Page Information:**

Not Applicable

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