Fenofibrate Capsules

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<td>01–May–2019</td>
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<td>Expert Committee</td>
<td>Chemical Medicines Monographs 2</td>
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In accordance with the Rules and Procedures of the 2015–2020 Council of Experts, the Chemical Medicines Monographs 2 Expert Committee has revised the Fenofibrate Capsules monograph. The purpose for the revision is to add Dissolution Test 6 to accommodate FDA-approved drug products with different dissolution tolerances than the existing dissolution tests.

- Dissolution Test 6 was validated using an Agilent Zorbax SB-C18 brand of L1 column. The typical retention time for fenofibrate is about 7.7 min.

The Fenofibrate Capsules Revision Bulletin supersedes the currently official monograph.

Should you have any questions, please contact Edith Chang, Senior Scientific Liaison–Team Leader (301-816-8392 or yec@usp.org).
Fenofibrate Capsules

**DEFINITION**
Fenofibrate Capsules contain NLT 90.0% and NMT 110.0% of the labeled amount of fenofibrate (C_{20}H_{28}ClO_{4}).

**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.

**Add the following:**
- **B.** The UV spectrum of the major peak of the **Sample solution** corresponds to that of the **Standard solution**, as obtained in the Assay.

**ASSAY**

**Change to read:**
- **Procedure**
  Use **Sample stock solution 2** for Capsules labeled to meet the requirements of **Dissolution Test 2**. For all other products, use **Sample stock solution 1**.

**Solution A:** 136 mg/L of monobasic potassium phosphate in water. Adjust with dilute phosphoric acid (1 in 10) to a pH of 2.9 ± 0.05.

**Mobile phase:** Methanol and **Solution A** (4:1)

**Standard solution:** 67 µg/mL of USP Fenofibrate RS in **Mobile phase**

**Sample stock solution 1:** Accurately weigh the contents of NLT 20 Capsules. Mix the contents, and transfer a weighed portion of the powder, equivalent to about 67 mg of fenofibrate, to a 100-µL volumetric flask. Add 80 mL of **Mobile phase**, sonicate for 10 min, stir for 15 min, and dilute with **Mobile phase** to volume.

**Sample stock solution 2** (for Capsules labeled to meet the requirements of **Dissolution Test 2**): Weigh the contents of NLT 20 Capsules. Mix the contents, melt in an oven at 80° for NLT 30 min, and homogenize. Allow the sample to solidify. Transfer a weighed portion of the sample, equivalent to about 67 mg of fenofibrate, to a 100-µL volumetric flask, dissolve in 30 mL of methanol with the aid of a mechanical shaker for NLT 4 h, and dilute with **Mobile phase** to volume.

**Sample solution:** Nominally 67 µg/mL of fenofibrate from the designated **Sample stock solution**, in **Mobile phase**. Pass a portion of this solution through a polyvinylidene difluoride (PVDF) filter of 0.45-µm pore size, discarding the first 5 mL.

**Chromatographic system**
(See **Chromatography** (621), **System Suitability**.)

**Mode:** LC

**Detector:** UV 285 nm

**Column:** 4.6-mm × 15-cm; 5-µm packing L1

**Flow rate:** 1 mL/min

**Injection volume:** 20 µL

**Run time:** NLT 1.5 times the retention time of the fenofibrate peak

**System suitability**
**Sample:** **Standard solution**

**Suitability requirements**
- ▲ USP 1-May-2019

- **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 fenofibrate (C_{20}H_{28}ClO_{4}) in the portion of Capsules taken:

\[
\text{Result} = \left(\frac{r_s}{r_U}\right) \times \left(\frac{C_s}{C_U}\right) \times 100
\]

- \(r_s\) = peak response from the **Sample solution**
- \(r_U\) = peak response from the **Standard solution**
- \(C_s\) = concentration of the **Standard solution** (mg/mL)
- \(C_U\) = nominal concentration of the **Sample solution** (µg/mL)

**Acceptance criteria:** 90.0%–110.0%

**PERFORMANCE TESTS**

**Change to read:**
- **Dissolution** (711)

  **Test 1**
  - **Medium:** 0.05 M sodium lauryl sulfate in water, 1000 mL, deaerated
  - **Apparatus 2:** 75 rpm
  - **Time:** 40 min
  - **Solution A and Mobile phase:** Prepare as directed in the Assay.
  - **Standard solution:** (0.001 × L) mg/mL of USP Fenofibrate RS in **Mobile phase**, where L is the label claim, in mg/Capsule
  - **Sample solution:** Pass a portion of the solution under test through a suitable PVDF filter of 0.45-µm pore size.

  **Chromatographic system**
  (See **Chromatography** (621), **System Suitability**.)

  **Mode:** LC

  **Detector:** UV 285 nm

  **Column:** 4.6-mm × 15-cm; 5-µm packing L1

  **Flow rate:** 1 mL/min

  **Injection volume:** 10 µL for Capsules labeled to contain 67 mg; 5 µL for Capsules labeled to contain 134 or 200 mg

  **System suitability**
  **Sample:** **Standard solution**

  **Suitability requirements**
  ▲ USP 1-May-2019

  **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 fenofibrate (C_{20}H_{28}ClO_{4}) dissolved:

  \[
  \text{Result} = \left(\frac{r_s}{r_U}\right) \times \left(\frac{C_s}{C_U}\right) \times V \times \left(\frac{1}{L}\right) \times 100
  \]

  - \(r_s\) = peak response from the **Sample solution**
  - \(r_U\) = peak response from the **Standard solution**
  - \(C_s\) = concentration of the **Standard solution** (mg/mL)
  - \(V\) = volume of **Medium**, 1000 mL
  - \(L\) = label claim (mg/Capsule)

  **Tolerances:** NLT 70% (Q) of the labeled amount of fenofibrate (C_{20}H_{28}ClO_{4}) dissolved.

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

  **Medium:** Phosphate buffer pH 6.8 ± 0.1 containing 0.1% pancreatin and 2% polysorbate 80; 900 mL, deaerated by vacuum
**2 Fenofibrate**

### Apparatus 2: 75 rpm with sinkers (see Dissolution (711), Figure 2a)

**Time:** 2 h

**Standard solution:** (L/1000) mg/mL of USP Fenofibrate RS in Medium, where L is the label claim in mg/Capsule. A volume of methanol, not exceeding 10%, can be used in the first dilution to solubilize fenofibrate.

**Sample solution:** Pass 20 mL of the solution under test through a suitable PVDF filter of 0.45-µm pore size, discarding the first 2 mL.

**Blank:** Medium

**Instrumental conditions**

(See Ultraviolet-Visible Spectroscopy (857),)

**Mode:** Spectrophotometry

**Detector:** UV 288 nm

**Path length:** 0.1-cm flow cell

**Analysis**

**Samples:** Standard solution and Sample solution

Calculate the percentage of the labeled amount of fenofibrate (C_{20}H_{31}ClO_{4}) dissolved:

\[
\text{Result} = (A_s/A_i) \times C_i \times V \times (1/L) \times 100
\]

- \(A_s\) = absorbance of the Sample solution
- \(A_i\) = absorbance of the Standard solution
- \(C_i\) = concentration of the Standard solution (mg/mL)
- \(V\) = volume of Medium, 900 mL
- \(L\) = label claim (mg/Capsule)

**Tolerances:** NLT 80% (Q) of the labeled amount of fenofibrate (C_{20}H_{31}ClO_{4}) is dissolved.

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

**Medium:** 0.72% sodium lauryl sulfate in water; 1000 mL, deaerated

**Apparatus 2:** 75 rpm, with sinkers with three prongs

**Time:** 30 min

**Standard solution:** (L/10) mg/mL of USP Fenofibrate RS in methanol, where L is the label claim in mg/Capsule. Transfer 10.0 mL of this solution to a 1000-mL volumetric flask, and dilute with Medium to volume.

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

**Instrumental conditions**

(See Ultraviolet-Visible Spectroscopy (857),)

**Mode:** Spectrophotometry

**Detector:** UV 291 nm

**Path length:** 0.1-cm flow cell

**Analysis**

**Samples:** Standard solution and Sample solution

Calculate the percentage of the labeled amount of fenofibrate (C_{20}H_{31}ClO_{4}) dissolved:

\[
\text{Result} = (A_s/A_i) \times C_i \times V \times (1/L) \times 100
\]

- \(A_s\) = absorbance of the Sample solution
- \(A_i\) = absorbance of the Standard solution
- \(C_i\) = concentration of the Standard solution (mg/mL)
- \(V\) = volume of Medium, 1000 mL
- \(L\) = label claim (mg/Capsule)

**Tolerances:** NLT 80% (Q) of the labeled amount of fenofibrate (C_{20}H_{31}ClO_{4}) is dissolved.

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

**Medium:** 0.05 M sodium lauryl sulfate in water; 1000 mL

**Apparatus 2:** 75 rpm, with helix sinkers or hoseclamp

**Times:** 30 min for products labeled to contain 67, 134, and 200 mg; 40 min for products labeled to contain 43 and 130 mg

**Standard stock solution:** 0.5 mg/mL of USP Fenofibrate RS in Medium prepared as follows. Dissolve a suitable quantity of USP Fenofibrate RS, taken in a suitable volumetric flask, in about 6% of the total volume of methanol, and dilute with Medium to volume.

**Standard solution:** Prepare solutions of USP Fenofibrate RS in Medium as per Table 1 from Standard stock solution.

### Table 1

<table>
<thead>
<tr>
<th>Capsule Strength (mg)</th>
<th>Concentration (mg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>67</td>
<td>0.065</td>
</tr>
<tr>
<td>130 and 134</td>
<td>0.13</td>
</tr>
<tr>
<td>200</td>
<td>0.2</td>
</tr>
<tr>
<td>43</td>
<td>0.045</td>
</tr>
</tbody>
</table>

**Sample solution:** Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size, discarding the first 3 mL of the filtrate.

**Instrumental conditions**

(See Ultraviolet-Visible Spectroscopy (857),)

**Mode:** Spectrophotometry

**Detector:** UV 291 nm

**Path length:** 0.1-cm flow cell

**Analysis**

**Samples:** Standard solution and Sample solution

Calculate the percentage of the labeled amount of fenofibrate (C_{20}H_{31}ClO_{4}) dissolved:

\[
\text{Result} = (A_s/A_i) \times C_i \times V \times (1/L) \times 100
\]

- \(A_s\) = absorbance of the Sample solution
- \(A_i\) = absorbance of the Standard solution
- \(C_i\) = concentration of the Standard solution (mg/mL)
- \(V\) = volume of Medium, 1000 mL
- \(L\) = label claim (mg/Capsule)

**Tolerances:** NLT 80% (Q) of the labeled amount of fenofibrate (C_{20}H_{31}ClO_{4}) is dissolved.

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

**Medium:** 0.025 M sodium lauryl sulfate in water; 1000 mL, deaerated

**Apparatus 2:** 75 rpm, with suitable sinkers

**Time:** 20 min

**Standard stock solution:** 0.5 mg/mL of USP Fenofibrate RS in methanol. Sonicate if necessary.

**Standard solution:** 12.5 µg/mL of USP Fenofibrate RS in methanol. Sonicate if necessary.

**Sample solution:** Pass a portion of the solution under test through a suitable filter of 0.45-µm pore size and discard the first few milliliters. Dilute with Medium, if necessary.

### Table 2

<table>
<thead>
<tr>
<th>Capsule Strength (mg)</th>
<th>Concentration (µg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>12.0</td>
</tr>
</tbody>
</table>
### Table 2 (continued)

<table>
<thead>
<tr>
<th>Capsule Strength (mg)</th>
<th>Concentration (µg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90</td>
<td>13.5</td>
</tr>
</tbody>
</table>

**Instrumental conditions**  
(See Ultraviolet-Visible Spectroscopy (857).)  
**Mode:** Spectrophotometry  
**Detector:** UV 290 nm  
**Blank:** Medium  
**Analysis**  
**Samples:** Standard solution and Sample solution  
Calculate the percentage of the labeled amount of fenofibrate \((C_{20}H_{21}ClO_4)\) dissolved:

\[
\text{Result} = \left(\frac{A_i}{A_j}\right) \times \left(\frac{C_j}{L}\right) \times D \times V \times 100
\]

\(A_i\) = absorbance of the Sample solution  
\(A_j\) = absorbance of the Standard solution  
\(C_j\) = concentration of the Standard solution (mg/mL)  
\(L\) = label claim (mg/Capsule)  
\(D\) = dilution factor for the Sample solution  
\(V\) = volume of Medium, 1000 mL

**Tolerances:** NLT 80% \((Q)\) of the labeled amount of fenofibrate \((C_{20}H_{21}ClO_4)\) is dissolved.  

**Test 6:** If the product complies with this test, the labeling indicates that it meets USP Dissolution Test 6.  
**Dissolution Test 6** is suitable for products labeled to contain 200 mg of fenofibrate.  
**Medium, Solution A, Mobile phase, and System suitability:** Proceed as directed in Test 1.  
**Apparatus 2:** 75 rpm, with suitable sinkers  
**Time:** 60 min  
**Standard solution:** 0.2 mg/mL of USP Fenofibrate RS prepared as follows. Transfer a suitable amount of USP Fenofibrate RS into a suitable volumetric flask. Add methanol to 2% of the total volume of the flask and sonicate to dissolve. Dilute with Medium to volume.  
**Sample solution:** Pass a portion of the solution under test through a suitable PVDF filter of 0.45-µm pore size. Discard the first few milliliters of filtrate.  
**Chromatographic system:** Proceed as directed in Test 1 except for Run time.  
**Run time:** NLT 2 times the retention time of the fenofibrate  
**Analysis**  
**Samples:** Standard solution and Sample solution  
Calculate the percentage of the labeled amount of fenofibrate \((C_{20}H_{21}ClO_4)\) dissolved:

\[
\text{Result} = \left(\frac{r_i}{r_j}\right) \times \left(\frac{C_j}{L}\right) \times V \times (1/L) \times 100
\]

\(r_i\) = peak response of fenofibrate from the Sample solution  
\(r_j\) = peak response of fenofibrate from the Standard solution  
\(C_j\) = concentration of USP Fenofibrate RS in the Standard solution (mg/mL)  
\(V\) = volume of Medium, 1000 mL  
\(L\) = label claim (mg/Capsule)  

**Tolerances:** NLT 80% \((Q)\) of the labeled amount of fenofibrate \((C_{20}H_{21}ClO_4)\) is dissolved.  

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**Change to read:**

- **Uniformity of Dosage Units** (905);  
  - Meet the requirements.  

**Procedure for content uniformity**  
**Solution A, Mobile phase, Standard solution, Chromatographic system, System suitability, and Analysis:** Proceed as directed in the Assay, except to prepare the Sample stock solution and Sample solution as follows.  
**Sample stock solution:** Place 1 Capsule in a suitable volumetric flask, add Solution A to 10%–20% of the final volume, and stir for 20 min to disintegrate the Capsule. Fill the flask to about 80% with methanol, sonicate for 10 min, and stir for 15 min. Dilute with methanol to volume to obtain a solution having a known concentration of about 0.4–0.7 mg/mL of fenofibrate, based on the label claim.  
**Sample solution:** Nominally 60–70 µg/mL of fenofibrate, from the Sample stock solution, in Mobile phase. Pass a portion of this solution through a PVDF filter of 0.45-µm pore size, discarding the first 3 mL.  

**IMPURITIES**

**Change to read:**

- **Organic Impurities**  
  - Use Sample solution 2 for Capsules labeled to meet the requirements of Dissolution Test 2. For all other products, use Sample solution 1.  
  - Solution A: 136 mg/L of potassium phosphate. Adjust with dilute phosphoric acid (1 in 10) to a pH of 2.9 ± 0.05.  
  - Mobile phase: Methanol and Solution A (4:1)  
  - System suitability solution: 0.67 mg/mL of USP Fenofibrate RS and 3.35 µg/mL of USP Fenofibrate Related Compound B RS in Mobile phase  
  - Standard solution: 3.35 µg/mL of USP Fenofibrate RS and 3.35 µg/mL of USP Fenofibrate Related Compound B RS in Mobile phase  
  - Sensitivity solution: 0.67 µg/mL of USP Fenofibrate RS and 0.67 µg/mL of USP Fenofibrate Related Compound B RS in Mobile phase  
  - Sample solution 1: Nominally 0.67 mg/mL of fenofibrate prepared as follows. Accurately weigh the contents of NLT 20 Capsules. Mix the contents, and transfer a weighed portion of the powder, equivalent to about 67 mg of fenofibrate, to a 100-mL volumetric flask. Add 80 mL of Mobile phase, sonicate for 10 min, stir for 15 min, and dilute with Mobile phase to volume. Pass a portion of this solution through a PVDF filter of 0.45-µm pore size, discarding the first 3 mL.  
  - Sample solution 2 (for Capsules labeled to meet the requirements of Dissolution Test 2): Nominally 0.67 mg/mL of fenofibrate prepared as follows. Weigh the contents of NLT 20 Capsules. Mix the contents, melt in an oven at 80°C for NLT 30 min, and homogenize. Allow the sample to solidify. Transfer a weighed portion of the sample, equivalent to about 67 mg of fenofibrate, to a 100-mL volumetric flask, dissolve in 30 mL of methanol with the aid of a mechanical shaker for NLT 4 h, and dilute with Mobile phase to volume. Pass through a PVDF filter of 0.45-µm pore size, discarding the first 1–2 mL.  

**Chromatographic system**  
(See Chromatography (621), System Suitability.)  
**Mode:** LC  
**Detector:** UV 285 nm  
**Column:** 4.6-mm × 15-cm; 5-µm packing L1

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Flow rate: 1 mL/min
Injection volume: 20 µL
Run time: NLT 3 times the retention time of the fenofibrate peak

System suitability
Samples: System suitability solution, Standard solution, and Sensitivity solution

Suitability requirements
Resolution: NLT 3.0 between fenofibrate and fenofibrate related compound B, System suitability solution
Tailing factor: NMT 2.0 for fenofibrate related compound B, System suitability solution
Relative standard deviation: NMT 2.0%, Standard solution
Signal-to-noise ratio: NLT 10 for the fenofibrate peak, Sensitivity solution

Analysis
Samples: Standard solution and designated Sample solution

Calculate the percentage of fenofibrate related compound B in the portion of Capsules taken:

\[
\text{Result} = \left( \frac{r_U}{r_S} \right) \times \left( \frac{C_S}{C_U} \right) \times 100
\]

\(r_U\) = peak response of fenofibrate related compound B from the Sample solution
\(r_S\) = peak response of fenofibrate from the Standard solution
\(C_S\) = concentration of fenofibrate related compound B in the Standard solution (mg/mL)
\(C_U\) = nominal concentration of fenofibrate in the Sample solution (mg/mL)

Calculate the percentage of any unspecified impurity in the portion of Capsules taken:

\[
\text{Result} = \left( \frac{r_U}{r_S} \right) \times \left( \frac{C_S}{C_U} \right) \times 100
\]

\(r_U\) = peak response of each unspecified impurity from the Sample solution
\(r_S\) = peak response of fenofibrate from the Standard solution
\(C_S\) = concentration of fenofibrate in the Standard solution (mg/mL)
\(C_U\) = nominal concentration of fenofibrate in the Sample solution (mg/mL)

Acceptance criteria
Individual impurities: NMT 0.5% for fenofibrate related compound B; NMT 0.2% for any other unspecified impurity
Total impurities: NMT 2.0%

ADDITIONAL REQUIREMENTS
• Packaging and Storage: Preserve in well-closed containers, and store at controlled room temperature.
• Labeling: When more than one Dissolution test is given, the labeling states the test used only if Test 1 is not used.
• USP Reference Standards (11)
USP Fenofibrate RS
USP Fenofibrate Related Compound B RS
2-[4-(4-Chlorobenzoyl)phenoxy]-2-methylpropanoic acid, or fenofibric acid.
\(C_{17}H_{15}ClO_4\) 318.75

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