**Coix lacryma-jobi Seed Powder**

**Final Authorized Version 1.0**

**Coix lacryma-jobi Seed Powder**

**DEFINITION**
The article consists of the dried ripe caryopsis, freed from the shell, of *Coix lacryma-jobi* var. *ma-yuen* (Rom.Caill.) Stapf (Family Poaceae, alt. Gramineae), collected in the fall, reduced to a fine or very fine powder. It contains NLT 3.5% of triglycerides calculated as the sum of trilinolein (C$_{57}$H$_{98}$O$_{6}$), 1,2-dilinoleoyl-3-palmitin (C$_{55}$H$_{100}$O$_{6}$), 1,2-dilinoleoyl-3-olein (C$_{57}$H$_{100}$O$_{6}$), 1-palmitoyl-2-oleoyl-3-linolein (C$_{55}$H$_{100}$O$_{6}$), 1,2-dioleoyl-3-linolein (C$_{57}$H$_{102}$O$_{6}$), 1,2-dioleoyl-3-palmitin (C$_{55}$H$_{102}$O$_{6}$), and triolein (C$_{57}$H$_{104}$O$_{6}$), on the dried basis.

**POTENTIAL CONFOUNGING MATERIALS**

*Coix lacryma-jobi* L.

*Coix puellarum* Balansa

Other varities of *Coix lacryma-jobi*, such as *Coix lacryma-jobi* var. *stenocarpa* Olive.

*Hordeum vulgare* L.

*Sorghum bicolor* (L.) Moench

**CONSTITUENTS OF INTEREST**

**Triglycerides:** Trilinolein, 1,2-dilinoleoyl-3-palmitin, 1,2-dilinoleoyl-3-olein, 1-palmitoyl-2-oleoyl-3-linolein, 1,2-dioleoyl-3-linolein, 1,2-dioleoyl-3-palmitin, and triolein

**Fatty acid:** Oleic acid

**IDENTIFICATION**

**• A. BOTANICAL CHARACTERISTICS**

**Macroscopic:** Pale whitish

**Microscopic:** Starch granules numerous and often in clumps, simple granule subround or polyhedral, 2–20 μm in diameter with stellate, y-shaped hilium; compound granule rare, usually consisting of 2–3 simple granules.

**• B. HPTLC (See High-Performance Thin-Layer Chromatography Procedure for Identification of Articles of Botanical Origin <203>.)**

**Standard solution A:** 2 mg/mL of USP Oleic Acid RS in methanol

**Standard solution B:** 40 mg/mL (about 50 μL/mL) of USP Coix Seed Oil Extract RS in methylene chloride

**Sample solution:** Sonicate 0.5 g of *Coix lacryma-jobi* Seed Powder in 5 mL of methylene chloride for 10 min. Centrifuge at 5000 rpm for 5 min and use the supernatant.

**Chromatographic system**
**Test conditions** and **System suitability:** See *Identification of Fixed Oils by Thin-Layer Chromatography* <202>, *Method II* [3].

**Analysis**

**Samples:** *Standard solution A, Standard solution B, and Sample solution*

Apply the Samples as bands to a suitable HPTLC plate and dry in air. Develop in a saturated chamber up to 7 cm, remove the plate from the chamber, and dry in air. Treat the plate with *Spray reagent*, heat at 120° for 3 min, and examine in white light.

**Acceptance criteria:** The Sample solution exhibits, in the upper half, a band corresponding in *R*<sub>F</sub> and color to the band due to oleic acid in Standard solution A. The Sample solution exhibits additional bands corresponding to similar bands in Standard solution B; these include one band right above oleic acid with the same color as it, a cluster of four blue bands in the lower half, and some faint bands in the middle-third.

• **C. HPLC**

**Analysis:** Proceed as directed in the Assay for Content of Triglycerides.

**Acceptance criteria:** The Sample solution exhibits a peak with a retention time corresponding to triolein in Standard solution A; the peaks due to trilinolein, 1,2-dilinoleoyl-3-palmitin, 1,2-dilinoleoyl-3-olein, 1-palmitoyl-2-oleoyl-3-linolein, 1,2-dioleoyl-3-linolein, and 1,2-dioleoyl-3-palmitin are at retention times corresponding to the same triglycerides in Standard solution B and the reference chromatogram provided with the lot of USP Coix Seed Oil Extract RS being used. 1,2-Dilinoleoyl-3-olein and 1,2-dioleoyl-3-linolein have the most intense peaks; 1-palmitoyl-2-oleoyl-3-linolein and triolein have medium intensity peaks; trilinolein, 1,2-dilinoleoyl-3-palmitin, and 1,2-dioleoyl-3-palmitin have weak intense peaks (distinction from *Sorghum bicolor* (L) Moench, which has similar triglycerides peaks but all of them are weak intensity; and *Hordeum vulgare* L., which has four triglycerides peaks and all of them are weak intensity).

**ASSAY**

• **Content of Triglycerides**

**Mobile phase:** Methanol

**Standard solution A:** 0.20 mg/mL of USP Triolein RS in methanol. Sonicate to dissolve.

**Standard solutions:** Dilute Standard solution A with methanol to obtain solutions of 0.0125, 0.025, 0.05, 0.10, and 0.20 mg/mL of USP Triolein RS. Pass each through a membrane filter of 0.45-μm or finer pore size.

**Standard solution B:** 0.5 mg/mL of USP Coix Seed Oil Extract RS in methanol

**Sample solution:** Accurately transfer about 500 mg of *Coix lacryma-jobi* Seed Powder into a suitable flask and add 35 mL of methanol. Sonicate for 30 min and filter with the aid of vacuum into a vacuum flask. Rinse the flask and the residue left in the flask with 5 mL of methanol, and wash the residue and paper on the filter using the rinsing. Repeat the rinse and wash procedure one more time. Transfer the filtrate into a 50-mL volumetric flask. Wash the vacuum flask with methanol, combine the washing into the volumetric flask, dilute with methanol to volume, and mix. Before injection, pass through a membrane filter of 0.45-μm or finer pore size, and discard the first portion of the filtrate. [**Note**—The filter and vacuum flask should be dry.]

**Chromatographic system**

(See *Chromatography* <621>, System Suitability.)

**Mode:** LC

**Detector:** Evaporative light-scattering. [**Note**—The parameters of the detector are adjusted to achieve the best signal-to-noise ratio, according to manufacturer recommendations.]

**Column:** 3.0-mm × 10-cm; 2.7-μm packing L7 (similar to AMT HALO™ C8 or Agilent Poroshell 120 EC
Column temperature: 20°
Flow rate: 0.3 mL/min
Injection volume: 5 µL

System suitability

Samples: Standard solution A, Standard solutions, and Standard solution B

Suitability requirements

Chromatogram similarity: The chromatogram of Standard solution B is similar to the reference chromatogram provided with the lot of USP Coix Seed Oil Extract RS being used.

Resolution: NLT 1.5 between the peak of 1,2-dilinoleoyl-3-olein and the small peak following it, Standard solution B

Tailing factor: NMT 1.5 for triolein peak, Standard solution A

Relative standard deviation: NMT 5.0% for the triolein peak, Standard solution A

Signal-to-noise ratio: NLT 15 for the triolein peak, Standard solutions of 0.0125 mg/mL

Correlation coefficient: NLT 0.995 for the regression line as determined in Analysis, Standard solutions

Analysis

Samples: Standard solutions, Standard solution B, and Sample solution

Using the chromatograms of Standard solutions and Standard solution B, and the reference chromatogram provided with the lot of USP Coix Seed Oil Extract RS being used, identify the retention times of the peaks corresponding to trilinolein, 1,2-dilinoleoyl-3-palmitin, 1,2-dilinoleoyl-3-olein, 1-palmitoyl-2-oleoyl-3-linolein, 1,2-dioleoyl-3-linolein, 1,2-dioleoyl-3-palmitin, and triolein in the Sample solution. [Note—See Table 1 for the approximate relative retention times.]

Table 1

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Approximate Relative Retention Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trilinolein</td>
<td>0.61</td>
</tr>
<tr>
<td>1,2-Dilinoleoyl-3-palmitin</td>
<td>0.67</td>
</tr>
<tr>
<td>1,2-Dilinoleoyl-3-olein</td>
<td>0.71</td>
</tr>
<tr>
<td>1-Palmitoyl-2-oleoyl-3-linolein</td>
<td>0.79</td>
</tr>
<tr>
<td>1,2-Dioleoyl-3-linolein</td>
<td>0.84</td>
</tr>
<tr>
<td>1,2-Dioleoyl-3-palmitin</td>
<td>0.94</td>
</tr>
<tr>
<td>Triolein</td>
<td>1.00</td>
</tr>
</tbody>
</table>

Plot the logarithms of peak responses versus the logarithms of concentrations, in mg/mL, of triolein from the Standard solutions, and determine the regression line using a least-squares analysis; or, establish a linear regression equation using a least-squares analysis according to the logarithms of the peak responses versus logarithms of concentrations, in mg/mL, of triolein from the Standard solutions. Determine the concentration, C, in mg/mL, of the relevant analyte in the Sample solution by regression line or linear regression equation.

Separately calculate the percentages of trilinolein, 1,2-dilinoleoyl-3-palmitin, 1,2-dilinoleoyl-3-olein, 1-palmitoyl-2-oleoyl-3-linolein, 1,2-dioleoyl-3-linolein, 1,2-dioleoyl-3-palmitin, and triolein in the portion of Coix lacryma-jobi Seed Powder taken:
\[ \text{Result} = C \times \left( \frac{V}{W} \right) \times 100 \]

- \( C \) = concentration of the relevant analyte in the \textit{Sample solution} as determined above (mg/mL)
- \( V \) = volume of the \textit{Sample solution} (mL)
- \( W \) = weight of \textit{Coix lacryma-jobi} Seed Powder taken to prepare the \textit{Sample solution} (mg)

Add the percentages of trilinolein, 1,2-dilinoleoyl-3-palmitin, 1,2-dilinoleoyl-3-olein, 1-palmitoyl-2-oleoyl-3-linolein, 1,2-dioleoyl-3-linolein, 1,2-dioleoyl-3-palmitin, and triolein.

\textbf{Acceptance criteria:} NLT 3.5\% on the dried basis

\section*{CONTAMINANTS}

\textbf{Test for Zearalenone}

\textbf{Mobile phase:} Acetonitrile and water (1:1)

\textbf{Standard stock solution:} 250 ng/mL of zearalenone in methanol

\textbf{Standard solution:} Accurately transfer 1 mL of \textit{Standard stock solution} to a 10-mL volumetric flask and add methanol to volume.

\textbf{Sample solution:} Accurately transfer about 20 g of \textit{Coix lacryma-jobi} Seed Powder to a suitable centrifuge tube. Add 4.0 g of sodium chloride, then accurately add 100 mL of 90\% acetonitrile. Mix for 2 min with a high speed disperser (not lower than 11000 rpm), and centrifuge for 5 min (4000 rpm). Immediately pipet 10 mL of the supernatant into a 50-mL volumetric flask, add water to volume, mix, and centrifuge. Pipet 10 mL of the supernatant onto the Immunoaffinity Column (ZearalaTest™), at a flow rate of 3 mL/min. Wash the column with 10 mL of water at a flow rate of 6 mL/min, let the column run dry, and discard the water elution. Wash the column with 1.5 mL of methanol at a flow rate of 1 mL/min, collect the methanol elution into a 2-mL volumetric flask, let the column run dry, add methanol to volume, and mix.

\textbf{Chromatographic system}

(See \textit{Chromatography <621>}, \textit{System Suitability}.)

\textbf{Mode:} LC
\textbf{Detector:} Fluorescence detector set at excitation wavelength (Ex) 232 nm and emission wavelength (Em) 460 nm.
\textbf{Column:} 4.6-mm × 15-cm; 5-µm packing L1 (similar to Inertsil ODS-3)
\textbf{Column temperature:} 30°
\textbf{Flow rate:} 1.0 mL/min
\textbf{Injection volume:} 20 µL of \textit{Sample solution}

\textbf{System suitability}

\textbf{Sample:} \textit{Standard solution}

\textbf{Suitability requirements}

\textbf{Theoretical plates:} NLT 10000
\textbf{Relative standard deviation:} NMT 10.0\% for the zearalenone peak
\textbf{Correlation coefficient:} NLT 0.999 for the regression line as determined in the \textit{Analysis}

\textbf{Analysis}

\textbf{Samples:} \textit{Standard solution} and \textit{Sample solution}

Inject the \textit{Standard solution} with volumes of 5, 10, 15, 20, and 25 µL, and separately measure the peak areas of each injection. Plot the peak responses versus the amount, in ng, of zearalenone in the \textit{Standard solution} injections, and determine the regression line using a least-squares analysis.
Using the chromatogram of the *Standard solution* identify the retention time of the peak corresponding to zearalenone in the *Sample solution.*

From the graph, determine the content, \( C \), in ng, of zearalenone in the *Sample solution.*

Calculate the content, in ng/g, of zearalenone in the portion of *Coix lacryma-jobi* Seed Powder taken:

\[
\text{Result} = 5000 \times \left( \frac{C}{W} \right)
\]

\( C \) = content of zearalenone as determined above (ng)

\( W \) = weight of *Coix lacryma-jobi* Seed Powder taken to prepare the *Sample solution* (g)

**Acceptance criteria:** NMT 60 ng/g on the dried basis

- **Articles of Botanical Origin**<561> [5] *Limits of Elemental Impurities* [5]: Meets the requirements
- **Articles of Botanical Origin**<61> [6] *General Method for Pesticide Residues Analysis* [6]: Meets the requirements

- **Microbial Enumeration Tests**<61>: [7][8] The total aerobic bacterial count does not exceed \( 10^5 \) cfu/g, the total combined molds and yeasts count does not exceed \( 10^3 \) cfu/g, and the bile-tolerant Gram-negative bacteria does not exceed \( 10^3 \) cfu/g.
- **Tests for Specified Microorganisms**<62>: [8][9] Meets the requirements of the tests for the absence of *Salmonella* species and *Escherichia coli*
- **Articles of Botanical Origin**<561>, [6] *Test for Aflatoxins* [6]: Meets the requirements

**Specific Tests**

- **Articles of Botanical Origin**<561>, [6] *Alcohol-Soluble Extractives, Method 1* [6]: Using absolute alcohol, NLT 5.5%
- **Loss on Drying**<731> [9]
  - **Sample:** 2.0 g of *Coix lacryma-jobi* Seed Powder
  - **Analysis:** Dry the Sample at 105° for 5 h.
  - **Acceptance criteria:** NMT 15.0%
  - **Analysis:** 2.0 g of *Coix lacryma-jobi* Seed Powder
  - **Acceptance criteria:** NMT 3.0%

**Additional Requirements**

- **Packaging and Storage:** Preserve in well-closed containers, protected from light and moisture, and store at 4°–10°.
- **Labeling:** The label states the Latin binomial and the part(s) of the plant contained in the article.
- **USP Reference Standards**<11> [20]
  - USP Coix Seed Oil Extract RS
  - USP *Oleic Acid* [2] RS
  - USP Triolein RS

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