# PHARMACOPOEIAL DISCUSSION GROUP

# CORRECTION CODE: E-27 NAME: METHYL PARAHYDROXYBENZOATE (Correction 2 to revision 1 of the sign-off document signed 10 June 2009)

## Item to be corrected:

- Addition of CAS numbers: [99-76-3]
- Appearance of solution/color: addition of comparison with alcohol

Attribute	EP	JP	USP
Definition	+	+	+
Identification A	+	+	+
(melting point)*			
Identification B	+	+	+
(IR)			
Appearance of	+	+	, +
solution/color	·		
Acidity	+	+	+
Related	+	+	+
substances**			
Sulphated ash	+	+	+
Assay	+	+	+

\* Melting point: listed in JP as a test and not as part of identification

\*\* Related substances: JP uses the term "relative response factor" instead of "correction factor"

# Legend

+ will adopt and implement

- will not stipulate

# Non-harmonised attributes

Characters, Storage

## Local requirements

Ph. Eur.	JP	USP
Second identification	Related substances: test for	none
(melting point, TLC)	required detectability, system	
	repeatability	
	Heavy metals (20 ppm)	
	Assay: column temperature	

## **Reagents and reference materials**

Each pharmacopoeia will adapt the text to take account of local reference materials and reagent specifications.

1-10 Km

# **European Pharmacopoeia**

Signature 🗲

Name

VIELCE COMME

Date

22-20-2020

Japanese Pharmacopoeia

Signature

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Name

Date

Date

for 7. Yoshida Haruhio Oluda

16 Dec / 2020

**United States Pharmacopeia** 

JLT. Mm KEVIN MODRE

19-NOU-2020

# **E27 - METHYL PARAHYDROXYBENZOATE**



C<sub>8</sub>H<sub>8</sub>O<sub>3</sub> [99-76-3]

DEFINITION Methyl 4-hydroxybenzoate. *Content:* 98.0 per cent to 102.0 per cent.

# **IDENTIFICATION**

A. Melting point: 125 °C to 128 °C.

B. Infrared absorption spectrophotometry.

Record the infrared absorption spectrum of methyl parahydroxybenzoate and compare with the Reference Spectrum or the spectrum obtained with the Reference Standard: the transmission minima correspond in position and relative size.

#### TESTS

Solution S. Dissolve 1.0 g in *alcohol* and dilute to 10 ml with the same solvent.

Appearance of solution. Solution S is clear and not more intensely coloured than *alcohol* or the reference solution.

# Primary solutions:

- Ferric chloride primary solution: a 45.0 g/l solution of ferric chloride (FeCl<sub>3</sub>, 6H<sub>2</sub>O).
- Cobalt chloride primary solution: a 59.5 g/l solution of cobalt chloride (CoCl<sub>2</sub>, 6H<sub>2</sub>O).
- Copper sulphate primary solution: a 62.4 g/l solution of copper sulphate (CuSO<sub>4</sub>, 5H<sub>2</sub>O).

## Reference solution:

To 5.0 ml of cobalt chloride primary solution, 12.0 ml of ferric chloride primary solution and 2.0 ml of copper sulphate primary solution, add hydrochloric acid (10 g/l HCl) to make 1000.0 ml.

Acidity. To 2 ml of solution S add 3 ml of *alcohol*, 5 ml of *carbon dioxide-free water* and 0.1 ml of *bromocresol green solution*. Not more than 0.1 ml of 0.1 M sodium hydroxide is required to change the colour of the indicator to blue.

## Related substances. Liquid chromatography.

*Test solution.* Dissolve 50.0 mg of the sample to be examined in 2.5 ml of *methanol* and dilute to 50.0 ml with the mobile phase. Dilute 10.0 ml of this solution to 100.0 ml with the mobile phase.

Reference solution (a). Dissolve 5 mg each of 4-hydroxybenzoic acid R and the substance to be examined in the mobile phase and dilute to 100.0 ml with the same solvent. Dilute 1 ml of this solution to 10.0 ml with the mobile phase.

*Reference solution (b).* Dissolve 50.0 mg of *methyl parahydroxybenzoate CRS* in 2.5 ml of *methanol* and dilute to 50.0 ml with the mobile phase. Dilute 10.0 ml of this solution to 100.0 ml with the mobile phase.

*Reference solution (c).* Dilute 1.0 ml of the test solution to 20.0 ml with the mobile phase. Dilute 1.0 ml of this solution to 10.0 ml with the mobile phase.

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Column:

— *size:* l = 0.15 m,  $\emptyset = 4.6 \text{ mm}$ ;

--- stationary phase: octadecylsilyl silica gel for chromatography (5 μm).

Mobile phase: 6.8 g/l solution of potassium dihydrogen phosphate, methanol (35:65 V/V).

Flow rate: 1.3 ml/min.

Detection: 272 nm.

Injection: 10 µl of the test solution and reference solutions (a) and (c).

Run time: 5 times the retention time of methyl parahydroxybenzoate.

*Relative retention* with reference to methyl parahydroxybenzoate (retention time = about 2.3 min): 4-hydroxybenzoic acid = about 0.6.

# System suitability:

- *resolution:* minimum of 2.0 between the peaks due to 4-hydroxybenzoic acid and methyl parahydroxybenzoate in the chromatogram obtained with reference solution (a).

# Limits:

- *correction factor*: for the calculation of content, multiply the peak area of 4-hydroxybenzoic acid by 1.4;

- 4-hydroxybenzoic acid: not more than the area of the principal peak in the chromatogram obtained with reference solution (c) (0.5 per cent);

- *unspecified impurities*: for each impurity, not more than the area of the principal peak in the chromatogram obtained with reference solution (c) (0.5 per cent);

- *total:* not more than twice the area of the principal peak in the chromatogram obtained with reference solution (c) (1.0 per cent);

- disregard limit: 0.2 times the area of the principal peak in the chromatogram obtained with reference solution (c) (0.1 per cent).

Sulphated ash: maximum 0.1 per cent, determined on 1.0 g.

## ASSAY

Liquid chromatography as described in the test for related substances with the following modification.

Injection: test solution and reference solution (b).

## System suitability:

- *repeatability:* maximum relative standard deviation of 0.85 per cent after 6 injections of reference solution (b).

Calculate the percentage content of  $C_8H_8O_3$  in the sample to be examined from the peak areas in the chromatograms obtained with test solution and reference solution (b) and the declared content of *methyl parahydroxybenzoate CRS*.

## REAGENTS

## Bromocresol green solution.

Dissolve 50 mg of *bromocresol green* in 0.72 ml of 0.1 *M sodium hydroxide* and 20 ml of *alcohol* and dilute to 100 ml with *water*.

*Test for sensitivity.* To 0.2 ml of the bromocresol green solution add 100 ml of *carbon dioxide-free water*. The solution is blue. Not more than 0.2 ml of 0.02 M hydrochloric acid is required to change the colour to yellow.

Colour change: pH 3.6 (yellow) to pH 5.2 (blue).

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