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Krma: metode vzorčenja in analize - Določevanje karbadoksa in olakvindoksa s HPLC/UV

Animal feeding stuffs: Methods of sampling and analysis - Determination of carbadox and olaquindox by HPLC/UV

Futtermittel - Probenahme- und Untersuchungsverfahren - Bestimmung von Carbadox und Olaquindox mittels HPLC/UVFANDARD PREVIEW

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Aliments des animaux - Méthodes d'échantillonnage et d'analyse - Détermination des teneurs en carbadox et olaquindox par CLHP/UV<sub>2017</sub>

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#### **English Version**

### Animal feeding stuffs: Methods of sampling and analysis - Determination of carbadox and olaquindox by HPLC/UV

Aliments des animaux - Méthodes d'échantillonnage et d'analyse - Détermination des teneurs en carbadox et olaquindox par CLHP/UV Futtermittel - Probenahme- und Untersuchungsverfahren - Bestimmung von Carbadox und Olaquindox mittels HPLC/UV

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EUROPEAN COMMITTEE FOR STANDARDIZATION COMITÉ EUROPÉEN DE NORMALISATION EUROPÄISCHES KOMITEE FÜR NORMUNG

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#### **European foreword**

This document (EN 16930:2017) has been prepared by Technical Committee CEN/TC 327 "Animal feeding stuffs - Methods of sampling and analysis", the secretariat of which is held by NEN.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by January 2018, and conflicting national standards shall be withdrawn at the latest by January 2018.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. CEN [and/or CENELEC] shall not be held responsible for identifying any or all such patent rights.

This document has been prepared under a mandate given to CEN by the European Commission and the European Free Trade Association.

WARNING — The use of this protocol involves hazardous materials, operations and equipment. This protocol does not purport to address all the safety problems associated with its use. It is the responsibility of the user of this protocol to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

According to the CEN-CENELEC Internal Regulations, the national standards organisations of the following countries are bound to implement this European Standard: Austria, Belgium, Bulgaria, Croatia, Cyprus, Czech Republic, Denmark, Estonia, Finland, Former Yugoslav Republic of Macedonia, France, Germany, Greece, Hungary, Iceland, Ireland, Italy, Latvia, Lithuania, Luxembourg, Malta, Netherlands, Norway, Poland, Portugal, Romania, Serbia, Slovakia, Slovenia, Spain, Sweden, Switzerland, Turkey and the United Kingdomstandards.iteh.ai/catalog/standards/sist/f68515fd-3dba-4bc1-9229-

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#### 1 Scope

This European Standard specifies a high performance liquid chromatographic – UV detection (HPLC-UV) method for the simultaneous determination of two growth promoters Carbadox and Olaquindox contents in compound feeds and raw materials at levels ranging from the limit of quantification to 100 mg/kg.

The limit of quantification of the method has been demonstrated to be lower than 3 mg/kg for olaquindox and 4 mg/kg for carbadox.

#### 2 Normative references

The following documents, in whole or in part, is normatively referenced in this document and is indispensable for its application. For undated reference, the latest edition of the referenced document (including any amendments) applies.

EN ISO 6498, Animal feeding stuffs - Guidelines for sample preparation (ISO 6498)

#### 3 Principle

The two growth promoters are extracted from the sample with a mixture of methanol and water 1:1, v:v. The extract of animal feeds is purified through a short open glass aluminium oxide column or by an alumina Solid Phase Extraction (SPE).

The final extract is analysed by reversed phase HPLC with UV detection at a wavelength of 375 nm. Alternatively, for confirmation purposes, a Diode Array Detector (DAD) can be used.

The presence of furazolidone can interfere with the determination of carbadox.

If a DAD is used, many other veter<u>inary drugs\_or7</u>feed additives can be detected (ronidazole, meticlorpindol, nitrofurazone, dimetridazola furaltadon, sulphonamides...), but their signals do not interfere with the target ones.

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In some special feeds, matrix interfering peaks may be present, very close to the carbadox peak.

#### 4 Reagents and materials

WARNING — Persons using this European Standard should be familiar with normal laboratory practice. This standard does not purport to address all of the safety problems, if any, associated with its use. It is the responsibility of the user to establish appropriate safety and health practices and to ensure compliance with any regulatory conditions.

- a) This method requires the use of solutions of Carbadox and Olaquindox. These substances were used as growth-promoting feed additives for piglets. Carbadox and Olaquindox are chemotherapeutics belonging to the quinoxaline group. They are suspected to be teratogen and mutagen. Avoid inhalation of and exposure to the toxic standard materials and solutions;
- b) These 2 growth promoters are subject to light degradation. Protect analytical work adequately from daylight, and keep standard solutions protected from light by using amber glassware, amber vials or aluminium foil.

Use only reagents as analytical grade at least unless otherwise stated.

- **4.1 Water**, demineralized or deionized or at least equivalent
- 4.2 Methanol
- 4.3 Acetonitrile
- 4.4 Glacial acetic acid, minimum purity 96 %
- 4.5 Ammonium acetate water free salt, CH<sub>3</sub>CO<sub>2</sub>NH<sub>4</sub>
- 4.6 Extraction solvent: methanol water mixture (1:1 v:v)

Combine equal volumes of methanol (4.2) and water (4.1). Mix well.

NOTE Only for 8.1 purposes, it is possible to combine equal volume of technical methanol (4.14) and water (4.1). It is essential to mix well.

#### **4.7 Ammonium acetate buffer,** 25 mM, pH = 4,35

Weigh 2,0 g of ammonium acetate (4.5) to the nearest 0,1 g, into a 1000 ml volumetric flask. Dissolve in 900 ml of water (4.1). Add 3,0 ml of glacial acetic acid (4.4). Adjust (5.1) the pH to pH = 4,35 with acetic acid (4.4), if necessary. Dilute to the mark with water (4.1) and mix well.

#### **4.8 Mobile phase for HPLC**: acetonitrile: buffer mixture (10:90; v:v)

Transfer 100 ml of acetonitrile (4.3) into a 1000 ml volumetric flask. Dilute to the mark with ammonium acetate buffer (4.7). Filter the eluent through a 0,45  $\mu$ m cellulose acetate membrane filter (5.9) using a solvent filtration system (5.2). If necessary, perform degassing for 5 min in an ultrasonic bath (5.3).

NOTE This mobile phase is suitable for the three recommended columns in 5.11.8 If another column is used, optimization of the mobile phase composition may be necessary before performing real analysis. See 8.4.1.

#### **4.9 Carbadox dissolving solvent**: methanol: acetonitrile mixture (1:1; v:v)

Combine equal volumes of methanol (4.2) and acetonitrile (4.3). Mix well.

#### 4.10 Reference standards

Guaranteed purity is required for each lot of reference standard.

- 4.10.1 Carbadox, 3- (2-quinoxalinyl methylene) carbazic acid methyl ester N,N'-dioxide
- 4.10.2 Olaquindox, (2-[N-2'-(hydroxyethyl) carbamoyl] 3-methyl quinoxaline) N,N'-dioxide

#### 4.11 Standard solutions

Protect all standard solutions from daily light

#### 4.11.1 Carbadox stock standard solution ca. 100 μg/ml

Weigh 25 mg of carbadox (4.10.1), to the nearest 0,1 mg, into a 250 ml amber volumetric flask. Dissolve in 200 ml of mixture (4.9). Mix well and place the flask in an ultrasonic bath (5.3) until total dissolution. Allow to cool down to room temperature, dilute to the mark with mixture (4.9) and mix well. Calculate the accurate concentration taking into account the purity of the reference standard (4.10.1).

Prepare fresh every month. Store in the dark at 0 °C to 8 °C (5.12).

#### 4.11.2 Olaquindox stock standard solution ca. 250 µg/ml

Weigh 50 mg of olaquindox (4.10.2), to the nearest 0,1 mg, into a 200 ml amber volumetric flask. Dissolve in about 190 ml of water (4.1). Mix well and place the flask in an ultrasonic bath (5.3) until total dissolution. Allow to cool down to room temperature. Dilute to the mark with water (4.1) and mix well. Calculate the accurate concentration taking into account the purity of the reference standard (4.10.2).

Prepare fresh every month. Store in the dark at 0 °C to 8 °C (5.12).

#### 4.11.3 Calibrations solutions

#### **4.11.3.1** Carbadox/olaquindox calibration solution ca. 10 μg/ml

Pipette 5 ml of the carbadox stock standard solution (4.11.1) and 2 ml of the olaquindox stock standard solution (4.11.2) in a 50 ml volumetric flask. Dilute to the mark with the extraction solvent (4.6) and mix well.

Prepare fresh for each series of samples.

#### 4.11.3.2 Carbadox/olaquindox calibration solution ca. 5 μg/ml

Pipette 5 ml of the carbadox stock standard solution (4.11.1) and 2 ml of the olaquindox stock standard solution (4.11.2) in a 100 ml volumetric flask. Dilute to the mark with the extraction solvent (4.6) and mix well.

Prepare fresh for each series of samples. DARD PREVIEW

#### 4.11.3.3 Carbadox/olaquindox calibration solution ca. 2,5 $\mu g/ml$

Pipette 5 ml of the carbadox stock standard solution (4.11.1) and 2 ml of the olaquindox stock standard solution (4.11.2) in a 200 ml volumetric flask. Pilute to the mark with the extraction solvent (4.6) and mix well.

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Prepare fresh for each series of samples.

#### 4.11.3.4 Carbadox/olaquindox calibration solution ca. 1 μg/ml

Pipette 5 ml of the Carbadox/olaquindox calibration solution ca.  $10 \,\mu\text{g/ml}$  (4.11.3.1) in a 50 ml volumetric flask. Dilute to the mark with the extraction solvent (4.6) and mix well.

Prepare fresh for each series of samples.

#### 4.11.3.5 Carbadox/olaquindox calibration solution ca. 0,5 μg/ml

Pipette 5 ml of the Carbadox/olaquindox calibration solution ca.  $5 \,\mu g/ml$  (4.11.3.2) in a 50 ml volumetric flask. Dilute to the mark with the extraction solvent (4.6) and mix well.

Prepare fresh for each series of samples.

#### 4.11.3.6 Carbadox/olaquindox calibration solution ca. 0,25 μg/ml

Pipette 5 ml of the Carbadox/olaquindox calibration solution ca. 2,5  $\mu$ g/ml (4.11.3.3) in a 50 ml volumetric flask. Dilute to the mark with the extraction solvent (4.6) and mix well.

Prepare fresh for each series of samples.

- 4.12 Neutral aluminium oxide, Brockmann activity I
- **4.13 Neutral aluminium oxide SPE cartridge,** 2 cm<sup>3</sup>, 1850 mg
- 4.14 Technical methanol
- 5 Apparatus

Usual laboratory apparatus and, in particular, the following:

- 5.1 pH meter
- 5.2 Solvent filtration system, suitable for 0,45 µm membrane filters
- 5.3 Ultrasonic bath
- 5.4 Rotary on mechanical shaker
- 5.5 Centrifuge
- 5.6 Glass wool
- **5.7 Glass column for chromatography,** length 200 mm to 400 mm; internal diameter 10 mm, restricted at the end and fitted with a wad of glass wool (5.6) or equivalent column
- 5.8 Filter papers or glass microfibre filter

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- 5.9 Cellulose acetate membrane filters of pore size 0,45 µm
- 5.10 PVDF syringe filters of pore size 0.45 µm and adaptable syringes

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- 5.11 HPLC system
- **5.11.1 pump.** capable of maintaining a volume flow rate of 0.4 ml to 1.5 ml min<sup>-1</sup>
- **5.11.2 Column heater** set at 30 °C
- **5.11.3 Injection system,** with a loop suitable for 10 µl to 50 µl injections
- **5.11.4 UV detector,** suitable for measurements at a wavelength of 375 nm
- **5.11.5 Diode array detector** for confirmation purposes
- 5.11.6 Recorder or data acquisition system
- 5.11.7 Guard column, silica bounded C8
- **5.11.8 LC analytical column,**  $250 \times 3$  mm,  $5 \mu m$  particle size, packed with Lichrosorb  $C_8$  or Lichrospher  $C_{18}$  or equivalent column.

NOTE During the method validation, these recommended LC columns have proved to be fit for purpose, especially with samples containing interfering peaks close to the carbadox zone.

#### 5.12 Refrigerator or refrigerated chamber at a temperature set between 0 °C and 8 °C

#### Sampling

It is important that the laboratory receives a sample that is truly representative and has not been damaged or changed during transport or storage.

Sampling is not part of the method specified in this European Standard. Recommended sampling methods are given in:

- Annex I of the Commission Regulation (EU) No152/2009 amended by Commission Regulation (EU) No 691/2013 [1]
- EN ISO 6497 [12].

#### 7 Preparation of test sample

#### 7.1 General

Prepare the test sample in accordance with EN ISO 6498.

#### 7.2 Laboratory sample

Grind the laboratory sample (usually 50 g) so that is passes completely through a sieve with 1 mm apertures. Mix thoroughly.eh STANDARD PREVIEW

#### 7.3 Test sample

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The test sample consists of a representative and homogenized aliquot of the ground laboratory sample of at least 20 g. SIST EN 16930:2017

7.4 Test portion https://standards.iteh.ai/catalog/standards/sist/f68515fd-3dba-4bc1-9229-9761120d91ee/sist-en-16930-2017

Accurately weigh 10,0 g to the nearest 0,1 g of the thoroughly mixed test sample into a 250 ml conical flask. Note down the mass expressed in g. Submit it to the analysis procedure (8).

#### 8 Procedure

#### 8.1 Extraction of feeding stuffs containing 0,5 mg/kg to 100 mg/kg of growth promoters

Add 100,0 ml of extraction solvent (4.6) to the test portion (7.4), stopper and shake vigorously for 1 h or 2 h on the rotary shaker (5.4).

#### 8.2 Filtration

Filter the solution obtained in 8.1 through a paper or glass micro fibre filter (5.8).

Alternatively centrifugation (5.5) is possible, if filtration is too long, for example.

Use the filtrate or the supernatant for the purification step.

#### 8.3 Purification

#### 8.3.1 General

Apply 8.3.2 or 8.3.3.