

SLOVENSKI STANDARD oSIST prEN 17550:2020

01-november-2020

Krma: metode vzorčenja in analize - Določevanje karotenoidov v krmnih mešanicah in premiksih s tekočinsko kromatografijo visoke ločljivosti z ultravijolično (UV) detekcijo (HPLC-UV)

Animal feeding stuffs: Methods of sampling and analysis - Determination of carotenoids in animal compound feed and premixtures by high performance liquid chromatography - UV detection (HPLC-UV)

Futtermittel - Probenahme- und Untersuchungsverfahren - Bestimmung von Carotinoiden in Mischfuttermitteln und Vormischungen für Tiere mittels Reverse Hochdruckphase-Flüssigchromatographie-UV Detektion (RP-HPLC-UV)

oSIST prEN 17550:2020

Aliments des animaux - Méthodes d'échantillonnage et d'analyse - Détermination de la teneur en caroténoïdes des aliments composés et des prémélanges pour animaux par chromatographie liquide à haute performance couplée à une détection UV (CLHP-UV)

Ta slovenski standard je istoveten z: prEN 17550

ICS:

65.120 Krmila Animal feeding stuffs

71.040.50 Fizikalnokemijske analitske Physicochemical methods of

metode analysis

oSIST prEN 17550:2020 en,fr,de

oSIST prEN 17550:2020

iTeh STANDARD PREVIEW (standards.iteh.ai)

oSIST prEN 17550:2020 https://standards.iteh.ai/catalog/standards/sist/27ac68a6-b27b-4b8c-a185-75e7348fece0/osist-pren-17550-2020

EUROPEAN STANDARD NORME EUROPÉENNE EUROPÄISCHE NORM

DRAFT prEN 17550

September 2020

ICS 65.120; 71.040.50

English Version

Animal feeding stuffs: Methods of sampling and analysis - Determination of carotenoids in animal compound feed and premixtures by high performance liquid chromatography - UV detection (HPLC-UV)

Aliments des animaux - Méthodes d'échantillonnage et d'analyse - Détermination de la teneur en caroténoïdes des aliments composés et des prémélanges pour animaux par chromatographie liquide à haute performance couplée à une détection UV (CLHP-UV) Futtermittel - Probenahme- und
Untersuchungsverfahren - Bestimmung von
Carotinoiden in Mischfuttermitteln und
Vormischungen für Tiere mittels Reverse
Hochdruckphase-Flüssigchromatographie-UV
Detektion (RP-HPLC-UV)

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European foreword

This document (prEN 17550:2020) 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 document is currently submitted to the CEN Enquiry.

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

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1 Scope

This analytical procedure specifies a reverse phase high performance liquid chromatographic with UV detection (RP-HPLC-UV) method for the simultaneous determination of four authorized carotenoids in fish compound feed, namely astaxanthin (AXN), canthaxanthin (CXN), adonirubin (ADR) and astaxanthin dimethyldisuccinate (AXN DMDS), and of six authorized carotenoids in poultry feed, namely canthaxanthin (CXN); capsanthin (CSN), ethyl ester of beta-apo-8'-carotenoic acid (BACARE), citranaxanthin (CIXN), lutein (LUT) and zeaxanthin (ZEA) at levels ranging from ca. 2 to ca. 4 500 mg/kg (depending on the carotenoid). Beta-carotene (BCAR), authorized in compound feed for all animal species, was also added to the scope. The analytical procedure is fit for the purpose of quantitation of declared carotenoids and labelling confirmation. The procedure applies to natural and synthetic feed additives.

Xanthophyll esters like those of lutein, zeaxanthin and capsanthin that might be present in feed materials are not authorized feed additives and therefore not part of the scope of this method.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

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

3 Terms and definitions the STANDARD PREVIEW

No terms and definitions are listed in this document.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

- IEC Electropedia: available at http://www.electropedia.org/ 5e/348/cce0/osist-pren-17530-2020
- ISO Online browsing platform: available at https://www.iso.org/obp

4 Principle

The carotenoids are first disclosed through an enzymatic reaction at 50 °C. The samples are extracted with acetone by means of a pressurized liquid extraction instrument or by liquid solid extraction. The extracts are centrifuged and analysed by reverse phase HPLC with UV or Diode Array Detection. A common isosbestic wavelength of 410 nm is selected for the determination of the target analytes, thus ensuring that the various isomers of each of the carotenoids have similar absorbance coefficients. The quantitation is performed through external calibration.

NOTE The maximum contents of the carotenoids as established by the European regulations for the authorization of feed additives ([1], [2]), are expressed in terms of the sum of the all-trans and cis isomers. Therefore, the areas of the corresponding isomers in the HPLC chromatogram have to be summed up prior to quantification of the individual carotenoids.

5 Reagents and materials

WARNING 1 — Carotenoids are subject to light degradation. Protect analytical work adequately from day light, and keep standard solutions protected from light by using amber glassware, amber vials or aluminium foil.

WARNING 2 — Avoid inhalation of and exposure to the toxic standard materials and solutions thereof. Work under fume hood when handling the solvents and solutions. Wear safety glasses and protective clothing.

WARNING 3 — Always wear a safety mask when handling Hydromatrix™.

Unless otherwise specified, use only reagents of recognized analytical grade.

5.1 Protease with the ability to release carotenoids from their encapsulated form

NOTE Suitable proteases are available. 1

- **5.2 Purified water**, e.g. Milli-Q or equivalent ²
- 5.3 Butylated hydroxytoluene BHT
- **5.4 High purity diatomaceous earth suitable for PLE,** e.g. Hydromatrix[™], bulk support material ²
- 5.5 Acetone, HPLC gradeh STANDARD PREVIEW
- 5.6 Acetone, spectroscopic grade and ards.iteh.ai)
- **5.7 Acetonitrile,** HPLC grade oSIST prEN 17550:2020 https://standards.iteh.ai/catalog/standards/sist/27ac68a6-b27b-4b8c-a185-
- 5.8 Methyl tert-butyl ether tBME, HPLC grade ren-17550-2020
- **5.9 Tetrahydrofurane stabilized with 250-350 ppm BHT**, HPLC grade
- **5.10** *n***-Hexane**, spectroscopic grade ³
- **5.11 Ethanol**, spectroscopic grade ³
- **5.12 Cyclohexane**, spectroscopic grade ³

¹ Alcalase® and Multifect PR 6L have been successfully used for the validation.

² Milli-Q, Hydromatrix[™], Alcalase® and Multifect PR 6L are examples of suitable products available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by CEN of these products.

³ The exact spectroscopic grade depends on the carotenoid for which the UV standardisation of the standard solution is performed (5.15.2).

5.13 Mobile phase for HPLC

5.13.1 Phase A: acetonitrile:methyl tert-butyl ether:water mixture 70:20:10; v:v:v, stabilized with 1 000 ppm BHT

Using a graduated cylinder (6.17), transfer 700 ml of acetonitrile (5.7) into a 1 000 ml bottle. Measure (6.17) and add 200 ml of methyl tert-butyl ether (5.8) and 100 ml water (5.2). Add 1,0 g of BHT (5.3). Perform mixing and degassing for 10 min in an ultrasonic bath (6.11). This mobile phase is stable for 28 days.

NOTE The retention time of the carotenoids is strongly influenced by slight differences in the composition of mobile phase A. The use of an HPLC quality control sample (9.1) is crucial for the correct signal allocation. Furthermore, the presence of non-targeted carotenoids might interfere with the signals of the targeted analytes when the composition of mobile phase A deviates from 5.13.1.

5.13.2 Phase B: acetonitrile:methyl tert-butyl ether mixture (70:30; v:v)

Using a graduated cylinder (6.17), transfer 700 ml of acetonitrile (5.7) into a 1 000 ml bottle. Measure (6.17) and add 300 ml methyl tert-butyl ether (5.8). Add 1,0 g of BHT (5.3). Perform mixing and degassing for 10 min in an ultrasonic bath (6.11). This mobile phase is stable for 28 days.

5.14 Reference standards

Guaranteed purity is required for each lot of reference standard:

- 5.14.1 Astaxanthin (AXN), purity ≥ 97% AND ARD PREVIEW
- 5.14.2 Canthaxanthin (CXN), purity **1979** mdards.iteh.ai)
- **5.14.3 Adonirubin** (ADR), purity ≥ 97 % <u>oSIST prEN 17550:2020</u>

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- 5.14.4 Astaxanthin dimethyldisuccinate (AXN DMDS), purity ≥95%
- **5.14.5 Capsanthin** (CSN), purity $\geq 95\%$
- **5.14.6 Ethyl ester of beta-apo-8'-carotenoic acid** (BACARE), purity ≥ 95 %
- **5.14.7 Lutein** (LUT), purity $\geq 95\%$
- **5.14.8 Citranaxanthin** (CIXN), purity $\geq 95\%$
- **5.14.9 Zeaxanthin** (ZEA), purity \geq 95 %
- **5.14.10 Beta-carotene** (BCAR), purity $\geq 95 \%$

5.15 Standard solutions

Protect all standard solutions from daily light.

5.15.1 Carotenoid stock standard solution, ca. 200 µg/ml

Prepare fresh and measure immediately.

NOTE Possible carotenoids are astaxanthin (AXN), canthaxanthin (CXN), adonirubin (ADR), astaxanthin dimethyldisuccinate (AXN DMDS), capsanthin (CSN), ethyl ester of beta-apo-8'-carotenoic acid (BACARE), citranaxanthin (CIXN), lutein (LUT), zeaxanthin (ZEA) and beta-carotene (BCAR).

5.15.1.1 Storage container containing more than 1,0 mg

When the storage container contains more than 1,0 mg of the selected carotenoid (see 5.14.1 or 5.14.2 or 5.14.3 or 5.14.4 or 5.14.5 or 5.14.6 or 5.14.7 or 5.14.8 or 5.14.9 or 5.14.10, depending on the carotenoid selected), make a 0,2 mg/ml standard solution by e.g. weighing 1,0 mg of this carotenoid (note down the weight of standard) into a 5 ml volumetric flask.

Dissolve and make up to the mark with THF/BHT (5.9). Mix well using a vortex mixer (6.27) and an ultrasonic bath (6.11). The accurate mass fraction needs to be standardized using a spectrophotometer (6.2) as described in detail in 5.15.2. Reserve a 1 ml portion for the isomerisation procedure (5.15.4), if needed. Reserve another aliquot for the identification of the analyte as described in paragraph 9.5.3.2 and store it in the freezer.

5.15.1.2 Storage container containing less than 1,0 mg

When the storage container specifies a nominal content equal to or less than 1,0 mg of the selected carotenoid (see 5.14.1 or 5.14.2 or 5.14.3 or 5.14.4 or 5.14.5 or 5.14.6 or 5.14.7 or 5.14.8 or 5.14.9 or 5.14.10, depending on the carotenoid selected), transfer quantitatively the whole content in a 5,0 ml volumetric flask. Dissolve and make up to the mark with THF/BHT (5.9). Mix well using a vortex mixer (6.27) and an ultrasonic bath (6.11). The accurate mass fraction needs to be standardized using the spectrophotometer (6.2) as described in detail in 5.15.2. Reserve a 1 ml portion for the isomerisation procedure (5.15.4), if needed. Reserve another aliquot for the identification of the analyte as described in paragraph 9.5.3.2 and store it in the freezer.

5.15.2 UV standardization of the standard solutions PEVEW

Pipette 100 μ l of the selected carotenoid stock standard solution (5.15.1) into a 10 ml volumetric flask (6.8) and make up to the mark with the appropriate solvent. The nominal value of the obtained solution is ca. 2 μ g/ml. Scan the spectrum of this solution from 300 nm to 550 nm and measure the absorption of this solution against the pure solvent, at the maximum, using the spectrophotometer (6.2). The maximum is solvent specific and is given as an approximate value in Annex A.

The exact content of the selected carotenoid is given by Formula (1).

Carotenoid
$$\mu$$
g/ml = $E_{\lambda max} \times 10~000/E^{1\%}_{1 cm}$ (1)

EXAMPLE For all-E AXN, the solvent suggested in the table in Annex A is n-hexane. For this solvent, the wavelength of measurement is approximately 470 nm and the $E^{1\%}$ 1cm (or $A^{1\%}$ 1cm) is 2 100.

Pipette with a suited pipette $100 \,\mu$ l of the astaxanthin stock standard solution (5.15.1) into a $10 \,\text{ml}$ volumetric flask and make up to the mark with n-hexane. The nominal value of the obtained standard measuring solution is ca. $2 \,\mu\text{g/ml}$. Measure the UV spectrum of this solution against pure n-hexane, using the spectrophotometer (6.2) set at the wavelength of maximum absorption (approx. 470 nm).

NOTE The solvent of the measured solution is not pure as there is a small presence of THF/BHT (5.9). However, we consider the effect negligible and we apply the same tabulated extinction coefficient given that the contribution of THF/BHT (5.9) is very small (1 %).

The content of AXN is given by Formula (2).

AXN
$$\mu g/ml = E_{max} \times 10\ 000/2\ 100$$
 (2)

A solution of the same concentration, $100~\mu l$ of the astaxanthin stock standard solution (5.15.1) pipetted into a 10~m l volumetric flask and made up to the mark with acetone (5.5), shall be injected simultaneously in the HPLC (see 5.15.3.1).

5.15.3 HPLC standard calibration curve

5.15.3.1 Carotenoid standard calibration curve

Pipette with a suited pipette $100 \mu l$ of carotenoid stock standard solution (5.15.1) into a $10 \mu l$ ml volumetric flask and make up to the mark with acetone (5.5).

5.15.4 Isomerisation

5.15.4.1 General

It is recommended to perform this step when ambiguity occurs regarding the presence of possible isomers.

5.15.4.2 Isomerisation of the standard solutions

The standards prepared according to 5.15.1.1 or 5.15.1.2 could be isomerised in order to have a profile of the all-trans and cis isomers of each carotenoid, which are present in the equilibrium phase.

Fill a LC vial with 1 ml standard. Close tightly and check that the vial is well sealed and the cap does not turn. Heat up the vial at about 80 °C for about 2 h to achieve the equilibrium between the isomers. A vials block heater (6.30) can be used for this purpose.

Inject a dilution of the isomerised standard (e.g. 1:200).

The isomerised standard is stable for several months at room temperature.

5.16 HPLC positive quality control sample (QCS)

Weigh approximately 0,8 mg AXN (5.14.1), 0,4 mg CXN (5.14.2) and 0,5 g BHT (5.3) in a round bottom flask (6.9). Add 200 ml of an acetone (5.5): water (5.2) 75:25 v:v solution. Fit a condenser (6.10) to the flask and immerse the flask in a heating mantle or heated bath (6.28) with a magnetic stirrer (6.29). Heat to boiling and allow to refluxing for about 1,5 h to 2 h.s.-pren-17550-2020

NOTE Alternatively, the mixture can be heated at 80°C in a pressure-resistant tube.

This reaction will cause the formation of a stable isomers ratio. Transfer to a 500 ml volumetric flask (6.8) and make up to volume with acetone (5.5). Mix well and transfer immediately in HPLC vials. The filling should be fast and each vial should be immediately closed to avoid evaporation. Store the vials at room temperature and away from light.

6 Apparatus

Usual laboratory apparatus and, in particular, the following:

- **6.1 HPLC system,** consisting of the following
- **6.1.1 Pump, pulse free,** capable of maintaining a volume flow rate from 0,1 ml/min to 2,0 ml/min
- **6.1.2** Injection system, manual or autosampler

If an autosampler is used, a cooled one is recommended although it was not used during the validation of the method.

6.1.3 UV/VIS detector, variable wavelength, suitable for reliable measurements at 410 nm, or UV/VIS photodiode array detector (DAD)

- 6.1.4 Computer data system
- **6.1.5** Analytical column, Supelco® Suplex pKb-100 5 μm, 250 mm × 4,6 mm or equivalent ⁴
- **6.1.6 Guard column,** 5 µm, 2 cm × 4 mm SUPELCOSIL[™] Suplex[™] pKb-100 Supelguard[™] Cartridge or equivalent 4
- **6.2 Spectrophotometer**, with 1 mm apertures
- 6.3 Grinding instrument
- **6.4 Sieve,** with 1 mm apertures
- **6.5 Balances,** one analytical, of 10 g capacity or greater with 0.1 mg readability, and one, of 100 g capacity or greater with 0.01 g readability
- **6.6 Polypropylene containers,** 100 ml with lids
- **6.7 Variable-volume positive displacement piston pipettes,** suitable for pipetting volumes ranging from 50 μ l to 100 μ l
- **6.8 Glass volumetric flasks** of, 5 ml, 10 ml, 150 ml, 500 ml and 1 000 ml
- 6.9 Round bottom flasks of 500 ml NDARD PREVIEW
- 6.10 Allihn condenser (standards.iteh.ai)
- **6.11 Ultrasonic bath,** temperature controlled, 17550:2020
- https://standards.iteh.ai/catalog/standards/sist/27ac68a6-b27b-4b8c-a185-6.12 Flat spatulas 75e7348fece0/osist-pren-17550-2020
- **6.13 PLE, pressurized solvent extraction system,** ASE 300 Dionex®, Büchi SpeedExtractor E-914 or equivalent ⁴
- **6.14 PLE cells,** suitable for the extraction unit used, 66 ml or larger
- **6.15 Cellulose filters** for PLE cells
- **6.16 PLE vials,** for the extraction unit used, at least 240 ml
- **6.17 Graduated cylinder** of 250 ml and 1 000 ml
- 6.18 Microcentrifuge safe-lock tubes, 1,8 ml, Eppendorf® tubes or equivalent 4
- 6.19 Microcentrifuge
- 6.20 1,5 ml HPLC glass vials

⁴ Supelco® Suplex pKb-100, SUPELCOSIL™ Suplex™ pKb-100 Supelguard™ Cartridge, ASE 300 Dionex®, Büchi SpeedExtractor E-914, Eppendorf® tubes and Techne Dri-Block Heater are examples of suitable products available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement by CEN of these products.

- 6.21 HPLC glass vials crimper
- 6.22 Centrifuge
- **6.23 Centrifuge tubes,** 50 ml or 100 ml
- 6.24 Ultraturrax mixer
- **6.25 PLE Funnels** for PLE cells
- 6.26 Powder funnels
- 6.27 Vortex mixer
- 6.28 Heating mantle or heated bath
- 6.29 Magnetic stirrer
- 6.30 Vials block heater, Techne Dri-Block Heater or equivalent 4

7 Sampling

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

Sampling is not part of the method specified in this document. A recommended sampling method is given in EN ISO 6497 [3].

8 Preparation of test sample

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8.1 General

Prepare the test sample in accordance with EN ISO 6498.

8.2 Laboratory sample

Grind the laboratory sample (usually 50 g) so that it passes completely through a sieve with 1 mm apertures (6.4) or until a fine paste is obtained. Mix thoroughly.

8.3 Test sample

The test sample consists of a representative and homogenized aliquot of the ground laboratory sample (8.2) of at least 10 g.

8.4 Test portion

Accurately weigh 5,0 g to the nearest 0,1 g of the thoroughly mixed test sample (8.3) into a 100 ml polypropylene container (6.6) for extraction procedure 9.3.1 or into a 50 ml or 100 ml centrifuge tube (6.23) for extraction procedure 9.3.2. Note down the mass expressed in g. Submit it to the analysis procedure (Clause 9).

9 Procedure

9.1 General

The complete procedure (9.2 to 9.4) should be applied to two test portions (8.4) of the same test sample in order to perform two independent and parallel determinations.

The use of quality control samples is recommended.

The QCS (5.16) shall be injected before and after the complete sequence samples each day of analysis, in order to assess the suitability of the HPLC/DAD system.

The QCS (5.16) contains two carotenoids:

- 1) CXN, a stable and more soluble carotenoid, and
- 2) AXN, a less stable and less soluble carotenoid.

NOTE Depending on the analytes to be determined a relevant carotenoid can be added to this mix.

The mixture is isomerised until a constant ratio of the isomers is reached. This solution, when protected from oxygen, is stable for several months.

9.2 Enzymatic disclosure

Add 0,2 g of BHT (5.3), 100 µl of enzyme (5.1) and 15 ml of purified water (5.2) to the test portion (8.4). Close tightly and shake vigorously to ensure that all the feed is permeated with water. Place in an ultrasonic bath (6.11) with the temperature set at 50 °C for 15 min to 20 min. Shake every 5 min.

9.3 Extraction

oSIST prEN 17550:2020 9.3.1 Extraction using PLE https://standards.iteh.ai/catalog/standards/sist/27ac68a6-b27b-4b8c-a185-

Remove the lid of the test portion after step 9.2, taking care of not losing sample drops. Add approximately 13 g of high purity diatomaceous earth suitable for PLE (5.4), close tightly and hand-shake vigorously until the wet feed sample looks all adsorbed on the diatomaceous earth beadlets and detaches well from the polypropylene container's wall.

NOTE Stomping the container on the table can help.

Open the containers carefully, remove the sample particles from the lid into the container using a flat spatula (6.12). Mix well with the spatula (6.12), making sure that there are no sample clusters left. Place two cellulose filters (6.15) on the bottom of the PLE cell (6.14). Transfer all the material in the PLE cell (6.14) using a funnel (6.25). Top up with diatomaceous earth (5.4) if needed. Close the cell and extract according to the parameters in Table 1.