

SLOVENSKI STANDARD oSIST prEN 16877:2015

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Krma - Metode vzorčenja in analize - Določevanje toksinov T-2 in HT-2, deoksinivalenola in zearalenona v sestavinah krme in krmni mešanici z LC-MS

Animal feeding stuffs - Methods of sampling and analysis - Determination of T-2 and HT-2 toxins, Deoxynivalenol and Zearalenone, in feed materials and compound feed by LC-MS

Futtermittel - Probenahme- und Untersuchungsverfahren - Bestimmung von T-2- und HT -2-Toxinen, Deoxynivalenol und Zearalenon in Einzelfuttermitteln und Mischfuttermitteln

mittels LC-MS

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Aliments des animaux - Méthodes d'échantillonnage et d'analyse - Dosage par LC-MS des toxines T-2 et HT-2, du déoxynivalénol et de la zéaralénone dans les matières premières pour aliments et les aliments composés

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Animal feeding stuffs - Methods of sampling and analysis - Determination of T-2 and HT-2 toxins, Deoxynivalenol and Zearalenone, in feed materials and compound feed by LC-MS

Aliments des animaux - Méthodes d'échantillonnage et d'analyse - Dosage par LC-MS des toxines T-2 et HT-2, du déoxynivalénol et de la zéaralénone dans les matières premières pour aliments et les aliments composés Futtermittel - Probenahme- und Untersuchungsverfahren -Bestimmung von T-2- und HT-2-Toxinen, Deoxynivalenol und Zearalenon in Einzelfuttermitteln und Mischfuttermitteln mittels LC-MS

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Foreword

This document (prEN 16877:2015) 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 method of analysis is applicable to the determination of HT-2 toxin (HT2) in the tested range of 22 μ g/kg to 178 μ g/kg, T-2 toxin (T2) in the tested range of 7 μ g/kg to 50 μ g/kg, Deoxynivalenol (DON) in the tested range of 88 μ g/kg to 559 μ g/kg, and Zearalenone (ZON) in the tested range of 14 μ g/kg to 430 μ g/kg in cereals and cereal-based compound animal feed. The actual working ranges may extend beyond the tested ranges. It is the responsibility of the laboratory to prove that the limit of quantitation (LOQ) for HT-2 and T-2 toxin is \leq 10 μ g/kg, for DON \leq 100 μ g/kg, and for ZON \leq 20 μ g/kg.

2 Normative references

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

EN ISO 3696, Water for analytical laboratory use — Specification and test methods (ISO 3696)

3 Principle

Two gram of finely ground and homogeneous test material is suspended in water. After addition of 16,0 mL ethyl acetate the sample is agitated for 30 min. Then sodium sulphate is added to facilitate phase separation and after 10 min to 20 min the sample is centrifuged to pellet particulate matter at the bottom of the extraction tube. The organic phase is transferred to a clean vial for possible storage. $500 \,\mu\text{L}$ of the organic phase, an equivalent of 1/16th of a gram of the test material, are mixed with stable-isotope labelled analogues of the analytes and evaporated to dryness in deactivated glass vials. After reconstitution of the dry extract with $250 \,\mu\text{L}$ of organic mobile phase modifier, addition of $250 \,\mu\text{L}$ of water, and thorough mixing the analytes are quantified with a Liquid Chromatography-Mass Spectrometry (LC-MS) system.

4 Reagents

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- **4.1 Water (deionized)** 4.1 Water (deionized) 4.1 Water (deionized
- 4.2 Water (LC-MS grade, double-distilled water or water of grade 1 as defined in EN ISO 3696)

4.3 Methanol (LC-MS grade)

WARNING — Methanol is hazardous and handling shall be carried out inside a fume cupboard. Appropriate safety equipment (lab coat, goggles, gloves) shall be worn.

4.4 Methanol (p.a.)

WARNING — Methanol is hazardous and handling shall be carried out inside a fume cupboard. Appropriate safety equipment (lab coat, goggles, gloves) shall be worn.

4.5 Ethyl acetate (p.a.)

WARNING — Ethyl acetate is hazardous and handling shall be carried out inside a fume cupboard. Appropriate safety equipment (lab coat, goggles, gloves) shall be worn.

4.6 Formic acid (98 % – 100 %, LC-MS grade)

WARNING — Formic acid is hazardous and handling shall be carried out inside a fume cupboard. Appropriate safety equipment (lab coat, goggles, gloves) shall be worn.

4.7 Acetonitrile (LC-MS grade)

WARNING — Acetonitrile is hazardous and handling shall be carried out inside a fume cupboard. Appropriate safety equipment (lab coat, goggles, gloves) shall be worn.

4.8 Sodium sulphate

anhydrous, granulated

4.9 Deoxynivalenol (DON)

WARNING — Deoxynivalenol is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.10 HT-2 toxin (HT2)

WARNING — HT-2 toxin is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.11 T-2 toxin (T2)

WARNING — T-2 toxin is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.12 Zearalenone (ZON)

WARNING — Zearalenone is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.13 ¹³C15-Deoxynivalenol (¹³C₁₅-DON)

WARNING — 13 C₁₅-Deoxynivalenol is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.14 ¹³C22-HT-2 toxin (¹³C₂₂-HT2)

WARNING — 13 C₂₂-HT-2 toxin is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.15 ¹³C24-T-2 toxin (¹³C₂₄-T2)

WARNING — 13 C₂₄-T-2 toxin is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.16 ¹³C₁₈-Zearalenone (¹³C₁₈-ZON)

WARNING — ¹³C₁₈-Zearalenone is highly toxic. Gloves and safety glasses shall be worn at all times and all standard and sample preparation stages shall be carried out in a fume cupboard.

4.17 Multitoxin stock solution

A mixture containing Deoxynivalenol (4.9), HT-2 toxin (4.10), T-2 toxin (4.11), and Zearalenone (4.12) in neat acetonitrile (4.7) at relevant concentrations.

When preparing this solution the certified purities of the mycotoxin reference materials need to be properly accounted for. In any case, the purities shall be \geq 95 %.

NOTE 1 3,2 μ g/mL DON, 0,5 μ g/mL HT-2 toxin, 0,3 μ g/mL T-2 toxin, and 0,3 μ g/mL ZON in neat acetonitrile have shown to work well. This solution is stable for three months in the dark at 2 °C –8 °C.

NOTE 2 Compare a new stock solution against the old one by adding 25 μ L of each into separate deactivated vials (5.6) and proceeding as described in "Test solution" (6.3).

NOTE 3 If 6.4"Spiking procedure" is executed at least 6 mL of the stock solution are needed.

4.18 Multitoxin working solution

Dilute Multitoxin stock solution (4.17) with Methanol (4.3) such that the resulting concentration in the working solution is applicable to the calibration range of the different compounds. Only prepare enough volume for one full calibration.

NOTE Adding 188 μ L of the Multitoxin stock solution to a 3 mL volumetric flask and making up to the mark with methanol will result in a solution containing 0,2 μ g/mL DON, 0,031 μ g/mL HT-2 toxin, 0,019 μ g/mL T-2 toxin, and 0,019 μ g/mL ZON in methanol/acetonitrile (94/6, v/v).

4.19 Multi internal standard (ISTD) stock solution

A mixture containing $^{13}C_{15}$ -DON (4.13), $^{13}C_{22}$ -HT-2 toxin (4.14), $^{13}C_{24}$ -T-2 toxin (4.15), and $^{13}C_{18}$ -ZON (4.16) in neat acetonitrile (4.7) at the same concentrations as the respective native compounds in the Multitoxin stock solution (4.17).

NOTE This solution is stable for three months in the dark at 2 °C – 8 °C.

4.20 Calibration

To six deactivated glass vials (5.6) add different volumes of the Multitoxin working solution (4.18) such that six equidistant calibration levels across the calibration range result. Proceed as described in 6.3. "Test solution".

NOTE Table 1 below shows example calibration levels using the solutions described in the notes above.

Once it has been shown that there is linearity the number of levels may be adjusted to local needs and requirements.

Volume of Multitoxin working Total mass of analyte solution (4.18) per vial [µL] [ng] DON HT-2 T-2 ZON 25 5 0,78 0,48 0,48 180 36 5,6 3,4 3,4 335 10 6,4 6,4 67 490 15 9,3 9,3 98 645 129 20 12 12 800 160 25 15 15

Table 1— Calibration solutions

4.21 Quality control material

An appropriate material with natural contamination or fortification of the tested mycotoxins which is sufficiently stable.

5 Apparatus

5.1 Mill

Single mill or multiple mills capable of comminuting test materials to particle sizes of < 500 μm.

5.2 Mixer

Capable of sufficiently homogenizing the comminuted test materials.

NOTE A tumble mixer that uses a folding action either through moving paddles or fins, or an end-over-end movement has shown to work well.

5.3 Conical polypropylen screw-cap centrifuge tubes 50 mL with caps

5.4 Volumetric flasks

3 mL, 5 mL, and 10 mL

5.5 Pipettors ITCh STANDARD PREVIEW

Adjustable 10 μL - 100 μL and adjustable 100 μL – 1 000 μL, properly calibrated.

5.6 Deactivated glass vials

5.7 Auto Liquid Sampler (ALS) vials

Of appropriate size for the Auto Liquid Sampler in use.

5.8 Shaker or Sonicator

5.9 Evaporator

Capable of maintaining a stable temperature in the range of 30 °C - 60 °C with a constant flow of dry nitrogen.

5.10 Centrifuge

Capable of generating a relative centrifugal force (RCF) of 3 000 g.

5.11 Syringe filter

Small internal volume, Nylon, Pore size: 0,2 µm Nylon.

5.12 LC-MS

5.12.1 Solvent delivery system

Capable of delivering a binary gradient at flow rates appropriate for the analytical column in use with sufficient accuracy.

5.12.2 Auto liquid sampler (ALS)

Capable of injecting an appropriate volume of injection solution with sufficient accuracy, cross-contamination below 0,1 %.

5.12.3 Analytical column

Capable of separating the four analytes with the following performance:

Peak asymmetry factor at 10 % height: 0.9 < As < 1.4; minimum apparent retention factor for any of the four analytes: $N \ge 1$ 200; minimum resolution between two adjacent analyte peaks: $Rs \ge 4$.

5.12.4 Mass spectrometer

An instrument capable of performing selected reaction monitoring (SRM) with a sufficiently wide dynamic range. Any ionization source giving sufficient yield may be employed.

5.13 Balance

Balance with readability d = 0,001 g or better.

6 Procedures

6.1 Sample preparation ANDARD PREVIEW

It is important that the laboratory receives a laboratory sample which is truly representative and has not been damaged or altered during transport or storage. Laboratory samples should be taken and prepared in accordance with European legislation where applicable [1] [2]. The laboratory sample should be finely ground and thoroughly mixed using a mill (5.1) and a mixer (5.2) or another process for which complete homogenization has been demonstrated before a test portion is removed for analysis.

The recommended way is to comminute the laboratory sample in several steps. Beginning with the totality of the laboratory sample each step consists of taking a representative aliquot of the previous step after sufficient homogenization. This aliquot is then comminuted to the next smaller particle size until a subsample of ca. 50 g of the final particle size is obtained. It is of utmost importance that the test portion is taken from a subsample which is sufficiently homogenous with a particle size of \leq 500 μ m. Care should be taken to not overheat the sample during this process.

In all instances everything should be at room temperature before any kind of manipulation takes place.

6.2 Extraction

Some of the steps described below are more critical for the accuracy of the results than others. These steps are marked as such and should be carried out with the necessary attention. A scale-up of the test portion size is deemed to be acceptable if such a need is assumed. In that case the amounts of added water, ethyl acetate, and sodium sulphate need to be increased at the same rate, for instance, scale-up by factor of 2:4 g test portion, 16 mL water, 32 mL ethyl acetate, 16 g sodium sulphate. In no way shall a scale-up be seen as replacement for proper sample preparation (6.1).

- For the test portion weigh 1,9 g to 2,1 g of the homogeneous sample into a conical polypropylene screw-cap tube (5.3), round and record the weight to the second decimal (the accuracy of this weight is critical for the accuracy of the final result).
- Add 7,2 mL to 8,8 mL of deionized water (4.1).

- Vortex thoroughly until test portion is completely suspended. Do not let this suspension stand for more than 15 min to prevent effects due to enzymatic activities.
- Add 16,0 mL of ethyl acetate (4.5, the accuracy of this volume is critical for the accuracy of the final result!).
- Extract for 27 min to 33 min in a sonicator or by vigorously shaking (5.8).
- Add between 7,2 g and 8,8 g of sodium sulphate (4.8).
- Instantly shake hard for 5 s.
- Let stand for 10 min to 20 min.
- Centrifuge (5.10) at RCF 3 000 g for at least 1 min to aid settlement of particulate matter and phase separation.
- If wanted for possible repeats: Transfer the extract (organic layer) into clean glass vial for storage of up to 7 days at 2 °C to 10 °C in the dark.
- Transfer 500 μL of the extract (organic layer) into a deactivated glass vial (5.6) for further processing (the accuracy of this volume is critical for the accuracy of the final result!).

6.3 Test solution

- Add 25 μL of the Multi ISTD stock solution (4.19) to the aliquot of the extract and/or the calibration solutions (4.20) (the accuracy of this volume is critical for the accuracy of the final result).
- Dry down the aliquot of the extract and/or the calibration solutions in an evaporator (5.9) with a gentle stream of dry nitrogen at 60 °C.
- Add 250 μ L of the organic mobile phase modifier to the dry residue for reconstitution.
- Vortex thoroughly for at least 10 s.
- Add 250 µL deionized water (4.1) to the reconstituted extract.
- Vortex thoroughly for at least 5 s.
- Transfer the test solution into an ALS vial (5.7); if solution is turbid it may be filtered through a syringe filter (5.11).

NOTE It has been shown that even very turbid samples can be injected without any negative effects to the life time of column and LC provided that appropriate in-line filters or guard columns are used.

6.4 Spiking procedure

If recovery needs to be determined execute the following in duplicate:

To three times 2 g of a material free of DON, HT2, T2, and ZON add three different volumes of the Multitoxin stock solution (4.17) such that 3 contamination levels across the calibration range result. Distribute the solutions evenly over the materials, mix to further distribute the spike, and leave for a minimum of 5 h to a maximum of 18 h. Proceed to 6.2 "Extraction" second step.

NOTE Addition of $360 \,\mu\text{L}$, $980 \,\mu\text{L}$, and $1600 \,\mu\text{L}$ of the Multitoxin stock solution (4.17) with the concentrations described in the note has been shown to work well.