
Krma - Določanje vsebnosti skupnega fosforja - Spektrofotometrična metoda (prevzet standard ISO 6491:1980 z metodo platnice)

Animal feeding stuffs - Determination of total phosphorus content - Spectrophotometric method

Aliments des animaux - Détermination de la teneur en phosphore total - Méthode spectrophotométrique

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SIST ISO 6491:1995

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Deskriptorji: prehrana živali, krmljenje živali, kemijska analiza, določanje vsebnosti, fosfor, spektrofotometrična analiza

ICS 71.040.40 * 65.120

Referenčna številka
SIST ISO 6491:1995 (en)

Nadaljevanje na straneh od II do III in 1 do 3

UVOD

Standard SIST ISO 6491, Krma - Določanje vsebnosti skupnega fosforja - Spektrofotometrična metoda, prva izdaja, 1995, ima status slovenskega standarda in je z metodo platnice prevzet mednarodni standard ISO 6491, Animal feeding stuffs - Determination of total phosphorus content - Spectrophotometric method, first edition, 1980-08-15.

PREDGOVOR

Mednarodni standard ISO 6491:1980 je pripravil tehnični odbor Mednarodne organizacije za standardizacijo ISO/TC 34 Kmetijski pridelki in živilski proizvodi.

Odločitev za prevzem tega standarda po metodi platnice je sprejela delovna skupina WG 10 Analitika krme v okviru tehničnega odbora USM/TC KŽP Kmetijski pridelki in živilski proizvodi.

Ta slovenski standard je dne 1995-05-08 odobril direktor USM.

ZVEZA S STANDARDI

S prevzemom tega mednarodnega standarda veljajo naslednje zveze:

- SIST ISO 6651:1995 (en) Krma - Določanje vsebnosti aflatoksina B₁
- SIST ISO 6654:1995 (en) Krma - Določanje vsebnosti sečnine
- SIST ISO 6866:1995 (en) Krma - Določanje vsebnosti prostega in skupnega gosipola
- SIST ISO 6870:1995 (en) Krma - Določanje vsebnosti zearalenona
- SIST ISO 5498:1995 (en) Kmetijski pridelki in živilski proizvodi - Določanje vsebnosti surove vlaknine - Splošna metoda
<https://standards.iteh.ai/catalog/standards/sist/5b40b5-c723-430b-803c-69d43aaa228f/sist-iso-6491-1995>
- SIST ISO 5983:1995 (en) Krma - Določanje vsebnosti dušika in izračun vsebnosti surovih beljakovin
- SIST ISO 5984:1995 (en) Krma - Določanje surovega pepela
- SIST ISO 5985:1995 (en) Krma - Določanje pepela, netopnega v klorovodikovi kislini
- SIST ISO 6490-1:1995 (en) Krma - Določanje vsebnosti kalcija - 1. del: Titrimetrična metoda
- SIST ISO 6490-2:1995 (en) Krma - Določanje vsebnosti kalcija - 2. del: Metoda atomske absorpcijske spektrometrije
- SIST ISO 6495:1995 (en) Krma - Določanje vsebnosti v vodi topnih kloridov
- SIST ISO 6496:1995 (en) Krma - Določanje vsebnosti vlage
- SIST ISO 5506:1995 (en) Sojini proizvodi - Določanje ureazne aktivnosti
- SIST ISO 6541:1995 (en) Kmetijski pridelki in živilski proizvodi - Določanje vsebnosti surove vlaknine - Modificirana Scharrerjeva metoda

OSNOVA ZA IZDAJO STANDARDA

- Prevzem standarda ISO 6491:1980

OPOMBI

- Povsod, kjer se v besedilu standarda uporablja izraz mednarodni standard , to pomeni v SIST ISO 6491:1995 slovenski standard .
- Uvod in predgovor nista sestavni del standarda.

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International Standard



6491

INTERNATIONAL ORGANIZATION FOR STANDARDIZATION • МЕЖДУНАРОДНАЯ ОРГАНИЗАЦИЯ ПО СТАНДАРТИЗАЦИИ • ORGANISATION INTERNATIONALE DE NORMALISATION

Animal feeding stuffs — Determination of total phosphorus content — Spectrophotometric method

Aliments des animaux — Détermination de la teneur en phosphore total — Méthode spectrophotométrique

First edition — 1980-08-15

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UDC 636.085/.087 : 543.42 : 546.18

Ref. No. ISO 6491-1980 (E)

Descriptors : animal feeding, animal nutrition, chemical analysis, determination of content, phosphorus, spectrophotometric analysis.

Price based on 3 pages

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards institutes (ISO member bodies). The work of developing International Standards is carried out through ISO technical committees. Every member body interested in a subject for which a technical committee has been set up has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work.

Draft International Standards adopted by the technical committees are circulated to the member bodies for approval before their acceptance as International Standards by the ISO Council.

International Standard ISO 6491 was developed by Technical Committee ISO/TC 34, *Agricultural food products*, and was circulated to the member bodies in February 1979.

It has been approved by the member bodies of the following countries :

SIST ISO 6491:1995
<https://standards.iteh.ai/catalog/standards/sist/3b40f33-c723-430b-803c-69d43aaa2222/iso-6491-1995>
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Australia	Hungary	Poland
Austria	India	Portugal
Brazil	Israel	Romania
Canada	Kenya	South Africa, Rep. of
Chile	Korea, Rep. of	Spain
Cyprus	Libyan Arab Jamahiriya	Thailand
Czechoslovakia	Malaysia	Turkey
Egypt, Arab Rep. of	Netherlands	United Kingdom
Ethiopia	New Zealand	Yugoslavia
France	Philippines	

No member body expressed disapproval of the document.

Animal feeding stuffs — Determination of total phosphorus content — Spectrophotometric method

1 Scope

This International Standard specifies a spectrophotometric method for the determination of the total phosphorus content of animal feeding stuffs.

2 Field of application

The method is applicable to all animal feeding stuffs.

It is particularly appropriate for the analysis of products low in phosphorus. In certain cases (products rich in phosphorus) a gravimetric method may be used.

3 Principle

Ashing of a test portion, either by dry combustion and dissolution in acid (in the case of organic feeding stuffs) or by acid digestion (in the case of mineral compounds and liquid feeding stuffs).

Treatment of this solution with molybdovanadate reagent and measurement of the absorbance of the yellow solution thus obtained, in a spectrophotometer, at 430 nm.

4 Reagents

All reagents shall be of recognized analytical quality. Distilled water or water of at least equivalent purity shall be used.

4.1 Calcium carbonate.

4.2 Hydrochloric acid, $c(\text{HCl}) \approx 6 \text{ mol/l}$.

4.3 Nitric acid, $c(\text{HNO}_3) \approx 1 \text{ mol/l}$.

4.4 Nitric acid, $\rho_{20} 1,38 \text{ g/ml}$.

4.5 Sulphuric acid, $\rho_{20} 1,84 \text{ g/ml}$.

4.6 Molybdovanadate reagent.

In a 1 000 ml volumetric flask, mix 200 ml of the ammonium heptamolybdate solution (4.6.1), 200 ml of the ammonium monovanadate solution (4.6.2) and 135 ml of the nitric acid (4.4). Make up to the mark with water.

4.6.1 Ammonium heptamolybdate solution.

Dissolve in hot water 100 g of ammonium heptamolybdate tetrahydrate $[(\text{NH}_4)_6\text{Mo}_7\text{O}_{24} \cdot 4\text{H}_2\text{O}]$. Add 10 ml of ammonia ($\rho_{20} 0,91 \text{ g/ml}$) and make up to 1 litre with water.

4.6.2 Ammonium monovanadate solution.

Dissolve 2,35 g of ammonium monovanadate (NH_4VO_3) in 400 ml of hot water.

Stirring constantly, slowly add 20 ml of diluted nitric acid [7 ml of the nitric acid (4.4) + 13 ml of water] and make up to 1 litre with water.

4.7 Phosphorus, standard solution containing 1 mg of phosphorus per millilitre.

In a 1 000 ml volumetric flask, dissolve 4,394 g of potassium dihydrogen phosphate (KH_2PO_4) previously dried at 103 °C, in water. Make up to the mark with water.

5 Apparatus

Usual laboratory apparatus, and in particular:

5.1 Ashing crucibles, of silica or porcelain.

5.2 Electric muffle-furnace, capable of being controlled at $550 \pm 20 \text{ }^\circ\text{C}$.

5.3 Kjeldahl flask, of capacity 250 ml.

5.4 One-mark volumetric flasks, of capacities 500 and 1 000 ml.

5.5 Spectrophotometer, fitted with 10 mm cells, suitable for measurements at 430 nm.

5.6 Glass test tubes, of capacity 25 to 30 ml, fitted with ground glass stoppers.

5.7 Sand bath.

5.8 Beaker, of capacity 250 ml.

5.9 Graduated pipettes.

5.10 Analytical balance.

6 Procedure

6.1 Test portion and preparation of the test solution

According to the nature of the sample, take a test portion and prepare the test solution as specified in 6.1.1 or 6.1.2.

6.1.1 Dry ashing (for samples containing organic substances and free from phosphates which give insoluble products on ashing)

Weigh about 2,5 g of the sample¹⁾ to the nearest 1 mg in an ashing crucible (5.1).

Mix the test portion thoroughly with 1 g of the calcium carbonate (4.1). Ash in the furnace (5.2) at 550 ± 20 °C until white or grey ash is obtained (a small quantity of carbon does not interfere).

Transfer the ash to the 250 ml beaker (5.8).

Add 20 ml of water and then hydrochloric acid (4.2) until effervescence ceases.

Add a further 10 ml of the hydrochloric acid (4.2).

Place the beaker on the sand bath (5.7) and evaporate to dryness to render the silica insoluble.

Allow to cool.

Add 10 ml of the nitric acid (4.3) to the residue and boil on the sand bath for 5 min, without evaporating to dryness.

Decant the liquid into a 500 ml volumetric flask (5.4), rinsing the beaker several times with hot water.

Leave to cool, make up to the mark with water, mix and filter.

6.1.2 Wet ashing (for mineral compounds and liquid feeding stuffs)

Weigh 1 g or more of the sample¹⁾ to the nearest 1 mg.

Place the test portion in the Kjeldahl flask (5.3), add 20 ml of the sulphuric acid (4.5), shake to impregnate the substance completely with acid and to prevent it from sticking to the wall of the flask, heat and keep at boiling point for 10 min.

Leave to cool slightly, add 2 ml of the nitric acid (4.4), heat gently, leave to cool slightly, add a little more nitric acid (4.4) and bring back to the boiling point.

Repeat this procedure until a colourless solution is obtained.

Cool, add a little water, and decant the liquid into a 500 ml volumetric flask (5.4), rinsing the Kjeldahl flask with hot water.

Leave to cool, make up to the mark with water, mix and filter.

6.2 Development of coloration and measurement of absorbance

Dilute an aliquot portion of the filtrate obtained (6.1.1 or 6.1.2) with water, to obtain a phosphorus concentration of not more than 40 µg/ml.

Transfer, by means of a pipette (5.9), 10 ml of this solution to a test tube (5.6) and add, by means of another pipette, 10 ml of the molybdovanadate reagent (4.6).

Mix and leave to stand for at least 10 min at 20 °C.

Transfer a portion of the solution obtained to a measuring cell and measure the absorbance in the spectrophotometer (5.5) at 430 nm, using as the reference liquid a solution obtained by adding 10 ml of molybdovanadate reagent (4.6) to 10 ml of water.

6.3 Number of determinations

Carry out two determinations on test portions taken from the same test sample.

6.4 Preparation of the calibration curve

6.4.1 Using the standard phosphorus solution (4.7), and by means of the graduated pipettes (5.9), prepare solutions containing respectively 5, 10, 20, 30 and 40 µg of phosphorus per millilitre.

6.4.2 Transfer, by means of a pipette (5.9), 10 ml of each of these solutions to a series of five test tubes (5.6) and add, to each by means of another pipette, 10 ml of the molybdovanadate reagent (4.6).

Mix and leave to stand for at least 10 min at 20 °C.

Measure the absorbance of each solution as specified in 6.2.

6.4.3 Draw the calibration curve by plotting the absorbances against the corresponding concentrations of phosphorus, in micrograms per millilitre, of the standard phosphorus solutions (6.4.1).

For concentrations between 0 and 40 µg/ml, the curve shall be linear.

1) The sampling and sample preparation of animal feeding stuffs will form the subject of future International Standards.