Product Code: EFZE3155

Clean-up of Commodity Extracts of Food and Feed Samples containing Zearalenon via Immunoaffinity Chromatography and Determination by HPLC-FLD

Principle:

This instruction of Zearalenon (ZON) determination in food and feed focuses on the enrichment step of extract using immunoaffinity column (IAC) and quantification with HPLC.

Accepted laboratory extraction methods could be maintained. Full performance of the IAC column is given if pronounced criteria regarding organic solvent tolerance, elution process of analyte and working range of column is followed.

Many pretreatment methods of Zearalenon determination in food and feed show low sensitivity because of interfering substances if problematic matrices are applied.

This method of content determination of Zearalenon combines the high selectivity of an immunoaffinity column (IAC) with its potential to concentrate elute and additional step of purification by HPLC column.

For the commodity extraction step a literature method is given. The given apparatus (e.g. HPLC system) might serve as example among other possibilities.

Extraction (Literature method given):

Assuming that 20g ground corn sample are extracted by a total of 50ml acetonitrile/water (90/10 v/v), as reported by Visconti et al.¹ If organic solvent proportion is varied the dilution of extract with PBS should be adapted accordingly in the enrichment step. On the other hand, if proportion of sample quantity and volume of extraction solvent is altered, calculation of gram equivalents must be corrected.

Enrichment Step IAC:

An aliquot of 1ml extract (see above, contains the quantity of Zearalenon of 0.4g sample) are diluted with 9ml PBS (pH=7.4) and then applied in a reservoir on top of the **SENSIColumn IAC Zearalenon 3ml** column.

To maintain full performance of the column, please take care that proportion of dilution buffer in the solution on top of the column is not too small. A proportion of 9% acetonitrile, resulting in this example enrichment, does not affect column performance.

The proportion of organic solvent of PBS diluted extract, which is applied on the column, should not exceed 15% methanol or 15% acetonitrile.

If organic solvent proportion lies above these limits, recovery rates are diminished. Increase of diluted extract volume by diluting extract with additional PBS, on the other hand, has almost no consequences to column performance.

If samples are to be prepared simultaneously, manifold of J.T. Baker for 12 samples has proven of value. Rate of flow through the affinity gel is 1 to 3 ml/min. In case of problematic matrices rate of flow should lie below 2ml/min.

<u>Caution!</u> Be aware that no big air bubbles are neither in the gel nor between gel and luer lock outlet of column which prevent a permanent flow or necessary exchange of matter.

Depending on application and on expected contents, larger or smaller extract aliquots can be applied. In such cases the sample calculation (see below) must be adapted.

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Wash:

After whole sample has passed through the gel the latter is washed with 20ml of deionized water. Remaining liquids in the gel are removed by applying either pressure from top of the column or underpressure from bottom.

Elution:

Sample reservoir on top of the **SENSIColumn IAC Zearalenon 3ml** column is removed and a appropriate vial is placed below the affinity column. The bound toxins are eluted by using a total of <u>1.5ml of methanol</u> as elution solvent. The elution process is performed in two steps to ensure complete release of analytes. First, a volume of 0.5ml elution solvent is applied. After that volume has passed through column half a minute is waited before the second portion of 1ml of elution solvent is eluted through the column. Remaining solvent solutions should be eluted by application of slight under- or overpressure. All methanolic fractions are unified to give the column eluate.

The column eluate may be injected into the HPLC directly or in case concentrations are low it may be concentrated by evaporation, e.g. using VLM evaporator at 50°C under a permanent stream of nitrogen.

<u>Caution:</u> To prevent loss of analyt during evaporation, it is recommended to use a keeper, e.g. 100µl deionized water or PBS.

The residue then is redissolved in HPLC solvent (e.g. 0.5ml) and an aliquot is finally injected into the system.

IAC Column Characteristics:

A) Working Range and Recovery Rates of SENSIColumn IAC Zearalenon 3ml Column:

Working Range of Column: Zero Contamination per Column:	5– 500ng ZON per IAC <5ng (LOD of HPLC-FLD method)
Guaranteed Recovery Rates within the Working	
Range ^(*) :	>85%

Recovery rates are confined to solvent content of diluted extract below 15% methanol or 15% acetonitrile (see details under Enrichment Step).

B) Cross Reactivities^(**) of SENSIColumn IAC Zearalenon 3ml Column:

Zearalenon:	100%	
α-Zearalanol: β-Zearalanol: α-Zearalenol: β-Zearalenol:	99% 94% 97% 86%	

(**)

Recovery rates if a total quantity of 500ng of Zearalenon, α -Zearalanol, β -Zearalanol, α -Zearalenol and β -Zearalenol (molar ratio of 1:1:1:1) is analyzed

C) Capacity^(***) of SENSIColumn IAC Zearalenon 3ml Column:

Maximum Column Capacity:	2.7µg ZON

An excess of ZON, e.g. 5µg, in a small volume of 2ml PBS is incubated with the IAC for 5 minutes; then the IAC is washed with 2ml PBS and the nonbonded fraction is analyzed. The difference of added analyt and nonbonded analyt equals maximum column capacity.

Analytical Method:

<u>HPLC:</u> Shimadzu; <u>Column</u>: Trentec Reprosil-Pur RP C18 120 ODS3 5µm; 125x3,0mm with guard column; <u>Mobile Phase A</u>: acetonitrile / deion. water (70/30, v/v); <u>Mobile Phase B</u>: 0.05 M sodium acetate (pH=8.6) / acetonitrile (65/35 v/v); <u>Isocratic conditions</u>: 0.01 min B 80%; <u>Flow Rate</u>: 0.5ml/min; <u>Time of Analysis</u>: 25min;

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Injector Volume: 100µl; Fluorescence-Detection: λ_{EX} [nm]: 305nm; λ_{EM} [nm]: 418nm. Temperature: Machine and eluents are at room temperature. Eluents are degassed with helium gas.

Example Sample Calculation of Zearalenon (ZON) content:

A) Calculation of Sample Gramm Equivalents per HPLC injection:

20g Sample		1ml Extract		0.1ml		0.08g
	х		Х	injector	=	Sample
50ml Extraction		0.5ml		volume		Equivalents
Solvent						•

B) Calculation of ZON contamination of examined commodity in ng/g:

# ng injected ZON	=	ng/g ZON in e.g. ground corn meal
Sample Equivalents [g]		

Buffer, Chemicals, Apparatus and Literature:

Phosphate Buffered Saline pH 7.4 (= PBS)

1.24g KH ₂ PO ₄	Dissolve in 1L deionized water. If necessary
7.27g K ₂ HPO ₄	adjust pH to 7.4 (\pm 0.3) with 1N NaOH or 1N HCl
8.76g NaCl	

Chemicals:

acetonitrile, HPLC grade
methanol, HPLC grade
acetic acid, 100% ultrapure

deionized water

Elution Solvent:

Methanol

Consumables:

SENSIColumn IAC Zearalenon 3ml

dipotassium hydrogenphosphate, >98%
potassium dihydrogenphosphate, >98%
sodium chloride

Evaporation:

•nitrogen gas 5.0 [Air Liquide M55763810] (to evaporate IAC-eluate)

Apparatus:

HPLC; Shimadzu; pump: LC-6A (2 pieces); auto sampler: SIL 6B; fluorescence detector: RF-10AXL; data handling: CLASS LC10

Evaporator (with tripod) [VLM EVA EC1-S]

Vacuum SPE Manifold (BAKER spe-24G Column Processor – process up to 24 samples) [J.T. Baker 7208]

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¹ "Determination of zearalenone in corn by means of immunoaffinity clean-up and high-performance liquid chromatography with fluorescence detection", Angelo Visconti, Michelangelo Pascale *Journal of Chromatography A*, 815 (1998) 133–140