Sorbent-Based Microextraction of Natural Toxins from Food Samples

Subjects: Biochemical Research Methods Contributor: Natalia Casado

Natural toxins are chemical substances that are not toxic to the organisms that produce them, but which can be a potential risk to human health when ingested through food. Thus, it is of high interest to develop advanced analytical methodologies to control the occurrence of these compounds in food products. Current trends in sample preparation involve moving towards "greener" approaches by scaling down analytical operations, miniaturizing the instruments and integrating new advanced materials as sorbents. The combination of these new materials with sorbent-based microextraction technologies enables the development of high-throughput sample preparation methods, which improve conventional extraction and clean-up procedures.

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1. Introduction

Natural toxins are chemical substances naturally produced by living organisms (animal, plants or microorganisms) that are not toxic to them, but which can be potential health hazards to humans when ingested through food. These substances may naturally occur in food endogenously (toxic compounds that are implicit constituents of food resulting from the metabolism of a genus, species or strain, e.g., glycoalkaloids in potato or tetradotoxin in pufferfish) or exogenously (toxic compounds resulting from the metabolism of living organisms that occur in food as contaminants as they are not intentionally added, e.g., mycotoxins produced by molds grown in different products and toxins produced by algae that may be accumulated in edible marine organisms) ^{[1][2]}. The World Health Organization (WHO) encourages national authorities to monitor the most relevant natural toxins in the food supply. In this context, natural toxins of exogenous origin have received the most attention because of their potential harmful health risks and their involvement as natural contaminants. With respect to international organisms, these natural toxins of exogenous origin can be grouped in mycotoxins, phycotoxins (or marine toxins) and plant alkaloids [1][3][4]. Mycotoxins are toxic metabolites produced by certain types of molds, which can grow on a large number of foodstuffs such as cereals, dried fruits, nuts and spices. Most of these mycotoxins are chemically stable and survive food processing. The most common are aflatoxins (B1, B2, G1, G2 and M1), ochratoxins (A, B and C), patulin and fusarium toxins (deoxynivalenol, nivalenol, T-2 toxin, HT-2 toxin, zearalenone and fumonisins) ^[5]. On the other hand, marine toxins are produced during blooms of particular naturally occurring microalgae species in the ocean and fresh water. Thus, these toxins can be retained and bioaccumulated in shellfish and fish or contaminate drinking water. Their intake can be a potential hazard to consumers, since they are not eliminated by cooking or freezing, and might cause several adverse effects ^[6]. Conversely, in recent years, awareness about alkaloids of plant origin, such as pyrrolizidine, tropane and opioid alkaloids, has raised because of their occurrence as contaminants in different food products and the lack of data and knowledge about their exposure through food. These alkaloids are secondary metabolites of some plants, which can grow in fields as weeds and contaminate food crops appearing throughout the production of plant-derived products and finally be ingested, being toxic to humans [4][7][8][9][10]

[11][12]. The control of all these exogenous natural toxins in food is of high importance since they can cause from mild disorders (headache, vomiting, diarrhea, etc.) to serious situations (neurological disorders, carcinogenic, teratogenic or/and mutagenic effects, hepatic and renal damage, etc.) and can even be lethal. Moreover, they may cause the appearance of chronic diseases due to their harmful effects after a long-term exposure at high levels ^{[1][2][3][4][5][6][7][8][9][10]} ^{[11][12]}. Therefore, food safety plays an essential role in reducing the risks related to the presence of harmful substances in food in order to protect consumers. In fact, the WHO in collaboration with the European Food Safety Authority (EFSA), the Food and Agriculture Organization (FAO) and the Codex Alimentarius Commission have established a legislation for mycotoxins and marine toxins ^{[13][14]}, whereas pyrrolizidine, tropane and opioid alkaloids are in the process of being legislated, and at the moment only recommendations have been established for them ^{[15][16][17]}. In this sense, maximum residue limits (MRLs) for many of these natural toxins have been established in these guidelines to control the occurrence of these compounds in food ^{[13][18]}.

Nonetheless, to achieve these limits and ensure the health of consumers it is important to develop high-throughput, sensitive and selective analytical methods to determine in a feasible way the presence of these natural toxins in foodstuffs ^[19]. However, the analysis of these compounds in food samples constitutes a challenging task because of the extreme complexity of these matrices, which considerably hinders the selective extraction of the target analytes and decreases the sensitivity of the method ^[20]. Despite significant advances in analytical instrumentation, particularly with respect to the combination of mass spectrometry and chromatographic separation, these techniques are not sensitive enough for direct analysis of complex matrices. Therefore, sample preparation is still a crucial step in food analysis in order to achieve an effective isolation and/or preconcentration of the analytes and provide an adequate clean-up of matrix interferences prior to instrumental analysis ^[21].

For many years, liquid-liquid extraction (LLE) and solid-phase extraction (SPE) have been the most extensively used sample preparation techniques. Due to the inherent drawbacks of LLE (such as: time-consumption, limited ability to extract polar compounds, requirement of large volumes of solvents, etc.), SPE has become more popular, as it provides more efficient recoveries and lower solvent consumption than LLE ^[22]. Nevertheless, current trends in sample preparation involve moving towards "greener" approaches by scaling down analytical operations and miniaturizing the instruments ^[23] ^[24]. This has led in recent years to the development of different microextraction techniques for sample preparation procedures. In this sense, the SPE technique has been the axis of improving and creating even better and greener sorbent-based sample preparation techniques, which require less time and labor than SPE, such as: miniaturized solid-phase extraction (m-SPE), micro-dispersive solid-phase extraction (μ-dSPE), microextraction by packed sorbents (MEPS), pipette-tip solid-phase extraction (PT-SPE), solid-phase microextraction (SPME), stir-bar sorptive extraction (SBSE), and micro-solid-phase extraction (μ-SPE). These sorbent-based microextraction techniques have been proposed in recent years as an alternative to conventional sample preparation techniques to meet the Green Analytical Chemistry (GAC) requirements, as they involve advantages such as minimal solvent and sample consumption, fewer treatment steps, and reduction of waste generation ^[25]. Thus, they enable the development of cheaper, more cost-effective, and more environmentally friendly extraction and purification procedures.

On the other hand, the synthesis of new advanced materials for their application as sorbents in sample preparation has achieved considerable progress in the last decade, since these materials can play an important role in preconcentration processes and, in some cases, provide selective extraction of the target compounds ^{[20][21][23][26][27]}. Magnetic nanoparticles (MNPs), silica-based nanomaterials, metal-organic frameworks (MOFs), multiwalled carbon nanotubes

(MWCNTs) and graphene oxide (GO) are currently the most used materials for the extraction of natural toxins from food samples, as they present large surface area and advanced physicochemical properties that enhance the efficiency, selectivity and sensitivity of the analytical procedures ^{[21][27][28][29]}. Additionally, the combination of these new materials with microextraction technologies enables the development of high-throughput sample preparation methods, which provide the advantages of both strategies leading to meet the GAC requirements and improving conventional extraction and clean-up technologies ^{[23][30]}.

Some works in the literature have previously reviewed the determination of several natural toxins, such as phytotoxins ^[27] or mycotoxins ^[31], in food samples and other matrices. However, these works have just focused on one type of compounds but have not considered other natural toxins. On the other hand, other published reviews have addressed the development of new materials for their application to extract or detect chemical contaminants in order to ensure food safety ^[27][32][33].

2. Sorbent-Based Microextraction of Natural Toxins from Food Samples

The miniaturization of conventional sample preparation procedures has been proposed as an alternative for developing analytical methods with improved analytical characteristics (accuracy, precision, sensitivity, etc.) along with a decrease in sample and solvent consumption, reduction of hazardous reagents and wastes, and saving energy and time. As a result, new formats and configurations have arisen to carry out microextraction procedures, which overcome drawbacks of conventional techniques. Table 1 collects the most relevant works published in the last decade dealing with microextraction techniques based on sorbent-adsorption, which have been applied for the isolation of natural toxins from different food samples. In this sense, Solid-Phase Microextraction (SPME) has been the most popular [34][35][36][37][38]. However, procedures based on the dispersion of the sorbent material, such as micro-dispersive solid-phase extraction (µdSPE) and micro-solid-phase extraction (µ-SPE) have also been used [39][40][41]. All the works reviewed were performed for the analysis of mycotoxins (ochratoxins, aflatoxins, zearalenone, fumonisins and patulin) in different food matrices (mainly, wine, cereals and nuts). Only three of the methodologies developed in these articles perform the simultaneous determination of different types of mycotoxins [37][39][40], while the other works only described the individual determination of a specific analyte [34][35][36][38][41][42]. Concerning detection mode, mass spectrometry (MS) and fluorescence detection (FLD) were the techniques employed to detect these natural toxins (Table 1). Most of these works used MS detection, which is the most suitable technique to detect the presence of contaminants in food at trace levels thanks to its high sensitivity and to its structural elucidation capability, which enables the unequivocal identification and confirmation of the target analytes. In contrast, the FLD also provides high sensitivity and selectivity, but if the analytes do not show fluorescence it is necessary to carry out a derivatization process (pre-column or post-column derivatization) for their detection, which can sometimes be time consuming.

 Table 1. Application of sorbent-based microextraction techniques for isolation of natural toxins in food samples (2009–2019).

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
Cereal flours (2 g)	AF (B1, B2, G1, G2)	Extraction with 10 mL of MeOH/phosphate buffer (80/20, v/v, pH 5.8). Evaporation to dryness and reconstitution with 4 mL of phosphate buffer. An aliquot of the extract (2 mL) subject to microextraction.	SPME Sorbent: Commercial fibers Elution: 0.1 mL MeOH	HPLC- FLD	49–59	0.035- 0.2 µg/Kg	[<u>34]</u>
Nuts, cereals, dried fruits and spices (0.5 g)	AF (B1, B2, G1, G2)	Extraction with 1 mL of MeOH/H ₂ O (80/20, <i>v/v</i>). An aliquot of the extract (0.1 mL) mixed with 0.1 mL of 50 mM Tris buffer and brought to a total volume of 1 mL with H ₂ O before microextraction.	In-tube SPME * Sorbent: SUPEL- Q PLOT capillary	HPLC- MS	81– 109	0.0021- 0.0028 μg /L	[<u>35]</u>
Fruit juice and dried fruit (1 mL)	PAT	-	In-tube SPME * Sorbent: Carboxen-1006 PLOT capillary	HPLC- MS	> 92	0.023 μg /L	[36]
Nut and grain samples (0.5 g)	OTA, OTB	Extraction with 1 mL of MeOH/H ₂ O (80/20, <i>v/v</i>). Defatted with 3 mL hexane, supernatant discarded. An aliquot of the clean extract (0.1 mL) brought to a total volume of 1 mL with H ₂ O before microextraction.	In-tube SPME * Sorbent: Carboxen-1006 PLOT capillary	HPLC- MS	88	0.089- 0.092 μg /L	[<u>37]</u>
Wine (0.05 mL)	ΟΤΑ	-	In-tube SPME * Sorbent: Luna C18 particles	HPLC- MS/MS	61–73	0.02 μg/L	[<u>38</u>]

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
Powdered infant milk (3 mL) and	ZEN, α-ZAL, β-ZAL,	Extraction of milk samples with 0.15 mL acetic acid and 6 mL ACN. Evaporation up to 2.5 mL and reconstitution	μ-dSPE Sorbent: 80 mg of MWCNTs	HPLC-	77–	0.05– 2.02	[<u>39]</u>
mineral waters (50 mL)	u-zel, β-zel, ZAN	with H ₂ O to 25 mL, pH adjusted to 3.0 before microextraction.	Elution: 30 mL MeOH/Acetone (1/1, v/v)	103/103	120	μg/L	
Peach seed, milk powder, corn flour (0.2 g) and beer (0.2 mL)	AF (B1), OTB, T-2, OTA, ZEN	Microwave assisted extraction of solid samples with 0.2 g NaCl and 5 mL MeOH/H ₂ O (70/30, <i>v/v</i>). An aliquot of the extract (0.2 mL) brought to a total volume of 5 mL with H ₂ O before microextraction. Liquid samples diluted with H ₂ O up to 5 mL before microextraction.	μ-dSPE Sorbent: 12.5 μg zirconia nanoparticles Elution: 0.1 mL MeOH	UHPLC- MS/MS	84– 105	0.0022– 0.017 μg/L 0.0036– 0.033 μg/Kg	[40]
Coffee (10 g) and grape juice (10 mL)	ΟΤΑ	Extraction of coffee samples with 100 mL of carbonate. An aliquot of the extract (10 mL) adjusted to pH 1.5 before microextraction. Grape juice samples adjusted to pH 1.5 before microextraction.	µ-SPE Sorbent: 15 mg AFFINIMIP [™] OTA Elution: 0.25 mL MeOH/Acetic acid (98:2, <i>v/v</i>)	HPLC- FLD	91– 101	0.02– 0.06 µg/Kg	[<u>41</u>]
Wine (0.35 mL)	ΟΤΑ	-	MEPS Sorbent: 4 mg C18 sorbent	HPLC- FLD	76– 108	0.08 μg/L	[42]

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
			Elution: 0.05 mL				
			ACN/2% Acetic				
			Acid (90/10, v/v)				

 2. López de Cerain, A.; Gil, A.; Bello, J. Alimentos con sustancias tóxicas de origen natural: Plantas
 * Elution performed with mobile phase (online system); ACN; Acetonitrile; AF: Aflatoxin; F: Fumonisin; HPLC-FLD: High superiores alimenticias. In Toxicologia Alimentaria, 1st ed.; Cameán, A., Repetto, M., Eds.; Ediciones performance liquid chromatography. coupled to fluorescence; HPLC-MS/MS: High performance liquid chromatography coupled to fluorescence; HPLC-MS/MS: High performance liquid chromatography coupled to mass spectrometry; HPLC-MS: High performance liquid chromatography coupled to mass Becontani, anto in the Fard, Chais: Mailable Ching, packed sorbent; MWCNTs: Multiwalled carbon nanotubes; OTALTO://WANOXinefaacere.opanabu/sites/default/files/nefservep/blabseraser_nassets/sontamizants_tiskintherfeod_refault/files/nefservep/blabseraser_nassets/sontamizants_tiskintherfeod_refault.pdf Ultra fighs ferformatically and the second to fluorescence; UHPLC-MS: Ultra High performance liquid chremetagraphy, coupled to tandem mass spectrometry; ZAL: Zearalanol; ZAN: Zearalanone; ZEL: Zearalenol; ZEN: Zearales?nec.led?ppa.eu/no-dineare/serieve/selid-maare_seriere/iebinta?PiFiaMis/earabilegbensiantractions_en (accessed on

11 December 2019).

3. Integration of New Advanced Materials as Sorbents on Microextraction Techniques to Isolate Natural Toxins from Food Samples

6. Park, D.L.; Guzman-Perez, S.E.; Lopez-Garcia, R. Aquatic Biotoxins: Design and Implementation of

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the Va/aarletical WethEcolorSpecincence Netwe YourkiaNoarb/Selevits9988; decolumnee 15612 cepts. df 57772000 preparation is the choice of

the sorbent material. Depending on the analytes to be extracted, the sorbent material must have specific characteristics 7. EFSA: Occurrence of Pyrrolizidine Alkaloids in Food. Available online: that allow obtaining the highest extraction efficiency. In addition, using minimal amounts of sorbents is one of the http://www.efsa.europa.eu/en/supporting/pub/en-859 (accessed on 11 December 2019). requirements of the Green Analytical Chemistry (GAC) when developing an analytical procedure ^[29]. Thus, the sorbent AUEFSA Solentifio Apicionapprotectale and the post and the sole and th exthattp://www.enfsabeursnang.rnu/ma/efsaiguusnal/tpub/a386n/acceasenfuends1inPneeneverp20212) of analytical methods

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multiwalled carbon instruction of the extraction of the extraction of the extraction of

natural toxins from food products (Table 2). The advanced properties of these materials, such as their large surface area, 10. EFSA: Scientific Opinion on Pyrrolizidine Alkaloids in Food and Feed. Available online: low resistance to diffusion, fast sorption kinetics and large adsorptive capability make them very suitable for sample https://efsa.onlinelibrary.wiley.com/doi/abs/10.2903/j.efsa.2011.2406 (accessed on 11 December 2019). preparation, as they improve the efficiency, selectivity and sensitivity of the analytical procedures. Moreover, the

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can be improved and GAC requirements can be accomplished. In this sense, in the last decade, different new materials 12. EFSA: Risks for Human Health Related to the Presence of Pyrrolizidine Alkaloids in Honey, Tea, Herbal have been used to extract natural toxins from food products by their combination with different microextraction Infusions and Food Supplements. Available online: http://www.efsa.europa.eu/en/efsajournal/pub/4908 techniques, such as m-SPE, in-syringe SPE, PT-SPE, μ-dSPE, μ-MSPE, μ-SPE, SPME and SBSE (**Table 2**). They have (accessed on 11 December 2019). proved their efficiency in the extraction of several mycotoxins (mainly aflatoxins, ochratoxins, patulin and zearalenone)

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1	Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.	
1			Extraction with 25 mL of MeOH/H ₂ O (80/20, v/v). Evaporation of the						2019). Sampling
		AF (B1	methanolic fraction of an aliquot of the	m-SPE			0.012-		ssed on
2	Cereals (5 g)	(B2, B2, G1,	extract (15 mL). Addition of Britton- Robinson buffer (pH	Sorbent: 50 mg hyperbranched polymer	HPLC- FLD	83–103	0.120 μg/Kg	[<u>43]</u>	lications ion for
2		62)	5.2) up to 3 mL. An aliquot of the extract (2 mL) subject to microextraction.	Elution: 0.2 mL ACN					r organic 1, C.M.,
2				m-SPE					samples
2	Apple juice (1 mL)	PAT	-	Sorbent: 30 mg CD-based polymers	HPLC- DAD	n.p.	n.p.	[<u>44]</u>	egies in d new
2	in proce			Elution: 1 mL Diethyl ether/ACN (4/1, <i>v/v</i>)					s, S.; A 2019,

http://www.fao.org/input/download/standards/13794/CXP_074e_2014.pdf (accessed on 11 December

2	Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.	traction. A., Eds.;
2	Apple juice (1 mL)	PAT	Dilution with 1 mL of H ₂ O before microextraction.	m-SPE Sorbent: 50 mg SiO ₂ maleicpolymer@MIP Elution: 5 mL de acidified ACN	HPLC- DAD	82–98	8.6 μg/L	[<u>45]</u>	ecent
2	Apple,								19, 277,
2 3 3	apple juice, hawthorn, hawthorn juice, mixed juice, wines and tomato (10 q)	PAT	Extraction with 10 mL of ACN, 4 mg MgSO ₄ and 1 g NaCl. An aliquot of the extract (1 mL) evaporated to dryness and reconstituted with 1 mL H_2O before microextraction.	m-SPE Sorbent: 30 mg dual dummy-MIP Elution: 3 mL MeOH	HPLC- MS/MS	81–106	0.05– 0.2 μg/Kg	[<u>46]</u>	lvances : . TrAC
	0,								12–33.
3	Bell pepper, rice and	F (B1, B2,	Extraction with 6 mL ACN/H ₂ O (84/16, <i>v/v</i>). An aliquot of the extract (1 mL) evaporated to dryness	m-SPE Sorbent: 20 mg MIP	HPLC-	62–86	4.5–44	[<u>47</u>]	arly ew. Anal aterials
3	corn flakes (1 g)	B3)	and reconstituted with 1 mL ACN/H ₂ O (90/10, <i>v/v</i>) before microextraction.	Elution: 1 mL MeOH/Acetic acid (95/5, <i>v/v</i>)	MS/MS		μg/Kg		ırs by งไ
3								[48]	−d
	Maize, barley and oat (5 a)	T-2	Extraction with 25 mL of ACN/H ₂ O (84/16, <i>v/v</i>). For oat samples.	m-SPE Sorbent: 50 mg MIP	HPLC- MS/MS	60–73	0.4–0.6 μg/Kg	ريجير	graphy–
C	our (0 9)		after the solid-liquid extraction, the extract was additionally defatted with 10 mL of	Elution: 3 mL MeOH/Acetic acid (95/5, v/v)					samples J.

 Saito, κ.; ικεucni, κ.; καταοκα, Η. Determination of ocnratoxins in nuts and grain samples by in-tube solid-phase microextraction coupled with liquid chromatography–mass spectrometry. J. Chromatogr. A 2012, 1220, 1–6.

3	Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.	se
3			hexane. An aliquot of the sample extracts (1 mL) evaporated to dryness and						er netry
4			reconstituted with 1 mL MeOH/H ₂ O (20/80, <i>v/v</i>) before microextraction.						Yang, d using
4		AF (B1,	Extraction with 5 ml						o-solid
4		MI), OTA, ZEN,	ACN with 0.1% formic acid. Supernatant of	m-SPE					e based
4	Milk (1 mL)	α- ZEL, β-	the extract evaporated to dryness and reconstituted with 0.5	Sorbent: 10 mg rGO/Au Elution: 5 mL	UHPLC- MS/MS	70–111	0.01– 0.07 ng/mL	[<u>49</u>]	h non- n. Acta
4		zel, ZAN, α-	v/v) and diluted up to 5 mL with 5 mL of H ₂ O before microextraction.	MeOH/ACN/Formic acid (50/49/1, <i>v/v/v</i>)					
4		ZAL, β-ZAL							action of
4		AF	Extraction with 10 mL ACN/H ₂ O (75/25, <i>v/v</i>).	In syringe SPE			0.00		, A.; Yan ed solid-
4	Soy-based foods (2 g)	(B1, B2, G1, G2)	Diluted up to 50 mL with 10% NaCl aqueous solution before microextraction.	Sorbent: 30 mg 3DG@Fe ₃ O ₄ Elution: 0.7 mL MeOH	HPLC- FLD	83–103	0.09– 0.15 μg/Kg	[<u>50</u>]	solid- tam.
4	Soy-based	AF (B1.	Extraction with 10 mL	In syringe SPE	HPLC-	76–101	0.09–	[<u>51</u>]	is and
4		B2, G1,	Diluted up to 50 mL with 7% NaCl aqueous	Sorbent: PU/GO nanofibers	. 20		μg/Kg		e and liquid
		G2)		Elution: 0.75 mL MeOH					I Chem.

50. Nouri, N.; Sereshti, H.; ⊢aranani, A. Graphene-coated magnetic-sheet solid-phase extraction followed by high-performance liquid chromatography with fluorescence detection for the determination of aflatoxins B1, B2, G1, and G2 in soy-based samples. J. Sep. Sci. 2018, 41, 3258–3266.

5	Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.	_
5			solution before microextraction.						e one- nt
5	Maize (5 g)	AF (B1, B2, G1, G2)	Extraction with 20 mL ACN/H ₂ O (80/20, <i>v/v</i>). Evaporation to dryness and reconstituted with 0.1 mL MeOH. Diluted up to 10 mL with H ₂ O before microextraction.	In syringe SPE Sorbent: 15 mg β-CDPG Elution: 2 mL MeOH/DCM (2/1, <i>v</i> / <i>v</i>)	HPLC- FLD	91–105	0.0075– 0.030 µg/Kg	[52]	press. Dipette is. xtraction natogr. /
5		YTX,		PT-SPF					;ore-
5		OA, DTX (1),	Extraction with 9 mL MeOH. An aliquot of	Sorbent: 2 mg graphene					nilk and
5	Shellfish (0.2 g)	GYM, SPX (1), PTX (2), AZA (1)	GYM,the extract (0.1 mL)SPXevaporated to dryness(1),and reconstituted withPTX0.2 mL H2O before(2),microextraction.AZA(1)	Elution: 2 mL ACN with 0.5% ammonium hydroxide (for basic conditions) or with 0.5% formic acid (for acid conditions)	HPLC- MS/MS	78–90	0.1–1.5 μg/Kg	[<u>53]</u>	oodstuf
5			Extraction with						for
5	Peanut (50 g)	AF (B1, B2, G1,	MeOH/H ₂ O (80/20, v/v). An aliquot of the extract (8 mL) diluted	μ-dSPE Sorbent: 5 mg GO	HPLC- FLD	85–101	0.08– 0.65 µg/Kg	[<u>54</u>]	
		G2)	with H ₂ O before microextraction.	Elution: 2 mL MeOH					294,
6	Milk and yogurt (1.5 mL)	ZEN, α- ZEL,	Extraction of milk samples with 3 mL ACN and 0.075 mL	μ-MSPE Sorbent: 80 mg	HPLC- MS/MS	70–120	0.21– 4.77 μg/L	[55]	_
6		β- ZEL, ZAN,	acetic acid. Evaporation of the supernatant until 1.5	Fe ₃ O ₄ @pDA Elution: 8 mL MeOH		07.050			hene- olution

62. Wang, Y.; Wen, Y.; Ling, Y.C. Graphene oxide-based magnetic solid phase extraction combined with high performance liquid chromatography for determination of patulin in apple juice. Food Anal. Methods

6	Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.	f
-		α- ZAL,	mL and diluted with H_2O up to 25 mL, pH						_
6		β-ZAL	adjusted to 7 before microextraction. Extraction of yogurt samples with 4.5 mL						omoic }
6			and 0.075 mL acetic acid. The rest of the procedure the same as						fied n
6			for milk samples.						orio,
6	Mineral and tap water (25 mL)	ZEN, α- ZEL, β- ZEL, ZAN, α- ZAL, β-ZAL	Adjustment of pH to 7 before microextraction.	μ-MSPE Sorbent: 60 mg Fe ₃ O ₄ @pDA NPs Elution: 6 mL MeOH	HPLC- MS/MS	70–119	0.02– 1.1 μg/L	[<u>56]</u>	, 294, ıflatoxins larly / Acta
6	Red wine (50 mL)	AF (B1, B2, G1,	-	μ-MSPE Sorbent: 4.4 mg PD-MNPs Elution: 0.25 ACN/MeOH	HPLC- MS/MS	97–108	0.0012– 0.0031 μg/L	[<u>57</u>]	gnetic C.
7		G2)		(1/1, <i>v</i> / <i>v</i>)					ognition
7	Milk and		Extraction with 5 mL hexane and 5 mL	μ-MSPE					[.] bent for J.
7	dairy products (5 mL)	AF (M1)	MeOH/2 mM NaCl aqueous solution (8/2, <i>v/v</i>) before microextraction.	Sorbent: 8 mg AMNPs Elution: 2 mL DCM/MeOH/Acetic acid (80/19/1, v/v/v)	HPLC- FLD	97–116	0.2 ng/L	[<u>58</u>]	/ith solid- ≀togr. A

73. Es nagni, Z., Sorayaei, H., Samaui, F., Masroumia, M., Bakherau, Z. Fabrication of a nover nanocomposite based on sol–gel process for hollow fiber-solid phase microextraction of aflatoxins: B1 and B2, in cereals combined with high performane liquid chromatography–diode array detection. J. Chromatogr. B 2011, 879, 3034–3040.

7	Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.	nolithic ins.
7	Shellfish (2 g)	AZA (1, 2, 3), OA, DTX (1, 2)	Extraction with 10 mL MeOH/H ₂ O (4/1, <i>v/v</i>). The supernatant mixed with 2 mL hexane, evaporated until 1 mL and addition of 4 mL of H ₂ O before microextraction.	μ-MSPE Sorbent: 50 mg MMM Elution: 2 mL Formic acid/MeOH (5/95, <i>v/v</i>)	UHPLC- MS/MS	83–119	0.4–1.0 µg/Kg	[<u>59]</u>	traction '.
	Maize (6 g)	ZEN, α- ZEL, β- ZEL, ZAN, α- ZAL, β-ZAL	Extraction with 24 mL of ACN/H ₂ O (75/25, <i>v</i> / <i>v</i>). The extract diluted up to 25 mL with H ₂ O before microextraction.	μ-MSPE Sorbent: 5 mg MNPs- MWCNT-nanoC18 Elution: 1 mL ACN	HPLC- MS	92–98	0.6–1.0 μg/mL	[<u>60</u>]	
	Rice, wheat and sesame (50 g)	AF (B1, B2, G1, G2)	Extraction of rice and wheat samples with 200 mL Acetone/H ₂ O (50/50, v/v). Elimination of the acetone fraction before microextraction. Extraction of sesame samples with 100 mL hexane and 200 mL Acetone/H ₂ O (50/50, v/v). The rest of the procedure the same as for rice and wheat samples.	μ-MSPE Sorbent: 10 mg MGNP Elution: 2 mL Acetone/H ₂ O (1/1, <i>v/v</i>)	HPLC- FLD	64–122	0.025– 0.075 μg/Kg	[<u>61</u>]	

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
Apple juice (5 g)	PAT	Extraction with 5 mL ethyl acetate/hexane (96/4, <i>v/v</i>), 1 g NaH ₂ PO ₄ and 5 g Na ₂ SO ₄ . An aliquot of the organic phase (3 mL) mixed with 0.02 mL acetic acid, evaporated to dryness and reconstituted with 2 mL H ₂ O at pH 6.2 before microextraction.	μ-MSPE Sorbent: 30 mg MGO Elution: 1 mL ACN	HPLC- UV	69–83	2.3 µg/Кg	[62]
Milk (20 mL)	AF (B1, B2, G1, G2)	-	μ-MSPE Sorbent: 90 mg M/ZIF-8 Elution: 1 mL ACN/DCM (1/1, <i>v/v</i>)	UHPLC- MS/MS	79–102	2.3–8.1 ng/L	<u>[63]</u>
Seafood (5 g)	DA	Extraction with 20 mL MeOH/H ₂ O (1/1, <i>v/v</i>). The resultant sample extract subjected to microextraction.	μ-MSPE Sorbent: 1 mg Fe ₃ O ₄ SPs@ZIF8/Zn ²⁺ Elution: 0.4 mL 3 mM histidine solution	HPLC- MS/MS	93–102	0.2 ng/L	[<u>64]</u>
Shellfish samples (5 g)	DA	Extraction with 20 mL MeOH/H ₂ O (1/1, <i>v/v</i>). The resultant sample extract brought to a total volume of 25 mL with MeOH/H ₂ O (1/1, <i>v/v</i>) before microextraction.	μ-MSPE Sorbent: 1 mg Fe ₃ O ₄ @SiO ₂ @UiO-6 Elution: 1.5 mL ACN with 20% acetic acid	HPLC- MS/MS	91–107	1.45 μg/L	[<u>65</u>]

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
Beer (6 mL)	DON, ZEN, AF (B1, B2, G1, G2), F (B1)	Clean-up with a C18 sorbent. An aliquot of the clean sample (0.1 mL) evaporated to dryness and reconstituted with 0.48 mL ACN/H ₂ O/acetic acid (49/50/1, <i>v/v/v</i>) before microextraction.	μ-MSPE Sorbent: 25 mg MNM Elution: 0.5 mL ACN/H ₂ O/acetic acid (79/20/1, <i>v/v/v</i>)	UHPLC- MS/MS	87	n.p.	[<u>66</u>]
Corn (25 g)	AF (B1, B2, G1)	Extraction with 5 g NaCl and 125 mL MeOH/H ₂ O (7/3, <i>v/v</i>). An aliquot of the extract (15 mL) mixed with 45 mL of PBS before microextraction.	μ-MSPE Sorbent: 80 mg MNPC Elution: 1.2 mL ACN/H ₂ O (6/4, <i>v/v</i>).	HPLC- FLD HPLC- MS/MS	75–99	0.05– 0.07 μg/L	[<u>67]</u>
Tea leaves and corn (5 g)	AF (B1, B2, G1, G2)	Extraction with 10 mL ACN/H ₂ O (60/40, <i>v/v</i>). 5 mL of the extract subjected to microextraction.	μ-MSPE Sorbent: 10 mg MMIP Elution: 1 mL ACN/formic acid (95/5, <i>v/v</i>).	UHPLC- MS/MS	76–95	0.05– 0.1 µg/Kg	[68]
Rice (25 g) and wine (20 mL)	ОТА, ОТВ, ОТС	Extraction of rice samples with 100 mL ACN/H ₂ O (60/40, <i>v/v</i>) before microextraction. Wine samples diluted up to 25 mL with a solution of 2.5 M NaCl and 0.24 M NaHCO ₃ before microextraction.	μ-MSPE Sorbent: 15 mg Fe ₃ O ₄ @PDA MIPs Elution: 1 mL ACN	HPLC- FLD	71–88	0.0018– 0.018 μg/Kg	[69]

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
Grape juice	ΟΤΑ	-	μ-MSPE Sorbent: 5 mg MMIP Elution: -	UV-vis	97	0.374 mg/L	[<u>70</u>]
Coffee (10 g) and cereals (5 g)	ΟΤΑ	Extraction with 10 mL 1% carbonate aqueous solution. Sample extract adjusted to pH 1.5 before microextraction.	μ-SPE Sorbent: 10 mg LTL Elution: 0.4 mL MeOH	HPLC- FLD	92–101	0.09– 0.3 µg/Kg	[<u>71</u>]
Cheese (0.05 g)	ΟΤΑ	-	SPME Sorbent: Carbon-tape fiber Elution: 0.15 mL MeOH	HPLC- MS/MS	93	1.5 µg/L	[72]
Rice and wheat (10 g)	AF (B1, B2)	Extraction with 1 g NaCl and 100 mL MeOH/H ₂ O (80/20, <i>v</i> / <i>v</i>). Evaporation of the methanolic fraction of the extract and diluted with 40 mL H ₂ O. An aliquot of the extract (25 mL) subject to microextraction.	SPME Sorbent: 50 mg CNT Elution: 2 mL MeOH	HPLC- DAD	47–103	0.061– 0.074 μg/L	[73]
Rice (2 g)	AF (B1), ZAN, STEH	Extraction with 10 mL ACN/MeOH/H ₂ O (51/9/40, v/v/v), 1.5 g MgSO ₄ and 0.5 g NaCl. Evaporation to dryness and	SPME in-tube * Sorbent: MAA-co-DVB Elution:-	HPLC- PDA	78–103	0.69– 2.03 µg/Kg	[<u>74</u>]

Food Matrix (Amount)	Analytes	Sample Pretreatment	Microextraction Technique	Analysis	Recovery (%)	LOD	Ref.
-		reconstituted with 3 mL					
		0.1% TFA/ACN (99/1,					
		v/v) before					
		microextraction.					
Milk (1 g) and baby foods (3 g)		Extraction of milk					
		samples with 3 mL 1%					
		formic acid solution.					
		Supernatant discarded					
		and solid residue					
		extracted with 6 mL					
		chloroform.					
		Evaporation to dryness					
	AF	and reconstitution with	SBSE				
	(B1,	4 mL H ₂ O before					
	B2,	microextraction. Baby	Sorbent: 0.5 g MMIP-SB	HPLC-	30 60	0.3-1.0	[<u>75</u>]
	G1,	food samples		MS/MS	39-00	ng/Kg	
	G2,	dissolved with 1%	Elution: 3 mL MeOH/acetic				
	M1)	formic acid solution.	acid (75/25, <i>v/v</i>)				
		Supernatant discarded					
		and solid residue					
		extracted with 18 mL					
		chloroform.					
		Evaporation to dryness					
		and reconstitution with					
		$6 \text{ mL H}_2\text{O}$ before					
		microextraction.					

* Elution performed with mobile phase (online system); ACN: Acetonitrile; AF: Aflatoxin; AMNPs: Aptamer-functionalized magnetic nanoparticles; AZA: Azaspiracid; CD: Cyclodextrin; CNT: Carbon nanotube; DA: Domoic acid; DAD: Diode array detector; DCM: Dichloromethane; DON: Deoxynivalenol; DTX: Dinophysistoxin; F: Fumonisin; Fe₃O₄ SPs@ZIF8/Zn²⁺: Modified magnetic zeolite imidazolate framework-8; Fe₃O₄@PDA MIPs: Magnetic polydopamine-based molecularly imprinted polymer; Fe₃O₄@pDA NPs: Core_shell polydopamine magnetic nanoparticles; Fe₃O₄@SiO₂@UiO-6: Magnetite@silica core-shell magnetic microspheres; FLD: Fluorescence; GO: Graphene oxide; GYM: Gymnodimine; HPLC: High performance liquid chromatography; LTL: Zeolites linde type; M/ZIF-8: Magnetic zeolite imidazolate framework-8; MAA-co-DVB: Methacrylic acid-co-divinyl-benzene; MeOH: Methanol; MEPS: Microextraction by packed sorbent; MGNP: Magnetic-graphene nanoparticles; MGO: Magnetic graphene oxide; MIP: Molecular imprinted polymer;

MMIP: Magnetic molecularly imprinted polymer; MMIP-SB: Magnetic molecularly imprinted stir-bars; MMM: Magnetic mesoporous microspheres; MNM: Magnetic nanostructured materials; MNPC: Magnetic nanoporous carbon; MNPs: Magnetic nanoparticles; MS: Mass spectrometry; MS/MS: Tandem mass spectrometry; m-SPE: Miniaturized solid phase extraction; MWCNTs: Multiwalled carbon nanotubes; n.p.: Not provide; OA: Okadaic acid; OTA: Ochratoxin A; OTB: Ochratoxin B; OTC: Ochratoxin C; PAT: Patulin; PBS: Phosphate buffer saline; PDA: Photodiode array; PD-MNPs: Polydopamine magnetic nanoparticles; PT-SPE: Pipette-tip solid phase extraction; PTX2: Pectenotoxin-2; PU: Polyurethane; rGO: Reduced Graphene oxide; SBSE: Stir-bar sorptive extraction; SPE: Solid-phase extraction; SPME: Solid-phase microextraction; SPX1: Spirolides-1; STEH: Sterigmatocystin; TFA: Trifluoroacetic acid; T-2: T-2 toxin; UHPLC: Ultra high performance liquid chromatography; UV/vis: Ultraviolet/visible; YTX: Yessotoxins; ZAL: Zearalanol; ZAN: Zearalanone; ZEL: Zearalenol; ZEN: Zearalenone; β -CDPG: β -cyclodextrin supported on porous graphene nanohybrid; μ -dSPE: Micro-dispersive solid-phase extraction; μ -SPE: Micro-magnetic solid-phase extraction; μ -SPE: Micro-solid-phase extraction; 3DG@Fe₃O₄: Magnetic three-dimensional graphene sorbent.