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### Co-occurrence of aflatoxins and ochratoxin A in spices commercialized in Italy

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# UNIVERSITÀ DEGLI STUDI DI TORINO

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22	Co-occurrence of aflatoxins and ochratoxin A in spices commercialized in Italy
23	
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#### 40 ABSTRACT

41

42 A total of 130 spice samples coming from India, China, South America, USA, Northern Africa, Europe and Sub-Saharan Africa were collected in different stores of Northern Italy. 43 44 They were analyzed for aflatoxins (AFs: AFB<sub>1</sub>, AFB<sub>2</sub>, AFG<sub>1</sub>, AFG<sub>2</sub>) and ochratoxin A (OTA) content by liquid chromatography with mass spectroscopy and positive electrospray 45 ionization (LC/ESI-MS/MS), and HPLC with fluorescence detector (FLD), respectively. The 46 47 analysis showed that 20 (15.4%) and 31 (23,8%) out of 130 samples were contaminated with 48 AFs and OTA, respectively. A low level of total AFs contamination was found in the positive samples, the average concentration was 0.64 ng g<sup>-1</sup>, far below the maximum threshold 49 admitted by the European legislation (5 ng  $g^{-1}$  for AFB<sub>1</sub>, and 10 ng  $g^{-1}$  for total aflatoxins 50 (AFB<sub>1</sub>, AFB<sub>2</sub>, AFG<sub>1</sub> and AFG<sub>2</sub>). A higher incidence of OTA was found in chili (60.0%) more 51 than in pepper (13.3%), ranging from 2.16 to 16.35 ng  $g^{-1}$ , and from 1.61 to 15.85 ng  $g^{-1}$ , 52 respectively. Moreover, three spice samples (2.3%) contaminated by OTA trespassed the 53 54 threshold admitted by the European Regulation (EC, 2010). The co-occurrence of OTA and AFs in spices was detected in 6 out of 130 samples (4.6%), ranging from 1.61 to 15.85 ng  $g^{-1}$ 55 and from 0.57 to 3.19 ng  $g^{-1}$ , respectively. 56

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<sup>58</sup> Keywords: aflatoxins, ochratoxin A, chili, Italy, pepper.

#### 60 1. Introduction

61

52 Spices are widely used all over the world for food preparation, to increase the flavour and 53 aroma, and also in the past as food preservative. Considering the global trade value, the most 54 cultivated spice crops are pepper, capsicum, nutmeg, cumin and cinnamon. Spices are 55 commercially produced in a relatively low number of countries. India is the most important 56 spice producer (74% of the world market), followed by Bangladesh (6%), Turkey (5%) and 57 China (5%) (FAOSTAT, 2010).

The fungal growth and development on spices are favoured by environmental conditions, 68 such as temperature and humidity, and by poor manufacturing conditions in the production 69 70 region. Moulds decrease quality and quantity of food production, and may also create 71 potential risk for human and animal health, due to the production of secondary metabolites, 72 called mycotoxins. Some mycotoxins, like AFs and OTA, are produced by species of 73 Aspergillus and Penicillium. These mould species can develop in post-harvest, during drying 74 and storage, particularly when good storage practices are not adopted (Scott, 1984). 75 Aflatoxins (AFs, sum of aflatoxins  $B_1$ ,  $B_2$ ,  $G_1$ ,  $G_2$ ) and ochratoxin A (OTA) can be found in 76 several types of food commodities, including spices (Wangikar, Dwivedi, Sinha, Sharma, & Telang, 2005). 77

AFs are difuranocoumarin compounds primarily produced by certain species of *Aspergillus: A. flavus, A. parasiticus, A. nomius,* and *A. tamarri* (Yabe, Nakamura, & Hamasaki, 1999). *A. flavus* and *A. parasiticus* are mainly producers of AFs: aflatoxin B<sub>1</sub> (AFB<sub>1</sub>) ,B<sub>2</sub> (AFB<sub>2</sub>), G<sub>1</sub>
(AFG<sub>1</sub>) and G<sub>2</sub> (AFG<sub>2</sub>) (Varga, Frisvad, & Samson, 2011). Occurrence of AFs contaminations
on several agricultural products, e.g. maize, wheat, rice, spices, dried fruits and hazelnuts, are
worldwide reported (Grajewski, Blajet-Kosicka, Twaruzek, & Kosicki, 2012; Jackson,
Kudupoje, & Yiannikouris, 2012; Prelle, Spadaro, Garibaldi, & Gullino, 2012). AFs have

85 been clearly identified as toxic, mutagenic, teratogenic, and carcinogenic compounds. AFB<sub>1</sub> is 86 the most potent carcinogenic compound found in nature (Castells, Marin, Sanchis, & Ramos, 87 2008). The International Agency for Research of Cancer (IARC, 1993) has classified AFB<sub>1</sub> as 88 a human carcinogen (Group I). OTA is a mycotoxin primarily produced by some strains of 89 Aspergillus belonging to the sections Circumdati and Nigri. In particular, it is produced in 90 warm and tropical climates by Aspergillus ochraceus, and in temperate countries by 91 Penicillium verrucosum. (Tittlemier, Varga, Scott, & Krska, 2011; Varga, Kevei, Rinyu, 92 Teren, & Kozakiewicz, 1996). OTA is usually found in different foods, such as cereals, 93 spices, coffee, wine and dried fruit (Coronel, Marin, Cano-Sancho, Ramos, & Sanchis, 2012; Spadaro, Lore, Garibaldi, & Gullino, 2010). OTA has been identified as nephrotoxic, 94 95 hepatotoxic, immunotoxic and teratogenic, and classified in the group 2B by IARC (1993) as 96 a possible carcinogen for humans.

Due to favourable conditions in tropical climates, both mycotoxins can co-contaminate some
typologies of spices, not only in the field, but also during drying and storage.

99 The European Union fixed a maximum admitted level of 5 ng g<sup>-1</sup> for AFB<sub>1</sub>, and 10 ng g<sup>-1</sup> for 100 total AFs (AFB<sub>1</sub>, AFB<sub>2</sub>, AFG<sub>1</sub> and AFG<sub>2</sub>) intended for chili, chili powder, paprika, white and 101 black pepper, nutmeg, turmeric, ginger and spice mixtures containing one or more of the 102 above-mentioned spices (European Commission, 2010). In the case of OTA, the maximum 103 admitted level for the same spices listed before has been decreased from 30  $\mu$ g kg<sup>-1</sup> to 15  $\mu$ g 104 kg<sup>-1</sup> since 1/7/2012 (European Commission, 2012).

105 Despite many papers reported the co-occurrence of AFs and OTA content in spices from 106 different countries, such as Turkey (Ozbey & Kabak, 2012) and Malaysia (Jalili & Jinap, 107 2012), only one paper reported the AFs content in 28 spice samples marketed in Italy 108 (Romagnoli, Menna, Gruppioni, & Bergamini, 2007).

109 In Italy, the consumption of spices is low (around 50%), compared to other European 110 countries, such as France, Germany or Spain, but growing quickly, due to the recent 111 immigration from extra-European countries and to the diffusion of ethnical restaurants 112 (Dalpozzo, 2011).

The aim of this study was to analyse the co-occurrence of AFs and OTA in a large number of spice samples collected in different stores of Northern Italy. Two efficient and simple methods of extraction, purification and analysis were validated on several spices: AFs were simultaneous detected by liquid chromatography-tandem mass spectroscopy with electrospray ionization (LC/ESI-MS/MS), and OTA by liquid fluorescence detector (FLD).

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## 2. Materials and methods

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121 2.1. Chemicals and reagents

122 HPLC grade acetonitrile and LC-MS grade methanol, formic acid and acetic acid were 123 purchased from Sigma-Aldrich (St Louis, MO, USA). AFs (AFB1, AFB2, AFG1, AFG2) and 124 OTA analytical standards were purchased from Sigma-Aldrich, and dissolved in methanol and 125 acetonitrile, respectively, to prepare working standard solutions at the concentration of 10 µg 1<sup>-1</sup>. All standard solutions were stored in the dark at 4°C. NaCl, KCl, Na<sub>2</sub>HPO<sub>4</sub>, KH<sub>2</sub>PO<sub>4</sub>, 126 127 polyethylene glycol (PEG), ammonium formate, NaHCO<sub>3</sub> and Tween 20 were purchased 128 from Merck (Merck, Darmstadt, Germany). AflaClean select and OtaClean select 129 immunoaffinity columns were obtained from LCTech (Dorfen, Germany). HPLC eluents 130 were degassed for 5 minutes and filtered through mixed cellulose ester 0.22 µm-filters 131 (Advantec MFS, Pleasanton, CA, USA) before use.

132

133 2.2. Samples

A total of 130 spices coming from India, China, South America, USA, Northern Africa, Europe and Sub-Saharan area were purchased randomly in North-Western Italy stores, from September 2011 to February 2012. Due to high different typologies of spices, samples were divided in: 30 samples of pepper, 25 of hot pepper and paprika, 21 of mixed spices (such as curry and food seasoning), and 54 other spices including 15 different types of spices (cinnamon, cloves, ginger, juniper, poppy seeds, coriander, fennel, vanilla, mustard, turmeric, nutmeg, sesame, cardamom, anise, dill).

141 Two hundred mg of each sample were stored in plastic bags, in the dark, at low relative 142 humidity and 4°C before the analysis. All samples, except powder spices, were pulverized 143 using a food processor, until homogeneous. Spice sampling was done in accordance with 144 sampling provision described on European regulation No 401/2006.

145

#### 146 2.3. LC-MS/MS and HPLC apparatus

147 Liquid chromatography coupled with mass spectrometry was used to detect aflatoxins in spice 148 samples and to confirm OTA absence in spices used for validation method. 1260 Agilent 149 Technologies consisting of binary LC pump and a vacuum degasser; connected with a Varian 150 autosampler Model 410 Prostar (Hansen Way, CA, USA) equipped a 100 µL loop was used 151 as liquid chromatograph and was coupled to a triple quadrupole mass spectrometer Varian 152 310-MS. The analytical column used for LC separation was a Zorbax Eclipse Plus C18 (100 153 mm x 4.6 mm, 3.8 µm particle size, Agilent). The chromatographic conditions were: column 154 temperature: 45°C; mobile phase consisting of eluent A (water with 20 mM ammonium formate buffer at pH 3.35) and eluent B (methanol), using a flow rate of 0.3 ml min<sup>-1</sup>. A 155 156 gradient elution was applied as follows: 0-5 min, 50% A / 50% B - 30% A / 70% B; 5-10 157 min, 30% A / 70% B – 10% A / 90% B; 10-11 min, 10% A / 90% B, and 11-15 min, 10% A / 158 90% B – 50% A / 50% B. Two minutes of post run was applied. The injection volume was 10

159 µl.

The triple quadrupole mass spectrometer was operated in the positive electrospray ionization 160 161 mode (ESI<sup>+</sup>). The ionization source conditions were: needle voltage of 2.5 kV, capillary 162 voltage of 60-77 V, source temperature of 50 °C, desolvation temperature of 350°C, cone gas 163 flow rate of 50 psi, desolvation gas flow rate of 50 psi with nitrogen. Multiple reaction 164 monitoring (MRM) mode of operation was used. The [M+H]<sup>+</sup> ions of AFs were used as 165 parent ions. The most intense daughter ions, resulting from collision-induced dissociation 166 with argon, were used to detect and quantify AFs content. The argon pressure was set at 1.8 167 psi. The most intense daughter ions detected were: m/z 284.9 at 14 eV of collision energy 168 (CE) for AFB<sub>1</sub>, m/z 258.9 at 22 eV CE for AFB<sub>2</sub>, m/z 242.9 at 18 eV CE for AFG<sub>1</sub>, m/z 245 169 at 24 eV CE for AFG<sub>2</sub>, m/z 358 at 18 eV CE and m/z 239 at 30 eV CE for OTA.

170 OTA was detected with a HPLC apparatus 1100 series Agilent equipped with G1311 171 quaternary pump, G1379 degasser, G1313A autosampler, G1316A column thermostat and 172 G1321A FLD – Fluorescence Detector. The mobile phase consisted in an isocratic mixture of 173 acetonitrile:water:acetic acid (49:49:2) for 15 min. Sample (30  $\mu$ L) was injected into the 174 analytical column Synergi 4u Hydro-RP (250 mm x 4.6 mm, Phenomenex) and detected using 175 333 and 460 nm as wavelengths for excitation and emission, respectively.

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177

#### 178 2.4. Aflatoxins extraction and clean up

In the first part of this study, concerning the validation of extraction method, two samples for each matrix, which confirmed to be aflatoxin free, were used as follows: one aliquot of the sample was analysed as such, whilst other aliquots were spiked with a known concentration of mycotoxin standard.

183 The method for AFs extraction from hazelnut, described in a previously published paper

184 (Prelle et al., 2012), was used with slight modification. For 25 g of sample, 5 g of NaCl and 185 125 ml of extraction solution, methanol: water (80:20) were added, and left for 2 h on a 186 shaker apparatus at 165 rpm. To eliminate the solid phase from the extraction solution, the 187 sample was filtered, first, through a Whatman No. 4 filter paper, and subsequently by using a 188 Whatman CA 0.45 µm syringe filter. A 10 ml aliquot of filtrate was diluted 1:4 in phosphate 189 buffer solution (PBS) and centrifuged at 4,000 rpm. To clean up the samples, immunoaffinity 190 columns (IAC) were used, by loading 10 ml of diluted sample. IAC were washed with 10 ml 191 of PBS solution and then with 10 ml of water and air dried. AFs were eluted with 3 ml of 192 methanol into an amber glass vial. The elute was evaporated at 65°C under air flow and 1 ml 193 of mobile phase was added to the precipitate and vortexed until dissolved. All samples were 194 analysed in triplicate.

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#### 196 2.5. Ochratoxin A extraction and clean up

197 The absence of OTA, avoiding positive matrix effect of fluorescence signal, was confirmed 198 by analysis through LC-MS/MS. The samples (4g) were put in a 50 mL centrifuge tube with 199 20 mL of extraction solution, methanol:water (80:20), and left for 5 minutes in ultrasonic 200 apparatus at 25°C. To separate the solid sample from the extraction solution, centrifugation 201 for 15 min at 6,000 rpm was applied. After centrifugation, the solution was diluted 1:4 with 202 PBS solution and filtered by cellulose acetate 0.45 µm syringe filter. Ten ml of diluted and 203 filtered solution were loaded into IAC and, then, 10 ml of washing solution (2.5% NaCl and 204 0.25% NaHCO<sub>3</sub>) and 10 ml of ultrapure water were added into column. Before eluting with 3 205 ml of methanol into an amber glass vial, the column was air dried. The elute was evaporated 206 at 65°C under air flow and 1 ml of eluent was added to the precipitate and vortexed until 207 dissolved. All samples were analyzed in triplicate.

210 Validation of AFs and OTA methods was optimized studying apparent recovery rate (R%), 211 limits of detection (LOD) and quantification (LOQ), based on the IUPAC definition (Thompson, Ellison, & Wood, 2002), and, to validate the extraction method and 212 213 chromatographic performances, capacity factor (k'), repeatability of recovery (RSD) and 214 capacity factor (RSDk') were calculated. Following the guidelines of Commission Decision 215 2002/657/EC, these parameters were validated. The recovery (R%), RSD, LOD and LOQ 216 were determined on pepper, chili and four spices chosen by "others" (cinnamon, clove, 217 nutmeg and sesame) and two spice by "mixed" group (curry and food seasoning). Samples were spiked with four concentrations of standard solutions of  $AFB_1$  (0.5, 5, 10, 25 ng ml<sup>-1</sup>) 218 for pepper, cinnamon and seasoning food and (1, 5, 10, 25 ng ml<sup>-1</sup>) for chili, clove, nutmeg, 219 sesame and curry. Validation levels for total AFs were obtained spiking at 3, 5, 10, 25 ng ml<sup>-</sup> 220 <sup>1</sup>), and for OTA at 5, 10, 25, 50 ng ml<sup>-1</sup> with standard solutions. RSDk' and k' were 221 222 calculated, by analysing spiked cinnamon, clove, food seasoning and pepper samples at 10 ng ml<sup>-1</sup> of AFB<sub>1</sub>, total AFs, and OTA. Each test was performed three times. 223

Limits of detection (LOD) and quantification (LOQ) of each method for all mycotoxins were assessed. LOD was defined as three times the electronic baseline noise and LOQ as ten times the level of the baseline noise. The baseline noise was obtained with a blank sample for each matrix processed following the tested procedures. LOD and LOQ were calculated, analysing six blank samples and for total AFs were obtained by meaning the values of each mycotoxin. The recovery was calculated, using a protocol presented by (Matuszewski, Constanzer, & Chavez-Eng, 2003).

Capacity factors (k'), intended as a measure of the time the sample component resides in thestationary phase relative to the time in the mobile phase, were used to assure reliability and

repeatability of chromatographic analysis and were calculated for each mycotoxin (IUPAC,1997).

235

#### 236 2.7. Statistical analysis

237 Normal distribution of toxin contents, means, standard errors and validation data were 238 analysed with SPSS software (SPSS Institute, Inc, 2000, Version 18.0). The calibration curves 239 used for quantification were calculated by least-squares method. Samples with a concentration 240 of AFs and OTA higher that LOD were considered positives, whilst samples with 241 concentrations lower than LOD were considered negatives. Mean AFs and OTA 242 concentrations were calculated only on the positive samples higher than the LOQ. The 243 Kruskal-Wallis test was used to compare the mean AF and OTA levels among the different 244 typologies (pepper, chili, mixed and others), and geographical origins of spice samples, while 245 the Mann-Whitney test was used to compare the mean OTA and AFs levels in pepper/chili 246 samples and grain/powder of pepper and chili samples, using the null hypothesis that the 247 levels were not different.

248

249 3. **Results and discussion** 

250

This study represents the first report about monitoring of AFs and OTA co-occurrence in a large number of spice samples marketed in Italy. Two extraction and detection methods for AFs and OTA were validated on chili, pepper, cinnamon, nutmeg, food seasoning and curry matrices by LC/ESI-MS/MS and by FLD, respectively.

Due to the limited consumption, and consequently limited availability of spices in Italy, with the exception of pepper and chili, we grouped the spices in four categories: pepper, chili, others and mixed. The geographical origins of the spices analysed are shown in table 1. In

agreement with a FAO report (FAOSTAT, 2012) about the main spice producers, most of the
analysed samples were coming from India (74), followed by North Africa (33), and China
(11).

261

#### 262 *3.1. Analytical performance*

263 The analytical methods were validated considering: linearity, apparent recovery rate (R%), 264 capacity factor (k'), repeatability of recovery (RSD) and capacity factor (RSDk), limits of 265 detection (LOD) and quantification (LOQ) for AFs and OTA methods. Data of performed 266 analytical methods are summarized in Table 2. Due to complexity of chili and pepper matrix, LOD and LOQ obtained for OTA detection, 0.86 ng  $g^{-1}$  and 2.61 ng  $g^{-1}$  for pepper, and 0.70 267 ng  $g^{-1}$  and 2.13 ng  $g^{-1}$  for chili, respectively, were higher than other spices tested. These 268 results could be explained by presence of matrix compounds which interfere with analytical 269 270 signal increasing baseline noise (Prelle, Spadaro, Denca, Garibaldi & Gullino, 2013). OTA recovery rate ranged from 70.8 % at 25 ng  $g^{-1}$  of sesame to 102.1 % at 10 ng  $g^{-1}$  of nutmeg. 271 272 According to Castagnaro et al. (2006), we could explain these recovery results obtained as 273 interaction between several interferences on specific identification antigen-antibody present 274 into IAC. In particular, the specific antibodies identification could be reduced by the formation 275 of open-ring OTA at alkaline pH and by cross-reaction due to the presence of nonchlorinated 276 analogue OTB (Castegnaro, Tozlovanu, Wild, Molinie, Sylla, & Pfohl-Leszkowicz, 2006). Repeatability of OTA measurements, ranging from 0.3% for curry to 18.2% for food 277 278 seasoning, resulting in accordance with requirement established by EU regulation. On the 279 contrary the values of LOD and LOQ for chili and pepper for aflatoxin detection, ranging from 0.08 ng  $g^{-1}$  of AFB1 to 0.41 of AFs, respectively, for pepper and from 0.17 ng  $g^{-1}$  of 280 281 AFB1 to 0.71 of AFs, respectively, for chili resulted less influenced by matrix interferences on analytical signal than OTA detection. Recovery values for AFs and AFB1 were above of 282

283 70% for all spices tested, despite the matrix complexity of some kind of spice, such as curry 284 and food seasoning, composed by different quantity of other spices. In the other hand the k' 285 and RSDk'values obtained reported in Table 3, demonstrated the high repeatability and 286 hardiness (RSDk'% < 0.1) of optimized chromatographic method, despite different spices 287 analysed.

288

289 *3.2. Mycotoxin occurrence* 

AFs were found in 20 (15.4%) out of 130 samples analysed with levels ranging from 0.59 to 5.38 ng g<sup>-1</sup>. The average concentration was 0.96 ng g<sup>-1</sup>. Five samples of each typology of spice (pepper, chili, mixed and others) resulted positive to one or more AFs. The AFs contamination range was between 0.59 and 3.68 ng g<sup>-1</sup>, with 11 (8.5%) samples contaminated by AFB<sub>1</sub>, with a mean on contaminated samples of 0.31 ng g<sup>-1</sup>, 14 (4.6%) by AFB<sub>2</sub> (contamination mean: 1.07 ng g<sup>-1</sup>), 3 (2.3%) by AFG<sub>1</sub> (0.42 ng g<sup>-1</sup>), and 7 (5.4%) by AFG<sub>2</sub> (2.26 ng g<sup>-1</sup>) (Table 4).

297 No sample was contaminated by AFs above the maximum admitted threshold established by 298 the European legislation (European Commission, 2010). In fact, the highest AFB<sub>1</sub> level detected was 1.95 ng g<sup>-1</sup>, 2.5 times lower than the maximum threshold specific for AFB<sub>1</sub> In 299 300 contrast with the results obtained by Romagnoli et al. (2007) on monitoring of AFs on spices 301 purchased in Italy, we detected a lower incidence of contamination and a lower level of AFs. Although pepper and chili are usually more susceptible substrates to mycotoxins 302 303 contamination (Reddy, Mayi, Reddy, Thirumala-Devi, & Reddy, 2001), our results showed 304 no differences among the spice typologies analysed. The low amount of AF content detected 305 in our study agrees with the results reported by similar studies ((Jalili & Jinap, 2012; Ozbey & 306 Kabak, 2012). The incidence of positive samples was slightly higher in the mixed and chili samples, 23.8% and 20.0%, respectively, compared to pepper (16.6%) and other (9.26%) 307

308 spices. The Kruskal-Wallis test showed that the level of contamination of AFs in different 309 countries were not statistically different (p=0.586). Average AF content in pepper and chili 310 samples were calculated with Mann-Whitney test, and resulted not statistically different 311 (p=0.248).

Considering the OTA contamination in the spice samples, the incidence resulted higher than AFs: 23.8% spices were positive to OTA, while only 15.4% to AFs. The values of OTA contamination ranged from LOD to 19.06 ng g<sup>-1</sup>. The contamination OTA mean on positive samples was 6.18 ng g<sup>-1</sup>. Fifteen out of 25 (60%) chili samples were contaminated at levels ranging from 2.16 to 16.35 ng g<sup>-1</sup>, 7 out of 21 (33.3%) mixed samples at levels ranging from 1.84 to 19.06 ng g<sup>-1</sup>. Pepper (13.3%) and other samples (9.3%) resulted positive ranging from 1.61 to 15.85 and from 2.21 to 11.08, respectively (Table 4).

Similarly to AFs, results of Kruskal-Wallis test showed no significant differences among the geographical origins of the samples. The Mann-Whitney test, instead, reported significant differences on OTA contamination levels between chili and pepper (p<0.05). Significant differences were also noticed for grain and powder chili (p<0.05). Chili powder showed higher OTA content (78.9%) than chili grains, probably, due to the higher contact surface presented to the microbial population, which can favor the mycotoxin release.

The current monitoring showed a higher incidence of OTA than AFs in spice samples marketed in Italy; moreover, concentration levels of OTA detected in three (2.3%) samples out of 130 trespassed the threshold admitted by the European Regulation (EC, 2010).

The co-occurrence of OTA and AFs in spices was detected in 6 out of 130 samples(4.6%), in particular in chili, pepper and curry samples. OTA and AFs concentration ranged from 1.61 to 15.85 ng g<sup>-1</sup> and 0.57 to 3.19 ng g<sup>-1</sup>, respectively. Several studies were performed over the last decade on the co-occurrence of AFs and OTA in spices. Most of the studies highlighted that the reason for a high mycotoxin contamination is the susceptibility of spices to fungal 333 contamination, and consequently to mycotoxin production, connected to environmental 334 conditions, such as high humidity and temperature (Patharajan et al., 2011), of the producer 335 countries located in tropical and sub-tropical areas. Regarding the co-occurrence of AFs and 336 OTA in same substrate, Sedmikova, Reisnerova, Dufkova, Barta, & Jilek (2001) 337 demonstrated that OTA could increase the mutagenic activity of AFB<sub>1</sub>. In agreement with our 338 results, most of previous work reported high incidence of AFs and OTA in pepper and chili. 339 In particular, during the last decade, AFs and OTA were monitored in over 1,500 spice 340 samples, resulting in a contamination incidence ranging from 30 to 70%. Despite the wide 341 range of spices present, attention mainly focused on pepper and chili, due to their high 342 demand in the market worldwide and to their susceptibility to fungal contamination, 343 especially during the drying processes (Shundo, de Almeida, Alaburda, Lamardo, Navas, 344 Ruvieri, et al., 2009). According to the available literature and to our results, chili and pepper 345 are more frequently contaminated by AFs and OTA than other spices, such as cumin or 346 cinnamon. This is the first study on AFs and OTA co-occurrence in a large number of spice 347 samples marketed in Italy. Future studies will also monitor the fungal population present on 348 the spices, which could be related to the AFs and OTA contamination, with microbiological 349 an molecular tools (Spadaro, Patharajan, Karthikeyan, Lorè, Garibaldi, & Gullino, 2011).

350

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- 456 Tables

# **Table 1.** Geographical origin of spice samples.

	Spices					
Origin	Pepper (n)	Chili (n)	Mixed (n)	Others (n)		
South America	2	2	0	0		
India	25	15	8	26		
North Africa	2	2	13	16		
Europe	0	0	0	1		
China	1	5	0	5		
Sub-Saharan Africa	0	1	0	5		
USA	0	0	0	1		
Total samples (n)	30	25	21	54		

Analytes	Matrix	Fortification	Recovery (n=3)	RSD (%)	LOD (ng g-1)	LOQ (ng g-1)
		(ng mi)	(%)	(11-3)	( <b>n=6</b> )	( <b>n=6</b> )
Aflatoxin B <sub>1</sub>	pepper	0.5, 5, 10, 25	91.8, 97.8, 99.6, 99.7	3.8, 0.6, 2.8, 3.5	0.08	0.28
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	pepper	3, 5, 10, 25	79.1, 82.8, 82.5, 83.3	25.6, 23.5, 25.0, 25.4	0.12	0.41
Aflatoxin B <sub>1</sub>	chili	1, 5, 10, 25	100.6, 97.3, 93.4, 90.5	1.8, 2.0, 2.9, 2.7	0.17	0.55
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	chili	3, 5, 10, 25	78.04, 83.4, 85.5, 88.0	1.9, 1.1, 2.5, 1.2	0.21	0.71
Aflatoxin B <sub>1</sub>	cinnamon	0.5, 5, 10, 25	99.8, 99.8, 99.6, 99.9	0.1, 2.7, 3.1, 3.0	0.12	0.41
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	cinnamon	3, 5, 10, 25	80.5, 81.5, 81.9, 83.1	24.8, 25.5, 25.8, 23.7	0.18	0.61
Aflatoxin B <sub>1</sub>	nutmeg	1, 5, 10, 25	79.0, 77.4, 77.6, 82.1	12.3, 14.3, 8.8, 8.9	0.2	0.66
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	nutmeg	3, 5, 10, 25	76.9, 80.3, 89.7, 87.4	9.1, 5.9, 12.7, 7.3	0.65	2.2
Aflatoxin B <sub>1</sub>	clove	1, 5, 10, 25	81.6, 92.7, 90.3, 85.1	13.3, 16.9, 12.4, 10.5	0.17	0.58
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	clove	3, 5, 10, 25	83.9, 96.6, 87.3, 81.2	15.6, 13.0, 9.4, 11.4	0.57	1.94
Aflatoxin B <sub>1</sub>	sesame	1, 5, 10, 25	73.2, 73.0, 77.8, 83.6	5.9, 3.2, 7.3, 5.6	0.19	0.62
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	sesame	3, 5, 10, 25	82.7, 81.1, 77.3, 81.9	20.3, 13.2, 10.4, 12.6	0.33	1.12
Aflatoxin B <sub>1</sub>	food seasoning	0.5, 5, 10, 25	97.2, 97.6, 87.8, 81.8	4.7, 1.1, 2.9, 1.8	0.13	0.45
Aflatoxins $B_1$ , $B_2$ , $G_1$ , $G_2$	food seasoning	3, 5, 10, 25	92.8, 93.1, 94.3, 83.8	18.6, 6.3, 4.7, 5.1	0.26	0.88
Aflatoxin B <sub>1</sub>	curry	1, 5, 10, 25	71.6, 74.7, 79.8, 80.3	19.6, 17.7, 15.9, 15.3	0.25	0.83
Aflatoxins $B_1$ , $B_2$ , $G_1$ ,	curry	3, 5, 10, 25	73.0, 71.6, 78.6, 79.4	17.3, 17.1, 18.0, 17.9	0.77	2.62

**Table 2.** Recovery of AFs and OTA spiked into pepper and cinnamon samples.

$G_2$						
OTA	pepper	5, 10, 25, 50	91.1, 94.8, 97.8, 98.7	2.1, 2.9, 4.8, 2.1	0.86	2.61
OTA	chili	5, 10, 25, 50	81.1, 79.3, 88.4, 87.8	4.7, 8.6, 4.5, 6.7	0.70	2.13
OTA	cinnamon	5, 10, 25, 50	97.4, 96.3, 99.2, 95.6	3.4, 4.5, 4.5, 3.6	0.03	0.09
OTA	nutmeg	5, 10, 25, 50	80.2, 102, 1, 84.6, 71.0	17.8, 7.8, 0.8, 0.4	0.03	0.10
OTA	clove	5, 10, 25, 50	78.4, 80.4, 75.3, 76.0	2.3, 17.1, 13.0, 0.68	0.02	0.07
OTA	sesame	5, 10, 25, 50	73.5, 74.5, 70.8, 75.4	6.0, 6.1, 6.9, 14.5	0.05	0.14
ΟΤΑ	food seasoning	5, 10, 25, 50	78.1, 78.2, 82.7, 77.3	1.8, 14.5, 18.2, 2.5	0.05	0.15
OTA	curry	5, 10, 25, 50	79.6, 72.9, 78.3, 71.2	0.3, 8.2, 9.4, 0.8	0.04	0.12

Analytes	Matrix	Capacity factor (k')	<b>RSDk'</b> (%)
AFs	pepper	38.1, 36.6, 34.1, 31.8	0.06, 0.07, 0.09, 0.1
OTA	pepper	5.2	0.04
AFs	cinnamon	38.4, 36.7, 34.3, 31.9	0.05, 0.07, 0.07, 0.09
OTA	cinnamon	5.2	0.05
AFs	food seasoning	38.0, 36.6, 34.3, 31.7	0.03, 0.04, 0.03, 0.06
OTA	food seasoning	5.2	0.03
AFs	clove	37.9, 36.2, 34.1, 31.0	0.09, 0.1, 0.1, 0.12
OTA	clove	5.3	0.07

**Table 3.** Analytical chromatographic performance.

			Number of positive samples (ng g <sup>-1</sup> )				
Mycotoxins	Positive/total (%)	Mean of contamination ± SD (ng g <sup>-1</sup> )	< LOD	LOD- 5.0	5.0-10.0	10.0- 15.0	>15.0
$AFB_1$	11/130 (8.5%)	0.31±0.003	8	3	-	-	-
$AFB_2$	6/130 (4.6%)	$1.07 \pm 0.086$	4	2	-	-	-
$AFG_1$	3/130 (2.3%)	$0.42\pm0025$	3	-	-	-	-
AFG <sub>2</sub>	7/130 (5.4%)	2.26±0.191	3	4	-	-	-
OTA	31/130 (23.8%)	6.18±0.639	-	15	11	2	3
OTA+AFs	6/130 (4.6%)	$0.85 \pm 0.030$					

464	Table 4. Occurrence and levels of AFs and OTA in the spices analysed.
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