Comparison of deoxynivalenol and zearalene concentration in conventional and organic cereal products in western Poland

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A – Research concept and design, B – Collection and/or assembly of data, C – Data analysis and interpretation,
D – Writing the article, E – Critical revision of the article, F – Final approval of article


INTRODUCTION

Food contamination with mycotoxins is a public health threat [1]. Mycotoxins are regarded as priority food contaminants by the Global Environmental Monitoring System/WHO Monitoring Programme on Food Pollutants (GEMS/Food 1995) [2]. They are usually thermostable and tend to survive during the transformation and processing of contaminated plants, and are not usually eliminated during cooking and sterilization [3].

Several factors can affect the occurrence of mycotoxins: harvesting, storage, transport of grain, and agricultural technology, are some of the basic factors. Therefore, the right approach and actions in this area can significantly reduce mould growth, as well as affect the presence of mycotoxins [4]. Because of their ability to bind to plasma proteins, mycotoxins can persist in organisms if the exposure is repeated or chronic [5]. Due to the variety of chemical and biosynthetic structures, mycotoxins have innumerable biological effects and some of these are classified as carcinogenic, immunosuppressive or estrogenic, potentially causing serious metabolic disturbances in humans [6].

DON are highly toxic mycotoxins [7] and occur in basic food products and commonly consumed products, such as cereals, milk and dairy products, dried fruit and legumes, coffee, wine and beer [8]. The toxicological effects and diseases associated with exposure to DON have recently been investigated; however, epidemiological studies are required to critically examine the potential connection between the consumption of large quantities of DON and the incidence of gastro-enteritis and chronic diseases.

ZEA (Zearalenone) and its metabolites can be found worldwide in a range of cereals, including corn, sorghum, wheat, rice, barley and oats. ZEA may also be consumed as a result of ingesting meat, milk and eggs from animals exposed to these disease toxins, or which received ZEA to promote their growth. ZEA and its metabolites show strong estrogenic activity, hence it is often referred to as mycogen. ZEA plays an important role in increasing the risk of hormone-dependent cancers [9].

MATERIALS AND METHOD

During the 2014–2015 timeframe, a total of 78 samples were examined, of which 20 were unprocessed cereals (maize, wheat, oat, barley), and 58 were cereal-based products (flour, groats, pasta). Among them, 39 samples were declared to be organic (10 cereals and 29 cereal-based products), while 39 samples were produced in a conventional manner (10 cereals and 29 cereal products). During the harvesting period (July–September) in 2014, unprocessed cereals were sampled from agricultural fields situated in western Poland. In 2015, cereal products were sampled in the form of packaged commercial
products produced by different domestic manufacturers available on the Polish market.

Samples of organically produced products had been labeled in accordance with Regulation 834/2007/EC on organic production and labeling of organic products, which repeals Regulation (EEC) No. 2092/91 [10]. These products also contain the national or private organic producers’ logo, the competent authority code and place of cultivation designation. Sampling and sample preparation were carried out in accordance with Commission Regulation 401/2006 [11] establishing sampling methods for the official control of mycotoxins levels in foodstuffs. Aggregate samples were taken of not less than 1 kg, connecting the 3 primary samples (a minimum of 500 g of each sample). Samples were stored in a cool, dry place, and transported to the laboratory within 48 hours.

The prepared test portions were ground to a 1.0 mm fine powder using an analytical mill and then stored in plastic bags at -4 °C until analysis. The mycotoxins were isolated from cereals and cereal products using Donprep® and EASIEXTRAT Zearalenon® immunoaffinity columns by R-Biopharm Rhône for extracting mycotoxins in accordance with the manufacturer’s procedures. The mycotoxins were analysed using high-performance liquid chromatography with fluorescence detection (HPLC-FLD).

**Determination of DON and ZEA content.** DON content was determined according to the National Institute of Hygiene Methodology – Determination of Fusarium toxins – deoxynivalenol (DON) in cereals and its products by high-performance liquid chromatography with purification using immunoaffinity columns.

The principle of the method is to extract a sample with a mixture of polyethylene glycol and water. The centrifuged and filtered extract is purified and concentrated in immunoenzyme columns. After elution with methanol and water, the methanol/aqueous solution is evaporated to dryness and dissolved in 9.5% methanol. The solution is injected into the chromatographic column. The measurement is carried out at a wavelength of 220 nm. The qualitative determination involves comparing the retention time of the standard and the sample tested. The quantitative determination involves reading the DON content in the test sample from the standard curve and making the appropriate calculation.

The DON content of the sample is expressed in μg kg⁻¹, calculated as:

\[
x = \frac{cxV3xV2}{mxV1xV3}
\]

X – DON content in the product [μg/kg];
C – DON content read from the curve [ng/cm³];
V1 – volume of extract extracted [cm³];
V2 – volume after dilution in PBS [cm³];
V3 – volume applied to the affinity column [cm³];
V4 – volume of eluate from the enzyme-linked column [cm³];
V5 – volume of extraction solvent [cm³];
m – weight [g].

Detection limit (LOD) – 5 μg kg⁻¹, limit of quantification (LQD) – 10 μg kg⁻¹, average recovery – 96%; the measurement is made using a fluorescence detector and an excitation wave of 276 nm, emission – 460 nm.

**Measurement conditions.** The solution was injected on a RP C18 250 x 4.6 mm column, 5μm, mobile phase composition: methanol 15%; water 85%, mobile phase flow – 1 cm³/min, injection volume – 100 μl, column temperature – 40°C, autosampler temperature – 4°C, wavelength – 220 nm.

ZEA was determined according to the National Institute of Hygiene Methodology – Determination of Fusarium toxins – zearalenon in cereals, and dairy products by high-performance liquid chromatography purification using immunoaffinity columns. The principle of the method is to extract the sample with 75% acetonitrile and water. The extract then filtered through filter paper, 20 cm² of the filtrate removed and mixed with 80 cm³ of saline (PBS), and pH 7.4 adjusted using HCl (0.1 mol/l). 25 cm³ of the solution was passed through a bed of EASIEXTRAT Zearalenon® enzyme immunoassays. Purification and concentration in the columns was performed according to R-Biopharm Rhône applications.

After elution, the acetonitrile-aqueous solution was injected into the chromatographic column. The ZEA quality assay consisted of comparing the retention time of the standard and the sample being tested, along with quantitative determination of the content of the mycotoxin tested in a sample from the standard curve the appropriate calculation was made.

ZEA content in the sample is expressed in μg kg⁻¹, calculated as:

\[
x = \frac{cxV2V4xV5}{mxV1xV3}
\]

X – ZEA content in the product [μg kg⁻¹];
C – ZEA content read from the calibration curve [ng/cm³];
V1 – volume of extract extracted [cm³];
V2 – volume after dilution in PBS [cm³];
V3 – volume applied to the affinity column [cm³];
V4 – volume of eluate from the enzyme-linked column [cm³];
V5 – volume of extraction solvent [cm³];
m – weight [g].

Detection limit (LOD) – 5 μg kg⁻¹, limit of quantification (LQD) – 10 μg kg⁻¹, average recovery – 96%; the measurement is made using a fluorescence detector and an excitation wave of 276 nm, emission – 460 nm.

Measurement conditions: RP C18 chromatographic column 250 x 4.6 mm, 5μm, mobile phase composition, acetonitrile 60% – water 40%, mobile phase flow – 1 cm³/min, injection volume – 100 μl, column temperature – 40°C, autosampler temperature – 4°C, emission wavelength – 460 nm, excitation wavelength – 276 nm.

**Statistical analysis.** The results were analysed statistically using STATISTICA v.10 software. The difference in mycotoxin concentration in the groups was assessed using the Mann-Whitney test. Assumed statistical significance level – p<0.05.

**RESULTS**

A total of 78 samples from organic and conventional systems were analyzed with HPLC. Occurrence of DON was detected in cereals from both production systems – organic and conventional. In cereals from the organic production system, the average content of DON was 285.25 ± 134.04 μg kg⁻¹ and was detected in 40% of the samples; range of contamination – 130–433 μg kg⁻¹. DON was detected in 70% of the cereal samples from conventional cultivation system, average content – 373.71 ± 171.20 μg kg⁻¹, range of contamination – 137–529 μg kg⁻¹.
In flour from organic farming, the average DON content was 213.80 ± 151.28 μg kg⁻¹ and detected in 50% of samples (range 104–459 μg kg⁻¹), in flour from conventional cultivation the average was 336.29 ± 188.90 μg kg⁻¹, level between 146–646 μg kg⁻¹, 70% of the flour samples were contaminated with the test mycotoxin.

The range of DON concentrations in samples of cereal products from organic and conventional cultivation was 102.00–465.00 μg kg⁻¹ and 137.00–792.00 μg kg⁻¹. It was detected in 26.3% and 31.6%, whereas the average concentrations of DON in cereal products was 199.60 ± 149.82 μg kg⁻¹ and 387.67 ± 250.24 μg kg⁻¹ in samples of cereal products from organic and conventional cultivations (Tab. 1).

There was no ZEA presence in cereals from either organic or conventional cultivation. Only one sample of flour from conventional cultivation was found to contain ZEA. ZEA was also found in one sample of cereal products from organic cultivation and 3 samples from conventional cultivation. Furthermore, ZEA was detected in 5.3 and 15.8%, whereas the average concentrations of ZEA in cereal products was 13,00 μg kg⁻¹ and 14,45 ± 2,76 μg kg⁻¹ in samples of cereal products from organic and conventional cultivations (Tab. 2).

**DISCUSSION**

Interest in organic food has increased worldwide in response to concerns about conventional farming practices, food safety, human health, animal welfare and the environment. In the EU, organic farming is an agricultural practice and modality of food production that combines favourable environmental and animal welfare standards and is supported by EU law [12, 10].

Today, many consumers prefer organic rather than conventional food because synthetic fungicides and mineral fertilizers are not used in organic production. It is generally believed that organic practices reduce the risk of plant infection [13], but at the same time there is an awareness that the worse use of fungicides may promote the presence of mycotoxins in ‘natural’ or ‘home’ chemicals – free products [14].

Data published insofar have shown that the presence of mycotoxins as frequent contaminants of different food products and feedstuffs greatly varies dependent on many parameters of influence, such as the type of cereal, weather conditions and among others [15, 16, 17, 18].

In this study, the occurrence of the mycotoxins in different unprocessed cereals cultivated in fields in Poland in an organic and conventional manner, and in cereal-based products available on the Polish market, was investigated. It was found that the average DON content was highest in samples of conventional cereal products and conventional flour – 387.67 and 336.29 μg kg⁻¹ respectively. It should be noted that the highest level of the mycotoxin being tested was recorded in a sample of pasta from conventional grains – 792 μg kg⁻¹. This level exceeded the permissible limit specified in the regulation [19].

Malmuaret et al. in France reported a co-occurrence of DON contamination of organic wheat was lower than that in conventional wheat (54.4% vs. 90.9%, respectively), but the medium and maximum levels of this mycotoxin were 2–3 times higher in cereal grains from organic farming than in the conventional (250 vs. 80 and 494 vs. 215 μg kg⁻¹, respectively) [20]. A study conducted in Germany between 1997–1998 [21] reported a greater contamination of DON wheat grains from organic farms. Malmuaret et al. [20] noted that more ‘conventional’ seeds were contaminated, although at a low level, while organic grains were less often contaminated, although in a few cases at a high level.

DON has proved to be a more common mycotoxic in cereal grains than ZEA. Many authors confirm the frequent presence of DON in cereals grown both in Poland and in other parts of the world. DON contamination was noted in a large percentage of samples tested in Russia, the USA, Bulgaria, China and Hungary, ranging from 50% – 96% [22, 23, 24]. DON levels in cereal grains increase in years with frequent rainfall and high humidity; for example, there was less pollution of rye with DON in Bavaria in 2000 and 2001, especially in 2001 [25]; and in 2003, despite the presence of Fusarium graminearum, DON was observed in only 18.5% of infected grains.

In a study in the UK, including 247 samples produced organically and 1,377 samples conventionally produced wheat grain, no significant differences in DON content were found in grain from the 2 cultivation systems; an average DON level of 230 μg/kg was recorded for organic

**Table 1. DON content in cereals, flour and cereal products [μg kg⁻¹]**

<table>
<thead>
<tr>
<th>Products</th>
<th>Organic</th>
<th></th>
<th>Conventional</th>
<th></th>
<th>Test Mann-Whitney p</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No./% of contaminated samples</td>
<td>SD</td>
<td>MIN–MAX</td>
<td>No./% of contaminated samples</td>
<td>SD</td>
</tr>
<tr>
<td>Cereals</td>
<td>4 (10); 40%</td>
<td>130–433</td>
<td>285.25 ± 134.04</td>
<td>7 (10); 70%</td>
<td>373.71 ± 171.20</td>
</tr>
<tr>
<td>Flours</td>
<td>5 (10); 50%</td>
<td>104–459</td>
<td>213.80 ± 151.38</td>
<td>7 (10); 70%</td>
<td>336.29 ± 188.90</td>
</tr>
<tr>
<td>Cereal products</td>
<td>5 (19); 26,3%</td>
<td>102–465</td>
<td>199.60 ± 149.82</td>
<td>6 (19); 31,6%</td>
<td>387.67 ± 250.24</td>
</tr>
</tbody>
</table>

**Table 2. ZEA content in cereals, flour and cereal products [μg kg⁻¹]**

<table>
<thead>
<tr>
<th>Products</th>
<th>Organic</th>
<th></th>
<th>Conventional</th>
<th></th>
<th>Test Mann-Whitney p</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No./% of contaminated samples</td>
<td>SD</td>
<td>MIN–MAX</td>
<td>No./% of contaminated samples</td>
<td>SD</td>
</tr>
<tr>
<td>Cereals</td>
<td>0 (10); 0%</td>
<td>0 (10); 0%</td>
<td>0 (10); 0%</td>
<td>0 (10); 0%</td>
<td>0 (10); 0%</td>
</tr>
<tr>
<td>Flours</td>
<td>0 (10); 0%</td>
<td>10±0.0</td>
<td>10±0.0</td>
<td>10±0.0</td>
<td>10±0.0</td>
</tr>
<tr>
<td>Cereal products</td>
<td>1 (19); 5.3%</td>
<td>13±0.0</td>
<td>13–13</td>
<td>3 (19); 15.8%</td>
<td>14.45 ± 2.76</td>
</tr>
</tbody>
</table>
substances and traditionally produced wheat [26]. That there are no differences in DON content in wheat from the 2 agricultural systems were also reported in studies of several other countries: France [27], Germany [28, 29], The Netherlands [30], Switzerland [31], Norway [32] and Spain [9]. The authors suggested that annual variability was more important for DON pollution than agricultural practices. The agricultural system did not affect DON contamination in 2010 in drier and warmer weather, significantly lower DON levels were found (340 μg/kg for organic production and 460 μg/kg for traditionally produced wheat grain) [18]. Lower DON levels were found in organic than in traditionally produced wheat in studies from Italy [33, 34], Belgium [35], Slovakia [36] and Norway [15].

Higher DON concentrations in organic farming samples (average 599 μg kg⁻¹) than in wheat from traditional agriculture (average 331 μg kg⁻¹) were reported by Kirinčič et al. [37]. A study of durum wheat from Italy reported lower levels of DON in organic than in conventionally produced grain [17]. However, in a Spanish study there was no difference in the content of DON in Durum wheat was detected in the 2 cultivation systems [16]. No differences were found in DON contamination in samples of barley grains produced by organic and conventional methods in Norway [15] and the UK [38]. In a study from Germany, there was no difference in the DON content of rye grain from two farming systems [39], while in Poland lower DON levels were found in organically produced rye than in conventional rye [40]. Most levels were well below the EU food limits [41]. Several authors have commented that weather conditions, crop year, location, crop rotation and tillage may be more important for the development of DON than crop type [15, 42, 16, 30, 17, 43, 18].

This study showed that the average DON content was highest in cereal processed samples (average 387.67 μg kg⁻¹), cereal grains (373.71 μg kg⁻¹ and flour (336.29 μg kg⁻¹) from conventional crops, compared to grains (285.25 μg kg⁻¹), flour (213.80 μg kg⁻¹) and cereal products (199.60 μg kg⁻¹) from organic farming. The acceptable limit of DON concentration was exceeded in one sample from conventional crops, but not in any samples from organic products. No statistically significant differences were established in DON content found in cereals or cereal-based products of both types (p<0.05). Stanisławczyk et al. [44] observed that the average content of DON in cereals from conventional crops was the highest in grains (127.95 μg kg⁻¹) and flour (127.36 μg kg⁻¹), compared to samples of groats and oat flakes, and the content of this mycotoxin was on a similar level – about 0.90 μg kg⁻¹. Studies by Mazurkiewicz et al. [45] did not show the permissible DON contents being exceeded in cereal grains. The average concentration of DON in cereal grains cultivated in the conventional manner was close to that obtained by Perkowski and Chełkowski [46] in studies carried out at the beginning of the 1990s. Birzele et al. [47] indicated greater contamination of DON grains from organic farming than conventional crops. Other studies carried out in Germany by the CVUA (Chemisches und Veterinaruntersuchungsamt Stuttgart) showed that the DON content in cereal products from organic farming was lower than in conventional farming products [48].

ZEA was not found in samples of the tested cereals. The current study shows the presence of ZEA in one sample of flour from conventional crops, and a significant difference in its occurrence in cereal products between two types of crops. The average ZEA content in cereal products was lower in organic farming products compared to conventional plant products (p<0.05). The maximum permissible ZEA content specified in the relevant European Regulation 1881/2006/EC was not exceeded in any of the products tested [19].

Stanisławczyk et al. [44] also did not find that contamination levels were exceeded, and the ZEA content in the samples they tested was significantly lower than in other mycotoxins content. The average content of mycotoxin in cereal products from conventional cultivation was 10 μg kg⁻¹. Much more varied results, although not exceeding the maximum levels, were obtained by Ghali et al. [49] who obtained different average ZEA contents in samples of wheat grains – 2.82 μg kg⁻¹ and barley – 15 μg kg⁻¹ from conventional crops. Solarska et al. [50] detected ZEA in 68.2% of cereal products from organic farming. In the studies of Lacko-Bartosova [36] and Vidal et al. [51], no differences were found in the content of ZEA between wheat produced by organic and conventional methods, while in wheat tests conducted by Meister in 2009 [43] showed lower concentrations of ZEA in samples from organic than in samples from conventional agriculture. In rye studies there was also no difference in ZEA concentration in samples from the 2 cultivation systems [40]. The relatively low sample number of studies, high variability, and the fact that ZEA is often below the detection level in a significant proportion of samples, reduces the possibility of statistical disclosure of the significant differences between organic and conventional agricultural practices.

CONCLUSIONS

In general, the mycotoxin contamination observed in organic cereals and cereal products does not differ significantly from what was observed in their conventional counterparts. It is likely that preventive measures used in organic farming are able to maintain mycotoxin contamination of organic cereals and organic cereal by-products at levels similar to those found in conventional cereals and cereal products, despite the fact that the fungicide is not handled. The findings of this study are important in terms of risk to human health, but they neither support nor reject the hypothesis that organic cereals are less contaminated than the conventional ones.

To reduce DON levels in products from organic and conventional farms, regular and increased monitoring supervising the most critical areas where DON contamination takes place, is required. In this respect, the control of pre-and post-harvest, storage and manufacturing processes, consisting of good agricultural practices, good storage practices, good manufacturing practices and hazard analysis critical control points, must abide within the standards of regulatory and monitoring organizations.

REFERENCES