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Determination of multi-mycotoxin occurrence in maize based porridges from selected regions of Tanzania by liquid chromatography tandem mass spectrometry (LC-MS/MS), a longitudinal study

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Abstract

Residents of certain areas of Tanzania are exposed to mycotoxins through the consumption of contaminated maize based foods. In this study, 101 maize based porridge samples were collected from villages of Nyabula, Kikelewa and Kigwa located in different agro-ecological zones of Tanzania. The samples were collected at three time points (time point 1, during maize harvest; time point 2, 6 months after harvest; time point 3, 12 months after harvest) over a 1-year period. Ultra-performance liquid chromatography tandem mass spectrometry (UPLC-MS/MS) was used to detect and quantify 9 mycotoxins: aflatoxin B₁ (AFB₁), aflatoxin B₂ (AFB₂), aflatoxin G₁ (AFG₁), aflatoxin G₂ (AFG₂), fumonisin B₁ (FB₁), fumonisin B₂ (FB₂), deoxynivalenol (DON), ochratoxin A (OTA) and zearaleneone (ZEN) in the samples following a QuEChERS extraction method. Eighty two percent of samples were co-contaminated with more than one group of mycotoxins. Fumonisins (FB₁+FB₂) had the highest percentage occurrence in all 101 samples (100%) whereas OTA had the lowest (5%). For all three villages the mean concentration of FB₁ was lowest in samples taken from time point 2. Conversely, in Kigwa village there was a distinct trend that AFB₁ mean concentration was highest in samples taken.
from time point 2. DON concentration did not differ greatly between time points but the percentage occurrence varied between villages, most notably in Kigwa where 0% of samples tested positive. ZEN occurrence and mean concentration was highest in Kikelelwa. The results suggest that mycotoxin contamination in maize can vary based on season and agro-ecological zones. The high occurrence of multiple mycotoxins found in maize porridge, a common weaning food in Tanzania, presents a potential increase in the risk of exposure and significant health implications in children.

1. Introduction

Mycotoxins are naturally occurring toxic secondary metabolites produced by filamentous fungi which can contaminate many kinds of agricultural products. Toxigenic fungi are capable of growing under a wide range of atmospheric conditions depending on the species and they can contaminate crops during pre-harvest, immediate post-harvest, storage, transport and processing (Bennett & Klich, 2003). Mycotoxins have been shown to contaminate a wide range of agricultural products including: cereals, nuts, fruit, spices and wine (Abia et al., 2013; Serra, Braga, & Venâncio, 2005; Van de Perre et al., 2014; Yogendrarajah, Van Poucke, De Meulenaer, & De Saeger, 2013). In the case of aflatoxins, they have also been detected in milk produced by cows that have consumed contaminated feed (Huang et al., 2013). Due to the ubiquitous presence of mycotoxins in both food and feed supply chains, and their association with various toxicological risks in both humans and animals, they have become a major economic and health concern.

More particularly, aflatoxin B₁ (AFB₁), aflatoxin B₂ (AFB₂), aflatoxin G₁ (AFG₁), aflatoxin G₂ (AFG₂), fumonisin B₁ (FB₁), fumonisin B₂ (FB₂), deoxynivalenol (DON), ochratoxin A (OTA) and zearaleneone (ZEN) have all been recognised global health, agriculture and trade concerns due to the high occurrence and associated health impacts of these mycotoxins that has been found around the world. However, due to socio-economic and environmental factors, developing countries tend to be more severely affected by the threat of mycotoxins, especially aflatoxins, than developed countries (Wild & Gong, 2010; Williams et al., 2004).

The growing recognition of the threat of mycotoxins has stimulated scientific research to better understand how exposure and toxicity impact human health. The toxicity varies among different types of mycotoxins but many have shown the capacity to be acutely toxic, carcinogenic, mutagenic and immunosuppressive (Bakirdere et al., 2012). The most fatal human aflatoxicosis outbreak occurred in
Kenya, 2004, with 317 recorded cases of acute hepatitis and 125 deaths (Nyikal et al., 2004). Public health officials discovered that this was linked to the consumption of AFB₁-contaminated maize and a case control study found that maize in case households had higher concentrations of aflatoxins compared to that of the maize from control households (Azziz-Baumgartner et al., 2005). The International Agency for Research on Cancer (IARC) classifies naturally occurring mixtures of aflatoxins (AFB₁+AFB₂+AFG₁+AFG₂) as Group 1 carcinogens (International Agency for Research on Cancer, 2002). Previous studies have also provided evidence that aflatoxins may cause immune suppression as a result of decreased protein synthesis, changes in enzymatic activity and changes in metabolism or cell cycles (Jiang et al., 2005; Jiang et al., 2008). A study in West African children reported a strong inverse correlation between the exposure of aflatoxin and body height increase (Gong et al., 2004).

Fumonisins, a group of mycotoxins produced by fungal species belonging to the *Fusarium* genus, have also been shown to have detrimental health effects. Fumonisins are widely distributed around the world and have been classified as possible carcinogens (Joint FAO/WHO Expert Committee on Food Additives, 2011). A study carried out in South Africa was able to demonstrate a positive correlation between fumonisin exposure and high incidences of human oesophageal cancer (Marasas, 2001). Fumonisin exposure has also been linked with increased occurrences of neural tube defects (Missmer et al., 2006). DON, another mycotoxin produced by fungal species within the *Fusarium* genus, has not yet been associated with any long-term health impacts in humans but animals with low dose chronic exposure to DON have shown that decreased growth and feed intake (Forsell, Witt, Tai, Jensen, & Pestka, 1986; Rotter, Thompson, Lessard, Trenholm, & Tryphonas, 1994). OTA, produced by *Aspergillus ochraceus*, is a common mycotoxin and a possible human carcinogen. Studies in animals have shown that it has the potential to be carcinogenic, immunosuppressive and neurotoxic (Álavarez, Gil, Ezpeleta, García-Jalón, & López de Cerain, 2004; Lioi, Santoro, Barbieri, Salzano, & Ursini, 2004; Schaaf et al., 2002). ZEN is produced by several fungi in the *Fusarium* genus including: *F. culmorum*, *F. graminearum* and *F. crookwellense*. It has estrogenic effects in pigs (Jiang et al., 2011) and suggested that it can trigger central precocious pubertal in human females (Massart, Meucci, Saggese, & Soldani, 2007).

Due to the toxic effect of mycotoxins in humans and animals it is important to develop analytical methods to detect them in food in order to facilitate their control and regulation. Liquid chromatography tandem mass spectrometry (LC-MS/MS) is an effective method of detection for mycotoxin analysis. In recent...
years, studies have demonstrated LC-MS/MS methods capable of sub ppb detection for multiple
mycotoxins in maize (Frenich, Vidal, Romero-Gonzalez, & Aguillera-Luiz, 2009; Liao et al., 2013;
Malachová, Sulyok, Beltrán, Berthiller, & Krska, 2014; Zachariasova et al., 2014). Analytical methods
for multiple mycotoxins should be selective for their target analytes, sensitive enough to detect toxins
at relatively low concentrations and efficient to ensure rapid and reliable analysis.

Recently a mycotoxin study carried out in Tanzania examined the extent of dietary exposure of AFB₁,
FB₁ and DON through the quantification of their respective biomarkers in serum and urine (Shirima et
al., 2013; Srey, Kimanya, Routledge, Shirima, & Gong, 2014). The study found that young children in
Tanzania are chronically exposed to AFB₁, FB₁ and DON through their diet. Urinary FB₁ was found to
be negatively associated with length for age Z-scores whilst the negative association between AF-Alb
and child growth did not reach statistical significance. In a recent study in Tanzania, maize kernels were
sampled from three districts and multi-mycotoxins were measured by LC-MS method (Kamala et al.,
2015). The study reported high occurrence of AFB₁ (50%) and FB₁ (73%). The food cooking process is
known to have varying impact on mycotoxin levels, therefore measuring the levels of mycotoxins in
cooked food can provide more close estimates of exposure than in maize flour.

This paper utilises a recently developed multi-mycotoxin detection method to determine the extent of
multi-mycotoxin contamination in the maize porridge, in order to build upon mycotoxin occurrence and
exposure data from previous studies; and to compare with the exposure biomarker data where possible.

2. Materials and Methods

2.1 Reagents and chemicals

Acetonitrile (LC-MS grade), ammonium hydroxide (≥25% in water), dimethyl sulfoxide (≥99.9%), formic
acid (≥98%), magnesium sulfate (anhydrous, ≥99.5%), methanol (LC-MS grade), mycotoxin standards
(AFBI, AFB₂, AFG₁, AFG₂, FB₁, FB₂, DON, ZEN and OTA), sodium chloride (≥99.0%) and Whatman®
Puradisc 4 syringe filters (0.2 µm, PTFE) were all acquired from Sigma-Aldrich (Poole, United Kingdom).
Each mycotoxin standard was separately dissolved in acetonitrile (0.2 mg/ml solution) and stored at -
20°C. Bondesil C₁₈ was acquired from Agilent Technologies (Waldbronn, Germany).

2.2 Study design and sampling
Cooked maize porridge samples were collected from households across three rural villages in Tanzania: Nyabula (Iringa region), Kikelelwa (Kilimanjaro region) and Kigwa (Tabora region), which are from different agro-ecological zones. The samples were collected at three time points over the period of a year: Time point 1 (June/July 2010, a maize harvesting season), time point 2 (January 2011, six months after maize was harvested) and time point 3 (June/July 2011, another maize harvesting season 12 months after time point 1). The cooked porridge samples were dried after collection. A total of 101 samples; 10 samples from each village at time points 1 and 2, 14 samples from Nyabula and Kikelelwa villages at time point 3 and 13 samples from Kigwa village at time point 3 were randomly selected. The samples were oven dried and kept frozen at -80°C until extraction for UPLC-MS/MS analysis. A blank maize flour sample was cooked into a porridge using the same recipe as the other Tanzanian porridge samples, oven dried and subsequently stored at -80°C until LC-MS/MS extraction for UPLC-MS/MS analysis.

2.3 Extraction procedure and UPLC-MS/MS analysis

A previously developed multi-mycotoxin UPLC-MS/MS method was adopted for the study. Nine mycotoxins of interest were quantified: AFB₁, AFB₂, AFG₁, AFG₂, FB₁, FB₂, DON, ZEN and OTA. Briefly, the LC-MS/MS method was developed on a Waters Acquity UPLC coupled to a Xevo TQ-S triple quadrupole mass spectrometer. Sample extraction method was based on QuEChERS method (Lacina et al., 2012). The quantification was achieved by interpolation from a standard curve prepared by spiking the blank matrix samples at 7 different levels with a mixture of mycotoxins before extraction. Calibrant solutions for matrix-matched calibration curves were prepared in blank matrix before extraction. Limits of detection (LOD) and quantification (LOQ) (S/N≥10) were previously determined in maize as: AFB₁ LOD: 0.05 ng/g, LOQ: 0.125 ng/g; AFB₂/AFG₁ LOD: 0.125 ng/g, LOQ: 0.25 ng/g; AFG₂ LOD: 0.25 ng/g, LOQ: 0.5 ng/g; DON LOD: 5 ng/g, LOQ: 12.5 ng/g; OTA LOD: 0.625 ng/g, LOQ: 1.25 ng/g; ZEN LOD: 2.5 ng/g, LOQ: 5 ng/g. Limits of detection and quantification were determined in wheat as 0.5 ng/g and 1.0 ng/g respectively for FB₁ and 0.2ng/g and 0.5ng/g respectively for FB₂.

2.4 Statistical analysis

Data was analysed using IBM SPSS Statistics for Windows, Software Version 21.0 (IBM Corp., 2012). Kruskal-Wallis tests were carried out to test for variance between villages within time points and
variance between time points within villages where applicable. A p value of <0.05 was considered statistically significant for this test and only positive samples >LOD were used. Spearman’s rank tests were carried out to test for correlation between the concentration of AFB$_1$, FB$_1$ and DON. A p value of <0.05 was considered statistically significant for this test.

3. Results

Descriptive statistics for all 101 maize porridge samples are displayed in Table 1a and 1b. Both FB$_1$ and FB$_2$ were detected in all 101 samples. The mean ratio of FB$_1$:FB$_2$ for Nyabula, Kikelelwa and Kigwa was 60:40, 60:40 and 64:36, respectively, with a mean ratio of 61:39 for all samples. Median FB$_1$ concentration was significantly higher in Kikelelwa (290.18 ng/g) and Kigwa (383.54 ng/g) than in Nyabula 60.14 ng/g) (p = 0.000). FB$_1$ concentration was lowest during time point 2 (6 months after harvest) in every village. Eleven percent of the 101 samples analysed showed fumonisin contamination levels greater than the maximum tolerable limit for total fumonisin in maize based foods intended for adult human consumption (FB$_1$+FB$_2$ ≥1000 ng/g) set by the European Commission (EC) (European Commission, 2006). Fifty-seven percent of samples also exceeded the EC limit for fumonisins in food products intended for infant consumption (FB$_1$+FB$_2$ ≥200 ng/g). At least one type of aflatoxin was detected in 50% of all samples. The mean ratios of AFB$_1$:AFB$_2$ and AFG$_1$:AFG$_2$ for all samples were both 90:10. The data from Kigwa village was chosen for statistical analysis due to the high (94%) occurrence of aflatoxins in comparison to Kikelelwa (27%) and Nyabula (24%).
### Table 1a

<table>
<thead>
<tr>
<th>Mycotoxin</th>
<th>Village/Region</th>
<th>Time point</th>
<th>Positive Samples (%)</th>
<th>Mean (ng/g)</th>
<th>Median (ng/g)</th>
<th>Range (ng/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nyabula/Iringa</td>
<td>1</td>
<td>10 (100)</td>
<td>158.06</td>
<td>109.64</td>
<td>41.70 - 375.64</td>
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<tr>
<td></td>
<td></td>
<td>2</td>
<td>10 (100)</td>
<td>44.96</td>
<td>46.54</td>
<td>13.01 - 65.53</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>14 (100)</td>
<td>87.18</td>
<td>60.14</td>
<td>12.87 - 421.36</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>34 (100)</td>
<td></td>
<td>95.61</td>
<td>60.14</td>
<td>12.87 - 421.36</td>
</tr>
<tr>
<td><strong>Fumonisin B₁</strong></td>
<td></td>
<td>1</td>
<td>10 (100)</td>
<td>555.74</td>
<td>393.41</td>
<td>218.19 - 1308.43</td>
</tr>
<tr>
<td></td>
<td>Kikelelwa/Kilimanjaro</td>
<td>2</td>
<td>10 (100)</td>
<td>159.38</td>
<td>82.10</td>
<td>12.65 - 564.10</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>14 (100)</td>
<td>438.14</td>
<td>311.63</td>
<td>36.02 - 1850.01</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>34 (100)</td>
<td></td>
<td>390.74</td>
<td>290.18</td>
<td>12.65 - 1850.01</td>
</tr>
<tr>
<td></td>
<td>Kigwa/Tabora</td>
<td>1</td>
<td>10 (100)</td>
<td>533.98</td>
<td>489.07</td>
<td>71.46 - 1206.09</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>10 (100)</td>
<td>299.26</td>
<td>309.83</td>
<td>12.65 - 554.57</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>13 (100)</td>
<td>394.84</td>
<td>328.50</td>
<td>57.08 - 1141.40</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>33 (100)</td>
<td></td>
<td>408.04</td>
<td>383.54</td>
<td>12.65 - 1206.09</td>
</tr>
<tr>
<td><strong>Fumonisin B₂</strong></td>
<td></td>
<td>1</td>
<td>10 (100)</td>
<td>110.42</td>
<td>80.56</td>
<td>27.67 - 261.89</td>
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<tr>
<td></td>
<td>Kikelelwa/Kilimanjaro</td>
<td>2</td>
<td>10 (100)</td>
<td>28.21</td>
<td>28.37</td>
<td>9.50 - 38.06</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>14 (100)</td>
<td>59.35</td>
<td>41.84</td>
<td>7.25 - 282.23</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>34 (100)</td>
<td></td>
<td>65.21</td>
<td>37.41</td>
<td>7.25 - 282.23</td>
</tr>
<tr>
<td></td>
<td>Nyabula/Iringa</td>
<td>1</td>
<td>10 (100)</td>
<td>298.68</td>
<td>255.38</td>
<td>23.57 - 790.05</td>
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<td></td>
<td></td>
<td>2</td>
<td>10 (100)</td>
<td>177.14</td>
<td>183.44</td>
<td>33.98 - 313.68</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>13 (100)</td>
<td>240.68</td>
<td>203.59</td>
<td>29.71 - 699.85</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>33 (100)</td>
<td></td>
<td>239.00</td>
<td>213.09</td>
<td>23.57 - 790.05</td>
</tr>
<tr>
<td><strong>Aflatoxin B₁</strong></td>
<td></td>
<td>1</td>
<td>4 (40)</td>
<td>7.16</td>
<td>0.45</td>
<td>0.15 - 27.60</td>
</tr>
<tr>
<td></td>
<td>Kikelelwa/Kilimanjaro</td>
<td>2</td>
<td>2 (20)</td>
<td>7.43</td>
<td>7.43</td>
<td>7.15 - 7.70</td>
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<td></td>
<td></td>
<td>3</td>
<td>4 (29)</td>
<td>0.34</td>
<td>0.33</td>
<td>0.20 - 0.50</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>8 (24)</td>
<td></td>
<td>4.49</td>
<td>0.43</td>
<td>0.15 - 27.60</td>
</tr>
<tr>
<td></td>
<td>Kigwa/Tabora</td>
<td>1</td>
<td>1 (10)</td>
<td>34.50</td>
<td>34.50</td>
<td>34.50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>0 (0)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>8 (57)</td>
<td>2.19</td>
<td>0.48</td>
<td>0.20 - 13.05</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>9 (27)</td>
<td></td>
<td>5.78</td>
<td>0.65</td>
<td>0.20 - 34.50</td>
</tr>
<tr>
<td></td>
<td>Kigwa/Tabora</td>
<td>1</td>
<td>10 (100)</td>
<td>4.05</td>
<td>1.15</td>
<td>0.40 - 13.55</td>
</tr>
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<td></td>
<td></td>
<td>2</td>
<td>10 (100)</td>
<td>10.21</td>
<td>5.95</td>
<td>0.55 - 25.80</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>11 (85)</td>
<td>0.67</td>
<td>0.68</td>
<td>0.20 - 1.55</td>
</tr>
<tr>
<td></td>
<td>All</td>
<td>31 (94)</td>
<td></td>
<td>4.73</td>
<td>0.95</td>
<td>0.20 - 25.80</td>
</tr>
</tbody>
</table>
## Table 1b

Levels of mycotoxin contamination in different villages and time points in Tanzania

<table>
<thead>
<tr>
<th>Mycotoxin</th>
<th>Village/Region</th>
<th>Time point</th>
<th>Positive Samples (%)</th>
<th>Mean (ng/g)</th>
<th>Median (ng/g)</th>
<th>Range (ng/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Aflatoxins (B&lt;sub&gt;1&lt;/sub&gt;+B&lt;sub&gt;2&lt;/sub&gt;+G&lt;sub&gt;1&lt;/sub&gt;+G&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>Nyabula/Iringa</td>
<td>1</td>
<td>4 (40)</td>
<td>8.26</td>
<td>0.80</td>
<td>0.15 - 31.30</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>2 (20)</td>
<td>14.48</td>
<td>14.48</td>
<td>14.05,14.90</td>
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<td></td>
<td></td>
<td>3</td>
<td>4 (29)</td>
<td>0.69</td>
<td>0.60</td>
<td>0.20 - 1.35</td>
</tr>
<tr>
<td></td>
<td></td>
<td>All</td>
<td>8 (24)</td>
<td>6.48</td>
<td>0.93</td>
<td>0.15 - 31.3</td>
</tr>
<tr>
<td></td>
<td>Kikelelwa/Kilimanjaro</td>
<td>1</td>
<td>1 (10)</td>
<td>39.70</td>
<td>39.70</td>
<td>39.70</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>0 (0)</td>
<td>-</td>
<td>-</td>
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<tr>
<td></td>
<td></td>
<td>3</td>
<td>8 (57)</td>
<td>3.23</td>
<td>1.10</td>
<td>0.25 - 14.9</td>
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<td></td>
<td></td>
<td>All</td>
<td>9 (27)</td>
<td>7.39</td>
<td>1.10</td>
<td>0.25 - 39.7</td>
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<tr>
<td>Deoxynivalenol</td>
<td>Nyabula/Iringa</td>
<td>1</td>
<td>1 (100)</td>
<td>6.27</td>
<td>1.65</td>
<td>0.40 - 22.7</td>
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<td></td>
<td></td>
<td>2</td>
<td>10 (100)</td>
<td>14.91</td>
<td>9.58</td>
<td>0.55 - 43.65</td>
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<td>11 (85)</td>
<td>1.03</td>
<td>1.10</td>
<td>0.20 - 2.40</td>
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<td></td>
<td></td>
<td>All</td>
<td>31 (94)</td>
<td>7.03</td>
<td>1.60</td>
<td>0.20 - 43.65</td>
</tr>
<tr>
<td>Zearalenone</td>
<td>Nyabula/Iringa</td>
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<td>0 (0)</td>
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<td>4 or more mycotoxins (%)</td>
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*Mycotoxins were organised into 5 groups for this co-occurrence analysis: 1. Total aflatoxins: AFB$_{1}$+AFB$_{2}$+AFG$_{1}$+AFG$_{2}$; 2. Total fumonisins: FB$_{1}$+FB$_{2}$; 3. DON; 4. ZEN; 5. OTA.

The highest concentration of AFB$_{1}$ in a single sample was found in Kikelelwa village at 34.50 ng/g but Kigwa and Nyabula village had more samples with aflatoxins detectable, and also had higher concentrations of aflatoxins; both villages had higher concentrations aflatoxins at time point 2 (during storage) than any other. The Tanzania regulatory limits for AFB$_{1}$ concentration (AFB$_{1}$ ≥5 ng/g) and total aflatoxins concentrations (total aflatoxins ≥10 ng/g) in maize, were exceeded in 14% and 12% of samples, respectively. DON was detected in 44% of all samples. Nyabula samples showed a successive increase in DON concentration between time points whereas Kikelelwa samples showed little difference. DON was not detected in any samples from Kigwa village. None of the analysed samples showed concentrations exceeded the EC limit for DON in maize-based foods intended for human consumption (DON ≥750 ng/g); however, 6% of samples were above the limit for infant food (DON ≥200 ng/g). ZEN was detected in 31% of all samples. ZEN concentration was highest in all three time points for Kikelelwa (mean: 62.14 ng/g; occurrence: 79%) however there was no significant difference between any of them ($p = 0.335$). Samples from Nyabula and Kigwa had a noticeably lower occurrence of ZEN (9% and 3% respectively). Only one sample exceeded the EC limit for ZEN in maize-based foods intended for human consumption (ZEN ≥200 ng/g). However, 23% of samples exceeded the limit for infant food (ZEN ≥20 ng/g). OTA was detected in 5% of all samples (mean: 3.80 ng/g, 3.80 ng/g, 3.80 ng/g).
median: 3.30 ng/g). Each sample that had detectable OTA exceeded the EC limit in maize products intended for both human adult and infant consumption (OTA ≥3.0 ng/g and >0.5 ng/g, respectively).

Overall, total aflatoxin concentration and total fumonisins were found to have a positive correlation by Spearman’s rank correlation (correlation coefficient: 0.254, \( p = 0.011 \)), while total aflatoxin was found to be negatively correlated to DON (correlation coefficient: -0.407, \( p = 0.000 \)). Total fumonisin and DON showed no statistically significant correlation (\( p = 0.919 \)). ZEN showed no statistically significant correlation with any of the other mycotoxins (\( p > 0.05 \)). Further data on co-occurrence can be seen in Table 2.

4. DISCUSSION

The LC-MS/MS analysis showed that the maize porridge samples collected from the three Tanzanian villages were subject to contamination of multiple mycotoxins. All three Tanzanian villages showed high occurrences of fumonisins (detected in 100% of samples) for all 101 samples. In all villages a lower fumonisin concentration was seen in samples taken at time point 2, 6 months after harvest, compared to samples taken at time point 1, during harvest. The reason for this difference is not immediately apparent. It is possible that as household food supplies from subsistence farming begin to dwindle during the dry season, and residents may be buying maize from other less contaminated areas. It has been suggested that certain strains of bacteria can be effective in reducing fumonisin levels in maize and that lactic acid, commonly produced by anaerobic bacteria, has a protective role against the growth of Fusarium species in stored maize (Benedetti, Nazzi, Locci, & Firrao, 2006). Potentially, the fungal species in the maize has been gradually degraded during storage by a bacterial agent. When all three time points were taken into account, Nyabula had the lowest FB\(_1\) mean concentration, followed by Kikelelwa. It is possible that the climate or seasonal weather variation in Nyabula may have had an influence in preventing the Fusarium growth in the maize. In terms of maize total aflatoxin concentration, Kigwa had the most positive samples for total aflatoxins, followed by Kikelelwa and Nyabula. Unlike fumonisins, maize aflatoxin concentration was highest in time point 2 compared to time points 1 and 3. This difference in concentrations between the two mycotoxins would suggest that the storage conditions are favoured by fungi that produce aflatoxins and field or harvest conditions at harvest are favoured by fungi that produce fumonisins. This also suggests that the overall climate and weather conditions of Tanzania are favourable for both aflatoxins and fumonisins.
It was found that Kikelelwa samples showed the highest frequency of DON in maize followed by Nyabula whilst DON was not detected in any samples from Kigwa. Kikelelwa has a temperature climate which is known to favour growth of DON producing fungi. Unlike aflatoxins and fumonisins, DON concentration between time points was not found to be statistically different. ZEN concentration showed a similar contamination pattern to DON; it was highest in Kikelelwa, followed by Nyabula and Kigwa, where it was only detected in <1% of samples. ZEN occurrence was highest in Kikelelwa village and lowest in Kigwa. The difference in ZEN concentration between time points was not found to be significant in any villages. OTA was the least prevalent of nine mycotoxins analysed. The low frequency meant that it was not possible to ascertain any discernable pattern in the contamination either between regions or time points.

There were several cases of multiple mycotoxins co-contamination in the analysed maize porridge samples. Mycotoxins were collated into five groups for co-contamination analysis: 1. Total aflatoxins: AFB₁ + AFB₂ + AFG₁ + AFG₂; 2. Total fumonisins: FB₁ + FB₂; 3. DON; 4. ZEN and 5. OTA. It was found that 82% of all samples contained two or more groups of mycotoxins, 36% contained three or more and 10% contained four or more. Samples taken from Kikelelwa contained the greatest proportion of co-contaminated samples with 79% containing three or more groups of mycotoxins. This is much greater in comparison to Nyabula and Kigwa where 21% and 6% of samples were co-contaminated with three or more groups of mycotoxins respectively. Certain mycotoxins also showed patterns of co-contamination in terms of their ratios to one another. The FB₁:FB₂ ratio of 60:40 stayed largely consistent for samples across all regions and time points. AFB₁:AFB₂ and AFG₁:AFG₂ both showed 90:10 ratios. Total aflatoxins and fumonisins showed statistically significant correlations with each other. Aflatoxins and fumonisins were found to have a positive correlation \( r = 0.254, p = 0.011 \) despite showing different general contamination patterns between time points. This suggests that, even though aflatoxin and fumonisins contamination may differ between seasons, the climate of Tanzania still promotes the growth of certain Aspergillus and Fusarium fungal species. Aflatoxins and DON were found to have a moderate negative correlation. This is most evident in Kigwa village which showed the highest aflatoxin contamination frequency but the lowest for DON. These two correlations highlight the need to further investigate and understand the nature of mycotoxin co-contamination.
Many of the samples exceeded EC regulatory limits for maize in both adult and infant food. Tanzania has official regulations for AFB₁ and total aflatoxins in maize but not for the other previously mentioned mycotoxins. Regulatory limits taken from European Commission Regulation No. 1881/2006, as amended, were used as a guideline for the other mycotoxins involved this study. Total aflatoxins and fumonisins were found to exceed the limits in 12% and 11% of samples, respectively. DON and ZEN were the two mycotoxins with the lowest number of samples over the limit at 0% and 1%, respectively. OTA was unique in that it was detected in only five samples but each of those exceeded the regulatory limit. The maize porridge was also compared against EC regulatory limits for food intended for infant consumption which are much lower than their adult counterparts. 50%, 57%, 6% and 23% of samples were over the infant limit for total aflatoxins, total fumonisins, DON and ZEN, respectively based on EC limits. These results show that the mycotoxins in these maize porridge samples are not just a health risk to infants and young children but also a serious risk to adults. Combine this with the fact that many of these samples are co-contaminated with multiple toxins and there is a very real possibility that a large number of these children are consuming food over the recommended limits of several different mycotoxins.

One of the applications of multi-mycotoxin food analysis such as this is to utilise the data in ongoing biomarker discovery and validation. The data in this study has been compared to biomarker data for samples taken from these villages at the same time points, to determine whether FB₁, AFB₁ and DON contamination in food are consistent with urinary FB₁ (uFB₁), aflatoxin albumin adduct (AF-Alb) and urinary DON (uDON) (Shirima et al., 2013; Srey et al., 2014). Generally speaking, uDON concentration in each Tanzanian village reflected the DON concentrations measured in maize food. The Kigwa maize samples showed no trace of DON and this was reflected in the uDON biomarker levels being the lowest out of the three villages. Table 1b shows the Kikelelewa median DON concentration was similar between time points 1 and 2 with a much lower level at time point 3, which is mirrored in the estimated DON intake from the biomarker study. The Nyabula median DON showed a constant rise from time point 1 through to 3 which is the same pattern as reported in the biomarker study (Srey et al. 2014).

The comparison of AFB₁ in maize porridge and AF-Alb biomarker in blood was not entirely consistent. The increase in the AFB₁ median concentration from time point 1 to 2 in Kigwa village was in a good agreement with a similar increase in AF-Alb from the same time points. However the maize and AF-
Alb data for time point 3 do not agree with each other – a decrease was observed in maize AFB$_1$
whereas AF-Alb stayed the same as time point 2 (Shirima et al., 2013). A possible explanation for this
is the increased consumption due to child growth and maize availability at harvest. Also the children
may have still been consuming the same AFB$_1$ contaminated maize from time point 2 which may still
affect the AF-Alb biomarker measurements.

The FB$_1$ found in this study was also compared against that of uFB$_1$ biomarker obtained from the same
village and time points. Nyabula village showed the lowest FB$_1$ contamination in maize but it was
Kikelelwa which showed the lowest FB$_1$ detected in urine. The lower exposure in Kikelelwa is
presumably due to lower FB$_1$ intake of children due to lower maize consumption but high other food
types than in the other two villages. It was observed that time point 2 consistently had the lowest FB$_1$
contamination in maize and this is in consistent with uFB$_1$ levels.

This study was compared against mycotoxin occurrence data in maize from Cameroon (Abia et al.,
2013), Nigeria (Ezekiel et al., 2014), Malawi (Matumba, Sulyok, Monjerezi, Biswick, & Krska, 2014), two
South African studies (Shephard et al., 2013; van der Westhuizen et al., 2010) and another Tanzanian
study with uncooked maize samples taken from the same Kikelelwa village at the same season
(Kimanya et al., 2014). All data sets showed high frequencies of FB$_1$ in maize for their respective areas.
The FB$_1$ median for Kikelelwa village determined from this study (290.18 ng/g) was similar to the FB$_1$
median from the Tanzanian study carried out in the same village (329 ng/g) (Kimanya et al., 2014). Both
Cameroon and Nigerian maize FB$_1$ levels were in a similar range as the levels in Kikelelwa and Kigwa
village. The two studies from South Africa and the one from Malawi all showed much greater FB$_1$ levels
compared to Tanzania. This comparison shows that FB$_1$ is heavily prevalent throughout Africa but the
levels of contamination not only vary between different countries but also within different areas of the
same country. The distribution of AFB$_1$ was mixed with the majority of areas showing detected
frequencies of ≤50%. The two studies featuring maize analysis from Kikelelwa village both showed
good agreement with each other. They both gave similar frequencies (29% and 24%, respectively) and
medians (0.65 ng/g and 1.27 ng/g, respectively). In comparison to Kigwa village (mean: 0.68 ng/g),
Cameroon and Nigeria showed relatively higher means (3.5 ng/g and 2.5 ng/g, respectively). The
distribution of DON between countries varied largely amongst countries especially if high DON
concentrations in Malawi Highlands (mean: 600 ng/g; max: 2328 ng/g) and compared against the lower
ones in Kikelelwa village, Tanzania (mean: 111.17 ng/g max: 410.90 ng/g). The frequency of ZEN was
relatively high for Kikelelwa of Tanzania when compared against Cameroon and South Africa. Nyabula and Kigwa of Tanzania showed lower ZEN concentrations comparable to Malawi and Nigeria. The limitation of comparing data between different countries is that mycotoxin contamination levels can vary between seasons and years, so conclusions about relative frequencies of contamination between different regions or countries cannot be drawn without multiple sampling over a period of time.

5. CONCLUSIONS

The UPLC-MS/MS method was successfully used to detect and quantify the nine mycotoxins of interest from the Tanzanian maize porridge samples. All three Tanzanian villages had a considerable mycotoxin contamination as 82% of samples contained two or more groups of mycotoxins. Fumonisins were by far the most prevalent mycotoxins as they were detected in all samples. FB$_1$ and FB$_2$ contamination was found to have a seasonal pattern, being lower following a period of storage. AFB$_1$ was detected primarily in Kigwa, with higher concentrations following storage. DON showed large regional variance, with complete absence from Kigwa village which matched up well the biomarker data in the same village. The data obtained for AFB$_1$, FB$_1$ and DON was comparable with exposure findings based on their corresponding biomarkers. Finally, the co-contamination data showed that both Tanzania adults and children from these villages are at risk from multiple mycotoxins in maize. It is not known to what extent co-contamination with multiple mycotoxins may contribute to health effects. This study highlights the need to understand the extent of mycotoxin co-contamination so that proper contamination control can be implemented in these vulnerable areas.

The authors declare that they have no actual or potential conflicts of interest.

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References


