In vivo formation of \( N \)-acyl-fumonisin \( B_1 \)

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Abstract Fumonisins are fungal toxins found in corn and in corn-based foods. Fumonisin \( B_1 \) (FB\( _1 \)) is the most common and is toxic to animals, causes cancer in rodents, and is a suspected risk factor for cancer and birth defects in humans. The hydrolyzed form of FB\( _1 \) (HFB\( _1 \)) also occurs in foods and is metabolized by rats to compounds collectively known as \( N \)-acyl-HFB\( _1 \) (also known as \( N \)-acyl-AP\( _1 \)). \( N \)-acyl-HFB\( _1 \) is structurally similar to ceramides, metabolites which have important structural and signaling functions in cells. FB\( _1 \) is \( N \)-acylated in vitro to ceramide-like metabolites which, like FB\( _1 \), are cytotoxic. However, metabolism of FB\( _1 \) and inhibition of ceramide synthase by its metabolites in vivo has not been demonstrated. Male rats were dosed ip with 0.5, 1, or 2 mg/kg body weight FB\( _1 \) on five consecutive days and the liver and kidney thereafter processed for chemical analysis. \( N \)-acyl derivatives of fumonisin \( B_1 \) were identified for the first time in these principal target organs of FB\( _1 \) in rats, at levels up to 0.4 nmol/g tissue using mass spectrometry. The \( N \)-acyl chain length of the metabolites varied in a tissue-dependent manner with \( C_{16} \) derivatives predominating in the kidney and \( C_{24} \) derivatives being prevalent in the liver. The toxicological significance of \( N \)-acyl-fumonisins is not known and warrants investigation.

Keywords Fumonisin · Metabolism · Mycotoxin · Biodistribution · \( N \)-acyl-fumonisins

Abbreviations

| Abbreviation | Definition |
|--------------|------------|
| FB\( _1 \) | Fumonisin \( B_1 \) |
| CerS | Ceramide synthase |
| ESI-MS/MS | Electrospray ionization-tandem mass spectrometry |
| HFB\( _1 \) | Hydrolyzed fumonisin \( B_1 \) |
| NAFB\( _1 \) | \( N \)-acyl-FB\( _1 \) |
| NAHFB\( _1 \) | \( N \)-acyl-HFB\( _1 \) |
| SL | Sphingolipid |
| TCA | Propane-1,2,3-tricarboxylic acid |

Introduction

Fumonisins are secondary metabolites of fungi, mainly \( Fusarium verticillioides \) and \( Fusarium proliferatum \). They were described for the first time by Gelderblom et al. (1988) who isolated and characterized fumonisin \( B_1 \) (FB\( _1 \)) by means of a biosassay based on the promotion of carcinogenesis in rat liver. Further studies demonstrated that FB\( _1 \), the most prevalent congener, is hepato- and nephrotoxic to rodents (Voss et al. 1989; Voss et al. 2001). In addition, several species-specific syndromes are caused by FB\( _1 \). These include neural tube defects in the LM/Bc mouse (Voss et al. 2009; Gelineau-van Waes et al. 2009; Gelineau-van Waes et al. 2005; Gelineau-van Waes et al. 2012), equine leukoencephalomalacia (Marasas et al. 1988), and pulmonary edema in pigs (Harrison et al. 1990). Fumonisins are structurally similar to the sphingoid bases...
sphingosine, and Wang et al. (1991) found that FB1 inhibits ceramide synthase (CerS), a critical enzyme in the de novo biosynthesis of ceramide and complex sphingolipids, and disrupts sphingolipid metabolism. These and subsequent studies have established disrupted sphingolipid metabolism as the mode of action of fumonisins (Bulder et al. 2012).

In a previous study, Humphf et al. (1998) found that hydrolyzed FB1 (HFB1), an alkaline hydrolysis product of FB1 that is present in some foods, is a substrate for CerS in rat liver microsomes (Humphf et al. 1998). Specifically, HFB1 substitutes for sphinganine or sphingosine so that CerS catalyzes the acylation of HFB1 at the primary amino group with fatty acids of various chain lengths to form ceramide analogs known as N-acyl-HFB1 (NAHFB1). HFB1 was also metabolized in vivo: NAHFB1 of various fatty acyl chain lengths were found in the liver and kidney of rats exposed to HFB1 (Seiferlein et al. 2007). Harrer et al. (2013), using an optimized mass spectrometry method, more recently demonstrated the in vitro formation of N-acyl-FB1 (NAFB1) in human cell lines which had been transfected for the overexpression of CerS. We now report the in vivo formation of NAFB1 in rats exposed to FB1.

Materials and methods

Animals and experimental design

The study protocol was approved by the Institutional Animal Care and Use Committee, Richard B. Russell Agricultural Research Center, Athens, GA. Male Sprague–Dawley rats (Harlan Industries, Indianapolis, IN, USA), 7 weeks of age at receipt, were individually housed in stainless steel, wire mesh cages in an environmentally controlled room having a 12-h light/dark cycle. Food (2019 Global Rodent Diet, Teklad, Madison, WI, USA) and fresh tap water were provided ad libitum. After a 1 week acclimation period, the animals were randomly assigned to five groups (n=2/group) having mean weights of 215 to 216 g (weight range of all animals, 211 to 220 g). For five consecutive days, they were observed, weighed, and given an intraperitoneal injection (ip) of 0 (vehicle), 0.5, 1.0, or 2.0 mg/kg body weight FB1 (provided by R. Eppley, US FDA), 1.0 mg/kg body weight HFB1 (positive control) (Voss et al. 2009), or vehicle (0.9 % physiological saline). These doses corresponded to 0.69, 1.38, and 2.77 μmol FB1/kg body weight and 2.47 μmol/kg body weight HFB1 per day. In order to compare the effects of FB1 and HFB1, dosages will be expressed as molar concentrations and, on this basis, the high dose of FB1 is roughly equivalent to that of the positive control dose of HFB1. Dosing solutions were sterile filtered and administered at a volume rate of 10 ml/kg body weight. The animals were fasted overnight prior to administration of the final dose. The rats were euthanized (CO2 inhalation and exsanguination) 60 to 90 min following the final dose administration and examined by necropsy. The kidney and liver specimens were fixed in 10 % neutral buffered formalin, processed, and microscopically evaluated without knowledge of the animal’s identity or treatment group. Three representative liver and kidney specimens (100 mg each) were also collected, immediately frozen, and stored (~80 °C) until processed, at which time the tissues were thawed and homogenized in distilled/deionized water (1 ml water/100 mg tissue). The homogenates were lyophilized before they were shipped by overnight courier for the analysis of fumonisin metabolites and sphingolipids.

Chemicals and reagents

Reverse phase columns were from Phenomenex (Aschaffenburg, Germany) and Varian (Darmstadt, Germany). All chemicals and solvents were analytical grade and purchased from Sigma-Aldrich (Steinheim, Germany). Sphingolipid standards are from Avanti Polar Lipids (Alabaster, USA). FB1, HFB1, NAFB1, NAHFB1, and the isotope-labeled standard FB1-d6 were isolated and synthesized with methods described earlier (Lukacs et al. 1996; Hübner et al. 2012; Harrer et al. 2013).

Quantification of NAFB1, NAHFB1, and sphingolipids

Lyophilized liver and kidney tissues (100 mg wet weight equivalents) were processed for the quantification of FB1, HFB1, NAFB1, NAHFB1, and sphingolipids. Preparation of the tissue samples was performed according to a previously published method (Harrer et al. 2013). For the quantification of FB1 and HFB1, a d6-labeled derivative of FB1 (FB1-d6) was used as the internal standard. Synthetic derivatives with a heptadecanoic fatty acid (C17:0) residue were used as internal standards for the quantification of N-acyl-fumonisins and ceramides (N-C17:0-FB1, N-C17:0-HFB1, and N-C17:0-ceramide). The sphingoid bases and their phosphate derivatives were quantified using synthetic derivatives with a C17-acyl-backbone: sphingosine (C17:1), sphingosine-1-phosphate (C17:1), sphinganine (C17:0), sphinganine-1-phosphate (C17:0) as internal standards. The standards were dissolved in methanol/chloroform (2:1, v/v) and then mixed with the tissue samples. The samples spiked with internal standard were kept at room temperature for 30 min before being processed further. The internal standard-spiked samples were first extracted with ethyl acetate, isopropanol, and water (60:30:10, v/v/v) and a second time using methanol, chloroform, and water (60:30:10, v/v/v). The extracts were combined, the solvents removed by evaporation, and the dried samples stored at ~80 °C till analyzed.
For the analysis, the samples were reconstituted in a mixture of water, methanol, and tetrahydrofuran (60:24:16, v/v/v). The NAFB1 and NAHFB1 derivatives having the following fatty acyl chain lengths were quantified by HPLC-MS/ MS: C16:0, C18:0, C20:0, C22:0, C24:0, C26:0. Furthermore, we also quantified the unsaturated derivatives C24:1 and C26:1. The results are given as the sum of all N-acyl derivatives of FB1 or HFB1. Quantitative results for each individual N-acyl derivative are given in Table S1 of the Supplementary Material.

Results and discussion

Histological examination of rat liver and kidney

Mild to moderate apoptotic, mitotic, and other effects consistent with fumonisin exposure (Bulder et al. 2012; Voss et al. 2001) were found in all rats given FB1. Dose-related differences in severity were not obvious, likely as a consequence of the small sample size of n=2/group and dosing regimen (levels and exposure route). In agreement with reports that HFB1 is less toxic than FB1 in rodents (Gelderblom et al. 1993; Collins et al. 2006; Howard et al. 2002; Voss et al. 2009), the microscopic appearance of the kidneys and liver from rats given 1.0 mg/kg body weight HFB1 per day could not be differentiated from that of tissues from the vehicle control group.

Tissue FB1 and HFB1 concentrations

In order to evaluate the metabolism of fumonisins, the concentrations of FB1, HFB1, and their metabolites were determined in the kidney and liver. No fumonisins or metabolites were detected in the animals given the vehicle (see Fig. 1). In the FB1-exposed groups, a dose-dependent increase in FB1 concentrations was found in the kidneys that ranged from 4 to up to 10 nmol/g tissue. The average concentrations of FB1 in the liver were 12- to 20-fold lower (see Fig. 1). This pattern of preferential accumulation of FB1 in the kidney is consistent with previously published data (Riley and Voss 2006) showing that the FB1 concentration (ng/g wet weight) in the kidney of rats fed with fumonisin-contaminated diet was approximately tenfold higher than that in the liver. Our findings are also consistent with those of a single-dose toxicokinetic study of FB1 (Martinez-Larranaga et al. 1999) in which area under the concentration–time curve (AUC) tissue/AUC plasma ratios of 29.9 and 2.03 were determined for the kidney and liver, respectively.

A different pattern was found for HFB1. Its molar concentration in the liver was about sixfold higher than the concentration in the kidney (1.78 nmol/g in the liver and 0.27 nmol/g in the kidney, see Fig. 1).

In vivo formation of NAFB1 and NAHFB1

N-acyl derivatives of fumonisins having various acyl chain lengths were found in animals treated with FB1 or HFB1, demonstrating for the first time the in vivo formation of N-acyl-FB1 in fumonisin’s two main target organs in rats (see Fig. 2). No metabolites have been observed in the vehicle-treated (control) animals.

The NAFB1 levels (expressed as the sum of all N-acyl derivatives) varied only slightly and ranged from 0.18 to 0.38 nmol/g. The amount of HFB1 metabolites in the liver was sixfold higher than that in the kidneys: the organs’ respective NAFB1 concentrations were 2.8 and 0.5 nmol/g tissue (see Fig. 2). Very low amounts of NAHFB1 (<0.1 nmol/g tissue) were also detected in the tissues of animals given FB1. Their source was not determined although it can be speculated that they resulted from absorption and metabolism of HFB1 that had been formed from hydrolysis of FB1 by the intestinal microbiota (Shephard et al. 1994 and 1995). However, the possibility that enzymatic hydrolysis of small amounts of absorbed FB1 provided a pool of HFB1 in the tissues cannot be discounted.

Total FB1 and HFB1 species in the kidney and liver

The tissue burdens of FB1, HFB1, and N-acyl metabolites in the kidney and liver after 5 days of exposure represented only a small fraction of the administered FB1 or HFB1. Differences in the accumulation in these target organs were apparent (see Fig. 3). Only 0.1 to 0.3 % of the total dose (TD) was recovered from the liver as unmetabolized FB1 while the amount found in the kidney was slightly higher, ranging from about 0.5 to slightly less than 0.8 % of TD. Compared to the high dose of FB1, a relatively high amount of unmetabolized HFB1, 0.7 % TD, accumulated in the liver and a relatively lower amount, <0.1 % TD of HFB1, in the kidney.

N-acyl-FB1 species in the liver ranged from 0.1 (low- and high-dose groups) to about 0.3 % TD (low-dose group) on a molar basis. Although low, these amounts represented 35 to 60 % of the total FB1 species in the liver (see Fig. 3). Only insignificant amounts of the FB1 species (<0.04 % TD and <10 % of FB1 species in the kidney) found in the kidneys were N-acyl metabolites. In contrast, N-acyl-HFB1 species in the liver accounted for 1.1 % TD, but, as was the case for FB1, this amount made up about 60 % of the total HFB1 species recovered in the liver. Similar to FB1 and its metabolites, the total amount of HFB1 species in the kidney was negligible and represented only 0.03 % TD. However, in contrast to the low contribution of metabolites to total renal FB1 species, 65 % of the HFB1 found in the kidney consisted of N-acyl metabolites.

The high dose of FB1 (2.77 μmol/kg body weight) and the HFB1 dose (2.47 μmol/kg body weight) were approximately equal, and, when comparing the results from these treatment
groups, it was apparent that the kidney accumulated FB1 much more readily than HFB1. The results also suggest that the kidney has only a limited capacity to metabolize both mycotoxins as less than 0.5 nmol/g of FB1, or HFB1 metabolites were found. The male rat kidney is recognized as being extremely sensitive to the apoptotic and other effects of FB1 (Howard et al. 2001; Dragnea et al. 2001). Nephrotoxicity has therefore served as a sentinel of exposure for applied studies (Burns et al. 2008; Vosse et al. 2011) and has also provided a benchmark dose for risk assessment (Bulder et al. 2012).

Tissue specificity of N-acylation of fumonisins

Different isoforms of CerS exist in the tissues of mammalian organisms. All are tissue specific, catalyze the N-acylation of sphinganine, and are specific for coenzyme A fatty acid cosubstrates having fatty acid residues that vary in length from C_{14} to C_{26} (Pewzner-Jung et al. 2006). We therefore analyzed the individual levels of ceramides and NAFAFB1 and NAHFB1 metabolites of different N-acyl chain lengths in the kidney and the liver of FB1- or HFB1-exposed rats. The amounts of ceramide species found in the two tissues were different with C_{16} derivatives being more predominant in the kidney. In contrast, CerS isoforms in the liver generated mainly C_{24}-ceramides (all quantitative results for each individual N-acyl derivative can be found in Table S1 the Supplementary Material). It is of interest that the ratio of N-C_{16}- to N-C_{24}-fumonisin derivatives in the kidney and liver was consistent with that of the ceramides found in each tissue (see Fig. 4). It can therefore be concluded that NAFAFB1 and NAHFB1 found in the liver and kidney were most likely products of reactions catalyzed by the CerS isoforms specific to those tissues. The gene expression of the CerS isoforms and

\[ \text{Fig. 1} \quad \text{The molar concentrations of FB1 and HFB1 in the kidney and liver of rats. The animals were treated by intraperitoneal injection of FB1 or HFB1 for 5 days with doses of 0.69, 1.38, and 2.77 \, \mu\text{mol FB1/kg body weight per day and 2.47 \, \mu\text{mol/kg body weight HFB1 per day. The dose of}} \]

\[ \text{HFB1 is the molar equivalent to the high dose group of FB1. FB1 and HFB1 in the tissues were quantified by HPLC-ESI-MS/MS. Values are means±S.D. (n=2). Note the difference in scale, y-axis. FB1 or HFB1 were not detected in tissues of vehicle-treated (control) rats.} \]

\[ \text{Fig. 2} \quad \text{The concentrations of NAFAFB1 and NAHFB1 in the kidney and liver of rats (expressed as the sum of all N-acyl derivatives). Animals were treated by intraperitoneal injection of FB1 or HFB1 for 5 days with doses of 0.69, 1.38, and 2.77 \, \mu\text{mol FB1/kg body weight per day or 2.47 \, \mu\text{mol/}} \]

\[ \text{kg body weight HFB1 per day. The dose of HFB1 is the molar equivalent to the high dose group of FB1. NAFAFB1 and NAHFB1 were analyzed by HPLC-ESI-MS/MS. Values are means±S.D. (n=2). Note the difference in scale, y-axis.} \]
the concordant ceramide pattern is well understood for the renal and hepatic tissue of mice. From the data of Laviad et al. (2008), it is seen that in the kidney of mice, the predominant ceramide species are the N-C16 derivatives and in the liver the N-C24 derivatives. Our data indicate that the ceramide pattern is consistent among mouse and rat tissues. However, there is no established knowledge about the chain length distribution and the gene expression of CerS isoforms in rats. Therefore, it could be interesting to verify these observations in further studies.

Sphingolipids

Inhibition of CerS by FB1 is the key event in fumonisins’ mode of action (Dragan et al., 2001, Bulder et al., 2012). Inhibition disrupts overall sphingolipid metabolism, resulting in increased concentrations of sphinganine, sphingosine, and their 1-phosphates and decreased levels of ceramide and complex sphingolipids in tissues in vivo (Voss et al. 2009) and in cultured cells (Wang et al. 1991). The levels of several sphingolipids were therefore analyzed in the samples, and, as expected (Riley and Voss 2006), the rats responded to FB1 with increased amounts of total sphingosine (sphingosine plus sphingosine-1-phosphate), total sphinganine (sphinganine plus sphinganine-1-phosphate) (see Fig. 5), and the individual sphingoid bases and 1-phosphate metabolites in the tissues (see Table 1). In contrast, treatment with HFB1 did not cause any significant changes in the levels of the sphingolipids compared to the vehicle-treated controls (see Fig. 5 and Table 1). The FB1 modified sphingolipid profiles in the liver and kidney differently. The two sphingoid bases and their 1-phosphates were increased somewhat in the liver and much more extensively in the kidney whereas tissue ceramide concentrations were decreased in both tissues, with levels in the liver tending to be slightly lower (see Fig. 5 and Table 1). A limited number of in vitro studies have shown that NAHB1 metabolites of HFB1 alter sphingolipid concentrations in various cell lines, including those overexpressing CerS isoforms (Seefelder et al. 2003; Harrer et al. 2013), and also were cytotoxic to IHKE human proximal tubule-derived cells (Seefelder et al. 2003) and other mammalian cell lines (Harrer et al. 2013). Whether or not NAFB1 and NAHB1 metabolites contribute to toxicity in vivo is not known and requires further investigation. That HFB1 treatment in this study did not significantly alter tissue sphingolipid...
concentrations, is consistent with reports that it exerts a significantly lesser effect on tissue sphingolipid profiles, and is less toxic to rats and mice than FB1 (Collins et al. 2006; Howard et al. 2002; Voss et al. 2009).

**Fig. 5** The levels of total sphinganine (Sa) plus sphinganine-1-phosphate (Sa1P), total sphingosine (So) plus sphingosine-1-phosphate (So1P), and ceramide in rat kidney and liver. Animals were treated by intraperitoneal injection with doses of 0.69, 1.38, and 2.77 μmol FB1/kg body weight per day and 2.47 μmol/kg body weight HFB1 per day. The dose of HFB1 is the approximate molar equivalent of the high dose of FB1. The tissue samples were extracted, and all analytes were determined by HPLC-ESI-MS/MS. Values are means±S.D. (n=2)

**Table 1** The levels of sphinganine (Sa), sphingosine (So), sphinganine-1-phosphate (Sa1P), and sphingosine-1-phosphate (So1P) and the Sa/So ratio in rat kidney and liver. Animals were treated by intraperitoneal injection with doses of 0.69, 1.38, and 2.77 μmol FB1/kg body weight per day and 2.47 μmol/kg body weight HFB1 per day. The dose of HFB1 is the approximate molar equivalent of the high dose of FB1. The tissue samples were extracted, and all analytes were determined by HPLC-ESI-MS/MS. Values are means±S.D. (n=2)

| Tissue | Substance | Dose | Sa nmol/g | So nmol/g | Sa/So ratio | Sa1P nmol/g | So1P nmol/g |
|--------|-----------|------|-----------|-----------|-------------|-------------|-------------|
| Kidney | Control   |      | 2±0.7     | 6±0.4     | 0.4±0.1     | 1.6±0.7     | 1.0±0.1     |
| Kidney | FB1       | Low  | 252±15.6  | 63±11.5   | 4.1±1.0     | 16.7±0.5    | 21.4±4.1    |
| Kidney | FB1       | Mid  | 146±11.3  | 59±8.3    | 2.5±0.2     | 11.3±1.5    | 20.8±0.7    |
| Kidney | FB1       | High | 207±79.9  | 56±6.2    | 3.8±1.8     | 19.9±1.0    | 21.6±4.5    |
| Kidney | HFB1      | High | 3±0.7     | 9±2.8     | 0.3±0.0     | 0.6±0.4     | 0.4±0.1     |
| Liver  | Control   |      | 3±0.3     | 3±0.6     | 1.0±0.1     | 0.2±0.0     | 0.3±0.0     |
| Liver  | FB1       | Low  | 68±27.6   | 16±3.3    | 4.2±0.9     | 0.2±0.2     | 0.4±0.2     |
| Liver  | FB1       | Mid  | 88±15.1   | 17±6.4    | 5.5±1.2     | 0.4±0.1     | 0.8±0.0     |
| Liver  | FB1       | High | 81±17.1   | 14±2.9    | 5.9±0.0     | 0.9±0.5     | 1.0±0.3     |
| Liver  | HFB1      | High | 2±1       | 3±0.2     | 0.5±0.4     | 0.1±0.1     | 0.2±0.0     |
Dosing route and relevance

Given the previous absence of evidence for in vivo FB₁ metabolism (other than conversion to HFB₁ by gut microflora) together with its low absorption after oral exposure (<5 % of dose) (reviewed by Bulder et al. 2012) and low conversion rate to NAFB₁ in vitro (Harrer et al. 2013), multiple high ip doses were used in this “proof of concept” study. The extent to which the NAFB₁ species are formed and accumulate in tissues after dietary FB₁ exposure in animals or humans is not known, but is almost certain to be manyfold lower than that found in this study. It should further be recognized that the high doses of FB₁ or HFB₁ possibly compromised tissue metabolic or transport systems, thereby influencing metabolite production and retention. This possibility is especially relevant for the kidney of rats given FB₁ because of the mild to moderate nephrotoxicity induced in these animals. Additional investigations following oral exposure to physiologically relevant doses are therefore needed to evaluate the role, if any, of NAFB₁ in FB₁ toxicity.

Conclusions

This experiment has to our knowledge revealed for the first time the metabolic conversion of FB₁ to a series of NAFB₁ species in vivo. Consistent with earlier reports, much more FB₁ was recovered from the kidneys than from the liver. Renal FB₁ was however almost exclusively unmetabolized whereas approximately half of the FB₁ in the liver consisted of NAFB₁. FB₁ exposure in animals or humans is not known, but is almost certain to be manyfold lower than that found in this study. It should further be recognized that the high doses of FB₁ or HFB₁ possibly compromised tissue metabolic or transport systems, thereby influencing metabolite production and retention. This possibility is especially relevant for the kidney of rats given FB₁ because of the mild to moderate nephrotoxicity induced in these animals. Additional investigations following oral exposure to physiologically relevant doses are therefore needed to evaluate the role, if any, of NAFB₁ in FB₁ toxicity.

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Conflict of interest The authors have declared no conflict of interest.

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