Cleavage of Zearalenone by *Trichosporon mycotoxinivorans* to a Novel Nonestrogenic Metabolite

Elisavet Vekiru, 1* Christian Hametner, 2 Rudolf Mitterbauer, 3 † Justyna Rechthaler, 4 Gerhard Adam, 3 Gerd Schatzmayr, 5 Rudolf Krška, 1 and Rainer Schuhmacher 1

Christian Doppler Laboratory for Mycotoxin Research, Department for Agrobiontotechnology (IFA Tulln), University of Natural Resources and Applied Life Sciences Vienna, Konrad Lorenz Str. 20, 3430 Tulln, Austria; 1 Institute of Applied Synthetic Chemistry, Vienna University of Technology, Getreidemarkt 9/163, 1060 Vienna, Austria; 2 Department of Applied Genetics and Cell Biology, University of Natural Resources and Applied Life Sciences Vienna, Muthgasse 18, 1190 Vienna, Austria; 3 University of Applied Sciences Wr. Neustadt, Konrad Lorenz Str. 10, 3430 Tulln, Austria; and Biomin Research Center, Technopark 1, 3430 Tulln, Austria

Received 19 June 2009/Accepted 22 January 2010

Zearalenone (ZON) is a potent estrogenic mycotoxin produced by several *Fusarium* species most frequently on maize and therefore can be found in food and animal feed. Since animal production performance is negatively affected by the presence of ZON, its detoxification in contaminated plant material or by-products of bioethanol production would be advantageous. Microbial biotransformation into nontoxic metabolites is one promising approach. In this study the main transformation product of ZON formed by the yeast *Trichosporon mycotoxinivorans* was identified and characterized by liquid chromatography-tandem mass spectrometry (LC-MS/MS) and LC-diode array detector (DAD) analysis. The metabolite, named ZOM-1, was purified, and its molecular formula, C_{35}H_{41}O_{7}, was established by time of flight MS (TOF MS) from the ions observed at m/z 351.1445 [M-H]⁻ and at m/z 375.1416 [M+Na]⁺. Employing nuclear magnetic resonance (NMR) spectroscopy, the novel ZON metabolite was finally identified as (5S)-5-{2,4-dihydroxy-6-[(1E)-5-hydroxypent-1-en-1-yl]benzoyl}oxy)hexanoic acid. The structure of ZOM-1 is characterized by an opening of the macrocyclic ring of ZON at the ketone group at C6. ZOM-1 did not show estrogenic activity in a sensitive yeast bioassay, even at a concentration 1,000-fold higher than that of ZON and did not interact with the human estrogen receptor in *in vitro* competitive binding assay.

Zearalenone (ZON) is the main member of a growing family of biologically important “resorcylic acid lactones” (RALs), which have been found in nature. ZON is produced by several *Fusarium* species, which colonize maize, barley, oat, wheat, and sorghum and tend to develop ZON during prolonged cool, wet growing and harvest seasons (38). Maize is the most frequently contaminated crop plant, and therefore, ZON can be found frequently in animal feeding stuff. Occurrence, toxicity, and metabolism data of ZON were summarized by the European Food Safety Authority (EFSA) (5) and in recent reviews (12, 38).

The potent xenohormone ZON leads to hyperestrogenism symptoms and in extreme cases to infertility problems, especially in pigs (15). Ovarian changes in pigs have been noted with toxin levels as low as of 50 µg/kg in the diet (1). Ruminants are more tolerant to ZON ingestion; however, hyperestrogenic syndrome, including restlessness, diarrhea, infertility, decreased milk yields, and abortion, have been well documented with cattle and sheep (4, 29).

Because widespread ZON contamination in feed can occur in problematic years, efficient ways to detoxify are desirable. The transformation of mycotoxins to nontoxic metabolites by pure cultures of microorganisms or by cell-free enzyme preparations (3) is an attractive possibility. Microbial metabolization of ZON to alpha-ZOL and beta-ZOL cannot be regarded as detoxification, because both ZOL products are still estrogenic (14). Also, formation of ZON-glucosides and -digluco-"sides (8, 17) and ZON-sulfate (7) cannot be considered true detoxification but rather formation of masked mycotoxins, because the conjugates may be hydrolyzed during digestion (11, 23), releasing ZON again (2).

As the estrogenic activity of ZON and its derivates can be explained by its chemical structure, which resembles natural estrogens (20), it can be expected that cleavage of the lactone undecyl ring system of ZON results in permanent detoxification.

El-Sharkawy and Abul-Hajj (9) were the first to report inactivation of ZON after opening of the lactone ring by *Gliocladium roseum*. This filamentous fungus was capable of metabolizing ZON in yields of 80 to 90%. Also Takahashi-Ando et al. (31) described the degradation reaction of ZON with *Clonostachys rosea* (synonym of *G. roseum*). A hydroxylase (encoded by a gene designated *ZHD101*) cleaves the lactone ring, and as recently proved (37; unpublished data) by subsequent decarboxylation of the intermediate acid, the compound 1-(3,5-dihydroxyphenyl)-10'-hydroxy-1'E-undecene-6'-one is formed. In contrast to ZON and 17β-estradiol, which showed potent estrogenic activity, this cleavage product did not show...
any estrogenic activity in the human breast cancer MCF-7 cell proliferation assay (16). Further details, e.g., on the conditions of the maximum activity of ZHD101 and its exploitation in genetically modified grains, can be found in later published work of this research group (32, 33).

Only a few authors reported the loss of estrogenicity in microbial metabolites of ZON, which is based on reactions other than cleavage of the lactone undecyl ring system. El-Sharkawy and Abul-Hajj demonstrated (10) that binding to rat uterine estrogen receptors requires a free 4-OH phenolic group (devoid of methylation or glycosylation). Loss of estrogenicity was, for instance, observed with 2,4-dimethoxy-ZON, one of the metabolites produced by Cunninghamella bainieri ATCC 9244B. Nevertheless, this rule cannot be generalized, as 8′-hydroxyzearalenone and 8′/H11032 bind to the estrogen receptor. Also, other authors reported that 8′-hydroxyzearalenone and 8′-epi-hydroxyzearalenone are nonestrogenic (13). However, so far, no practical application in feed or food detoxification has been found for the microorganisms producing these compounds.

It has been shown previously that the yeast Trichosporon mycotoxinivorans has a very high capability to degrade both ochratoxin A (OTA) and ZON (22, 26, 27). When T. mycotoxinivorans is used as a feed additive preparation, microbial degradation of the mycotoxins is assumed to take place in the gastrointestinal tract of the animal after consumption of contaminated feed. The protective effect of T. mycotoxinivorans against OTA toxicity has already been shown with broiler chicken (24).

In the present study we report the isolation, analytical characterization, and structure elucidation, as well as the evaluation, of the estrogenic activity of the main degradation product of ZON produced by T. mycotoxinivorans.

MATERIALS AND METHODS

Microbial cultivation of T. mycotoxinivorans and degradation of ZON. Erlenmeyer flasks with 30 ml yeast medium (10 g/liter glucose, 20 g/liter malt extract, 10 g/liter yeast extract, 5 g/liter peptone of casein) were inoculated with T. mycotoxinivorans (22) directly with aliquots from a culture stock stored at −80°C and incubated at 37°C and 200 rpm on an orbital shaker. After 48 h, biomass was harvested by centrifugation, resuspended in the same volume of sterile 0.9% NaCl containing 10 mg/liter ZON (Biopure Referenzsubstanzen GmbH, Tulln, Austria), and incubated under the same conditions. As controls, flasks containing solutions without ZON (matrix control) or without biomass (substrate control) were incubated in parallel. Samples (1.0 ml) were taken at given time points for up to 6 days and heat inactivated in glass vials for 5 min in a boiling water bath. Samples were stored frozen (−20°C) until analysis.

For preparative-scale production of the ZON metabolite ZOM-1, a total culture volume of 900 ml (150 ml per flask) was incubated under the conditions described above. Biomass was harvested, washed with 0.9% NaCl solution, dissolved in 900 ml minimal medium (MM) (27) containing 50 mg/liter of ZON, portioned in 50-ml volumes, placed in a 300-ml Erlenmeyer flask, and incubated at 35°C for 192 h. Heat inactivation in an autoclave followed for 10 min. Samples were stored frozen (−20°C) until analysis.

LC-MS/MS and LC-UV analysis. Liquid chromatography-tandem mass spectrometry (LC-MS/MS) analyses were performed with a QTrap LC-MS/MS system (Applied Biosystems, Foster City, CA), equipped with an electrospray ionization (ESI) source and an 1100 series high-pressure liquid chromatography (HPLC) system (Agilent Waldbronn, Germany), including an 1100 series diode array detector (DAD).

Enhanced mass spectra (EMS) and enhanced product ion (EPI) scans, as well as other MS operation modes, e.g., precursor ion scan (data not shown), were used to gain structural information about the molecules. For analysis, samples were clarified by centrifugation (Beckman GS-6; 10 min at 3,500 rpm), and the supernatant was transferred to HPLC vials for direct injection.

Chromatographic separation was achieved at 25°C using a gradient on an 250- by 3.0-mm column (by inner diameter [i.d.]), 5-μm-particle-size Phenomenex Luna C18(2) column (Phenomenex Inc., Torrance, CA), including an identical matrix guard column. The injection volume was 10 μl, while the flow rate was 0.5 ml/min. Based on the mobile phases A, H2O/HCOOH (50 μ/liter), and B, MeOH/H2O/HCOOH (50 μ/liter), the elution started at 15% solvent B, with a linear gradient to 100% solvent B from 0 to 20 min. After holding for 7 min, the initial conditions returned within 1 min and were held for reequilibration until the end of the run at 38 min.

The ESI source was operated at 400°C in negative ionization mode. EMS parameters in the negative ionization mode were as follows: curtain gas (CUR), 20 lb/in 2; nebulizer gas (GS1), 25 lb/in 2; auxiliary gas (GS2), 65 lb/in 2; ion spray voltage (IS), −4,200 V; declustering potential (DP), −30 V; entrance potential (EP), −10 V; collision energy (CE), −30 V; mass range, 100 to 850 amu; scan rate, 1,000 amu/s; and linear iontrap (LIT) fill time, 40 ms, where 1 lb/in 2 is 6.895 kPa. MS/MS spectra were recorded in the EPI mode with the following parameters: LIT fill time, 40 ms; scan rate, 1,000 amu/s; Q1 resolution, unit; Q0 trapping; yes; and MR pause, 5.0 ms. The collision energy and the declustering potential for m/z 351.1 were DP, −51 V; and CE, −35 eV.

Data acquired by the DAD were recorded at 220 and 270 nm.

ZOM-1 isolation and purification. Ethyl acetate was added at a ratio of 0.5:1 (vol/vol) to the clarified culture fluid, and extraction was repeated three times. The combined organic fractions were dried over anhydrous sodium sulfate (Na2SO4) and concentrated to dryness in a rotary evaporator, and the residue was reconstituted in ethyl acetate and transferred to a 4-ml screw vial in which it was vortexed again in a scheme to reconstitute up to 42.5 mg of crude residue. The crude residue was reconstituted in 2 ml of a mixture of MeOH/H2O (70:30) and cleared by centrifugation, and the supernatant was transferred to an HPLC vial.

ZOM-1 isolation was done at 25°C on a semi-preparative Phenomenex Luna C18(2) column (Phenomenex Inc., Torrance, CA), 250- by 100-mm length (by i.d.), 5-μm particle size, including a C18 security guard column (10.0 by 10.0 mm; 5 μm). The injection volume was 80 μl, while the flow rate was 3.0 ml/min. The mobile phases consisted of A (H2O/HCOOH [50 μ/liter]) and B (MeOH/ HCOOH [50 μ/liter]). Elution started isocratically at 65% B for 14 min, increased to 95% to wash the column, and returned to 65% B for column equilibration. Fractions belonging to ZOM-1 were collected after peak detection at 270 nm. After solvent evaporation using nitrogen gas flow and overnight lyophilization, the yield of isolated ZOM-1 was 16.7 mg.

ZOM-1 characterization via time of flight (TOF) MS. Accurate mass measurements of the purified ZOM-1 were performed on an ESI microOTOF mass spectrometer (Bruker Daltonik, Bremen, Germany) with flow injection.

Data acquisition was controlled by Bruker Daltonics microOTOF control version 1.1. Data evaluation was performed using the Generate Molecular Formula (GMF) software suite within Bruker Daltonics microOTOF DataAnalysis version 3.2. For assigning both accurate mass positions and isotopic patterns for the calculation of molecular formulae, the mass accuracy tolerance window of generated masses was set to 30 ppm in order to check how many hits are possible in the mass range of interest. Only molecular formulae of ions which yielded a “sigma value” of <0.02 (sigma is an indicator value for the isotopic pattern fit) were considered confident.

Calibration was done internally using sodium formate clusters [Na(NaCOOH)x]+ in positive ionization mode and formate adducts of sodium formate clusters in the form of [HCOO(NaCOOH)y]− in negative ionization mode. Samples containing approximately 1 μg/liter ZOM-1 in MeOH/H2O (1:4 vol/vol) and the calibration standard were introduced into the mass spectrometer via a Hamilton syringe at a flow rate of 180 μl/h, and measurements were carried out in the positive as well as negative ion mode using a scan range of m/z 50 to 1,000 and the following settings for both polarities: end plate offset, 500 V; capillary voltage, 4,500 V; nebulizer pressure (N2), 0.5 bar; gas flow (N2), 5 liter/min; dry temperature, 200°C; flight tube voltage value, 9,000 V; and reflector voltage, 1,300 V. The following transfer parameters were applied in positive ion mode: capillary exit at 120 V, skimmer 1 at 50 V, hexapole 1 at 24 V, skimmer 2 at 23 V, hexapole 2 at 21 V, hexapole RF at 150 voltage per pole (Vpp), transfer time at 45 μs, and pre puls storage at 5 μs. In the negative ion mode, the transfer parameters were capillary exit at −120 V, skimmer 1 at −40 V, skimmer 2 at −24 V, skimmer 2 at −21 V, hexapole RF at 120 Vpp, transfer time at 45 μs, and pre puls storage at 5 μs.

Nuclear magnetic resonance (NMR) spectroscopy of ZOM-1. For structure elucidation of ZOM-1, 1H, 13CAPT, 1H-1H correlation spectroscopy (COSY), 1H-13C heteronuclear single quantum correlation (HSQC), and 13C-13C hetero-
The ability of ZON and ZOM-1 to bind to the human estrogen receptor was assayed in vitro using the HitHunter EFC estrogen chemiluminescence assay kit (DiscoveRx/Amersham Biosciences) and the human estrogen receptor-α (Sigma) as previously described for zearalenone-4-glucoside (25). Luminescence of the β-galactosidase product was measured after 1.5 h of incubation with a luminometer (Victor 2; Wallac/Perkin Elmer, Monza, Italy).

RESULTS

T. mycotoxinivorans was incubated for 0 to 144 h in saline medium with 10 mg/liter ZON, and culture medium was analyzed by LC-MS/MS and LC-DAD. Saline was used in order to keep matrix effects in the subsequent analysis to facilitate the identification of newly formed compounds as easy as possible. Due to the substrate controls (ZON incubated in culture medium without T. mycotoxinivorans), loss of ZON due to adsorption or decomposition during incubation could be excluded. Comparison between DAD chromatograms after cultivation of T. mycotoxinivorans alone (culture controls) and cocultivation of T. mycotoxinivorans and ZON, both treated in the same manner, resulted in the identification of one main ZON-derived metabolite (ZOM-1) eluting at a retention time of 18.2 min. (Fig. 1). The increase of ZOM-1 concentration over incubation time is illustrated in Fig. 1 (inset). ZOM-1 concentration was estimated by assuming a UV response equal to that for ZON at the same concentration. This seemed to be feasible since no other major metabolites were detected in the samples, and the UV spectra of both ZON and ZOM-1 were similar (data not shown). After 48 h of incubation, 95% of ZON was metabolized. It was converted to ZOM-1, which appeared to be stable for several days, indicating that ZOM-1 constitutes a stable metabolization end product under the conditions applied.

The data of the LC-MS total ion chromatogram (TIC) obtained in the negative enhanced full scan mode (EMS) confirmed the formation of one main ZON metabolite at 18.2 min. After subtraction of the culture control TICs, the resulting full-scan mass spectrum of the putative ZOM-1 (Fig. 2a) was evaluated in detail. Initially this spectrum was difficult to interpret; however, after these data were combined with additional measurements in the positive ionization mode (showing [M + Na]⁺ at m/z 375 as the most abundant ion [data not shown]), a molar mass (M) of 352 g/mol was assumed for ZOM-1. The mass shift of 34 amu might indicate the addition of 2 hydrogen and 2 oxygen atoms to ZON. While the signal at m/z 351 was assigned to the deprotonated molecular ion [M–H]⁻, m/z 373 was assumed to originate from its sodium adduct [M–2H + Na]⁺. Since sodium adduct formation in negative ESI is rare but has been reported for aromatic acids (28), the signal at m/z 373 might correspond to the presence of a free carboxylic group in the ZON metabolite.

As demonstrated by the EPI spectrum in negative ionization of the deprotonated molecular ion [M–H]⁻ at m/z 351 (Fig. 2b), the signals at m/z 351 was assigned to the deprotonated molecular ion [M–H]⁻, m/z 373 was assumed to originate from its sodium adduct [M–2H + Na]⁺. Since sodium adduct formation in negative ESI is rare but has been reported for aromatic acids (28), the signal at m/z 373 might correspond to the presence of a free carboxylic group in the ZON metabolite.
and isolation, a putative molecular formula of C$_{18}$H$_{24}$O$_7$ was formed. After extraction and preparative HPLC purification lar formula of C$_{12}$H$_{13}$O$_5$, C$_{12}$H$_{11}$O$_4$, and C$_{18}$H$_{22}$NaO$_7$ for determined by NMR analysis. From 1H and 13C-APT spectra

m/z amu and formation of the ion at 

ilarities to the MS/MS spectrum of ZON, e.g., the loss of 114 

Highlighted are the deprotonated molecular ion [M-H]$^-$ at m/z 351 and further signals of its sodium and potassium adducts (m/z 373, m/z 389) as well as the molecular dimer ion at m/z 703 and further alkali-bridged dimer ions at m/z 725, 741, 747, and 763 (zoomed). (b) Structure of the ZON metabolite (ZOM-1) of T. mycotoxinivorans and consecutive numbering of the carbon atoms. Enhanced product ion data of its deprotonated molecule ion [M-H]$^-$ at m/z 351 and scheme of proposed major fragmentation pathways. MW indicates molecular weight.

For further characterization of ZOM-1, a large-scale incubation experiment with ZON and T. mycotoxinivorans was performed. After extraction and preparative HPLC purification and isolation, a putative molecular formula of C$_{18}$H$_{23}$O$_7$ was assigned to the purified ZOM-1 by TOF MS measurements. In detail, measurement in the negative ionization mode showed the formation of [M-H]$^-$ at m/z 351,1445. Evaluation with the GMF software revealed the molecular formula C$_{18}$H$_{23}$O$_7$. The relative deviation between the measured mass and the theoretical mass of this signal was 1.1 ppm, and the sigma factor was calculated to be 0.0082, indicating very good agreement between measured spectra and calculated molecular mass as well as isotope pattern. The spectrum also included the main fragments of [M-H]$^-$ at m/z 237 and 219 as well as an adduct formation at m/z 373. GMF evaluation suggested the molecular formula of C$_{12}$H$_{13}$O$_5$, C$_{12}$H$_{11}$O$_4$, and C$_{18}$H$_{22}$NaO$_7$ for these signals, respectively. This confirmed our early interpretation of m/z 373 as the sodium adduct [M+Na-H]$^-$ and supported our structural interpretation of the fragments at m/z 237 and 219, as shown in Fig. 2b. Measurement in the positive ionization mode (data not presented) showed only the formation of [M+Na]$^+$ at m/z 375,1416 and not of [M+H]$^+$. GMF evaluation suggested the molecular formula C$_{18}$H$_{22}$NaO$_7$ with an accuracy of $\pm 0.554$ ppm and a calculated sigma factor of 0.0043. The spectrum also included an [M-H + 2Na]$^+$ adduct formed at m/z 397,1231.

The complete molecular structure of ZOM-1 was finally determined by NMR analysis. From 1H and 13C-APT spectra (Table 1), it was evident by comparison that a substantial part of the ZON structure is preserved in ZOM-1, namely, the tetra-substituted aromatic ring, including the benzoic ester moiety and the olefinic double bond. This was proven by two-dimensional spectra (1H-1H COSY, 1H-13C HSQC, and 1H-13C HMBC), which also allowed the elucidation of the remaining part of the structure.

<table>
<thead>
<tr>
<th>Position</th>
<th>13C (ppm)$^a$</th>
<th>1H (ppm)$^a$</th>
<th>J (Hz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>104.1 (s)</td>
<td>6.28 (1H)</td>
<td>2.5 (d)</td>
</tr>
<tr>
<td>2</td>
<td>164.3 (s)</td>
<td>6.28 (1H)</td>
<td>2.5 (d)</td>
</tr>
<tr>
<td>3</td>
<td>101.6 (d)</td>
<td>6.22 (1H)</td>
<td>2.5 (d)</td>
</tr>
<tr>
<td>4</td>
<td>162.6 (s)</td>
<td>6.22 (1H)</td>
<td>2.5 (d)</td>
</tr>
<tr>
<td>5</td>
<td>108.2 (d)</td>
<td>6.38 (1H)</td>
<td>2.5 (d)</td>
</tr>
<tr>
<td>6</td>
<td>143.8 (s)</td>
<td>6.38 (1H)</td>
<td>2.5 (d)</td>
</tr>
<tr>
<td>1'</td>
<td>131.7 (d)</td>
<td>6.96 (1H)</td>
<td>15.5 (d), 1.5 (t)</td>
</tr>
<tr>
<td>2'</td>
<td>131.6 (d)</td>
<td>5.92 (1H)</td>
<td>15.4 (d), 6.9 (t)</td>
</tr>
<tr>
<td>3'</td>
<td>29.4 (t)</td>
<td>2.29 (2H)</td>
<td>7.6 (t), 6.9 (d), 1.5 (d)</td>
</tr>
<tr>
<td>4'</td>
<td>32.3 (t)</td>
<td>1.73 (2H)</td>
<td>(m)</td>
</tr>
<tr>
<td>5'</td>
<td>65.1 (t)</td>
<td>3.64 (2H)</td>
<td>6.6 (t)</td>
</tr>
<tr>
<td>6'</td>
<td>176.3 (s)</td>
<td>3.64 (2H)</td>
<td>6.6 (t)</td>
</tr>
<tr>
<td>7'</td>
<td>33.7 (t)</td>
<td>2.36 (2H)</td>
<td>6.5 (t)</td>
</tr>
<tr>
<td>8'</td>
<td>21.1 (t)</td>
<td>1.80–1.65 (2H)</td>
<td>(m)</td>
</tr>
<tr>
<td>9'</td>
<td>35.5 (t)</td>
<td>1.85–1.70 (2H)</td>
<td>(m)</td>
</tr>
<tr>
<td>10'</td>
<td>72.3 (d)</td>
<td>5.21 (1H)</td>
<td>(m)</td>
</tr>
<tr>
<td>11'</td>
<td>19.4 (q)</td>
<td>1.38 (3H)</td>
<td>6.3 (d)</td>
</tr>
<tr>
<td>12'</td>
<td>171.3 (s)</td>
<td>1.38 (3H)</td>
<td>6.3 (d)</td>
</tr>
</tbody>
</table>

$^a$ 13C and 1H NMR spectra were recorded at 100 MHz and 400 MHz, respectively. Chemical shifts in ppm are referenced to tetramethylsilane (TMS) as an internal standard. Spectra were recorded in CD$_3$OD.

$^b$ Multiplicity was determined from APT spectra.

$^c$ See Fig. 2b for numbering of the carbon atoms. Multiplicities (shown in parentheses) are abbreviated as follows: s, singlet; d, doublet; t, triplet; q, quartet; m, multiplet.
position 10', the methyl group 11' and the adjacent CH2 group (9') were easily identified, although the protons of the latter are part of an overlapped region of six signals between 1.90 and 1.60 ppm. Because of this overlap, the next members in the chain were identified via two- and three-bond C-H correlations, leading to two further CH2 groups 8' and 7' (21.1 and 33.7 ppm in the carbon spectrum). The chemical shift and coupling pattern of the 7' signals indicated a neighboring C—O fragment, which was assigned to the 176.3-ppm carbon signal via the HMBC spectrum. Its characteristic shift and the coupling pattern of the 7

In this work we studied the degradation of ZON by T. mycotoxinivorans, a basidiomycete yeast which is used as a microbial feed additive against mycotoxins. A nonestrogenic ZON metabolite (ZOM-1) was the main product of this ZON degradation. In order to facilitate metabolite identification and characterization, high concentrations of ZON (10 and 50 mg/liter) were used in a simple cultivation medium lacking some nutrients. Therefore, the incubation period had to be extended since under these conditions the ZON metabolizing activity of T. mycotoxinivorans is reduced. Moreover, additional metabolism experiments using complex culture media were carried out, which demonstrated a fast transformation of ZON to ZOM-1 and further currently uncharacterized metabolites in minor amounts (data not shown).

LC-MS/MS and TOF MS measurements demonstrated the structural similarity of ZOM-1 to ZON and revealed its fragment pattern and molecular formula. Subsequent NMR analysis of the purified metabolite confirmed its identity as (5S)-5-{(2,4-dihydroxy-6-[(1E)-5-hydroxypent-1-en-1-yl]benzoyl}oxy)hexanoic acid (Fig. 2b) in accordance with the molecular formula elucidated by mass spectrometry.

With the purified ZOM-1, an estrogenicity test was conducted. The growth of the indicator yeast strain YZRM7, as well as the positive-control strain YZGA376, on SC-His-Ura plates supplemented with either ZON or ZOM-1 is shown in Fig. 3A. Whereas a concentration of 3 nM ZON (~1 µg/liter ZON) allows growth of the estrogen-dependent bioassay strain YZRM7 due to activation of the estrogen-inducible URA3 gene, a 1,000-fold-higher concentration of the ZON metabolite does not. (B) Detection of the luminescent product of β-galactosidase activity released after 90 min of incubation. Competitive displacement of the estrogen receptor from the estrogen-linked donor peptide by ZON reconstitutes β-galactosidase activity, while the ZON metabolite is inactive.

DISCUSSION

In this work we studied the degradation of ZON by T. mycotoxinivorans, a basidiomycete yeast which is used as a microbial feed additive against mycotoxins. A nonestrogenic ZON metabolite (ZOM-1) was the main product of this ZON degradation. In order to facilitate metabolite identification and characterization, high concentrations of ZON (10 and 50 mg/liter) were used in a simple cultivation medium lacking some nutrients. Therefore, the incubation period had to be extended since under these conditions the ZON metabolizing activity of T. mycotoxinivorans is reduced. Moreover, additional metabolism experiments using complex culture media were carried out, which demonstrated a fast transformation of ZON to ZOM-1 and further currently uncharacterized metabolites in minor amounts (data not shown).

LC-MS/MS and TOF MS measurements demonstrated the structural similarity of ZOM-1 to ZON and revealed its fragment pattern and molecular formula. Subsequent NMR analysis of the purified metabolite confirmed its identity as (5S)-5-{(2,4-dihydroxy-6-[(1E)-5-hydroxypent-1-en-1-yl]benzoyl}oxy)hexanoic acid. The opening of the ring at the keto group of the macrocyclic ring of ZON and formation of a carboxy and hydroxy group is different from the hydrolysis of the preexisting lactone in ZON by Gliocladium/Clonostachys (followed by decarboxylation). The product ZOM-1 probably is formed by the proposed two-step mechanism illustrated in Fig. 4. First,
of the corresponding gene(s), it may also become feasible to develop enzymatic detoxification systems (36) or to engineer this detoxification pathway in other organisms.

ACKNOWLEDGMENTS

This work was supported by grants from the Christian Doppler Society and the Austrian Research Promotion Agency (FFG) and was done in cooperation with Biomin GmbH. We are grateful to Martin Täubel and Alexander Frank for their assistance with the degradation experiments. We also thank Dieter Moll for critically reading the manuscript.

REFERENCES


