STUDY OF AFLATOXIN B1-DESTROYING ACTIVITY OF Gliocladium roseum AND Trichoderma viride AND THEIR ANTAGONISM TOWARD TOXIGENIC Aspergillus flavus

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Abstract

Aflatoxin B1-destroying activity and antagonistic potential of Gliocladium roseum GRZ7 and Trichoderma viride TV35 strains isolated from natural substrates colonized by aflatoxigenic Aspergillus flavus were studied in vitro. Submerged cultures of G. roseum grown on liquid Czapek’s medium with casein hydrolysate (Czapek-CasH) at 28 °C and 200 rpm for 7 days were able to destroy 80-90 % of aflatoxin B1 (AFB1), which was added in the nutrient medium before inoculation. T. viride grown under the same conditions destroyed only 48 % of initial AFB1 during the same time of cultivation. The tested T. viride strain effectively suppressed the growth of toxigenic A. flavus strain A11 on Czapek-CasH agar. Co-cultivation of A11 with T. viride TV35 resulted in 64 % diminution of the average colony diameter of the aflatoxigenic strain. The strain GRZ7 of G. roseum was ineffective as an antagonist of A11. AFB1-destroying activity was detected in samples of high-molecular weight metabolites (> 5 kDa) isolated from culture liquid of G. roseum grown without AFB1. In addition, T. viride ability to degrade the mycotoxin was shown to be inducible. Obtained results were supposed to be of interest for further investigation on decontamination of feeds, which are contaminated with AFB1 or AFB1-producers.

Keywords: aflatoxin B1, biological decontamination, Gliocladium roseum, Trichoderma viride

Contamination of forage grains and plant materials with aflatoxins, the secondary metabolites of aspergillus fungi, is a serious problem. Because of hepatotoxicity, carcinogenicity and teratogenicity, aflatoxins are dangerous for the vast majority of warm-blooded organisms [1]. Feed contamination with aflatoxins can cause animal death or decreased production, and also lead to food stuff contamination [2]. Due to wide spread of aspergillas, their ability to evolve as facultative parasites on vegetating plants and, at the same time, to keep up saprophytic growth, it is almost impossible to completely exclude the contamination of grain and fodder grass by aflatoxin producers during harvesting and storage. Therefore, the approaches to solving the problem are mainly aimed at decontamination of raw materials [3, 4] or biocontrol of potentially aflatoxigenic Aspergillus species, in particular Aspergillus flavus, by microbial antagonists [5]. Due to the high stability of aflatoxins, their chemical or physical degradation is carried out under stringent conditions [6, 7]. This often reduces the feed quality making such treatment economically unprofitable. The biological method of decontamination [8-10] is based on the use of secondary metabolites of some plants toxic to A. flavus [11], and also on the search for natural inhibitors of aflatoxinogenesis or microorganism that could serve as a source of enzymes destroying aflatoxins or transform them to non-hazardous derivatives [12-16].

For example, when some micromycetes colonizing natural substrates are grown together with the toxigenic isolates of A. flavus, aflatoxin B1 (AFB1), typ-
ical for this fungus, decomposes in the culture liquid (CL) [17]. The enzymatic nature of the detoxifying activity of CL in a such biodestructor, *Phoma glomerata* (strain PG41), has been confirmed [18].

Here we first investigated the ability of two other micromycetes, *Gliocladium roseum* and *Trichoderma viride*, previously isolated from the toxigenic *A. flavus* consortium, to destroy AFB1 in view to assess the prospects of their use as sources of the AFB1-catabolizing enzymes or as antagonists of the of AFB1 producer.

**Technique.** Destroying activity of GRZ7 (*Gliocladium roseum* Bainier) and TV35 (*Trichoderma viride* Pers.) strains from the collection of Laboratory of Pathophysiology (All-Russian Research Institute of Phytopathology) was studied in 7-day cultures grown in liquid Czapek's medium with casein hydrolyzate (Czapek-CasH) at 28 °C and 200 rpm (Excella™ E-25/25R, New Brunswick Scientific Co., Inc., USA). Prior to fungi inoculation, AFB1 (5-10 μg/ml, Sigma™, USA) was added under sterile conditions. High molecular weight metabolites with nominally cut off molecular weight of > 5 kDa were isolated from CL of *G. roseum*, grown under the same conditions in the absence of toxin (intact CL), by precipitation of the filtrate with ammonium sulfate and ultrafiltration of aqueous solution of the precipitate. AFB1 was added to sterile samples of the high molecular weight fraction and the mixture was incubated at 27-28 °C for 3 days. Residual amounts of B1 in CL or in the sample of the fraction were determined using high performance liquid chromatography [18].

The effect of *T. viride* and *G. roseum* on the growth of toxigenic *A. flavus* (strain A11) was studied by the method of double cultures (co-cultivation on potato-glucose agar at 20-22 °C for 6 days). The minimum and maximum diameters were measured, and the area of the colonies was calculated based on the maximum diameter. Control cultures of strain A11 were grown under the same conditions in the absence of the putative antagonists.

Data were processed using Statistica 6.0 (StatSoft, USA). The significance of the differences between control and experimental values was confirmed by a t-test for independent variables (p ≤ 0.05). The table and the figure show the mean (M) and standard error of the mean (m). Each experiment included at least 6 replicates with 3-fold reproduction.

**Results.** A study of the dynamics of destruction of aflatoxin by *G. roseum* showed that after 3 days of its growth on medium with AFB1 the content of the latter in CL was halved, and by the end of cultivation it did not exceed 10-20 % of the initial amount (Fig., curve 4). A significant decrease in the AFB1 concentration also occurred after its incubation in the fraction of high-molecular metabolites isolated from the filtrate of intact CL of *G. roseum* (see Fig., curve 5). Thermal treatment of the fraction led to the loss of toxin-degrading activity (see Fig., curve 1).

These results suggest that extracellular enzymes of this biodestructor may participate in the decomposition of AFB1, similar to that in *P. glomerata* [18] or other ascomycetes [12, 19]. Earlier, it was found that *G. roseum* produces zearalenone-specific lactonase [20] which hydrolyses the lactone ring in ZEN, the mycotoxin of fusarium fungi dangerous for mammalians, thereby reducing ZEN toxicity [21]. The inhibition of most micromycetes by ZEN significantly reduces the possibility of using antagonistic fungi against toxic fusariums, which affect plants, but *G. roseum* is not sensitive to ZEN [20]. Like ZEN, aflatoxin B1 molecule contains the lactone ring which is associated with its toxicity, mutagenic and carcinogenic effects [22-25]. Aflatoxin-destroying activity that we have discovered in *G. roseum*, makes it even more attractive as a bioagent against fungi
that produce mycotoxins of different chemical nature.

Aflatoxin B1 (AFB1) biodegradation by the micromycetes Gliocadium roseum GRZ7 (4) and Trichoderma viride TV35 (2, 3) grown on the medium with aflatoxin, and intact (5) or heat inactivated (1) high molecular extracellular metabolites of G. roseum. Controls: for 2, 3, 4 — the AFB1 concentration in a nutrient medium not inoculated with micromycetes; for 1, 5 — the AFB1 amount immediately after its adding to analyzed fraction of metabolites (without incubation). Differences between the values in treated and control samples are statistically significant at P ≤ 0.05; the Y-errors bars show SEM for 3 experiments with 6 replicates per option in each.

The destroying activity of the studied T. viride strain was low. The utilization of AFB1 from the nutrient medium occurred slowly, and its concentration in the CL decreased not significantly as the fungus grew (see Fig., curve 2). However, the ability of the strain to destroy AFB1 in LC increased (curve 3) in case of using spore suspension of the fungus previously grown on the Czapek-CasH agar medium supplemented with AFB1 (0.9 μg/ml). When using such an inoculum, the proportion of the destructured toxin in the immersed culture increased by the end of the fermentation by 20 % (p = 0.03). Therefore, the AFB1-degrading activity which we detected in T. viride was inducible and could be increased by mutagenesis and selection of the most effective clones. Moreover, the tested T. viride strain proves to be an effective antagonist of A. flavus. When the toxigenic A11 strain was grown together with T. viride, the growth of the aflatoxin producer was significantly lower (Table) with a reduction in the colony size by 64 % on average.

**Radial growth of toxigenic Aspergillus flavus A11 colonies in single culture and under co-cultivation with Trichoderma viride TV35 (Tv) or Gliocadium roseum GRZ7 (Gr) (M±m)**

<table>
<thead>
<tr>
<th>Variant of cultivation</th>
<th>Average colony diameter, cm</th>
<th>Average colony area, cm²</th>
<th>Growth inhibition, % to control value</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>A11 (control)</td>
<td>6.90±0.03</td>
<td>7.10±0.00</td>
<td>39.50±0.00</td>
<td>0.005</td>
</tr>
<tr>
<td>A11 + Tv</td>
<td>3.33±0.09</td>
<td>4.26±0.06</td>
<td>14.25±0.56</td>
<td>63.9</td>
</tr>
<tr>
<td>A11 + Gr</td>
<td>6.20±0.06</td>
<td>6.89±0.08</td>
<td>37.27±0.12</td>
<td>5.6</td>
</tr>
</tbody>
</table>

Mycoparasite G. roseum did not noticeably influence the radial growth of A. flavus colonies as compared to the control, however, the differences were significant (Table), and in some cases the A11 strain colonies were overgrown with the G. roseum mycelium by the end of co-cultivation which could be a sign of hyperparasitism.

Thus, it has been found out that the studied strains of Tricho-derma viride and Gliocadium roseum, the antagonist species of many pathogenic fungi, competing with Aspergillus flavus for natural substrates, were different in their capacity to degrade the aflatoxin B1 in vitro and to inhibit the growth of aflatoxigenic A. flavus strain. Additionally, the inducibility of degrading activity, discovered by us, is of undoubted interest both in the theoretical and practical aspects. The obtained results can be used in the development of technologies for biological decontamination of feed contaminated with both aflatoxin B1 and its producer. In this case, G. roseum, apparently, can serve as a source of enzymes that decompose this mycotoxin, and T. viride can be a bioagent that inhibits the development of a producer.

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