Comparison of Soil Fungal Community Structure in Different Peanut Rotation Sequences Using Ribosomal Intergenic Spacer Analysis in Relation to Aflatoxin-Producing Fungi


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ABSTRACT


The present study focuses on determining soil fungal community structure in different peanut-cropping sequences by using a high-resolution DNA fingerprinting technique: ribosomal intergenic spacer analysis (RISA). This study was initiated to determine fungal community profiles in four peanut-cropping sequences (continuous peanut, 4 years of continuous bahiagrass followed by peanut, peanut-corn-cotton, and peanut-cotton rotations), with a special focus to evaluate whether the profiles under investigation may have also indicated microbial differences that could affect Aspergillus flavus populations. Results indicated 75% similarities among fungal communities from the same cropping sequences as well as with similar times of sampling. Polymerase chain reaction (PCR)-based detection of A. flavus directly from these soils was carried out using A. flavus-specific primers (FLA1 and FLA2) and also through quantitative estimation on A. flavus and A. parasiticus agar medium. Population levels of A. flavus in soil samples ranged from zero to \(1.2 \times 10^3\) CFU g\(^{-1}\) of soil (based on culturable methods); however, the fungus was not detected with A. flavus-specific primers. The minimum threshold limit at which these aflatoxin-producing fungi could be detected from the total soil genomic DNA was determined through artificial inoculation of samples with 10-fold increases in concentrations. The results indicated that a minimum population density of \(2.6 \times 10^6\) CFU g\(^{-1}\) of soil is required for PCR detection in our conditions. These results are useful in further determining the relative population levels of these fungi in peanut soils with other soil fungi. This is a new approach to understanding soil fungal communities and how they might change over time and under different rotation systems.

Additional keywords: aflatoxinogenic fungi.

Among different plant biotic stresses, diseases caused by soilborne fungal pathogens are important. The relative abundance of soilborne fungi in a given soil can play a defining role in the manifestation of soilborne diseases or may influence other organisms to induce plant diseases and, therefore, losses to crop production (4). Describing the soil microbial community in agricultural crops is an important step in understanding plant disease complexes and, ultimately, their management. Peanut is an important cash crop in the southeastern United States and is affected by many soilborne diseases as well as aflatoxin contamination. Aflatoxins, produced by Aspergillus flavus-group fungi, are known to be carcinogenic, hepatotoxic, and teratogenic (14,28). In soils in which peanut crops are produced, resident fungal communities vary in location and abundance (19,25). Determining the soil fungal community composition in peanut soils might be an important step not only in understanding disease complexes but also in the management of aflatoxigenic fungi, such as A. flavus-group fungi.

The relative occurrence of soilborne diseases is subject to change with different cropping histories (2). The rotation sequences that affect the relative disease occurrence could also affect the A. flavus group of fungi in peanut soils. However, short-term manipulation of a plant community through cropping sequence may not have a significant impact on soil microbial diversity (7). To arrive at a particular microbial community profile that would be beneficial to crop health, a long-term rotation or sequence of crops is desirable (12,22,26). Once a soil suppressive to preharvest aflatoxin contamination is achieved through such long-term crop rotations, then it is possible to concentrate on microbial interactions that make the soil suppressive to A. flavus populations. Investigations in crop–fungus interactions that reduced the A. flavus populations and subsequent identification of microbial groups that helps to reduce the population levels of these aflatoxin-producing fungi in peanut soils will be of much use in devising novel strategies for reduced risk in peanut.

Over recent years, it has become possible to investigate soil microbial community structure using culture-independent techniques. These methods include DNA fingerprinting techniques such as denaturing gradient gel electrophoresis, terminal restriction fragment length polymorphism, length heterogeneity polymerase chain reaction (PCR), ribosomal intergenic spacer analysis (RISA), cloning, and sequencing in addition to conventional microbial estimation through quantitative methods (6,21,27). These techniques provide rapid, lucid, reliable, and highly reproducible results for determining soil fungal community structures. These molecular techniques, RISA provides greater resolution and analysis of soil microbial diversity than other techniques (6). Use of RISA in determining fungal diversity in soils exploits the length polymorphism of the nuclear ribosomal DNA (rDNA) region that contains two internal transcribed spacers (ITSs) and the 5.8S rRNA gene (ITS1-5.8S-ITS2) (24). A better insight into these community structures with an efficient molecular technique such as RISA, with emphasis on aflatoxin-producing fungi, may...
provide a better understanding and subsequent management approaches to the aflatoxin problem.

In this study, the fungal communities from cropping sequences of peanut crops (P-P-P-P), 4 years of continuous bahiagrass followed by peanut (B-B-B-B-P), peanut-corn-cotton (P-Cn-Ct-P), and peanut-cotton (P-Ct-P-Ct) were compared. In addition to total fungal communities, the specific identification of *A. flavus* aflatoxin-producing fungi was addressed. Detection of *A. flavus* populations in soils was carried out using specific primers and through quantitative estimation. Quantification of minimum population loads of aflatoxin-producing fungi in soils, to determine levels needed for analysis of soil genomic DNA through molecular methods, was conducted.

**MATERIALS AND METHODS**

**Field site history and description.** The experimental site was located at the Wiregrass Research and Extension Center, Headland, AL. The site was established in 1988 with an objective of managing peanut diseases such as early leaf spot, late leaf spot, southern stem rot, and peanut root knot diseases (11). There are 34 peanut-cropping sequences at this site, of which four were selected for this study: P-P-P-P; B-B-B-B-P, a 5-year sequence that started most recently in 2003; Ct-P-Ct-P; and P-Cn-Ct-P (Table 1). The soil is Dothan fine sandy-loam (fine-loamy, siliceous, thermic Plinthic Palendults) type (<1% organic matter). The crops were supplemented with side-roll irrigation system as needed. The plots were arranged in a randomized complete block design with four replications.

**Soil sampling.** For each sample, five soil cores were collected from the pegging zone (up to 10-cm depth) during 2006, 2007, and 2008 from each plot, bulked, and mixed for a representative sample. Samples were collected three times: early season (June), midseason (August), and at peanut harvest (October). The samples were processed within 24 h for total soil genomic-DNA extraction and preserved genomic material was stored at −20°C for further downstream applications. The samples collected at the time of harvest were also subjected to enumeration of *A. flavus*-group populations as well as determining the minimum threshold population limit of the *A. flavus* group of fungi at which they are detectable in soils.

**DNA extraction and quantification.** Total genomic DNA was extracted from 10 g of each soil sample using the Power Mac Soil kit from MoBio (MoBio Laboratories Inc., Carlsbad, CA) following the manufacturer's instructions. Extracted DNA quality (260/280 ratio) and quantity was assessed by Nano Drop Spectrophotometer (Thermo Scientific). DNA dilutions were carried out to 5 ng µl⁻¹ and stored at −80°C for further downstream applications.

**Fungal community fingerprinting by RISA.** The length heterogeneity of the ITS1-5.8S-ITS2 region was exploited to characterize the fungal community. The primers used to amplify this region represent consensus sequences found at the 3’ end of the 18S genes in fungi (primer 2234C, 5’-ATATGCTTAAGTTCAGCGGGT-3’ and at the 5’ end of the 28S genes (primer 3126T, 5’-GTTTCCGTAGGGTTCCTAGCGAGCC-3’) (24). The reverse primer was labeled with IRD700 fluorochrome from Li-Cor (Lincoln, NE) at the 5’ end. Reaction mixtures (50 µl) for PCR contained 5 µl of 10× dilution buffer (20 mM Tris-HCl [pH 7.5]), 100 mM KCl, 15 mM MgCl₂, 1 mM dithiothreitol, 0.1 mM EDTA, 0.5% Tween 20 [vol/vol], 0.5% Nonidet P40 [vol/vol], and 50% glycerol [vol/vol], 2 µl of 10 mM dNTPs, 2 µl of 10 µM primer 2234C, 1 µl of 10 µM primer 3126T, 10 µl of 1 µM 2234C primer labeled with IRD700 fluorochrome from Li-Cor at the 5’ end, 0.2 µl of Taq polymerase (Promega Corp., Madison, WI), 10 µl of 25 mM MgCl₂, and 20 µl of template DNA (100 ng). PCR amplification was carried out in a Peltier Thermal Cycler (PTC-200, MJ Research) after a hot start at 94°C for 3 min; followed by 30 cycles consisting of 94°C for 45 s, 55°C for 1 min, and 68°C for 2 min; and final extension for 7 min at 68°C. Polyacrylamide gels were then prepared by mixing 30 µl of 5.5% Li-Cor polyacrylamide (KB plus-Li-Cor), 200 µl of 10% ammonium per sulfate, and 20 µl of N, N’, N”-tetramethylethylenediamine. The PCR product (5 µl) was then transferred to new tubes and, to this, 2.5 µl of stop buffer (Li-Cor, Blue Stop Solution) was added. The contents were thoroughly mixed and denatured at 98°C for 5 min before loading onto the polyacrylamide gel. The denatured PCR products (0.6 to 0.8 µl) were electrophoresed on the NEN Global Edition IR2 DNA Analyzer (Li-Cor) under denaturing conditions for 8 h at 1,500 V following the manufacturer’s instructions.

The RISA gel images were analyzed by the BIONUMERICS V. 5.0 (Applied Maths, TX) software program. Levels of similarity between fingerprints were calculated with the Pearson product-moment correlation coefficient (r) after the banding patterns were subjected to conversion, normalization, and background subtraction with mathematical algorithms. The program used binary data based on the presence or absence of particular band in the fingerprint profile pairs being compared. The dendrogram was constructed by using unweighted pair-group method with arithmetic mean. The multidimensional scaling analysis (MDSA) tool in BioNumerics was used to evaluate the similarities or dissimilarities between soil fungal communities. Cluster analysis was performed with the unweighted pair-group method using average linkages to determine the relatedness among the fungal communities of different peanut-cropping sequences (1).

**Enumeration and determination of minimum threshold population of *A. flavus* in soils.** Initially, the soil population levels of *A. flavus* and *A. parasiticus* were enumerated on *A. flavus* and *A. parasiticus* agar (AFPA) according to Pitt et al. (23). Soil samples with no traces of *A. flavus*-group fungi, as determined by plating on AFPA, were selected for this experiment. These *A. flavus*-free soil samples were augmented with *A. flavus* spore suspensions at different concentrations ranging from 2 × 10³ to 2 × 10⁸ CFU g⁻¹. Six soil samples (10 g each) were weighed and 5 ml of each spore concentration was added to individual soil samples in order to achieve *A. flavus* concentrations ranging from 2 × 10³ to 2 × 10⁸ CFU g⁻¹. The total genomic DNA was extracted immediately from these samples using the Power Mac Soil kit from MoBio following the manufacturer’s instructions. DNA was purified using protocol suggested by M. R. Liles (personal communication) for the determination of the minimum threshold level for *A. flavus* detection directly from the soil.

**PCR-based detection for *A. flavus* in soil.** The purified total soil genomic DNA was PCR amplified using the *A. flavus*-specific primers FLA1 (5’-GTAGGGTTCTAGCGAGCC-3’) and FLA2 (5’-GGAAAAGATGATTGAGCC-3’) (9). The concentration of *A. flavus* spore suspension that was added to the soil at which the fungal pathogen was detectable through PCR using *A. flavus*-specific primers was determined sequentially by increasing

<table>
<thead>
<tr>
<th>Crop rotation</th>
<th>2005</th>
<th>2006</th>
<th>2007</th>
<th>2008</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuous peanut (P-P-P-P)</td>
<td>Peanut</td>
<td>Peanut</td>
<td>Peanut</td>
<td>Peanut</td>
</tr>
<tr>
<td>Bahiagrass-peanut (B-B-B-B-P)</td>
<td>Bahiagrass</td>
<td>Peanut</td>
<td>Peanut</td>
<td>Peanut</td>
</tr>
<tr>
<td>Peanut-cotton (Ct-P-Ct-P)</td>
<td>Cotton</td>
<td>Peanut</td>
<td>Cotton</td>
<td>Peanut</td>
</tr>
<tr>
<td>Peanut-corn-cotton (P-Cn-Ct-P)</td>
<td>Peanut</td>
<td>Corn</td>
<td>Cotton</td>
<td>Peanut</td>
</tr>
</tbody>
</table>
or decreasing the spore concentration as needed. PCR reactions were performed in the Peltier Thermal Cycler (PTC-200; MJ Research). The PCR amplification protocol for *A. flavus* detection was as follows: 1 cycle of 5 min at 95°C; followed by 26 cycles of 30 s at 95°C, 30 s at 58°C, and 45 s at 72°C; and a final extension of 5 min at 72°C. The PCR products were run on 1% agarose gel electrophoresis and subjected to ethidium bromide staining.

Statistical analysis. Data were analyzed using analysis of variance (SAS Institute, Cary, NC). The PROC GLM procedure was used to separate the mean population levels of *A. flavus* over 3 years.

**RESULTS**

**RISA analysis of fungal communities.** Fungal RISA profiles showed similarities among communities from replicate plots of the same crop rotation (data not shown). In general, similarities of 55 to 75% were observed among plots of different cropping sequences in 2006 to 2007. Common bands were observed among all samples regardless of sampling time and cropping sequence. RISA banding patterns in P-P-P-P indicated that there was ≈75% similarity in these plots sampled in October 2006. In the case of B-B-B-B-P, 70% similarity was observed in plots sampled during June 2006. The banding patterns from these same plots showed a marginal increase in similarity as the months advanced; for instance, in BT-P, for the plots sampled in August 2007, ≈80% similarity was noticed, whereas the plots sampled in August and October 2006 had ≈75%. On the other hand, P-Cn-Ct-P rotation plots sampled in August 2006 clustered at 85% similarity and at ≈80% when sampled during August 2007.

MDSA of RISA community profiles revealed that fungal communities showed greater similarity based on the month of sampling than based on cropping sequence (Fig. 1A). This can be clearly seen in the tighter grouping of the August 2006 observations and few outliers in the case of other groupings. Lesser similarities were observed with respect to fungal composition cropping sequence (Fig. 1B), indicating that the impact of cropping sequence on soil fungal diversity is minimal.

**Enumeration and PCR-based detection of soil *A. flavus*.** *A. flavus* soil populations, determined at the time of harvest from each cropping sequence, indicated a marginal increase in population levels of *A. flavus* within cropping sequences of P-P-P-P and in Ct-P-Ct-P rotation. The population levels in these cropping sequences ranged from 1 × 10^3 to 1.2 × 10^3 CFU g⁻¹ of soil through 2006 to 2008. Populations remained constant in the P-Cn-Ct-P rotation through 2006 to 2008. A substantial decline in soil *A. flavus* populations was observed in the B-B-B-B-P cropping sequence in 2006 to 2008 (0.7 × 10^3 to 0.2 × 10^3 CFU g⁻¹ of soil) (Table 2).

Comparisons of mean *A. flavus* population levels over 3 years indicated that, among the four cropping sequences, *A. flavus* populations were significantly less in soils from B-B-B-B-P (0.2 × 10^3 CFU g⁻¹ of soil) compared with other treatments. The population levels in P-P-P-P, Ct-P-Ct-P, and P-Cn-Ct-P sequences were 1.2 × 10^3, 1.2 × 10^3, and 1.0 × 10^3 CFU g⁻¹ of soil, respectively; and no significant differences were observed among these three cropping sequences (Table 2). The minimum threshold limit at which *A. flavus* can be detected in peanut soils directly from soil genomic DNA with *A. flavus*-specific primers was found to be 2.6 × 10⁶ CFU g⁻¹ (Fig. 2).

**DISCUSSION**

Fungal RISA profiles with common bands in all cropping sequences in the present study may be attributed to the presence of common saprophytes in these soils. However, the phylogenies of these saprophytes in soils were not determined. The effect of particular plant species on the selection of resident soil microbial communities is well documented (7,8,10,29). This influence is due to differential root exudates into the rhizosphere. For this reason, the microbial diversity of a rhizosphere and nonrhizosphere soil differ greatly (15). In our study, however, sampling time had a more profound effect on similarities of these communities compared with crop. One probable explanation for this is that the experimental site is in a long-term rotation, and crop-management practices such as tillage, irrigation, and soil nutrient application year after year might be masking the effect of plant type on the soil microbial communities (5). Another reason could be that, at each sampling time, at least two of the four rotations had the same crop (e.g., peanut in two treatments in 2006). MDSA also revealed that there was no selection of fungal communities based on plant type because this analysis enables grouping of fungal communities based on presence or absence of bands and their intensity.

Overall, our results suggest that, though different RISA groups were detected based on sampling times and cropping sequences with peanut, a specific fungal community associated with these sequences could not be confirmed concretely. To ascertain the phylogeny of these fungal communities in different cropping sequences, 18S rDNA sequencing of fungi from the soil of these cropping sequences is an alternative for better understanding of the fungal communities associated with these crops in a peanut ecosystem.

Our study on the detection of aflatoxin-producing fungal populations in soils revealed that PCR-based methods using *A. flavus*-specific primers was found to be 2.6 × 10⁶ CFU g⁻¹ (Fig. 2).

**TABLE 2. Population levels of* Aspergillus flavus* group in soils from different peanut cropping sequences**

<table>
<thead>
<tr>
<th>Soil <em>A. flavus</em> population at harvest (x10³ CFU g⁻¹ of soil)</th>
<th>Rotation</th>
<th>2006</th>
<th>2007</th>
<th>2008</th>
<th>Mean*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuous peanut</td>
<td>1.0</td>
<td>1.0</td>
<td>1.2</td>
<td>1.1</td>
<td>a</td>
</tr>
<tr>
<td>Bahiagrass-peanut</td>
<td>0.7*</td>
<td>0.0</td>
<td>0.2*</td>
<td>0.3</td>
<td>b</td>
</tr>
<tr>
<td>Peanut-cotton</td>
<td>1.0*</td>
<td>1.1*</td>
<td>1.2</td>
<td>1.1</td>
<td>a</td>
</tr>
<tr>
<td>Peanut-corn-cotton</td>
<td>1.0*</td>
<td>0.8*</td>
<td>1.0</td>
<td>0.9</td>
<td>a</td>
</tr>
</tbody>
</table>

*Mean over 3 years; means followed by the same letter are not significantly different and the treatment means were separated using least significant difference (P = 0.01).

* Plots were planted to bahiagrass.

* Plots were planted to cotton.

* Plots were planted to corn.

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Fig. 2. Polymerase chain reaction with *Aspergillus flavus*-specific primers. Lane M = 100-bp marker, lane N = negative control, and lanes 1 to 6 are soil samples inoculated with *Aspergillus flavus* spores at 10⁴ to 10⁶ CFU/g of soil.
flavus-specific primers could not detect these fungi even at a concentration of 10⁴ CFU g⁻¹ soil. There are several possible reasons for this. One reason might be the presence of high amounts of PCR inhibitors in the soil metagenomic DNA such as humic acids (31). Separation of humic substances from a DNA sample is always a critical phase because both of these are acid macromolecules. Another reason could be that the DNA extraction method used in the present study might affect the yields of DNA from A. flavus and, subsequently, the obtained DNA might be less than the primer’s sensitivity limits. However, other extraction methods that might yield more A. flavus DNA can be more specific in identifying these populations from soils. Further studies on how to improve PCR-based detection of A. flavus from soil would be beneficial.

Soil enumeration studies indicated that using bahiagrass in rotation with peanut resulted in lower A. flavus population levels compared with other cropping sequences. Though we did not dig deeper in our study as to why bahiagrass rotations did not favor the A. flavus populations, earlier reports indicated that sod-based rotations, including bahiagrass, improved soil health parameters by improving soil water infiltration rates, allowing greater nutrient uptake, increasing earthworm population densities, increasing the abundance of beneficial microorganisms, and, ultimately, increasing organic matter content (16–18,20). Bahiagrass rotations might have contributed to soil organic matter at our study site, thereby increasing the antagonistic microflora and causing significant reductions in A. flavus population levels. In addition, in bahiagrass rotations in our study, more bands were observed in fingerprinting profiles (data not shown), indicating that an increase in soil microbial diversity is another contributing factor in lowering the A. flavus populations.

High population levels of A. flavus in P-P-P-P, Ctn-P-Ctn-P, and P-Ctn-P-Ctn-P rotations compared with the rotation that included bahiagrass, improved soil health parameters by improving soil water infiltration rates, allowing greater nutrient uptake, increasing earthworm population densities, increasing the abundance of beneficial microorganisms, and, ultimately, increasing organic matter content (16–18,20). Bahiagrass rotations might have contributed to soil organic matter at our study site, thereby increasing the antagonistic microflora and causing significant reductions in A. flavus population levels. In addition, in bahiagrass rotations in our study, more bands were observed in fingerprinting profiles (data not shown), indicating that an increase in soil microbial diversity is another contributing factor in lowering the A. flavus populations.

We thank O. Olivares-Fuster of Fisheries and Allied Aquacultures department for his assistance in this project.

LITERATURE CITED
