Introduction

In 2002 the National Program for the Augmentation of Forest Cover in Poland was revised. The Program anticipates afforestation of 1.5 million ha of marginal land (including post-agricultural and fallow land) and an increase in forest cover to 30% by 2020 and 33% by 2050. Considering soil and weather conditions, this marginal land is being planted with Scots pine (*Pinus sylvestris* L.), which is the dominant forest tree species in Poland (71%).

**Heterobasidion** species are considered to be the most economically important conifer forest pathogens in the northern hemisphere [1, 2]. Scots pine becomes seriously damaged by *Heterobasidion annosum* (Fr.) Bref. [3]. The fungus spreads through aerial basidiospores to stump surfaces and wounds, and by mycelia via root contacts from tree to tree. It causes root and butt rot, growth loss, and mortality of trees. In favourable conditions seedlings can become infected very soon after planting. In Europe and the USA *Heterobasidion* is responsible for the loss of $1.5-2 billion annually.

Original Research

Effect of Scots Pine Sawdust Amendment on Abundance and Diversity of Culturable Fungi in Soil

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Abstract

Scots pine and birch were planted in soils left fallow for 3, 6, or 15 years after arable cropping. We investigated the effects of Scots pine sawdust amendment, applied a year before planting, on the abundance and diversity of culturable soil fungi 4, 14, and 16 years after treatment. The treatment was intended to increase populations of fungi antagonistic to *Heterobasidion* and soil suppressiveness to tree root pathogens. Effects of treatment on fungal abundance were inconsistent; general or local, seasonal or continuous decreases or increases, often significant, were observed. There were, however, significant and continuous increases in frequency of antagonistic *Clonostachys* + *Trichoderma* and the mycorrhizal fungus *Oidiodendron* in treated soils compared with the control in all three fallow areas. Local and seasonal decreases in frequency of *Penicillium* + *Talaromyces*, *Pseudogymnoascus*, and entomopathogenic and nematophagous species were observed in treated soils. Abundance of fungi was moderately and negatively correlated with soil pH ($R^2 = -0.61$, $P<0.0001$). Abundance of *Clonostachys* + *Trichoderma* was moderately and positively correlated with mean annual temperature and positively correlated with total annual rainfall. Fresh sawdust, even applied undecomposed and without added mineral N (to aid microbial decomposition and plant growth), may be beneficial in sandy soil.

Keywords: *Heterobasidion*, *Oidiodendron*, organic matter, soil fungi, *Trichoderma*

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Damage is generally greater in the first or second generations of trees grown on post-agricultural or fallow land, which has the least organic matter; a deficiency in nitrogen, potassium, phosphorus, and magnesium; higher pH; increased soil compaction below the plough level; and poor capillarity for water [4]. The shortage of lignified tissues and a specific carbon and energy source for microbiota creates an unfavourable microbiological habitat and causes deficiency of microorganisms that are potentially competitive and antagonistic toward H. annosum and helpful in forest site-forming processes.

The aim of our study was to evaluate the abundance, diversity, and temporal and spatial distribution of culturable fungi in soils left fallow for different lengths of time, then amended with fresh Scots pine sawdust. The hypothesis was that the organic matter amendment would change the physical and chemical properties of the soil, which would in turn initiate and stimulate microbiological shifts that would increase the population of fungal antagonists of Heterobasidion and suppressiveness of soil to root pathogens. Studies made 4, 14, and 16 years after treatment were intended to show the temporal dynamics and diversity of microorganisms.

This study is part of a larger project on development and promotion of non-chemical methods of pest management as part of an integrated pest management strategy.

Material and Methods

Site Description, Treatments, and Sampling

The experiment was established in three areas (each 300 m², 1 km apart), that had been left fallow for 3, 6, and 15 years in Głęboki Bród Forest District in northeast Poland (geographic coordinates 53.983333, 23.3). In October 1995, four treatment plots and four control plots (each 25 m²) were designated at the four corners of each area. The first soil samples were collected in October 1995; six soil cores (400 g each) were taken from each corner of each plot. The upper and lower 5 cm of each core were discarded and a subsample (0.26 mg) of the soil-sand mixture was put into a petri dish and covered with liquid (50°C) Johnson-Martin’s agar (JMA). Thirty replicates were made from the three flasks for each treatment and each control. A subsample (0.26 mg) of the soil-sand mixture was put into a petri dish and covered with liquid (50°C) Johnson-Martin’s agar (JMA). Thirty replicates were made from the three flasks for each treatment and each control. All plates were incubated for 20 days at 25°C. All colonies on each plate were examined macro- and microscopically and distinguished on the basis of color, growth rate, hyphal characteristics, and sporulation. Colonies of each species were counted and representatives of fungi were identified by morphotyping on potato dextrose agar (PDA) and synthetic nutrient agar (SNA). Aspergillus and Penicillium species were identified on Czapek yeast autolysate agar (CYA) and 2% malt extract agar (MEA). Identification was made using mycological keys [5-8].

Abundance of fungi was defined as the number of colony forming units (cfu) in a sample. Frequency of an individual species or group of species was defined as proportion (%) of isolates in the total number of isolates. Diversity was defined as the number of species in a sample. A species, or group of related species of fungi, was considered as:

i) eudominant, with frequency >10%
ii) dominant, with frequency 5-10%
iii) subdominant, with frequency 2-5%
iv) recedent, with frequency 1-2%
v) subrecedent, with frequency <1% (in at least one treatment).

Fungi in the main dominance class were often grouped on the basis of their morphology and taxonomy.

Isolation and Identification of Fungi

The classical soil-dilution method was applied for qualitative and quantitative analyses of fungal communities in 1995 (when nucleic acid sequencing was not easily accessible), and 4, 14, and 16 years later (for direct comparison with results from 1995). The affiliation, activity, and microbiological interactions of many culturable soil fungi are already known and can be used for evaluation of the possible microbiological status of soil (i.e., comparing soil qualities and properties).

A subsample (1 g) of soil from each treatment was diluted in 149 g of sterile quartz sand in a flask, in a safety cabinet. Three replicates (= three flasks) of soil-sand mixture were prepared for each treatment and each control. A subsample (0.26 mg) of the soil-sand mixture was put into a petri dish covered with liquid (50°C) Johnson-Martin’s agar (JMA). Thirty replicates were made from the three flasks for each treatment and each control. All plates were incubated for 20 days at 25°C. All colonies on each plate were examined macro- and microscopically and distinguished on the basis of color, growth rate, hyphal characteristics, and sporulation. Colonies of each species were counted and representatives of fungi were identified by morphotyping on potato dextrose agar (PDA) and synthetic nutrient agar (SNA). Aspergillus and Penicillium species were identified on Czapek yeast autolysate agar (CYA) and 2% malt extract agar (MEA). Identification was made using mycological keys [5-8].

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Fungi in the main dominance class were often grouped on the basis of their morphology and taxonomy.
Table 1. Chemical properties of soils left fallow for different durations before (1995) and after (1999, 2009, 2011) sawdust amendment and tree planting.

<table>
<thead>
<tr>
<th>Soil characteristics</th>
<th>3 years fallow</th>
<th>6 years fallow</th>
<th>15 years fallow</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH in KCl</td>
<td>3.8 5.0 4.9 4.85* 4.92* 5.23 4.33</td>
<td>6.0 4.3 4.2 4.89* 4.15* 4.3 4.0</td>
<td>4.1 4.39* 4.18* 4.1 4.1</td>
</tr>
<tr>
<td></td>
<td>F=16.56 P=0.0066</td>
<td></td>
<td>F= 8.03 P=0.0471</td>
</tr>
<tr>
<td>pH in H2O</td>
<td>4.4 6.1 6.1 5.9 6.1 5.8 5.2</td>
<td>7.0 5.6 5.3 5.9 5.1 5.0 4.7</td>
<td>5.1 5.5 5.1 4.9 4.7</td>
</tr>
<tr>
<td></td>
<td>F=6.11 P=0.0484</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organic carbon (%)</td>
<td>0.87 1.48 1.22 1.11 0.93 1.16 1.53</td>
<td>0.98 1.81 1.49 1.76 1.12 2.03 2.02</td>
<td>1.20 1.89 1.27 2.39 2.04</td>
</tr>
<tr>
<td></td>
<td>F=0.8968 P=0.0032</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total nitrogen (%)</td>
<td>0.059 0.065 0.061 0.078 0.059 0.069 0.066</td>
<td>0.071 0.083 0.082 0.095 0.079 0.083* 0.098*</td>
<td>0.091 0.1 0.094 0.083* 0.088*</td>
</tr>
<tr>
<td></td>
<td>F=26.47 P=0.0358</td>
<td></td>
<td>F=25.00 P=0.0377</td>
</tr>
<tr>
<td>Organic nitrogen (%)</td>
<td>0.058 0.064 0.060 0.077 0.058 0.068 0.065</td>
<td>0.070 0.082 0.081 0.094 0.078 0.082 0.097</td>
<td>0.090 0.1 0.093 0.082 0.087</td>
</tr>
<tr>
<td></td>
<td>F=1.11 P=0.0484</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C:N</td>
<td>14.7 22.8 20.0 14.2 15.8 16.8 23.1</td>
<td>13.8 21.8 18.2 18.5 14.2 24.5 20.6</td>
<td>13.2 18.9 13.5 28.7 23.2</td>
</tr>
<tr>
<td>Extractable phosphorus P2O5 (mg/100 g)</td>
<td>10.7 3.2 2.2 3.7 3.1 6.4 5.1</td>
<td>2.4 2.8 2.5 3.9 3.7 6.7 5.5</td>
<td>1.1 3.8 3.5 3.2 3.3</td>
</tr>
<tr>
<td>Extractable potassium (mg/100 g)</td>
<td>1.2 3.0 2.4 3.8 4.1 4.8 3.7</td>
<td>5.0 2.3 2.5 5.1 3.5 4.3 4.3</td>
<td>4.0 2.6 1.4 3.7 2.9</td>
</tr>
<tr>
<td>Extractable magnesium (mg/100 g)</td>
<td>2.5 2.6 2.2 2.0 2.9 1.9 0.7</td>
<td>92.2 1.8 1.2 3.2 1.1 1.2 0.6</td>
<td>2.5 1.9 0.7 0.5 0.3</td>
</tr>
<tr>
<td>Extractable calcium (mg/100 g)</td>
<td>18.5 57.0 52.2 45.9 59.7 40.1 16.3</td>
<td>143.5 31.0 24.0 62.3 17.6 13.8 5.1</td>
<td>32.5 39.7 14.3 10.6 6.7</td>
</tr>
<tr>
<td>Extractable sodium (mg/100 g)</td>
<td>1.5 0.9 1.0 0.3 0.3 0.3 0.4</td>
<td>1.5 0.8 1.0 0.2 0.2 0.4 0.4</td>
<td>1.5 0.3 0.3 0.5 0.5</td>
</tr>
</tbody>
</table>

S – sawdust application  
C – control  
*a statistically significant difference between sawdust application (S) and control (C) in an individual year according to one-way ANOVA.
Statistics

Differences in soil chemical properties were analysed with one-way analysis of variance (ANOVA) using Statgraphics™ Centurion (Statpoint Technologies, Inc. Warrenton, VA, USA). Abundance and diversity of fungi were analysed with $X^2$ test. Diversity within and between fungal communities was compared using diversity indices calculated for each community [9]. Species richness was indicated by the total number of species in the community, Margalef’s index (DMg), Shannon’s diversity index ($H'$), and Simpson’s diversity index (D). Evenness and dominance were indicated by Shannon’s evenness index (E) and Berger-Parker’s index (d). Similarity between fungal communities in two systems was determined by calculating the qualitative Sorensen’s similarity index (CN). Relationships between soil chemical properties, temperature, rainfall, and abundances of fungi were estimated with Pearson’s correlation coefficient.

Results

Changes in Chemical Properties of Soils after Sawdust Amendment

In 1995 the soil left fallow for three years had a higher content of extractable phosphorus than the other soils, and the soil left fallow for six years had higher pH and higher content of extractable potassium, magnesium, and calcium (Table 1). Sawdust amendment usually resulted in slightly higher (usually not significant) pH, percentages of organic carbon and total and organic nitrogen, C:N ratio, and content of extractable phosphorus, potassium, magnesium, and calcium than in the control. Large differences between sites in 1995 tended to disappear after four years.

Fungal Community Structure

Totals of 221 species of fungi and three species of Oomycetes were recorded in the non-rhizosphere soil in all samples (Table 3). Fungi included mostly representatives of Ascomycota. Zygomycota (17 species) were rare and Basidomycota (1 species) occurred sporadically. Seven groups of fungi occurred with frequency $>10\%$ (eudominants) in at least one treatment. Eudominants, dominants, subdominants, recedents, and subrecedents included 12, 25, 46, 46, and 92 species, respectively.

Effects of sawdust amendment on abundance of fungi were not consistent. Abundances were less (significant or non-significant) in amended than in control plots in soil left fallow for three years (1999 sample) or six years (all samples), and were greater in amended plots in soil left for three years (2009 and 2011 samples) or 15 years (all samples). There was a significant interaction between duration of fallow and year of sample on abundance of fungi ($\chi^2$ test, $P\leq0.001$ or $P\leq0.05$).

The most common fungi before sawdust amendment included Penicillium admetezii, P. janczewskii, P. jenseni, Pseudogymnoascus roses, and Purpureocillium lilacinum. Frequencies of Oidiodendron and Trichoderma were greater (usually significantly) after sawdust amendment than in the control. Frequencies of Oidiodendron tended to increase throughout the experiment. Frequencies of Trichoderma tended to decrease after 1999. Trichoderma hamatum and T. harzianum were the most frequent species of Trichoderma. Local or seasonal effects of sawdust amendment on frequencies were observed for Acremonium + Sagenomella spp.; they were significantly more frequent than in controls after three years of fallow in 1999 sample, and after six years of fallow in the 2011 sample. Frequency of Penicillium + Talaromyces spp. was often significantly less after sawdust amendment throughout the experiment and tended to decrease over time. There were local or seasonal effects of sawdust amendment on frequency of Zygomycota (Absidia + Cunninghamella + Mortierella + Mucor + Umbelopsis spp.); they were significantly less frequent than in controls after six years of fallow in the 2009 and 2011 samples, and after 15 years of fallow in the 2009 sample. Pseudogymnoascus spp. varied between samples and was sometimes less frequent after sawdust amendment than in the control: after three years of fallow in the 1999 sample, after six years of fallow in the 2009 sample, and after 15 years of fallow in the 2011 sample. Entomopathogenic and nematophagous species were sometimes less frequent after sawdust amendment.

Diversity in the communities (measured as number of species) was often less after sawdust amendment than in controls but by only non-significant amounts. The relatively small number of fungal taxa and the infrequent occurrence of many taxa soon after treatment resulted in relatively small diversity indices based on species richness (DMg) and proportional abundance of species ($H'$). Generally, the greatest diversity occurred in the 1999 sample and in soil left fallow for three years (Table 4). Diversity decreased over time in soils that had been left fallow for three or 15 years.

Rare dominance of single taxa resulted in higher values for Shannon’s evenness index (E) and often lower values for Simpson’s dominance index (d). Generally, communities were more evenly distributed than in the control in soil left fallow for three years, in the 2009 sample, and in sawdust-treated plots. Sorensen’s qualitative similarity index (CN), when used for comparing fungal communities from treated plots and control in a particular year, showed that most similarity in species composition occurred in soils left fallow for six or 15 years and sampled in 2009.

There was moderate negative correlation between the abundance of fungi and soil pH ($R^2=0.61$, $P<0.0001$), and moderate positive correlation between the abundance of Trichoderma and mean annual temperature ($R^2=0.45$, $P<0.0001$). Stronger correlation was observed between abundance of Trichoderma and total annual rainfall ($R^2=0.85$, $P<0.0001$).
Table 2. Temperature and rainfall at Głęboki Bród.

<table>
<thead>
<tr>
<th></th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean seasonal (IV-X)</td>
<td>12.9°C</td>
<td>13.4°C</td>
<td>12.6°C</td>
<td>13.5°C</td>
</tr>
<tr>
<td>Mean annual temperature</td>
<td>6.8°C</td>
<td>7.7°C</td>
<td>6.8°C</td>
<td>7.3°C</td>
</tr>
<tr>
<td>Total seasonal (IV-X)</td>
<td>392 mm</td>
<td>418 mm</td>
<td>411 mm</td>
<td>461 mm</td>
</tr>
<tr>
<td>Total annual rainfall</td>
<td>584 mm</td>
<td>605 mm</td>
<td>597 mm</td>
<td>589 mm</td>
</tr>
<tr>
<td>Hydrothermal coefficient(K)</td>
<td>1.4</td>
<td>1.63</td>
<td>1.73</td>
<td>1.47</td>
</tr>
</tbody>
</table>

*1961-90, **1971-2000

Discussion

Scots pine sawdust has been used to improve forest nursery conditions in the production of containerized or bare-root tree seedlings. Sawdust that had been allowed to age was shown (with delayed nitrogen application) to benefit conifer seedlings [10], possibly favoring the growth of antagonists of soil-borne pathogens. Sawdust applied beneath Scots pine roots was thought to have contributed to better development of the assimilation organs [4]. Sawdust improves the physical properties of soil and decreases the toxicity of some soil chemical contaminants [11].

Many nurseries, particularly those using bare-root propagation, have used pine sawdust with success. However, inappropriate application of sawdust can be dangerous, and effects of sawdust on soil conditions – including chemical and microbiological properties – need to be fully investigated.

The fungi detected in the present studies are the main organic matter decomposers in acidic forest soils. Growth-based measurements have revealed fivefold less bacterial growth and fivefold more fungal growth in acidic forest soils (pH=4.5) compared to agricultural alkaline soils (pH=8.3), indicating approximately 25-30-fold greater fungal importance in the former [12]. All fungi detected in the present studies were classified as eudominants, dominants, subdominants, recedents, or subrecedents to indicate the scale of their functional potential.

The study showed that amending fallow soil with sawdust often caused either general or local, and seasonal or long-term decreases in the abundance and diversity of the soil fungi. Since similar decreases were also often observed in the control (only ploughed) plots the inhibiting effect of ploughing itself cannot be excluded. Tillage, including ploughing, disrupts hyphal networks and decreases hyphal viability, fungal reproduction and colonization of plant roots [13-15].

Generally, the decrease in abundance of soil fungi can also result from immobilization of nitrogen. Scots pine sawdust has only 0.2% nitrogen and a very high C:N ratio. Wood decomposers naturally occurring in soil require materials with 1-2% of nitrogen to proceed with degradation. Finding only 0.2% in sawdust, they utilize inorganic soil nitrogen. This leads to a shortage of inorganic soil nitrogen and less decomposition of organic residues with high C:N ratio, and this is followed consequently by decreased abundance of mycobiota. However, total nitrogen content was only occasionally slightly less in amended soils (left fallow for six or 15 years; 2011 sample) than in the non-amended control, which suggests that N immobilization was not a main reason for decreased fungal populations after treatment.

Scots pine sawdust has high cellulose (55%), and less lignin (30%) and hemicellulose (10%) content. Cellulolysis provides carbohydrates (the breakdown products) shortly after sawdust incorporation into soil. Theoretically, the abundance of carbohydrates should have stimulated the growth and reproduction of microorganisms [16]. However, the communities described here only rarely behaved according to such a scheme. This may be explained by short- or long-term suppression of certain species of fungi by:

i) alkalization of the treated soils
ii) higher concentration of CO₂ produced during decomposition of sawdust
iii) toxic compounds present in sawdust or produced during their degradation (i.e., soluble resins, tannins, terpenes, and biologically active phenolic monomers leached from wood particles during the winter)
iv) antibiotics produced by actinomycetes stimulated by sawdust
v) competition by bacteria that are the predominant degraders of lignocellulosic detritus
vi) competition by the dominant fungal species, including Trichoderma.

Trichoderma and Oidiodendron were more abundant in treated soils despite slight alkalinization caused by the sawdust amendment. A similar level of alkalinization was observed in two soil types, i.e., in sand (left fallow for three or six years) and lightly clayed sand (left fallow for 15 years) despite the greater buffering capacities of clayed soils [17]. The acidity of the soils after treatment was, however, still in the range favoured by Trichoderma (pH 5.5-7.50) [18]. The greater abundance of Trichoderma was often associated with smaller populations of Zygomycota and P. roseus. This may be partly because of their poor ability to decompose cellulose and hemicellulose.
<table>
<thead>
<tr>
<th>No.</th>
<th>Taxon</th>
<th>3 years fallow</th>
<th>6 years fallow</th>
<th>15 years fallow</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>S</td>
<td>C</td>
<td>S</td>
</tr>
<tr>
<td>1.</td>
<td><em>Humicola</em> spp. (&lt;sup&gt;1&lt;/sup&gt;)</td>
<td>2.6</td>
<td>0*</td>
<td>10.5*</td>
</tr>
<tr>
<td>2.</td>
<td><em>Absidia</em> + <em>Cunninghamella</em> + <em>Mortierella</em> + <em>Muco</em> + <em>Umbelopsis</em> spp.&lt;sup&gt;2&lt;/sup&gt;</td>
<td>9.2</td>
<td>8.1</td>
<td>6.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.6*</td>
<td>1.3*</td>
<td>8.0*</td>
</tr>
<tr>
<td>3.</td>
<td><em>Oidiodendron</em> spp.&lt;sup&gt;3&lt;/sup&gt;</td>
<td>1.0</td>
<td>2.6</td>
<td>6.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4.</td>
<td><em>Penicillium</em> + <em>Talaromyces</em> spp.&lt;sup&gt;4&lt;/sup&gt;</td>
<td>44.4</td>
<td>20.3</td>
<td>18.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>7.7*</td>
<td>18.2*</td>
<td>14.1</td>
</tr>
<tr>
<td>5.</td>
<td><em>Pseudogymnoascus</em> spp.&lt;sup&gt;5&lt;/sup&gt;</td>
<td>10.0</td>
<td>2.6</td>
<td>6.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>6.</td>
<td><em>Trichoderma</em> spp.&lt;sup&gt;6&lt;/sup&gt;</td>
<td>1.2</td>
<td>17.0*</td>
<td>2.5*</td>
</tr>
<tr>
<td>7.</td>
<td>Entomopathogenic and nematophagous species&lt;sup&gt;7&lt;/sup&gt;</td>
<td>0.3</td>
<td>0</td>
<td>3.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

**Eudominants (frequency > 10% in at least one treatment)**

**Frequency of eudominants**

69.2 | 48.7 | 48.6 | 82.4 | 72.1 | 87.1 | 90.8 | 51.8 | 94.5 | 87.3 | 86.4 | 86.7 | 77.7 | 88.5 | 71.9 | 85.7 | 86.6 | 85.0 | 90.4

**Dominants (frequency 5-10% in at least one treatment)**

**Frequency of dominants**

10.3 | 32.7 | 12.9 | 15.3 | 17.3 | 4.1 | 6.3 | 20.4 | 4.2 | 5.1 | 7.7 | 4.1 | 8.9 | 2.4 | 18.1 | 8.1 | 3.9 | 5.4 | 2.5
<table>
<thead>
<tr>
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<th>3 years fallow</th>
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<tr>
<td></td>
<td></td>
<td>S</td>
<td>C</td>
<td>S</td>
</tr>
</tbody>
</table>

Subdominants (frequency 2-5% in at least one treatment)

17. Alternaria sp. + Lewia sp.  
18. Aureobasidium pullulans (de Bary) G. Amadu  
19. Cadophora fastigiata Lagerb. & Melin  
20. Cladosporium spp.  
22. Chaetomium spp.  
23. Cylindrocarpon spp.  
25. Phialocephala botulispora (Cole & W.B. Kendr.) Grünig & T.N. Sieber  
27. Phoma spp. + Pyrenochaeta sp.  
28. Tolyphodium geodes W. Gams  
29. Truncatella truncata (Lév.) Steyaert  

**Frequency of subdominants**

**Frequency of recedents + subrecedents** (listed below)

**Abundance** – number of colony forming units in a sample

**Diversity** – number of species in a sample

S – sawdust application  
C – control  
*a statistically significant difference between sawdust application (S) and control (C) in an individual year according to χ² test at P ≤ 0.001 or at P ≤ 0.05.  
**belongs to Oomycota**
Table 3. Continued.

Explanations:

Humbicola fuscaotra Traen + H. grisea Traen


Oidiobolus cladosporioides Moraill + O. echinulatum G.L. Barron + O. griseum Robak + O. tenuissimum (Peck) S. Hughes + O. truncatum G.L. Barron


Paecilomyces marquandii (Massee) S. Hughes + P. cf. victoriae (Sylv.) A.H.S. Br. & G. Sm. + P. varioti Bainier

Sarocladium glaucum (W. Gams) Summerb. + S. strictum (W. Gams) Summerb. + Alternaria alternata (Fr.) Keissler + Leve illa infectoria (Fuckel) M.E. Barr & E.G. Simmons

Cladosporium cladosporioides (Fresen.) G.A. de Vries + C. herbarum (Pers.) Link

Clonostachys cardelabrum (Bonord.) Schroers + C. rosea (Link) Schroers, Samuels, Seifert & W. Gams

Chaetomium fucicola Cooke + Ch. homopilatum Omvick + Ch. truncatum sp.

Cylindrocarpon magnusianum Wollenw.+ Cylindrocarpon sp.


Philophora alba J.F.H. Beyma + P. bakubak (Laxa) Schol-Schwarz + P. cyclanimis J.F.H. Beyma


Subrecedents (frequency < 1% in at least one treatment) – Ascomycota spp., Basidiomycota spp., Bipolaris sorokiniana (Sacc.) Schoemaker, Boeberia exigua (Desm.) Eaveskamp, Gruyter & Verkley, Cladosphalophora caespitosa (Grove) Crous & Arzlanlou, Exophiala mansonii (Castell.) de Hoog, Fusicola aquaeductuum (Radlik. & Rabenh.) Gräfenhan, Seifert & Schroers, Geotrichum candidum Link, Gibellulopsis nigrescens (Pehlybr.) Zare, W. Gams & Summerb., Gliomastix macrospora Moustafah, Hormonema sp., Hyalosporaceae sp., Lasiadlae lasiosphaericae (W. Gams) Rábola & W. Gams, Leptosphaeria coniothyriomium (Fuckel) Sacc., Mariannae elegans (Corda) Samson, Mennonella echinata (Riv.) Galloway, Microdochium bolleyi (W. Gams & M. druidicale (Riv.) Galloway, Moniliforme arbusculum (Sopp.) Samson, N. Yilmaz, Frisvad + C. tropicum (Wallr.) S. Hughes, Pestotum sp., Phialocephala chlamydosporia (de Vries & Klien-Natrop) Gams, Verticillium nubilum Pethybr., non sporulating – 22 morphologically different cultures.

Recedents (frequency 1-2% in at least one treatment) – Arthrinium phaeospermum (Corda) B.M.Ellis, Candida albicans (Robin) Berkhout, Chrysosporium meridarium (Link) J.Carm., Ch. pannicola (Corda) v. Oorschot & Stalpers, Ch. tropicum J.W. Carmich., Chrysosporium sp., Cladorrhinum bulbillosum W. Gams & Mouch., Epicoccum nigrum Link, Gliomastix roseogrisea (S.B. Sakasena) Summerb. Gams, Monodictys putredinis (Wallr.) S. Hughes, Populaspora equi Shadomy & Dixon, P. nishigaharana Ts. Watan., Scytalidium japonicum Udawaga.Tominaga & T Hamaoka, Spoonrillous schenkii Hecton & C.F. Perkins, Torula sp., Trichocladium asperum Harz.,

<table>
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<th>Index</th>
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S – sawdust application  
C – control

Zygomycota are saprotrophs, with the potential to increase plant resistance to phytopathogens, while *P. roseus* is associated with ectomycorrhizae of conifers. These beneficial properties suggest that increased populations in soil would be useful.

The genus *Trichoderma* includes 89 species [19]. Many species are opportunistic avirulent plant symbionts and these are often considered for development as biocontrol agents against fungal phytopathogens [20]. *Trichoderma* occurs in soils at all latitudes and stimulation of its populations and activity, by manipulation of the soil environment (by amendment of organic matter), in one habitat may have general relevance.

A few species of *Trichoderma* detected in our studies (*T. harzianum, T. koningii, T. viride*) are known to restrict the growth of *H. annosum in vitro* and *in vivo* [3, 21]. *Trichoderma hamatum*, eudominant in the present study, is known for its antagonistic properties toward *Thanatephorus cucumeris* (A. B. Frank) Donk. Possible mechanisms involved in disease control include antibiosis, competition, mycoparasitism, and induction of plant defense responses. *Trichoderma harzianum*, another eudominant after sawdust amendment, can produce significant amounts of carbon dioxide, ethanol, water-soluble antibiotics and antifungal compounds, and fixed or volatile metabolites that are responsible for antibiosis, including the inhibition of growth and sporulation of some soil fungi.

An increase in *Trichoderma* populations in treated areas may increase the suppressiveness of soil toward *H. annosum*. Because of the preference of *Trichoderma* for higher temperatures [18, 21], more successful biocontrol effects may be expected in warmer years or in areas more exposed to solar radiation. Stimulation of growth of *Trichoderma* by moisture may be necessary in sandy soils, considering that the *Trichoderma* population size required for suppression was found to be $10^7$ conidia·g$^{-1}$ of soil in dry fine sandy loam and $10^8-10^9$ conidia·g$^{-1}$ of soil in moist clay loam. The beneficial effects of rainfall on the development of *Trichoderma* populations in sandy soils are supported by the correlation between abundance of *Trichoderma* and annual rainfall in the present study. More rainfall washes out the clay and humus, thus reducing the soil’s buffering capacity and increasing its acidification, which would generally stimulate the growth of fungi.

*Oidiodendron* is another generally beneficial fungus owing to its relationship with plants; some *Oidiodendron* species are ericoid mycorrhizal fungi. It is considered to be a cosmopolitan genus in soil and in/on cellulose substrates. In the present study, however, it was not always found in some fallow soils. Its population increased slowly (becoming greatest 14-16 years after sawdust treatment), partly as a result of stabilization of soil acidity (the fungus favours soil pH = 3.0-5.0) [22] and from the presence of *Vaccinium* on the site.

The application of sawdust did not increase the abundance of another *Heterobasidion* antagonist: *Penicillium adametzii* [23].

Since *Heterobasidion* infects trees continuously over a long period, antagonists are required to have long persistence in the ecosystem without the addition of nutrients or amendments. Increased populations of *Trichoderma* and *Oidiodendron* in sawdust-amended soils relative to non-treated controls, even 16 years after treatment, should ensure long-term suppressiveness of root pathogens in sandy soil.

**Conclusions**

Fresh sawdust may be beneficial as an amendment to sandy soil, even where applied without exposure to weathering and with no added mineral N (in quantities sufficient to meet the requirements of both sawdust-decomposing microorganisms and the development of transplanted plants).
Acknowledgements

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References