The Effect of Re-Planting Trees on Soil Microbial Communities in a Wildfire-Induced Subalpine Grassland

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Abstract: Wildfire often causes tremendous changes in ecosystems, particularly in subalpine and alpine areas, which are vulnerable due to severe climate conditions such as cold temperature and strong wind. This study aimed to clarify the effect of tree re-planting on ecosystem services such as the soil microbial community after several decades. We compared the re-planted forest and grassland with the mature forest as a reference in terms of soil microbial biomass C and N (C_{mic} and N_{mic}), enzyme activities, phospholipid fatty acids (PLFA) composition, and denaturing gradient gel electrophoresis (DGGE). The C_{mic} and N_{mic} did not differ among the grassland, re-planted forest and mature forest soil; however, ratios of C_{mic}/C_{org} and N_{mic}/N_{tot} decreased from the grassland to re-planted forest and mature forest soil. The total PLFAs and those attributed to bacteria and Gram-positive and Gram-negative bacteria did not differ between the re-planted forest and grassland soil. Principle component analysis of the PLFA content separated the grassland from re-planted forest and mature forest soil. Similarly, DGGE analysis revealed changes in both bacterial and fungal community structures with changes in vegetation. Our results suggest that the microbial community structure changes with the re-planting of trees after a fire event in this subalpine area. Recovery of the soil microbial community to the original state in a fire-damaged site in a subalpine area may require decades, even under a re-planted forest.

Keywords: wildfire; enzyme activities; microbial biomass; phospholipid fatty acid; reforestation; subalpine

1. Introduction

In forest ecosystems, fire alters the soil microbial community composition and activity in the form of direct heat-induced microbial mortality [1,2]. Furthermore, loss of forest by fire and grass invasion may alter nutrient cycling and quality of organic matter [3], which can indirectly affect the soil microbial community. Soil bacterial and fungal biomass is lower in fire-induced grassland areas than unaffected forest areas [4]. Post-fire tree replanting can accelerate the recovery of forest function in an ecosystem [5], thereby leading to greater recovery of the soil microbial community.

In Taiwan, grasslands are generally distributed above the tree line, at 3100 to 3400 m above sea level (asl). Approximately 50 years ago, a wildfire led to a vegetation shift from forest to grassland at Tatajia, the saddle (2600 to 2800 m asl) of Yushan, one of the highest mountains (3950 m) in East Asia.
This region provides a unique environment for observing long-term ecological changes in vegetation succession in a subalpine area.

Soil microorganisms are critical for ecosystem function and contribute substantially to organic matter decomposition and maintenance of soil structure [6,7]. Fungi and bacteria often occupy separate ecological niches in the soil environment and play critical roles in nutrient cycling [8,9]. Microbial biomass increases quickly in the early forest succession stage and then decreases and remains at a constant level with long-term forest restoration [10]. Hedo et al. [11] reported that post-fire silvicultural treatment did not have a direct effect on soil microbial properties and soil enzyme activities. However, little is known about the response of microbial communities to the succession of vegetation after fire [12]. A decade after a forest fire in a tropical rainforest, the dominant soil bacteria and actinobacteria did not differ between unburnt and burnt areas [13]. On the other hand, with the construction of 16S rRNA gene clone libraries, the diversity of bacterial communities was significantly lower in planted or natural forest than in fire-induced grassland soil in subalpine area after 3 decades [14]. Soil microbial biomass, activity and microbial community may increase with vegetation succession.

Analysis of phospholipid fatty acids (PLFA) is useful for determining the active soil microbial community structure [15]. It provides quantitative data on microbial communities [16]. As another simple, rapid and molecular technique, denaturing gradient gel electrophoresis (DGGE) can provide information on dominant microbial species such as bacteria, fungi, and target groups by the use of different primers in the same soil DNA extract [17].

With the hypothesis that replanting of trees in a wildfire-induced grassland might enhance the recovery of microbial communities and function, we examined microbial activities, biomass and community structure in a subalpine forest ecosystem after a fire disturbance using PLFA and DGGE.

2. Materials and Methods

2.1. Site and Soil Sampling

This study was conducted at Tatajia (23°28′ N, 120°54′ E) in central Taiwan, a subalpine area of Yushan (Mt. Jade). The site has an elevation 2600–2800 m, and mean annual precipitation of 4100 mm and mean annual temperature of 9.5 °C. The area is a long-term ecological study site of subalpine forest ecosystems in Taiwan. Soils are characterized as Humic Dystrudepts with low base saturation and strong acidity (pH 3.6–4.2) [18].

The study involved 3 vegetation types: grassland, re-planted forest, and natural forest. The study site was covered by natural hemlock and cypress forest before the wildfire. The grassland used as a control had formed from a wildfire event approximately 50 years ago in the forest area. The vegetation in the grassland included dwarf bamboo (*Yushania niitakayamensis* (Hayata) Keng f.) and alpine silver grass (*Miscanthus transmorrisonensis* Poaceae). The re-planted forest adjacent to the grassland plots was established by planting *Pinus taiwomensis* Hayata about 30 years ago. The *Pinus* plantation was weeded for three years in the early stage of establishment, after which no management was applied to the plantation. The non-disturbed natural forest used as a reference is dominated by Chinese hemlock (*Tsuga chinensis* (Franch.) Pritzel ex Diels.) with low density of Taiwan false cypress (*Chamaecyparis formosensis* Matsum.), spruce (*Picea morrisonicola* Hayata), and Armand’s pine (*Pinus armandi* Franch.). Natural and re-planted forest plots were separated by 200 m and were about 1 km away from grassland plots.

Each plot was demarcated into five subplots (50 × 50 m; replicates) for sampling. In each subplot, three samples were collected and mixed as a composite sample. The soil samples were collected in January 2010 and stored in an ice box for immediate transport to the laboratory. After removing undecomposed litter on the surface, soil samples were collected at 0–10 cm depth, which include a thin O horizon (Oa) and A horizon, using a metal auger (8 cm diam.). Visible detrital material, such as roots and litter, was hand-removed by sieving, and then the soil samples were ground to
pass through a 2-mm sieve. Soil samples were stored at 4 °C in the dark before analysis within one month. Subsamples were frozen at −20 °C immediately after sampling for analysis of PLFA. Biochemical analyses, including microbial biomass and enzymatic activities, were completed using fresh samples within 1 month after field sample collection. Another subsample was air-dried for soil chemical analysis.

2.2. Biochemical Assay

Organic C and total N in soil samples were determined using an NSC elemental analyzer (NA1500 Series 2, Fisons, Italy). Total P was determined using the Bray-1 method (Olsen and Sommers 1982). Soil subsamples were weighed and oven-dried for 72 h at 105 °C to determine the moisture content. Soil pH values of air-dried samples were measured using a combination of glass electrodes (soil: water ratio 1:2.5) [19]. Soil microbial biomass was analyzed using the chloroform fumigation extraction method [20]. Total organic C in the extracted solution was measured with a total organic C analyzer (Model 1010 O.I. Analytical, College Station, TX, USA) and converted to microbial biomass C (C_{mic}) assuming a conversion factor of 2.22 [21]. Microbial N (N_{mic}) was calculated from ninhydrin-reactive N released from the biomass and determined colorimetrically at 560 nm [22].

Urease activity (EC 3.5.1.5) was determined as described in Kandeler and Gerber [23]. The determination of phosphatase activity (EC 3.1.3.2) followed the method of Tabatabai and Bremner [24]. Cellulase activity (EC 3.2.1.4) and xylanase activity (EC 3.2.1.8) were determined by the method of Schinner and von Mersi [25].

Extraction and analysis of PLFA was as described in [26]. Lipids were extracted in a single-phase mixture of chloroform-methanol-citrate (1:2:0.8). Phospholipids were split into neutral and glycolipid phospholipids using a solid-phase extraction column and eluted with chloroform, acetone and methanol. Phospholipids then underwent methylation to form fatty acid methyl esters (FAMEs). FAMEs were identified, and their content was quantified by capillary gas chromatography (GC) with a flame ionization detector (Thermo Finnigan Trace chromatograph) as described in [27]. Fatty acid nomenclature followed Frostegård et al. [26]. The fatty acids i15:0, a15:0, 15:0, i16:0, 16:1ω7c, 17:0, i17:0, cy17:0, 18:1ω7c, and cy19:0 represent bacteria; 18:2ω6 fungi; i15:0, a15:0, i16:0, and i17:0 Gram-positive (G+) bacteria; 16:1ω7c, cy17:0, 18:1ω7c, and cy19:0 Gram-negative (G−) bacteria; and 10Me18:0 actinobacteria [28,29] (Zogg et al., 1997; Zelles 1999).

Soil DNA was extracted using the PowerSoil DNA isolation kit (MO BIO lab, Solana Beach, CA, USA). For soil bacteria, PCR conditions with primers F968-GC and R1401 [30] were as follows. The PCR mixtures (50 µL) contained 20 µL 2X Quick Taq HS DyeMix (Toyobo, Osaka, Japan), 10 pmol each primer, and 1 µL template DNA and deionized water to bring the final volume to 50 µL. The PCR program was 2 min 94 °C, followed by 34 cycles of 94 °C for 30 seconds, 61 °C for 30 seconds and 68 °C for 1 min. For soil fungi, PCR conditions with primer NS1 and GC fungi [31] were as follows. The PCR mixtures (50 µL) contained 20 µL 2X Quick Taq HS DyeMix (Toyobo), 10 pmol each primer, and 1 µL template DNA and deionized water to bring the final volume to 50 µL. The PCR program was 2 min 94 °C, followed by 32 cycles of 94 °C for 30 seconds, 52 °C for 30 seconds and 68 °C for 1 min.

The PCR products were analyzed using DGGE as described [32]. The DGGE gel comprised 6% polyacrylamide gel and 50% to 70% denaturants for bacterial analysis and 7% polyacrylamide and 20% to 45% denaturants for fungal analysis. The electrophoresis was performed for 18 h at 50 V in 1× TAE buffer at a constant temperature of 58 °C for bacteria and 60 °C for fungi with a Dcode Universal Detection system (Bio-Rad Laboratories, Hercules, CA, USA). The gel was stained with Gelstar (20 µL of Gelstar in 200 mL of 1× TAE buffer) for 30 min, and then DGGE band profiles were visualized under UV light. Digital image capturing involved the use of a Gel Doc XR gel imaging system (Bio-Rad). The obtained DGGE fingerprints were analyzed for similarities between microbial communities by the Dice similarity index based on the unweighted pair group method with arithmetic averages (UPGAMA) with Quantity One 4.5.1 (Bio-Rad, Hercules, CA, USA).
2.3. Statistical Analysis

Data from biological and chemical analyses were converted based on the oven-dried mass. Statistical analysis involved the use of Student’s t-test to evaluate significance of difference between the means of the three vegetation systems. A normality test (Shapiro-Wilk test) for all variables was performed before the test, and non-normal distributed data were transformed logarithmically. Principal component analysis (PCA) was used to test biochemical properties and relative concentrations (mole %) of individual fatty acids; relationships between vegetation types and individual PLFA were estimated using 2-D PC plots. Correlation between microbial parameters and soil properties involved Pearson’s correlation coefficient. Statistical analyses involved SPSS v12.0 (SPSS Inc., Chicago, IL, USA). P < 0.05 was considered statistically significant.

3. Results

3.1. Soil Properties and Microbial Biomass

Soil pH in the re-planted forest and grassland was similar and higher than in the mature forest (Table 1). Soil organic C and total N were higher in the mature than re-planted forest and grassland soil, with no significant difference between the re-planted forest and grassland soil. Soil total phosphorus did not show any significant difference among the three types of vegetation, although it was higher in the mature forest soil than in the re-planted forest and grassland soil. Soil microbial biomass C and N did not significantly differ among the three types of vegetation; however, the ratios of soil C/Corg and N/Ntot decreased from the grassland to the re-planted forest and mature forest.

Table 1. Soil chemical properties and microbial biomass characteristics in the three vegetation zones.

<table>
<thead>
<tr>
<th>Vegetation</th>
<th>pH</th>
<th>Organic C (g kg(^{-1}))</th>
<th>Total N (g kg(^{-1}))</th>
<th>Total P (mg kg(^{-1}))</th>
<th>C(_{\text{mic}}) (mg kg(^{-1}))</th>
<th>N(_{\text{mic}}) (mg kg(^{-1}))</th>
<th>C(_{\text{mic}})/Corg (%)</th>
<th>N(_{\text{mic}})/Ntot (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grassland</td>
<td>4.14 a</td>
<td>106 c</td>
<td>7.93 c</td>
<td>606 a</td>
<td>5905 a</td>
<td>587 a</td>
<td>5.19 a</td>
<td>7.43 a</td>
</tr>
<tr>
<td>Re-planted forest</td>
<td>4.20 a</td>
<td>131 b</td>
<td>9.82 b</td>
<td>574 a</td>
<td>4792 a</td>
<td>557 a</td>
<td>3.65 b</td>
<td>5.67 b</td>
</tr>
<tr>
<td>Mature forest</td>
<td>3.56 b</td>
<td>472 a</td>
<td>25.0 a</td>
<td>650 a</td>
<td>6165 a</td>
<td>648 a</td>
<td>1.30 c</td>
<td>2.88 c</td>
</tr>
</tbody>
</table>

C\(_{\text{mic}}\): microbial biomass C; N\(_{\text{mic}}\): microbial biomass N. In each column, different letters after the values indicate significant differences between treatments (Student’s t-test, \(p < 0.05\)).

3.2. Soil Enzyme Activities

Cellulase, xylanase and phosphatase activities were significantly higher in the mature forest soil than other soils, with no significant difference between the re-planted forest and grassland soil (Table 2). Urease activity was lowest in the re-planted forest soil, intermediate in the mature forest soil, and highest in the grassland soil (\(p < 0.05\)).

Table 2. Soil enzymatic activities in the three types of vegetation.

<table>
<thead>
<tr>
<th>Vegetation</th>
<th>Cellulase (mg Glucose g(^{-1}) d(^{-1}))</th>
<th>Xylanase (mg Glucose g(^{-1}) d(^{-1}))</th>
<th>Urease (mmole NH(_4^+)-N g(^{-1}) h(^{-1}))</th>
<th>Acid Phosphatase (µg Nitrophenol g(^{-1}) h(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grassland</td>
<td>2.786 b</td>
<td>5.252 b</td>
<td>77.89 a</td>
<td>1919 b</td>
</tr>
<tr>
<td>Re-planted forest</td>
<td>0.867 c</td>
<td>5.167 b</td>
<td>17.52 c</td>
<td>2189 b</td>
</tr>
<tr>
<td>Mature forest</td>
<td>5.608 a</td>
<td>10.98 a</td>
<td>41.90 b</td>
<td>2897 a</td>
</tr>
</tbody>
</table>

In each column, different letters after the values indicate significant difference between treatments. (Student’s t-test, \(p < 0.05\)).

3.3. PLFA Biomarkers

The planting of trees in the grassland affected PLFA biomarkers (Table 3). The total PLFA content was higher in the re-planted forest than the grassland soil, though it was still lower than the mature forest. The levels of both fungal and arbuscular mycorrhizal fungal PLFA biomarkers were higher in
the re-planted forest than the grassland soil. Bacterial biomass did not differ between the re-planted forest and grassland soil, nor did the PLFA levels of biomarkers for actinobacteria. The ratios of G+/G− and fungi/bacteria were higher but not significantly so in the re-planted forest compared with the grassland soil. Although the planting of trees increased the total PLFA and biomarker levels for fungi and arbuscular mycorrhizal fungi compared to the grassland, the levels of these microbial groups were still lower than those in the mature forest.

Table 3. Content of phospholipid acid biomarkers of the total microbial community and individual groups in soils under three types of vegetation.

<table>
<thead>
<tr>
<th>Vegetation</th>
<th>Total PLFA (nmol g⁻¹ soil)</th>
<th>Bacteria</th>
<th>Fungi</th>
<th>VAM Fungi</th>
<th>Actinobacteria</th>
<th>G+ Bacteria</th>
<th>G− Bacteria</th>
<th>G+/G− Fungi</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grassland</td>
<td>389 c</td>
<td>202 b</td>
<td>18.3 b</td>
<td>14.7 c</td>
<td>31.6 b</td>
<td>43.7 b</td>
<td>91.1 b</td>
<td>0.461 c</td>
</tr>
<tr>
<td>Re-planted forest</td>
<td>508 b</td>
<td>223 b</td>
<td>37.4 a</td>
<td>18.6 b</td>
<td>37.4 b</td>
<td>51.7 b</td>
<td>94.8 b</td>
<td>0.568 b</td>
</tr>
<tr>
<td>Mature forest</td>
<td>769 a</td>
<td>371 a</td>
<td>52.3 a</td>
<td>22.5 a</td>
<td>53.7 a</td>
<td>99.2 a</td>
<td>131 a</td>
<td>0.755 a</td>
</tr>
</tbody>
</table>

PLFA: phospholipid fatty acids; VAM, vesicular arbuscular mycorrhizal. In each column, different letters after the values indicate significant difference between treatments. (Student’s t-test, p < 0.05).

3.4. Principal Component Analysis

The first two PCA components explained 71.5% of the variability in the soil biochemical parameters. The first factor (PC axis 1), which explained 53.6% of the variance, was mainly positively associated with soil organic C, total N and enzymatic activities and negatively associated with soil pH, C_mic/C_Org and N_mic/N_tot (Figure 1). The second factor (PC axis 2), which explained 19.7% of the variance, was associated with microbial biomass. The soil biochemical PCA plot (Figure 1a) separated the mature forest from the planted forest and grassland.

![Figure 1](image_url)  
**Figure 1.** Plots of (a) the first 2 principal components (PCs) and (b) corresponding loading values in principal component analysis of the chemical and biochemical properties in the three types of vegetation.
Soil microbial communities, as analyzed by the PCA of the PLFA levels, significantly differed among the three types of vegetation and could be divided into three clusters: mature forest, re-planted forest, and grassland. The first and second principal components (PC1, PC2) accounted for 70% of the PLFA variability (Figure 2). The PC1 differentiated the mature forest from the other vegetation, and PC2 differentiated the grassland from the re-planted forest. High positive loadings for G+ bacteria (i15:0, a15:0, i16:0, 10Me16:0 and 10Me17:0), high negative loadings for G− bacteria (cy17:0 and 18:1ω7c), and positive loading for fungi (18:2ω6c) contributed to the separation of the mature forest from other types of vegetation along the PC1 axis.

![Figure 2](image)

**Figure 2.** Plots of (a) the first 2 principal components (PCs) and (b) corresponding loading values in the principal component analysis of the mole% of microbial phospholipid fatty acids in three types of vegetation.

### 3.5. DGGE Analysis

Dendrograms of genetic similarity obtained by DGGE of bacterial communities in the soil samples clustered differently for different types of vegetation (Figure 3). For soil bacterial communities, the re-planted forest and grassland were in one group, whereas in the other group was the mature forest.

Similarly, cluster analysis of fungal communities showed two main clusters for the three types of vegetation (Figure 4). Re-planted forest and grassland soil clustered together, being separate from the mature forest soil.
3.6. Correlation of Soil Properties and Microbial Communities

Soil organic C was closely correlated with soil levels of enzyme, with the exception of urease and phosphatase, and microbial community composition (Table 4). Total N content was highly correlated with the organic C content; the correlation of the total N content with soil enzymes and microbial community composition not as high as that with organic C. $C_{mic}$ was not correlated with soil enzymes or microbial community composition. Soil pH was negatively correlated with cellulase and xylanase activity and microbial community composition.
Table 4. Pearson’s correlation coefficients for soil properties, soil enzyme and microbial variables.

<table>
<thead>
<tr>
<th></th>
<th>Cellulase</th>
<th>Xylanase</th>
<th>Urease</th>
<th>Phosphatase</th>
<th>Total PLFA</th>
<th>Bacteria</th>
<th>Gram-Positive Bacteria (G+)</th>
<th>Gram-Negative Bacteria (G−)</th>
<th>G+/G−</th>
<th>Fungi</th>
<th>VAM Fungi</th>
<th>Bacteria/Fungi</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>0.790 ***</td>
<td>0.849 ***</td>
<td>−0.216</td>
<td>0.432</td>
<td>0.897 ***</td>
<td>0.867 ***</td>
<td>0.876 ***</td>
<td>0.732 **</td>
<td>0.903 **</td>
<td>0.753 **</td>
<td>0.776 **</td>
<td>0.517 *</td>
</tr>
<tr>
<td>N</td>
<td>0.569 *</td>
<td>0.719 **</td>
<td>−0.158</td>
<td>0.269</td>
<td>0.650 **</td>
<td>0.564 *</td>
<td>0.598 *</td>
<td>0.390</td>
<td>0.762 **</td>
<td>0.633 *</td>
<td>0.505</td>
<td>0.529 *</td>
</tr>
<tr>
<td>P</td>
<td>0.163</td>
<td>0.160</td>
<td>0.134</td>
<td>0.114</td>
<td>0.379</td>
<td>0.359</td>
<td>0.341</td>
<td>0.329</td>
<td>0.292</td>
<td>0.378</td>
<td>0.294</td>
<td>0.213</td>
</tr>
<tr>
<td>C/N</td>
<td>0.682 **</td>
<td>0.641 *</td>
<td>−0.178</td>
<td>0.426</td>
<td>0.710 **</td>
<td>0.796 ***</td>
<td>0.775 **</td>
<td>0.804 ***</td>
<td>0.611</td>
<td>0.398</td>
<td>0.733 **</td>
<td>0.116</td>
</tr>
<tr>
<td>C&lt;sub&gt;mic&lt;/sub&gt;</td>
<td>0.485</td>
<td>0.362</td>
<td>0.166</td>
<td>0.319</td>
<td>0.486</td>
<td>0.409</td>
<td>0.399</td>
<td>0.309</td>
<td>0.394</td>
<td>0.657 **</td>
<td>0.279</td>
<td>0.511</td>
</tr>
<tr>
<td>N&lt;sub&gt;mic&lt;/sub&gt;</td>
<td>0.422</td>
<td>0.363</td>
<td>0.117</td>
<td>0.197</td>
<td>0.461</td>
<td>0.393</td>
<td>0.385</td>
<td>0.300</td>
<td>0.391</td>
<td>0.592 *</td>
<td>0.279</td>
<td>0.462</td>
</tr>
<tr>
<td>C&lt;sub&gt;mic&lt;/sub&gt;/C&lt;sub&gt;org&lt;/sub&gt;</td>
<td>−0.587 *</td>
<td>−0.733 **</td>
<td>0.455</td>
<td>−0.388</td>
<td>−0.844 ***</td>
<td>−0.804 ***</td>
<td>−0.821 ***</td>
<td>−0.663 **</td>
<td>−0.886 **</td>
<td>−0.627 *</td>
<td>−0.820 **</td>
<td>−0.458</td>
</tr>
<tr>
<td>N&lt;sub&gt;mic&lt;/sub&gt;/N&lt;sub&gt;tot&lt;/sub&gt;</td>
<td>−0.517 *</td>
<td>−0.694 **</td>
<td>0.396</td>
<td>−0.306</td>
<td>−0.767 **</td>
<td>−0.701 **</td>
<td>−0.723 **</td>
<td>−0.543 *</td>
<td>−0.828 **</td>
<td>−0.619 *</td>
<td>−0.710 **</td>
<td>−0.489</td>
</tr>
<tr>
<td>Moisture</td>
<td>0.368</td>
<td>0.196</td>
<td>0.312</td>
<td>0.083</td>
<td>0.289</td>
<td>0.251</td>
<td>0.239</td>
<td>0.203</td>
<td>0.181</td>
<td>0.451</td>
<td>0.062</td>
<td>0.311</td>
</tr>
<tr>
<td>pH</td>
<td>0.440</td>
<td>0.597 *</td>
<td>−0.265</td>
<td>0.356</td>
<td>0.790 ***</td>
<td>0.683 **</td>
<td>0.696 **</td>
<td>0.487</td>
<td>0.805 **</td>
<td>0.752 **</td>
<td>0.701 **</td>
<td>0.602 *</td>
</tr>
<tr>
<td>Cellulose</td>
<td>0.886 ***</td>
<td>0.866 ***</td>
<td>0.062</td>
<td>0.541 *</td>
<td>0.666 **</td>
<td>0.699 **</td>
<td>0.683 **</td>
<td>0.679 **</td>
<td>0.591</td>
<td>0.580 *</td>
<td>0.514 *</td>
<td>0.318</td>
</tr>
<tr>
<td>Xylanase</td>
<td>0.886 ***</td>
<td>0.035</td>
<td>0.473</td>
<td>0.651 **</td>
<td>0.676 **</td>
<td>0.683 **</td>
<td>0.602 *</td>
<td>0.681 **</td>
<td>0.489</td>
<td>0.577 *</td>
<td>0.265</td>
<td></td>
</tr>
<tr>
<td>Urease</td>
<td>0.062</td>
<td>0.035</td>
<td>1</td>
<td>−0.557 *</td>
<td>−0.369</td>
<td>−0.199</td>
<td>−0.211</td>
<td>−0.025</td>
<td>−0.433</td>
<td>−0.346</td>
<td>−0.432</td>
<td>−0.575 *</td>
</tr>
<tr>
<td>phosphatase</td>
<td>0.541 *</td>
<td>0.473</td>
<td>−0.557</td>
<td>1</td>
<td>0.394</td>
<td>0.278</td>
<td>0.260</td>
<td>0.184</td>
<td>0.365</td>
<td>0.550 *</td>
<td>0.341</td>
<td>0.553 *</td>
</tr>
<tr>
<td>Total PLFA</td>
<td>0.666 **</td>
<td>0.651 **</td>
<td>−0.369</td>
<td>0.394</td>
<td>1</td>
<td>0.959 ***</td>
<td>0.961 ***</td>
<td>0.946 ***</td>
<td>0.936 **</td>
<td>0.806 **</td>
<td>0.919 ***</td>
<td>0.556 *</td>
</tr>
</tbody>
</table>

VAM: vesicular arbuscular mycorrhizal. * p < 0.05, ** p < 0.01, and *** p < 0.001.
4. Discussion

The planting of trees can have direct effects on soil properties [33,34], mostly because of increased soil organic matter (SOM) through the supply of litter and root exudates. In our study of a Pinus plantation in central Taiwan (2600–2800 m asl) after a fire event, the trend we found in soil SOM, total N, microbial activities and microbial community structure may be related to the vegetation and recovery history.

The result of the PCA carried out with soil biochemical parameters clearly indicated a differentiation of chemical properties and soil enzyme activities in soils under different types of vegetation (Figure 1). Soil pH was more acidic in the mature forest than in the re-planted forest and grassland in our study area, which may be attributed to accumulation of more organic acids under the older forest [35]. We found the soil organic C content increased from the grassland to the mature forest in our study site. SOM is highly correlated with the structure of the microbial community of forest soils because of the quantity and quality of organic matter input [36,37]. Soil microbial biomass is significantly decreased after fire events in coniferous forests [38,39]. Furthermore, microbial biomass in forest soils takes decades to recover to the original state once it has been depleted [40]. In our previous study at this site, both levels of microbial biomass C\textsubscript{mic} and ratios of fungal to bacterial biomass were higher in the mature forest than the grassland soil [4], but after 15 years of succession at this study site, microbial biomass C\textsubscript{mic} and N\textsubscript{mic} did not differ among the vegetation zones. Our current study suggests that the establishment of a young forest, the Pinus plantation, caused the change in soil C\textsubscript{mic}/C\textsubscript{org} and N\textsubscript{mic}/N\textsubscript{tot} ratios. The soil C\textsubscript{mic}/C\textsubscript{org} ratio may reflect the potential for soil organic mineralization after fresh input of organic materials [41]. The grassland soil in our study site showed the greatest potential for mineralization, with a high C\textsubscript{mic}/C\textsubscript{org} ratio. Furthermore, the re-planted forest gradually provided more recalcitrant SOM into soil and decreased the C\textsubscript{mic}/C\textsubscript{org} ratio. The re-planted forest still had a higher C\textsubscript{mic}/C\textsubscript{org} ratio than the mature forest, which may indicate that the re-planted forest is still under succession.

Recovery of microbial biomass in burned forest–grassland ecosystems is linked to plant community recovery, providing an important interaction between plants and soil microbes [42]. In general, enzyme activities in soil are closely related to the content of organic matter [43] and C\textsubscript{mic} [44]. High SOM usually sustains high C\textsubscript{mic} and enzyme activities [45]. Xylanase is produced and released mostly by fungi in the forest environment [46]. Ali et al. [47] showed that in coniferous tree species such as Pinus taeda L., ectomycorrhizal fungi produce extracellular phosphatase to improve phosphorus uptake. In our study, the mature forest soil had higher fungal biomass than other soils and provided more xylanase and phosphatase (Table 3). In addition, fungal biomass, particularly of vesicular arbuscular mycorrhizal (VAM) fungi, is a good predictor of phosphatase in forest soil [48] which coincides with our results, with high fungal biomass in the mature forest soil (Tables 1 and 3). On the other hand, acid phosphatase activity decreased in soil with severe wildfires [49], which is consistent with our results in the grassland and re-planted forest.

Although soil microbial biomass (both C\textsubscript{mic} and N\textsubscript{mic}) was similar among the three types of vegetation, urease activity was significantly higher in the grassland than in the re-planted and mature forest. Increased communities of gramineous and herbaceous species of grasses more likely provide higher inputs of labile C and N than do perennial forest species. Scott and Binkley [50] showed that the quality of substrate (low lignin to N ratios) provided by existing plants can directly influence N turnover rates.

Our PCA of PLFA revealed different microbial communities in soils under different vegetation (Figure 2). Other studies have also shown that associated changes in vegetation rather than direct changes in soil properties may lead to shifts in the soil microbial community structure [51,52].

We found no differences in bacterial biomarker features between grassland and re-planted forest soil. However, fungal and VAM levels were higher in the re-planted forest than the grassland soil. Fungal communities play a dominant role in fresh organic matter decomposition [53]. Djukic et al. [54] showed similar results with higher amounts of fungi in forest soils than in grassland soils.
Fungal biomass contributes considerably to total biomass in coniferous forest soils [55]. By comparison, high availability of labile substrate (root exudates), for which bacteria are effective competitors, may constrain fungal biomass in grassland sites [54]. Allison et al. [56] found that C inputs in grasslands were mainly derived from root exudates: the rapidly metabolized C limits the development of fungi because fungi use more recalcitrant sources of C.

VAM fungi play an important role in improving plant growth and uptake of nutrients, especially phosphorus [57]. PLFA 16:1ω5 is present in VAM fungi [58] and in G− bacteria [59]. Jonasson et al. [60] indicated that 20% to 30% of total organic P is immobilized by microbes, and Thoms et al. [61] showed a high association of soil P content and PLFA 16:1ω5. The alterations in the 16:1ω5 pattern we found support results from previous studies showing higher extractable P in mature forest than other vegetation soils (Table 1).

The ratio of fungi to bacteria was increased in the re-planted forest soil (Table 3). The increased ratio of fungi to bacteria can be an important indicator of the amount and composition of litter that enters soils because fungi are the dominant decomposers of plant cell-wall polymers in the litter [62].

In this study, the switch from grassland to forest ecosystems was critical for the source of SOM because these ecosystems differ in the quantity and quality of dead plant biomass input. Although bacterial biomass was similar in the grassland and re-planted forest soils, fungal biomass increases from grassland to re-planted and mature forests. Our previous study found that both soil ergosterol and the respiration rate of fungi to bacteria decreased from the forest to grassland soil [4]. In this study, the ratio of fungi to bacteria PLFA patterns was similar in both the mature forest and re-planted forest soil, which suggests the recovery of the microbial community structure by reforestation after wildfire events.

The composition of the soil microbial community was mainly controlled by the pH and ratio of C to N of the substrate [63]. Bacterial biomass has been reported to increase in soils with a lower C to N ratio [64]. Ingham et al. [65] stated that grasslands more strongly dominated by bacteria than a coniferous forest, which supports our results. In grasslands, fast-growing plant species, especially those with highly branched fine root systems, supply large quantities of exudates [66], which are favored by bacteria.

A low ratio of G+/G− bacteria, found in grassland soils, may be due to better growth of G− bacteria under substrate-rich conditions [27]. By contrast, with progressive succession from a grassland to a forest system, the detritus food webs become more complex and recalcitrant. Slow-growing specialists, such as G+ bacteria, are more competitive than G− bacteria in resource-limited areas [67] because of effective cell metabolism and effective use of recalcitrant substances such as cellulose and lignin in a coniferous environment [68,69]. This situation is similar to the decrease in the G+/G− ratio for the soils in a bamboo-invaded cedar forest, as herbaceous litter favors the growth of G− bacteria [70].

Our dendrograms of soil bacteria and fungi from PCR-DGGE showed primary differences in the microbial communities related to the succession of vegetation (Figures 3 and 4). By using 16S rRNA gene clone libraries for this same study site, Lin et al. [14] found that the soil bacterial communities differed between the grassland and the mature forest 30 years after the fire. These findings support our results. The Pinus plantation can reform soil microbial communities. Although the microbial community structure was similar in grassland and re-planted forest soil, our DGGE findings agreed with the PCA results of PLFA, showing a distinct composition of bacterial and fungal communities with forest succession.

5. Conclusions

Our results suggest substantial changes in microbial community structure with the re-planting of trees in a wildfire-induced grassland. The plantation changed the fraction of the labile form of SOM, with a decrease in the ratio of C_{mic}/C_{org} in the re-planted forest soil. Microbial biomass (PLFA content) increased from the grassland to the re-planted forest soil, accompanied by increases in fungi and VAM.
fungi but not the bacterial content. The shift in vegetation with the succession of the fire-induced grassland to the forest led to an increased ratio of G+/G− bacteria. DGGE cluster analysis revealed changes in bacterial and fungal community structure associated with the Pinus plantation. Therefore, the planting of trees 30 years ago after a fire in a subalpine area changed the soil microbial activities and community structure, but recovery of the soil microbial community to the original state may need much more time.

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