Ectomycorrhizal fungal communities in endangered *Pinus amamiana* forests

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Abstract

Interactions between trees and ectomycorrhizal (ECM) fungi are critical for the growth and survival of both partners. However, ECM symbiosis in endangered trees has hardly been explored, complicating conservation efforts. Here, we evaluated resident ECM roots and soil spore banks of ECM fungi from endangered *Pinus amamiana* forests on Yakushima and Tanegashima Islands, Kagoshima Prefecture, Japan. Soil samples were collected from remaining four forests in the two islands. The resident ECM roots in soil samples were subjected to molecular identification. Soil spore banks of ECM fungi were analyzed via bio-assays using a range of host seedlings (*P. amamiana*, *P. parviflora*, *P. densiflora* and *Casta-nopsis sieboldii*) for 6–8 months. In all remaining *P. amamiana* forests, we discovered a new *Rhizopogon* species (*Rhizopogon* sp.1), the sequence of which has no match among numerous *Rhizopogon* sequences deposited in the international sequence database. Host identification of the resident ECM roots confirmed that *Rhizopogon* sp.1 was associated only with *P. amamiana*. *Rhizopogon* sp.1 was far more dominant in soil spore banks than in resident ECM roots, and its presence was confirmed in nearly all soil samples examined across the major remaining populations. While *Rhizopogon* sp.1 did not completely lose compatibility to other pine species, its infection rate in the bioassays was highest in the original host, *P. amamiana*, the performance of which was improved by the infection. These results indicate that *Rhizopogon* sp.1 is very likely to have a close ecological relationship with endangered *P. amamiana*, probably due to a long co-evolutionary period on isolated islands, and to play the key role in seedling establishment after disturbance. We may need to identify and utilize such key ECM fungi to conserve endangered trees practically.

Introduction

Forests have disappeared and deteriorated all over the world due to habitat destruction and environmental changes caused by human activity [1]. As a result, many tree species are threatened with extinction [2]. In particular, the dominant trees in a forest play critical roles in primary production and ecosystem structuring, directly and/or indirectly supporting many other organisms. Thus, the extinction of a dominant tree species can have a serious impact on
biodiversity in the entire forest ecosystem, and for this reason, conservation measures are urgently needed for dominant trees threatened with extinction.

Dominant trees in temperate forests, such as species of the Fagaceae and Pinaceae, are associated with ectomycorrhizal (ECM) fungi and depend on them for soil nutrients. Without compatible ECM fungi, these trees cannot grow normally [3]. Moreover, it has become increasingly clear that the species composition and distribution of ECM fungi in soil affect the establishment of tree seedlings [4, 5]. Therefore, knowledge of the ECM fungi associated with an endangered tree species could be key to its conservation.

There are two main infection pathways used by ECM fungi in nature: the mycelium (mycelial network) extending from the existing ECM roots (e.g. [6]) and spores dispersed from fruiting bodies [7]. Both types of infection are ubiquitous in soils of developed forests, where host roots can be easily infected with ECM fungi. Mycelial networks are the predominant infection pathway in less disturbed forests, while spores that have accumulated in the soil, called soil spore banks, are the primary source of ECM infection for regenerating seedlings after disturbances that eliminate existing trees [8, 9]. Therefore, pioneer trees that require disturbance for regeneration depend on both soil spore banks, at the seedling stage, and mycelial networks, at the mature stage.

The species composition of ECM fungi often differs between spore banks and resident ECM roots, with soil spore banks usually composed of fewer ECM fungal species [8, 10–12]. Especially in pine forests, soil spore banks are dominated by pine-specific Rhizopogon [8, 10, 13–15], because their spores have a longer life span and greater environmental resistance than do those of other ECM fungi [16–19]. All Rhizopogon species produce hypogeous sporocarps and depend on animal ingestion for spore dispersal [14, 20]. Thus, gene flow of Rhizopogon species is more restricted than fungi producing epigeous sporocarps and is easily inhibited by geographical barriers [21]. This limited gene flow leads to genetic differentiation among isolated Rhizopogon populations [21–23] and eventually to speciation in geographically isolated areas [24, 25]. While most pine species are successful pioneer colonizers after disturbances throughout the northern hemisphere [20, 26], establishment of their seedlings may be sustained by unique Rhizopogon species that evolved locally.

The genus _Pinus_ consists of two subgenera, four sections and over 100 extant species, and it is the largest genus of conifers and the most widespread genus of trees in the Northern Hemisphere [27]. While some pine species have wide distribution ranges over a continent, many have been restricted to small isolated areas [27] as a result of long biogeographic history since the Early Cretaceous [27–29] and competition with broadleaf trees. Species with small distribution ranges are prone to extinction caused by environmental and demographic stochasticity in combination with inbreeding depression [30]. In fact, 19 pine species are classified as threatened (12 endangered and 7 vulnerable species) [2].

_Pinus amamiana_ is endemic to Yakushima and Tanegashima Islands, Kagoshima Prefecture, Japan [27] and is classified as “Endangered A3ce; B1ab(iii,v)+2ab(iii,v)” on the IUCN Red List and as “Vulnerable” by the Japanese government [31]. The abundance of _P. amamiana_ has been drastically reduced by timber harvesting and pine wilt disease, and only 1,500 and 100 individuals persist in Yakushima and Tanegashima Islands, respectively [31]. Although we know nothing about the ECM fungi colonizing this endangered pine species, _P. amamiana_ may be associated with unique ECM fungi that coevolved locally and may depend on these fungi for existence.

ECM fungi of endangered trees have long been overlooked, and thus available information on these fungi is quite limited. However, our recent study in forests of _Pseudotsuga japonica_ (Pinaceae), another endangered conifer species in Japan, found that soil spore banks are dominated by _Rhizopogon togasawariana_ that coevolved with the endangered host for more than 30
million years [12]. Notably, this fungus was completely absent in resident tree roots [32]. Because *P. japonica* is a light-demanding pioneer tree species, its seedlings are rarely found on the dark forest floor [33, 34]. These findings strongly suggest that both *P. japonica* and *R. togasawariana* depend on disturbance for regenerating their populations, and that *R. togasawariana* plays the key role in the establishment of seedlings during regeneration. The existence of this ECM fungus specific to the endangered tree species and its potential role in establishment of endangered tree seedlings have critical implications for conservation; yet, it is difficult to generalize the results obtained from this single case. Similar studies in *Pinus*, which includes far more endangered species than those of *Pseudotsuga*, would further improve our understanding of ECM symbiosis on endangered trees.

In the present study, our objective was to characterize the ECM fungal communities of both soil spore banks and resident ECM roots in remaining *P. amamiana* forests. Specifically, we examined the following hypotheses: (1) *P. amamiana* is associated with some host specific ECM fungi that have coevolved with this endangered pine, and (2) these fungi are more frequently detected in soil spore banks than in resident ECM roots as *Rhizopogon* species in other ecosystems. We believe that the results of this study can provide key information for *in situ* conservation of *P. amamiana* and has many important implications for the conservation of other endangered trees.

**Materials and methods**

**Study sites**

This study was conducted in four *P. amamiana* forests, i.e. Hirauchi (site 1) and Banri (site 2) on Yakushima Island, and Wasedagawa (site 3) and Injo (site 4) on Tanegashima Island, Kagoshima Prefecture, Japan (Table 1, S1 Appendix). Sampling permission for the site 1, 2 and 3 was issued by Yakushima forest office, and it for site 4 was issued by the owner of the land. These forests harbor the majority of remaining *P. amamiana* populations. At the Yakushima sites, *P. amamiana* is usually found on steep slopes or ridges with other ECM trees (*Castanopsis sieboldii, Lithocarpus edulis, Quercus acuta, Q. phillyracea, Q. salicina, Tsuga sieboldii*). At the Tanegashima sites, this tree is usually found on hilly land (site 3) or by the seaside (site 4) along with other ECM trees (*C. sieboldii, L. edulis, P. thunbergii, Q. salicina*). In addition to ECM trees, these forests also contain non-ECM sub-trees (i.e. *Cleyera japonica, Cryptomeria japonica, Distylium racemosum, Elaeocarpus japonicas, Ilex pedunculosa, Morella rubra, Myrsine seguittii, Pieris japonica, Prunus sp., Rhaphiolepis indica var. umbellate, Rhododendron tashiroi, Syzygium buxifolium, Ternstroemia gymnanthera*). The annual mean temperature ranged from 17.9 to 20.4˚C, with abundant precipitation (Table 1).

| Site                | Elevation (m) | Mean annual precipitation (mm) | Annual mean temperature (˚C) | Site area (ha) | Latitude/longitude | Mean DBH of *Pseudotsuga japonica* (cm) |
|---------------------|---------------|--------------------------------|-----------------------------|----------------|--------------------|----------------------------------------|
| Hirauchi (site 1)   | 363–514       | 3114                           | 20.4                        | 1.12           | N: 30˚14’ 49.7–57.6”, E: 130˚30’ 14.9–18.5” | 44.2 (n = 41) |
| Ohko Forestry Road  | 635–748       | 2412                           | -                           | 0.54           | N: 30˚19’ 40.1–53.4”, E: 130˚24’ 37.7–42.2” | 38.4 (n = 21) |
| Banri (site 2)      | 71–96         | 2941                           | 17.9                        | 0.33           | N: 30˚36’ 36.9–39.2”, E: 131˚01’ 43.4–45.4” | 19.7 (n = 32) |
| Wasedagawa (site 3) | 87–103        | 2941                           | 17.9                        | 0.24           | N: 30˚34’ 23.1–27.9”, E: 131˚01’ 27.1–31.0” | 25.8 (n = 10) |

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Sampling
In September 2014, 26, 21 and 32 pairs of soil samples were collected near randomly selected *P. amamiana* trees at sites 1, 2 and 3, respectively. In November 2015, an additional 15 and 10 soil samples were collected at sites 1 and 4, respectively. The number of samples collected depended on the number of surviving trees at each site. Selected trees were >5 m apart. Two soil samples (5 × 5 × 10 cm depth), one for resident ECM root analysis and the other for soil spore bank analysis, were collected near each selected tree. Geographical positions were recorded using a GPS (GPSPMAP62SJ, Garmin, Olathe, KS, USA). Samples were placed separately in plastic bags and kept at 4˚C until further analyses.

Bioassays
Bioassay experiments were performed to assess the soil spore banks following an established method [11, 12]. In these experiments, only soil samples collected in 2014 were used. After removing roots, organic debris and small stones, the soil samples were air-dried for 2–8 months (S2 Appendix). The bioassay containers were 50-mL centrifuge tubes (Ina Optika Inc., Osaka, Japan), with two drainage holes near the bottom blocked by a cotton ball to prevent soil loss. Each tube was filled with approximately 40 mL of the air-dried soil sample.

We used four tree species, *P. amamiana*, *P. parviflora*, *P. densiflora* and *Castanopsis sieboldii* (Fagaceae), to evaluate the host specificity of soil propagule banks, particularly the specificity of the *Rhizopogon* spp., which are expected in spore banks. While *P. parviflora* belongs to the same subgenus (*Strobus*) as that of *P. amamiana*, *P. densiflora* belongs to the subgenus *Pinus*. *C. sieboldii* is a broadleaf tree species that is predominant in warm-temperate forests of the region. We were able to use only 46 *P. amamiana* seeds for all bioassay tubes because of the limited seed availability and low germination rate of this endangered species.

Seeds of all tree species were soaked in tap water for 48 h, surface-sterilized using a 5% sodium hypochlorite solution for >10 min, and rinsed with tap water. To induce germination, surface-sterilized *P. amamiana* and *P. parviflora* seeds were placed on sterilized, well-moistened peat moss in an incubator (25˚C for 1 month and then 5˚C for 2 months), while *P. densiflora* and *C. sieboldii* seeds were placed on sterilized Shibanome soil (fine red granular soil of Pleistocene volcanic origin) in an incubator (25˚C). For each soil-host combination, one germinated seed was planted and covered with sterilized Shibanome soil in a bioassay tube, for a total of 283 seedlings (S2 Appendix). To monitor airborne spore contamination of ECM fungi, we prepared a control treatment using 10 tubes containing an autoclaved, randomly selected soil sample (per host). Bioassay seedlings were watered with tap water (2–3 mL per seedling) every 3–5 days depending on soil conditions and were harvested after 6–9 months of growth in a growth chamber (MLR-351; SANYO Inc. Tokyo, Japan) set to 25˚C with 16 h of light (15 fluorescent lamps: 20,000 lx) and then to 20˚C for 8 h of dark (S2 Appendix).

Identification of ectomycorrhizal fungi
Roots of resident trees collected from field soil samples and bioassay seedlings were gently washed with tap water. ECM root tips were classified into morphotypes under a dissecting microscope based on their surface color, texture and emanating hyphae, as described in previous studies [35, 36]. For molecular identification of ECM fungi, triplicate ECM root tips, if available, were selected for each morphotype in a soil sample, then placed individually into 2.0-mL tubes. A total of 682 root tips from ECM were used for molecular identification on resident trees. In addition, 208, 246, 257 and 191 ECM root tips from the bioassays of the *P. amamiana*, *P. parviflora*, *P. densiflora* and *C. sieboldii*, respectively, were subjected to DNA analysis.
Molecular identification of ECM fungi was performed following Murata et al. [12]. Briefly, each ECM tip sample was placed in a 2.0-mL tube containing a zirconia ball and then pulverized using a bead beater. Total DNA was extracted using a modified cetyl trimethylammonium bromide method [35]. Internal transcribed spacer regions (ITS1-5.8S-ITS2) of ribosomal DNA were amplified using the ITS1F or ITS0F-T forward primer and several reverse primers (ITS4, ITS4Cg, LB-W [37–40] depending on amplification success. Platinum® Multiplex PCR Master Mix (Applied Biosystems, Foster City, CA, USA) was used for polymerase chain reaction (PCR).

The amplified products were purified and subjected to direct sequencing (3730xl DNA Analyzer; Applied Biosystems) using ITS1 as the sequencing primer. For poorly sequenced samples, ITS4 was also used as a sequencing primer. The obtained high-quality sequences were grouped into molecular operational taxonomic units (MOTUs) with ≥ 97% ITS sequence similarity (including 5.8S regions) using ATGC software (ver. 7.0; GENETYX Corp., Tokyo, Japan). Species identity was assigned based on the results of BLAST searches against known taxa in international sequence databases (DNA Data Bank of Japan [DDBJ]/EMBL/GenBank). Potential PCR chimeras were removed manually before further analyses, based on inconsistent BLAST results between segments within each MOTU sequence, e.g., between the initial and last parts (about 100 bp each) of the sequence. If the ITS similarity with a described species from herbarium specimens was > 97%, we used that species name for the MOTU. When the similarity to known species was 90–97% or < 90%, we identified MOTUs at the genus and family level, respectively. One exception was C. geophilum, which was identified by amplifying the Cenococcum-specific primer (ITS4Cg) and confirmed by sequencing randomly selected samples. Although several different MOTUs of C. geophilum were found (< 97% ITS sequence similarity), they were treated as one MOTU in this study, because no taxonomic agreement was reached among the many previous studies of ECM fungal communities (e.g., [36, 41]). Representative sequences of individual MOTUs were deposited in the DDBJ under accession numbers LC315810–LC315919.

To confirm the identity of host species in all molecular samples, the plastid trnL or rbcL region of plant DNA was amplified using primers trnC (5’-cgaaatcgtagacgctacg-3’) or rbcLa-F (5’-atgtcaccacaaacagagactaaagc-3’) in combination with trnD (5’-ggggatagaggctagactaaagc-3’) or rbcLa-R (5’-gttaaatcaagtctccaccccg-3’). All the amplicons were purified and subjected to direct sequencing using trnC or rbcLa-F as the sequencing primer. As with species identification of ECM fungi, the identity of host species was assigned based on the results of BLAST searches against known taxa in international sequence databases.

**Statistical analyses**

The frequency of an ECM fungal species was defined as the number of bioassay seedlings colonized by each ECM fungal species or, for resident trees, the number of soil samples containing each ECM fungal species. The frequencies of major fungal species were compared between different bioassay host species using Fisher’s exact test implemented in SPSS (ver. 11.5; SPSS Japan Inc., Tokyo, Japan). Statistical significance was set at α = 0.05. Estimate S software (ver. 8.2; [42]) was used to calculate the minimum number of total species richness indicators (Jackknife 2).

A fungal community matrix was built for the major host groups (P. amamiana, T. sieboldii and Fagaceae) at each site. Community compositional similarity was visualized using non-metric dimensional scaling (NMDS), implemented in the ‘vegan’ package of R software [43], using the Bray-Curtis distance (Manhattan distance for NMDS based on the unit of soil
samples for resident ECM roots) and 999 permutations. Differences in community composition between resident ECM roots and assayed spore banks were determined using Adonis (permutation multivariate analysis of variance; [44]) in ‘vegan’, again using the Bray-Curtis distance and 999 permutations [45]. Data dispersions of the communities among groups were analyzed by the Betadisper test (Permutational analysis of multivariate dispersions; [44]) in ‘vegan’.

Results

ECM fungal communities of Pinus amamiana forests

Of the 104 soil samples collected from the four study sites, 44 contained P. amamiana ECM roots. Other dominant host taxa in the belowground ECM tips were T. sieboldii and Fagaceae, which were found in 9 and 58 soil samples, respectively (Table 2). As a minor host, the subgenus Pinus was confirmed in one soil sample. No ECM roots were found in 15 soil samples.

Of the 101 putative ECM fungi that were identified in the four sites, 42 species were found on P. amamiana roots (Table 2, S3 Appendix). Thirteen and sixty-one ECM fungi were found on T. sieboldii and Fagaceae roots, respectively. The rarefaction curves indicated that neither the observed ECM fungal richness on P. amamiana nor richness on all hosts approached an asymptote with the maximum sample size (Fig 1), indicating that additional fungal species would be found with additional sampling effort. The richness estimator Jackknife2 showed that at least 229 ECM fungal species should inhabit these forests (Fig 1). The Jackknife2 richness estimator for ECM fungi on P. amamiana was 102 (Fig 1). The observed ECM fungal richness Jackknife2 estimators on T. sieboldii and Fagaceae trees were 31 and 141, respectively.

The total ECM fungal community was composed of a few common species and a large number of rare species. Only eight ECM fungal taxa appeared in five or more soil samples. In contrast, 73 taxa were found only once (i.e., singletons). Cenococcum geophilum was the most frequent taxon, found in 39% of the soil samples. The occurrence frequencies of Russulaceae (48%), Boletaceae (36%), Clavulinaceae (17%), Rhizopogonaceae (14%) and Thelephoraceae (13%) were also high at the family level. Russulaceae (30 spp.) was the most species-rich ECM fungal lineage, followed by Boletaceae (19 spp.), Clavulinaceae (eight spp.) and Thelephoraceae (six spp.), while only two species of Rhizopogonaceae were found (Table 2).

Only Rhizopogon sp.1 was found at all sites (Table 2). In addition, Boletaceae sp.1, Clavulinaceae sp.3, Phylloporus sp.1 and Thelephoraceae sp.1, among the major ECM fungal species with >5% relative frequencies, were only found on Yakushima Island (sites 1 and 2). In contrast, none of the major ECM fungi were exclusive to Tanegashima Island (sites 3 and 4).

Cenococcum geophilum, Clavulinaceae sp.1, Clavulinaceae sp.3 and Phylloporus sp.1 were found in all of the host groups (two conifer species and Fagaceae, Table 2). In contrast, Rhizopogon sp.1 and Thelephoraceae sp.1 were only found in P. amamiana, with a >5% relative frequency, while Boletaceae sp.2, Elaphomyces sp.1, Lactarius sp.1 and Phylloporus sp.2 were only found in Fagaceae. ECM fungal communities were separated by host groups (Fig 2, S4 Appendix), with statistical significance, as determined by the Adonis test \(\text{pseudo-F}_{3,4} = 2.06, R^2 = 0.26, P < 0.01\) in combination with the Betadisper test \(F_{3,7} = 3.241, P = 0.101\). The effect of site was also significant by the Adonis test \(\text{pseudo-F}_{3,4} = 2.56, R^2 = 0.49, P < 0.01\), yet it was affected by the difference in data dispersion (Betadisper test: \(F_{3,6} = 8.484, P = 0.014\)).

Bioassay experiments

By the end of the growth period, three, fifteen, two and seven seedling mortalities occurred among the P. amamiana, P. parviflora, P. densiflora and C. sieboldii soil samples, respectively (S5 Appendix). ECM formation was observed in 100%, 81.3%, 75.3% and 80.6% of seedlings in P. amamiana, P. parviflora, P. densiflora and C. sieboldii, respectively. Five, four, nine and
Table 2. Ectomycorrhizal (ECM) fungal species detected on resident tree roots from *Pinus amamiana* forests. The number of soil samples containing each ECM fungus is shown, along with sequence accession numbers and BLAST results.

| Experiment types | Among-forests comparison | Host | DDBJ accession No. | Seq. length (bp) | Best BLAST match |
|------------------|--------------------------|------|-------------------|-----------------|----------------|
| Sites            | 1 2 3 4                  |      |                   |                 |                |
| ECM fungi        |                          |      |                   |                 |                |
| *Amanita* sp.1   | 0 1 0 0                  | T    | 559               | KP711844        | 98  96         |
| *Amanita* sp.2   | 1 0 0 0                  | P    | 639               | AB922858        | 99  99         |
| *Amanita* sp.3   | 0 1 0 0                  | P    | 688               | KC424544        | 100 99        |
| *Amanita* sp.4   | 0 0 1 0                  | F    | 203               | JO991635        | 100 96        |
| *Amanitaceae* sp.1 | 1 0 0 0                | F    | 393               | KP276311        | 89  89         |
| *Austroboletus* sp.1 | 1 0 0 0               | F, T | 616               | JQ991650        | 91  99         |
| *Boletellus* aurocontextus | 0 0 1 1              | F, P | 798               | AB898004        | 100 99        |
| *Boletus* sp.1   | 1 1 0 0                  | F    | 639               | AB973752        | 100 99        |
| *Boletus* sp.2   | 0 0 1 0                  | F    | 694               | KX756398        | 99  90         |
| *Boletus* sp.3   | 0 0 1 0                  | -    | 644               | AB973811        | 99  90         |
| *Boletaceae* sp.1 | 1 5 0 0               | F, P, T | 439             | AB972824        | 99  100        |
| *Boletaceae* sp.2 | 1 0 4 0               | F    | 794               | KM198314        | 93  93         |
| *Boletaceae* sp.3 | 0 1 0 0               | F    | 692               | KJ676960        | 100 92        |
| *Boletaceae* sp.4 | 0 1 0 0               | F    | 262               | KC552019        | 100 94        |
| *Boletaceae* sp.5 | 0 1 1 0               | F    | 806               | KC551993        | 100 99        |
| *Boletaceae* sp.6 | 0 2 0 0               | P    | 765               | AB973727        | 100 99        |
| *Boletaceae* sp.7 | 0 0 1 0               | P    | 630               | AB509871        | 59  99         |
| *Boletaceae* sp.8 | 1 0 0 0               | P    | 652               | KM595001        | 99  92         |
| *Boletaceae* sp.9 | 0 1 0 0               | F    | 572               | JQ991917        | 100 100       |
| *Boletaceae* sp.10 | 0 0 0 1             | F    | 675               | JF273511        | 100 99        |
| *Cantharellaceae* sp1 | 1 0 0 0              | F, T | 565               | KT200524        | 84  93         |
| *Cenococcum geophilum* | 12 6 15 0           | F, P, T | -               | -               | -             |
| *Ceratobasidiaceae* sp.1 | 0 1 0 1       | F, P, T | 584             | AB605643        | 100 100       |
| *Ceratobasidiaceae* sp.2 | 0 1 0 0       | P    | 597               | AB605649        | 98  96         |
| *Ceratobasidium* sp.1 | 0 0 0 1     | P    | 612               | AB303058        | 100 99        |
| *Ceratobasidium* sp.2 | 0 0 0 1     | P    | 602               | JQ991676        | 93  98         |
| *Clavulina* sp.1 | 0 0 0 1               | F    | 569               | JF273519        | 100 99        |
| *Clavulinaeae* sp.1 | 2 0 0 0           | F, P, T | 641             | AM412265        | 100 89        |
| *Clavulinaceae* sp.2 | 2 0 0 0           | F, P, T | 711             | AB807913        | 92  99        |
| *Clavulinaceae* sp.3 | 5 0 0 0           | F, P, T | 403             | AB807910        | 94  99        |
| *Clavulinaceae* sp.4 | 1 0 0 0           | P    | 655               | KC876295        | 100 87        |
| *Clavulinaceae* sp.5 | 0 1 0 0           | P    | 592               | AB848424        | 100 100       |
| *Clavulinaceae* sp.6 | 0 1 0 0           | F    | 235               | KC876295        | 100 89        |
| *Clavulinaceae* sp.7 | 1 0 0 0           | P    | 601               | FR731325        | 100 88        |
| *Coltricia* sp.1 | 1 0 0 0               | P    | 268               | KU360702        | 100 95        |
| *Cortinarius* sp.1 | 0 0 2 0             | F, P | 737               | HQ604690        | 100 93        |
| *Cortinarius* sp.2 | 0 0 1 0             | F    | 616               | AB848438        | 100 99        |
| *Cortinarius* sp.3 | 0 0 1 0             | F    | 660               | LC096896        | 100 92        |
| *Cortinarius* sp.4 | 0 1 0 0             | F    | 780               | AB973753        | 100 99        |
| *Cortinarius* sp.5 | 0 0 0 1             | F, P | 507               | HQ285384        | 100 96        |
| *Craterellus* sp.1 | 0 1 0 0             | F    | 237               | JQ991672        | 99  94        |
| *Elaphomyces* sp.1 | 1 0 3 0             | F    | 535               | JQ991717        | 92  95        |
| *Elaphomyces* sp.2 | 0 1 0 0             | F    | 417               | KX165351        | 98  97        |
| *Elaphomyctaceae* sp.1 | 0 0 1 0           | F    | 337               | JQ991901        | 99  98        |

(Continued)
| Experiment types | Among-forests comparison | Host | DDBJ accession No. | Seq. length (bp) | Best BLAST match |
|------------------|--------------------------|------|--------------------|-----------------|------------------|
| ECM fungi        |                          |      |                    |                 |                  |
| Entoloma sp.1    | 0 0 1 0 F                | 554  | KP403072           | 100             | 97               |
| Hydnellum sp.1   | 0 0 1 0 F                | 602  | EU293832           | 84              | 92               |
| Hydnellum sp.2   | 0 0 1 0 P                | 601  | KM403015           | 71              | 97               |
| Hydnum sp.1      | 1 0 0 0 P                | 555  | KU612573           | 94              | 99               |
| Hydnum sp.2      | 0 0 1 0 F                | 569  | AB906676           | 100             | 100              |
| Hydnum sp.3      | 1 0 0 0 P                | 392  | AB251813           | 100             | 100              |
| Hymenochaetaceae sp.1 | 0 1 0 0 P | 704  | JC991687           | 91              | 89               |
| Hymenochaetaceae sp.2 | 1 0 0 0 P | 710  | KM594987           | 99              | 91               |
| Inocybe sp.1     | 1 0 0 0 T                | 528  | AM882711           | 100             | 95               |
| Lactarius sp.1   | 2 1 0 0 F                | 625  | AB777482           | 100             | 99               |
| Lactarius sp.2   | 0 0 2 0 F, P             | 654  | LC096473           | 100             | 99               |
| Lactarius sp.3   | 1 0 0 0 F                | 638  | JC991640           | 96              | 99               |
| Lactarius sp.4   | 0 1 0 0 T                | 656  | GQ268638           | 100             | 98               |
| Lactarius sp.5   | 0 1 0 0 F                | 573  | AB973742           | 100             | 99               |
| Lactarius sp.6   | 0 1 0 0 F                | 543  | JC991763           | 97              | 94               |
| Pezizaceae sp.1  | 0 1 0 0 F                | 565  | JN102406           | 99              | 94               |
| Pezizaceae sp.2  | 0 0 0 1 F                | 360  | JC991767           | 100             | 95               |
| Phylloporus sp.1 | 7 1 0 0 F, P, T          | 835  | DQ533980           | 99              | 91               |
| Phylloporus sp.2 | 0 1 2 0 F                | 775  | AB973776           | 82              | 99               |
| Phylloporus sp.3 | 0 1 1 0 F                | 765  | JC967270           | 82              | 96               |
| Rhizopogon sp.1  | 5 3 1 2 P                | 754  | LC216339           |                 |                  |
| Rhizopogon sp.2  | 0 0 0 1 Ps               | 286  | AB923020           | 99              | 99               |
| Rossbeevera griseovelutina | 0 1 0 0 F | 505  | KC551986           | 100             | 99               |
| Russula sp.1     | 3 0 6 0 F, P             | 768  | AB507012           | 84              | 99               |
| Russula sp.2     | 0 0 2 0 F                | 600  | JX556180           | 99              | 99               |
| Russula sp.3     | 0 0 1 0 F, P             | 625  | JX987768           | 99              | 99               |
| Russula sp.4     | 2 0 0 0 F, P             | 608  | LC096863           | 100             | 99               |
| Russula sp.5     | 1 0 0 0 F                | 641  | JC991802           | 94              | 99               |
| Russula sp.6     | 0 0 1 0 F                | 687  | AB507006           | 88              | 99               |
| Russula sp.7     | 1 0 0 0 F                | 712  | LC096934           | 100             | 96               |
| Russula sp.9     | 1 0 0 0 F                | 488  | AB509875           | 82              | 99               |
| Russula sp.10    | 0 1 0 0 P                | 630  | AB594957           | 94              | 99               |
| Russula sp.11    | 0 0 1 0 -                | 526  | EF634135           | 100             | 90               |
| Russula sp.12    | 0 0 1 0 F                | 573  | AB629011           | 100             | 99               |
| Russula sp.13    | 0 0 1 0 -                | 535  | AB636110           | 97              | 99               |
| Russula sp.14    | 0 0 1 1 F, P             | 718  | LC096811           | 100             | 100              |
| Russula sp.15    | 1 1 0 1 F, P             | 736  | KJ769295           | 99              | 99               |
| Russula sp.16    | 1 0 0 0 -                | 502  | JN129409           | 100             | 96               |
| Russula sp.17    | 0 0 0 1 F                | 608  | AB291756           | 98              | 99               |
| Russula sp.18    | 0 0 0 1 F                | 603  | JC991823           | 97              | 99               |
| Russula sp.19    | 1 0 0 0 F                | 615  | LT602972           | 99              | 97               |
| Russula sp.20    | 1 0 0 0 F                | 614  | JC991790           | 99              | 99               |
| Russula sp.21    | 1 0 0 0 P                | 594  | AB769909           | 100             | 95               |
| Russula sp.23    | 1 0 0 0 F                | 599  | AB218154           | 100             | 99               |

(Continued)
seven ECM fungi were identified from 208 *P. amamiana*, 246 *P. parviflora*, 257 *P. densiflora* and 191 *C. sieboldii* DNA samples, respectively (Table 3 and S3 and S5 Appendices). None of the control seedlings had ECM roots.

In bioassays of the subgenus *Strobus*, six ECM fungal species were found, of which three were shared between *P. amamiana* and *P. parviflora* (Table 3, Fig 3). Nine and seven ECM fungal species were detected in the *P. densiflora* and *C. sieboldii* bioassays, respectively. The estimated species richness (Jackknife2) was 7.9, 7.9, 20.7 and 13.8 in *P. amamiana*, *P. parviflora*, *P. densiflora* and *C. sieboldii*, respectively.

*Rhizopogon* sp.1 was the most dominant ECM fungal species detected in the *P. amamiana* bioassay, found in 40 of 43 (93% frequency) soil samples (Fig 3). This fungus was also found in *P. parviflora* and *P. densiflora* samples but at lower frequencies, 67% and 31%, respectively (Fig 3). *Rhizopogon* sp.1 was not found in any of the *C. sieboldii* bioassays.

*Cenococcum geophilum* was the most common fungal species in the *P. densiflora* and *C. sieboldii* bioassays (Fig 3). Among the 16 ECM fungal species found in bioassays, only *C. geophilum* was shared among the four host species (Fig 3).

Soil spore bank communities were clearly separated by host and site using NMDS ordination (Fig 4), with statistical significance determined by the Adonis test (host: *pseudo-F*₁,₈ = 12.08, *R²* = 0.69, *P* < 0.01; site: *pseudo-F*₂,₆ = 5.12, *R²* = 0.2, *P* < 0.01) in combination with the Betadisper test (host: *F*₁,₈ = 1.351, *P* = 0.325; site: *F*₂,₉ = 1.819, *P* = 0.206). A significant difference (S6 Appendix; *pseudo-F*₁,₁₈ = 5.76, *R²* = 0.24, *P* < 0.01) was found between soil propagule banks and resident ECM fungal communities on mature trees in these same three forests, however, it was affected by significant difference in data dispersion among the groups (Betadisper, *F*₁,₁₈ = 44.257, *P* < 0.001).

**Discussion**

We found *Rhizopogon* sp.1 in all remaining *P. amamiana* forests. Host identification of the resident ECM roots confirmed that *Rhizopogon* sp.1 was associated only with the endangered
species *P. amamiana*. In soil spore banks, *Rhizopogon* sp.1 was far more dominant than in resident ECM roots and was identified in nearly all soil samples examined (93%) among the major remaining *P. amamiana* populations. These findings generally support both our hypotheses, confirming the existence of an ECM fungus specific to the endangered pine species in its natural settings and the predominance of this fungus in soil spore banks [12]. *Rhizopogon* sp.1 is very likely to have a close ecological relationship with the endangered *P. amamiana*, probably due to long co-evolutionary periods on isolated islands.

Dominant ECM fungi in soil spore banks generally contribute to establishment of tree seedlings, especially for pine trees regenerating after a disturbance [46–48]. Thus, the *Rhizopogon* sp.1 found in this study likely also contributes to seedling establishment of the endangered *P. amamiana* after a disturbance [49, 50]. Although no other native pine species are distributed in the remaining forests of *P. amamiana*, some broadleaf trees could compete with *P. amamiana* during regeneration. It should be noted here that the dominant broadleaf tree *C. sieboldii* was not compatible with *Rhizopogon* sp.1 in the bioassay experiments. Alternatively, *C. sieboldii* bioassay seedlings were dominated by the true generalist *C. geophilum*, sclerotia of which

Fig 1. Sample-based rarefaction curves for ectomycorrhizal (ECM) fungi found in *Pinus amamiana* forests. Black circles and triangles represent observed ECM fungal species richness for all host species and for *P. amamiana*, respectively. Jackknife2 minimal species richness estimates of ECM fungi are also shown for all host species (white circles) and for *P. amamiana* (white triangles).

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exist in almost all forests investigated (e.g. [8, 11, 12, 15]). Murata & Nara [51] suggested that *C. geophilum* competes with other ECM fungi, thus affecting the infection rates of other soil spore bank fungi. While ECM symbiosis is a prerequisite for trees to grow and survive in nature [3], controlling ECM fungal communities may be difficult, especially after forest development. Eliminating other ECM fungi from soil spore banks in disturbed sites via treatments, such as heating [12, 52–54] or filtering [51], may allow an increase in the relative frequency of *Rhizopogon* sp.1 and eventually increase regeneration of the endangered *P. amamiana* preferentially.

We found no sequences that match *Rhizopogon* sp.1 in an international nucleotide sequence database (DNA Data Bank of Japan [DDBJ]/EMBL/GenBank), potentially indicating its endemism to these islands. In fact, based on the morphological characteristics of the sporocarps and phylogenetic relationships with other described *Rhizopogon* species, we are now describing it as a new species, which does not belong to any subgenus proposed by Grubisha et al. [55] (Sugiyama et al. under revision). The existence of a *Rhizopogon* fungus specific to an endangered tree was documented in our recent study of *Pseudotsuga* species in Japan and

**Fig 2.** Non-metric multidimensional scaling (NMDS) depicting ECM fungal communities of resident trees in four endangered *Pinus amamiana* forests. Stress = 0.089. White, gray and black symbols indicate the communities on *P. amamiana*, *Tsuga sieboldii* and Fagaceae, respectively. Circles, diamonds, squares and triangles represent communities at sites 1, 2, 3 and 4, respectively. 

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China; i.e., *P. japonica* is associated with *R. togasawariana* [12, 32] and *P. sinensis* with an undescribed *Rhizopogon* species [56]. Here, we first confirmed the existence of a *Rhizopogon* fungus specifically associated with endangered *Pinus*, which is a far larger genus than *Pseudotsuga* and includes far more endangered species. Given the predominance of the host-specific fungus in soil spore banks and its potential roles in seedling regeneration, such fungi should be explored further in other endangered *Pinus* species that have been geographically isolated for long periods. Without identifying such key ECM fungi, conservation of endangered pine species will be difficult.

*Rhizopogon* sp.1 was associated solely with the endangered *P. amamiana* under natural settings. This fungus was also compatible with *P. parviflora* and *P. densiflora* in bioassays, but the colonization frequency was reduced significantly with increasing phylogenetic distance. *P. amamiana* is closely related to *P. armandii* var. *armandii* and *P. armandii* var. *mastersiana* [57, 58], belonging to the same clade of subsection *Strobus* found in East Asian subtropical areas [29, 57], as the divergence of *P. amamiana* from *P. parviflora* or *P. densiflora* dates approximately 6.4 Mya or 85 Mya, respectively [29]. The partial compatibility with these distantly related pine species may indicate that the isolation period has not been long enough for compatibility to be lost completely. In previous studies of *R. togasawariana* associated with *Pseudotsuga japonica*, colonization in *P. densiflora* was totally absent in a similar bioassay, but it was compatible with North American *Pseudotsuga menziesii* [12]. *R. togasawariana* belongs to the subgenus *Villosulii*, which is specific to *Pseudotsuga*, and its origin precedes the migration of the host to Asia approximately 34 Mya. Although *Rhizopogon* is regarded as a host-specific ECM fungal lineage [59], its compatibility is not strict within the same host genus. Yet, the compatibility with non-native hosts may not be fully functional in terms of nutrient transfer and host performance.

### Table 3. Ectomycorrhizal (ECM) fungal species detected from soil spore banks in *Pinus amamiana* forests by bioassay experiments using four host species.

| Experiment types | Among-forests comparison | DDBJ accession No. | Seq. length (bp) | Best BLAST match |
|------------------|--------------------------|---------------------|-----------------|-----------------|
| Bioassay hosts a | Pa | Pp | Pd | C |
| Sites            | 1  | 2  | 3  | 1  | 2  | 3  | 1  | 2  | 3  | 1  | 2  | 3  | 1  | 2  | 3  |
| Atheliaceae sp.1 | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 652 | AB456674 | 84  |
| *Cenococcum geophilum* | 6  | 2  | 10 | 8  | 8  | 18 | 14 | 8  | 29 | 14 | 16 | 26 | -   | -   | -   |
| Clavulinaceae sp.3 | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 473 | JQ991682 | 100  |
| Craterellus sp.2 | 1  | 0  | 0  | 1  | 0  | 0  | 1  | 0  | 1  | 0  | 0  | 0  | 516 | AB973729 | 100  |
| Elaphomyces sp.2 | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 1  | 0  | 0  | 0  | 0  | 417 | KX165351 | 98   |
| Laccaria vinaceavellanea | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 1  | 0  | 0  | 0  | 0  | 480 | KJ609167 | 100  |
| Pezizaceae sp.3 | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 3  | 539 | AB571493 | 98   |
| *Rhizopogon* sp.1 | 17 | 6  | 17 | 4  | 16 | 22 | 3  | 5  | 16 | 0  | 0  | 0  | 754 | LC216339 | -   |
| *Rhizopogon* sp.2 | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 286 | AB923020 | 99   |
| *Russula* sp.25 | 1  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 1  | 541 | AB253521 | 100  |
| *Suillus* bovinus | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 537 | HE814113 | 100  |
| Thelephoraceae sp.7 | 2  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 1  | 0  | 608 | JX456851 | 100  |
| Thelephoraceae sp.8 | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 0  | 1  | 597 | GU907787 | 100  |

**a** Pa, Pp, Pd and C indicate *Pinus amamiana*, *P. parviflora*, *P. densiflora* and *Castanopsis sieboldii*, respectively.

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Another interesting finding of this study is that *P. amamiana* was not associated with any *Suillus* species, either in ECM roots of resident trees or in soil spore banks. *Suillus* produces epigeous sporocarps for spore dispersal by wind but is a phylogenetic sister to *Rhizopogon* [55]. While the life span of *Suillus* spores is shorter than that of *Rhizopogon*, it is frequently recorded in soil spore banks [20], probably because of its strong spore dispersal abilities [60, 61] and good germination rates [62]. Some *Suillus* species are specific to, or prefer, the subgenus *Strobus* within the genus *Pinus* [63–65], and these fungi have wide intercontinental distributions. For example, *Suillus spraguei* is found in *Strobus* pine forests from North America to East Asia [63]. In Japan, *S. spraguei* is associated with *P. parviflora, P. pumila* and *P. koraiensis* [64, 66], all of which belong to the subgenus *Strobus*. In addition, *S. spraguei* and some closely related species are associated with *P. armandii*, which is closely related to *P. amamiana* [63,
The nearest pine population belonging to *Strobus, P. parviflora*, is located in the Takaku-mayama mountains on the main island of Kyushu, which is its southern population limit [68], and this population is located >100 km away from the remaining *P. amamiana* forests by sea. Therefore, the absence of *Suillus* species in the remaining *P. amamiana* forests may indicate the difficulty of airborne spore dispersal from distant *Strobus* forests together with local extinction on Yakushima and Tanegashima Islands.

In the bioassay experiment, it was difficult to characterize the effect of ECM infection on the initial growth of seedlings, because the experimental period was too short and the bioassay tubes were too small to evaluate the growth of *P. amamiana*. Therefore, we did not compare the growth of seedlings infected with ECM fungi with that of control plants in the bioassays. Instead, seedlings infected with *Rhizopogon* sp.1 and uninfected seedlings were raised for 1 year in a separate bioassay experiment, after which growth was compared. Seedlings infected with *Rhizopogon* sp.1 exhibited increased growth compared with control seedlings, with marginal statistical significance (S7 Appendix) and the difference would become much larger with the growth period. Moreover, *P. amamiana* seedlings colonized by *Rhizopogon* sp.1 are much more tolerant to transplantation than are uncolonized seedlings (S8 Appendix). Thus, even in a nursery setting, the application of *Rhizopogon* sp.1 is a promising approach to producing good seedlings for transplantation to conserved areas.

Supporting information

S1 Appendix. Location of four study sites in Japan.
(PDF)

S2 Appendix. Air-dried periods, growth periods and number of sample of each tree species
in bioassay experiment.
(PDF)

S3 Appendix. Blast results of ectomycorrhizal fungal species identified in this study using
UNITE database.
(PDF)

S4 Appendix. Non-metric multidimensional scaling (NMDS) depicting ECM fungal com-
munities of resident trees in four endangered *Pinus amamiana* forests based on individual
soil samples. Stress = 0.142. White, gray and black symbols indicate the communities on Fag-
ceae, *P. amamiana* and *Tsuga sieboldii*, respectively. Circles, squares, diamonds and triangles
represent communities at sites 1, 2, 3 and 4, respectively. The effects of both host and site were
significant (Adonis, *P*<0.01) after confirming data variance among the groups was not signifi-
cant (Betadisper, *P>*0.05).
(PDF)

S5 Appendix. Summary of bioassays using soil from three endangered *Pinus amamiana*
forests.
(PDF)

S6 Appendix. Comparison of ectomycorrhizal fungal communities between soil propagule
banks and resident trees in endangered *Pinus amamiana* forests using non-metric dimen-
sional scaling (NMDS). Each symbol represents a community in each host per site.
Stress = 0.134. White and black symbols indicate the communities assayed with resident trees
and soil propagule banks, respectively.
(PDF)
S7 Appendix. In the bioassay experiment different from this study (unpublished data), the number of leaves and tree height of seedlings infected with *Rhizopogon* sp.1 and seedlings not infected with ECM fungi (control) grown for 1 year. Significant probability between *Rhizopogon* sp.1 and Control of leaf number and tree height by T test was p = 0.08 and p = 0.06, respectively. The bar shows the standard error.

(S7)

S8 Appendix. From a bioassay experiment separate from this study (unpublished data), transplantation of seedlings infected with *Rhizopogon* sp.1 (upper) and uninfected seedlings (lower) to new pots after 1 year of growth.

(S8)

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