1	1.	Title:
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2	Mycorrhizal associations in woody plant species at the Mt. Usu volcano, Japan.
3	2. Informative title
4	Mycorrhizal associations in woody plant species at volcano Usu.
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19	

1 **Title:**

2 Mycorrhizal associations in woody plant species at the Mt. Usu volcano, Japan.

3

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6

7 Abstract

8 We investigated the association between ectomycorrhizal (ECM) and arbuscular mycorrhizal (AM) fungi and pioneer woody plant species in areas devastated by the 9 10 eruption of Mt. Usu, Japan, in 2000. We observed 8 woody plant species at the research site, most of which were associated with ECM and/or AM fungi. In particular, 11 dominant woody plant species Populus maximowiczii, Salix hultenii var. angustifolia 12 13 and Salix sachalinensis were consistently associated with ECM fungi and erratically 14 associated with AM fungi. We found 1 to 6 morphotypes in the roots of each ECM host and on average 2 in the roots of each seedling, indicating low ECM fungal diversity. 15 ECM colonization ranged from 17 to 42% of root tips. Using morphotyping and 16 molecular analyses, 15 ECM fungi were identified. ECM fungi differed greatly 17 18 between hosts. However, Laccaria amethystea, Hebeloma mesophaeum, Thelephora 19 terrestris and other Thelephoraceae had high relative colonization, constituting the

1	majority of the ECM colonization in the roots of each plant species. These ECM fungi
2	may be important for the establishment of pioneer woody plant species and further
3	revegetation at Mt. Usu volcano.
4	
5	Key words;
6	Mycorrhizal association, Ectomycorrhizal fungi, Woody plant, Disturbed area, Volcano.
7	
8	Introduction
9	Woody plant species invade and become established in devastated areas immediately
10	following volcanic eruption, despite the presence of environmental stresses such as
11	low soil nutrients, instability of the soil surface and drought (Goto 1937; Yoshii 1942;
12	Tsuyuzaki 1987). These woody plant species, called pioneer species, contribute to
13	vegetation recovery by facilitating the establishment of later seral vegetation (Walker
14	and del Moral 2003).
15	Ectomycorrhizal (ECM) hosts such as the Salicaceae often dominate areas
16	devastated by volcanic eruption (Goto 1937; Yoshii 1942; Tsuyuzaki 1987). The
17	dominant woody plant species at our Mt. Usu study site are Salix sachalinensis Fr.
18	Schm., Salix hultenii var. angustifolia Kimura and Populus maximowiczii A. Henry,
19	which belong to a family usually colonized by ECM and arbuscular mycorrhizal (AM)

1	fungi. These species are considered to be significant for future reforestation. ECM
2	fungi enhance the growth of host plant species: recent studies have revealed
3	coinoculation with various ECM fungi can alter host growth and nutrient acquisition
4	(Reddy and Natarajan 1997; Baxter and Dighton 2001). Thus, the composition of the
5	ECM fungal community influences establishment of host plant species and to
6	understand the effect of ECM associations on growth and survival of host plants, it is
7	important to know which species comprise a given community.
8	Although few studies have examined ECM associations in woody plant species
9	established in devastated areas, efforts have been made to describe the ECM fungi
10	involved in primary succession. Jumpponen et al. (2002) investigated the
11	chronosequence of ECM fungi occurring at the front of the Lyman Glacier. They noted
12	that the occurrence of ECM fungal sporocarps varies according to the time since
13	deglaciation, indicating an early and late stage model for succession. Allen et al.
14	(1992) noted that several years after the last eruption of Mt. St. Helens, several woody
15	plant species were associated with ECM fungi. Yang et al. (1998) investigated the
16	occurrence of ECM morphotypes in Larix kaempferi (Lamb.) Carr. at the Mt. Koma
17	volcano, Japan. They demonstrated that, as with litter accumulation and soil conditions,
18	the composition of ECM morphotypes varied with elevation, emphasizing the
19	importance of ECM diversity for survival and growth of seedlings. Recently, molecular

analyses have been applied to mycorrhizal research in order to differentiate and 1 2 identify ECM fungi. Using polymerase chain reaction (PCR) amplification of the internal transcribed spacer (ITS) region of fungal nuclear ribosomal DNA (rDNA), 3 4 morphologically similar ectomycorrhizae can be distinguished and identified by their restriction fragment length polymorphism (RFLP) patterns and sequencing, 5 respectively. Nara et al. (2003a, b) used both conventional morphotyping and 6 7 molecular analyses to reveal the presence of ECM flora in the roots of Salix reinii 8 Franch. et Savat. and demonstrate succession in underground ECM fungi from a volcanic desert on Mt. Fuji. Ashkannejhad and Horton (2006) investigated the ECM 9 10 flora of Pinus contorta var. contorta seedlings on coastal sand dunes. However, little is known about ECM flora in areas devastated by volcanoes, particularly in the period 11 immediately following cessation of volcanic activity. 12 Using morphotyping and molecular analyses we investigated: 1) the status of 13 mycorrhizal associations in seedlings of woody plant species; and 2) the underground 14 ECM fungal community associated with pioneer woody plant species in areas 15

16 devastated by the 2000 eruption of Mt. Usu.

17

18 Materials and Methods

19 Mt. Usu $(42^{\circ} 32' \text{ N}, 140^{\circ} 50' \text{ E}; 773.1 \text{ m asl})$ is an active volcano located in

southwest Hokkaido, Japan (Fig. 1) that has erupted repeatedly since 1663. It erupted 1 2 again on March 31, 2000, 22 years after the previous eruption. A number of small craters formed at the foot of the Nishiyama and Konpira areas, and were accompanied 3 by the accumulation of a considerable amount of volcanic debris. Ejection of debris 4 subsided in autumn 2000, but the effects of thermal activity such as elevated soil 5 temperatures, as well as the emission of noxious gases, continued near the K-A, K-B 6 and N-B craters. Prior to the 2000 eruption, there was a natural secondary forest 7 8 comprising broadleaf species such as Betula spp., Acer spp. Quercus spp. and Magnolia spp., and a partially planted forest of L. kaempferi and Abies sachalinensis 9 10 (Fr. Schm.) Masters. However, the deposition of 1-3m of volcanic debris (fine volcanic ash and pumice) devastated ca. 71ha of forest around the craters. This study was 11 conducted in the devastated area around the N-A crater and at the foot of the 12 13 Nishiyama area, where it appeared that volcanic activity had ceased as we observed no emissions of volcanic gases or elevation of soil temperature. In 2004, 15 woody plant 14 species had established near the Nishiyama area craters, reaching a total density of 15 1038ha⁻¹. The mean growth rate of the dominant species (S. sachalinensis) was ca. 16 10cm year⁻¹. Thus, conditions at present remain unfavorable for the establishment of 17 18 woody plant species. In 2004, climatic data from the Sapporo meteorological station at Date (42° 30' N, 140° 54' E; 84.7m asl), indicated a mean annual precipitation of 19

835mm and annual temperature of 8.9°C, ranging between -12.0 and 30.8°C
 (December to August, respectively).

3

4 Sampling procedure

In May 2004, we established a 4-ha research site encompassing several craters in which no trees had survived the 2000 eruption and where all understory vegetation had disappeared due to the deposition of volcanic debris (> 1m). From June to September 2004, we randomly selected 1-12 seedlings from each woody plant species and sampled their lateral roots, which extended from the soil surface to a depth of 15cm.

10

11 ECM and AM associations

We investigated the ECM and AM associations in each woody plant species with > 612 13 seedlings. Adhering soil was separated from the roots by soaking and careful washing 14 of samples in tap water. Appearance and the presence of a mantle and Hartig net was used to identify ECMs under differential interference microscopy (400-1000x 15 magnification). AMs were identified using the staining procedure described by Phillips 16 and Hayman (1970), with some modifications. Roots were rinsed with distilled water, 17 18 cleared with 10% KOH for 80 min at 80°C, bleached in 0.5% H₂O₂ for 10-20 min at 60°C, acidified in 1% HCl at room temperature (ca. 15-20°C), and stained using 0.05% 19

trypan blue in lactophenol for 15 min at 80°C. AM colonization was identified by the
 presence of vesicles or arbuscules, as well as internal hyphae.

3

4 Determination of mycorrhizal colonization

5 We focused on ECM hosts and investigated their underground ECM fungal flora. 6 The overall morphologies of ECMs were observed under stereoscopic microscopy. 7 ECMs from each woody plant species were classified into morphological groups and 8 divided into two subsamples: one was placed in FAA solution (formaldehyde: acetic 9 acid: ethyl alcohol: distilled water = 1:1:9:9) for microscopic investigation and the 10 other stored at -80°C for DNA extraction.

ECM abundance was estimated as the proportion of each morphotype relative to the total ECM. Frequency was estimated as the proportion of seedlings colonized by one morphotype relative to all seedlings.

14

15 DNA extraction, PCR amplification and RFLP

The samples contained one ECM root tip of each morphotype from each seedling; 3 to 5 samples from each morphotype identified in a given woody plant species were categorized individually by PCR-RFLP. ECM fungal DNA was extracted from 5-10mg ground, lyophilized tissue using the DNeasy Plant Mini kit (QIAGEN, USA) according

1	to the manufacturer's instructions. The ITS region, including the 5.8S rDNA, was
2	amplified using a specific primer for higher fungi (ITS1-f; Gardes and Bruns 1993) and
3	a universal primer (ITS4; White et al. 1990). The following PCR amplification
4	conditions were used: 94°C for 3 min, followed by 30 cycles of 94°C for 1 min, 50°C
5	for 1 min and 72°C for 3 min, then a final extension at 72°C for 10 min (Landeweert et
6	al. 2005).
7	Single enzyme digests using <i>Hin</i> fI and <i>Alu</i> I were performed on PCR products from 3
8	to 5 ECM root tips of each ECM morphotype. Using 2.5% agarose gel electrophoresis,
9	we determined the quality and quantity of the PCR products, as well as the size of
10	restriction fragments. Band lengths were calculated using KiloACE
11	(http://www.nih.go.jp/%7Ejun/cgi-bin/kiloace.pl).

12

13 Sequencing

We used the primer ITS1f to sequence samples of each PCR product arising from different ECM morphotypes and exhibiting differences in RFLP analysis. Sequencing reactions were performed using the BigDye Terminator v3.1/1.1 Cycle Sequencing Kit (Applied Biosystems, USA), followed by ethanol precipitation and analysis with an ABI Auto Sequencer 310 (Applied Biosystems, USA). ECM sequences were compared with the GenBank database at the DNA Data Bank of Japan (DDBJ) using the BLAST program, and species names were assigned to BLAST matches exhibiting > 95%
 homology.

3

4 Sporocarps of ECM fungi

Although no ECM fungal sporocarps were found during preliminary work at the 5 6 study site, we identified 9 taxa in the area that contained surviving mature trees and herbaceous plants (Obase et al. 2005). These were identified microscopically as 7 8 Laccaria sp., Inocybe nitidiuscula (Britzelm.) Sacc., Inocybe dulcamara (Pers. Albertini and Schweinitz) P. Kumm., Hebeloma crustuliniforme (Bull. Fr.) Quel., 9 10 Hebeloma mesophaeum complex, Hebeloma sp., Suillus laricinus (Berk. in Hook.) O. Kuntze, Suillus grevillei (Klotzsch Fr.) Singer and Scleroderma bovista Fr. In order to 11 perform alignments between above- and below-ground fungal sequences, we extracted 12 and sequenced DNA from these ECM sporocarps. The procedures for DNA extraction, 13 14 PCR amplification and sequencing were as described above, except that the ratio of DNA template to sterilized distilled water was altered from 9:16 to 1:24 in the PCR 15 procedure. 16

17

18 **Results**

19 Mycorrhizal association

We observed 8 woody plant species at the research site (Table 1) and observed ECM 1 2 colonization in almost all seedlings of Betula platyphylla Sukatchev var. japonica (Miq.) Hara, Quercus crispula Blume, P. maximowiczii, S. hultenii var. angustifolia, 3 Salix integra Thunb. and S. sachalinensis. AM colonization was detected in the roots 4 of B. platyphylla var. japonica, P. maximowiczii, Q. crispula, S. hultenii var. 5 angustifolia, S. sachalinensis, Acer mono Maxim. var. marmoratum (Nichols.) Hara f. 6 dissectum (Wesmael) Rehder and Rosa multiflora Thunb. We also observed AM and 7 8 ECM co-colonization in some seedlings of B. platyphylla var. japonica, P. maximowiczii, Q. crispula and S. hultenii var. angustifolia, but the association 9 10 frequency of the former was lower than that of the latter. In general, we observed association with ECM and/or AM fungi in most woody plant species and with the 11 12 exception of A. mono var. marmoratum f. dissectum, we found that the dominant 13 woody plant species were associated consistently with ECM fungi and erratically with AM fungi. 14

15

16 ECM morphotype and colonization

We found between 1 and 6 morphotypes in the roots of each ECM host (Table 2) and 17 to 42% of all root tips were colonized by ECM fungi. On average, 2 morphotypes 19 were observed in the roots of each seedling, except for those of *Q. crispula* and *B.* 1 *platyphylla* var. *japonica*, which had 1 and 4, respectively.

2

3 PCR-RFLP patterns and genetic identification

4	Although DNA amplification using the primers ITS1f and ITS4 resulted in nearly
5	100% amplification of PCR products, some types produced multiple PCR products,
6	possibly because of the presence of other fungi within or around the root tissues. As
7	some samples could not be determined by RFLP analysis alone, we digested the most
8	well-separated and abundant PCR products from each sample with HinfI and AluI, and
9	thus categorized each ECM morphotype (Table 3). With the exception of Lk-2, Ss-2
10	and $Bp-2$, the banding patterns were identical from different samples within each ECM
11	morphotype.
12	Alignment of these sequences with those from GenBank resulted in potential
13	matches for 15 ECM fungi (Table 4). Sequences of 2 ECM morphotypes matched the H.
14	mesopaeum complex and S. bovista sporocarps that were observed in the preliminary

15 study (Obase et al. 2005).

16

17 Colonization by each ECM morphotype

The ECM fungal flora differed between hosts (Table 4) and 11 of the 15 fungi were
observed in the roots of only one host. However, *Laccaria amethystea*, *H. mesophaeum*,

Thelephora terrestris Fr. and Thelephoraceae 1 were observed in the roots of 2, 3, 4
 and 4 ECM hosts, respectively. These fungi were abundant and represented most of the
 ECM colonization in the roots of each woody plant species.

4

5 Discussion

6 In 2000, the study site was strongly disturbed by the eruption, which resulted in the loss of nearly all the plant species that had colonized the site before 2000. Thus, almost 7 8 all seedlings were new recruits that had become established independently on the new substrate. Although the deposition of a thick layer of new volcanic debris around 9 10 craters must presumably make it problematic for woody plant species to associate with mycorrhizal fungi, such associations were nonetheless observed in the roots of 11 newly-recruited seedlings. Following volcanic eruptions, AM and ECM associations 12 reestablish immediately (Allen et al. 1992), and are of major importance to the primary 13 14 succession of plant species in volcanic areas (Titus and Tsuyuzaki 2002; Fujiyoshi et al. 2005; Tsuyuzaki et al. 2005). 15

At the time of eruption, the scale of disturbance in our study area was relatively small (ca. 71ha) and the surrounding forest edge recovered quickly. It would appear that there was a rapid recovery of, or minimal damage to the fungal flora at the forest edge, as Obase et al. (2005) identified a variety of fungal species by investigating sporocarp occurrence. The speed of recovery of both vegetation and fungal flora in
 these edge areas, as well as their proximity to the study site, both play a role in the
 recruitment of mycorrhizal inocula to the devastated area.

Almost all seedlings of the dominant woody plant species P. maximowiczii, S. 4 hultenii var. angustifolia and S. sachalinensis exhibited ECM fungal associations. In 5 2002, only two years after the volcanic eruption, a preliminary study revealed the 6 7 presence of ECM colonization in the roots of Salix. ECM and AM fungi both colonize 8 the Salicaceae (Harley and Harley 1987). In the present study, we observed a very low percentage of AM colonization, with less than half of the seedlings exhibiting an AM 9 10 association. Thus, it appears that AM fungi represent a relatively insignificant factor in the establishment of Salicaceae seedlings, compared to ECM fungi. In a study on Mt. 11 Fuji, Nara (2006) reported a strong relationship between established S. reinii 12 13 individuals and ECM fungal association, but found only rare associations with AM 14 fungi. In contrast, in about 50% the seedlings of A. mono var. marmoratum f. dissectum associated with AM fungi and no ECM fungi were observed in present study. Acer spp. 15 16 have been observed with AM, ECM or non-mycorrhizal associations (Harley and Harley 1987). Under different environmental conditions, some seedlings of A. mono 17 18 var. marmoratum f. dissectum associated with AM fungi but others did not form 19 mycorrhizal associations (unpublished data). Thus, it seems A. mono var. marmoratum f. *dissectum* intrinsically forms erratic relationships with AM fungi during seedling
 stage, that also appeared in primary succession.

3 Analysis of RFLP and sequence data derived from root materials demonstrated that 3 Salicaceae woody plant species that are dominant in the study area harbored 9 ECM 4 fungal taxa, with one woody plant species alone containing 3 to 5 ECM fungal taxa. 5 6 These numbers are very low compared to the high ECM fungal diversity in temperate 7 and boreal forests (Horton and Bruns 2001). In the roots of Salix repens L. established 8 in sand dunes, 78 ECM fungal species were recorded as sporocarps (van der Heijden 1999). Nara et al. (2003a, b) reported 23 ECM species as sporocarps and 21 ECM 9 10 species in the roots of S. reinii established in the volcanic desert on Mt. Fuji. However, in a 6-year-old plantation, a study on the ECM community associated with Salix 11 viminalis L. and Salix dasyclados Wimm. identified only 4 and 7 ECM taxa, 12 respectively (Püttsepp et al. 2004). In addition, Nara et al. (2003a, b) observed only 5 13 14 ECM taxa in young S. reinii seedlings. As the seedlings investigated in the present study were young, their age and isolation from mature trees will have had an influence 15 on the diversity of their ECM communities. Ashkannejhad and Horton (2006) revealed 16 that both the ECM diversity per seedling and the total number of ECM fungi was lower 17 18 in isolated dunes than in forests. They also showed that some ECM fungi found in the 19 forest also colonized seedlings in sand dunes. Thus, it appears that isolated seedlings that are undergoing primary succession are only able to associate with a limited range
 of early-stage ECM fungi.

3 We observed Hebeloma, Laccaria, Thelephoraceae species consistently in the roots of the dominant woody plant species P. maximowiczii, S. hultenii var. angustifolia and 4 S. sachalinensis in the study area. These ECM fungi are well known colonizers of 5 plants in disturbed or primary habitats (e.g. Nara et al. 2003a, b; Trowbridge and 6 Jumpponen 2004) and may be important for the establishment of pioneer woody plant 7 8 species, as well as the further revegetation of Mt. Usu. In general, however, the role played by mycorrhizal fungi in the growth and survival of seedlings of woody plant 9 10 species remains unclear, since their interactions may vary according to the combination of species and environmental conditions. In the future, it would be useful to examine 11 the effects of inoculation of these mycorrhizal fungi in the field. 12

13

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14	
15	Table and Figure Legends
16	Table 1. Frequencies (F) of ECM and AM associations with woody plant species,
17	observed at a research site on the Mt. Usu volcano, Hokkaido, Japan in 2004.
18	*Non, Non-mycorrhizal; ECM, ectomycorrhizal; and AM, arbuscular mycorrhizal.
19	Table 2. Number of ECM morphotypes, mean number of ECM morphotypes per

1	seedling and percentage of all types of ECM colonization (Ec) in the roots of woody
2	plant species on the Mt. Usu volcano, Hokkaido, Japan.
3	*Standard deviations are indicated.
4	Table 3. ECM fungi detected according to type and best BLAST match.
5	*Ss, S. sachalinensis; Pm, P. maximowiczii; Sh, S. hultenii var. angustifolia; Si, S.
6	integra; Qc, Q. crispula; Bp, B. platyphylla var. japonica; Bm, B. maximowicziana;
7	and <i>Lk</i> , <i>L. kaempferi</i> .
8	**The assignment of two names for one ECM type indicates that some ECM types
9	were identified initially as identical but were differentiated later by PCR-RFLP.
10	*** n.d., not detected; + not cleaved.
11	Table 4. Percentage of colonization (Ec)** and frequencies (F) of each ECM fungus
12	observed in ECM hosts*** established at the study site on Mt. Usu, Hokkaido, Japan.*
13	Percentage of colonization and frequencies of these fungi are obscured because two
14	fungal species were included in one ECM type.
15	** Mean and standard deviations (in parenthesis) are presented.
16	***See Table 3.
17	Fig. 1. Location of study site on Mt. Usu, Hokkaido, Japan.
18	
19	

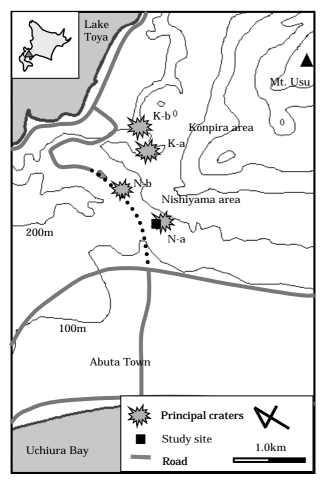


Fig. 1 Location of study site on Mt. Usu, Hokkaido, Japan

Table 1 The frequency (F) of ECM and AM association with
woody plant species, observed at a research site on Mt. Us
 $\mathbf{1}$,
Hokkaido, Japan in 2004.

Woody plant species	M	F	
Woody plant species	Mycorrhiza*	ECM	AM2
Betula platyphylla var. japonica	ECM, AM	6/6	3/6
Populus maximowiczii	ECM, AM	9/9	4/9
Quercus crispula	ECM, AM	5/6	1/6 ³
<i>Salix hultenii</i> var. <i>angustifolia</i>	ECM, AM	9/9	3/9
Salix integra	ECM	6/6	0/6
Salix sachalinensis	ECM, AM	12/12	1/12
Acer mono	AM	0/6	3/6
Rosa multiflora	AM	0/6	4/65

*ECM ectomycorrhizal, AM arbuscular mycorrhizal

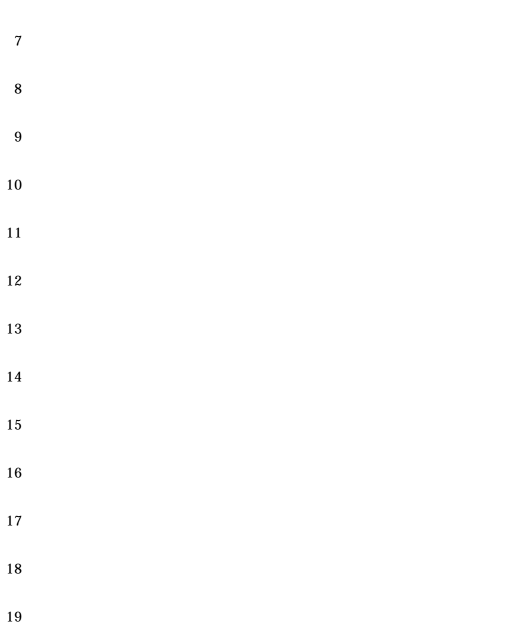


Table 2 The number of ECM morphotypes, the average number of ECM morphotypes per seedling and the percentage of all types of ECM colonization in the roots of woody plant species on Mt. Usu, Hokkaido, Japan

	ECM	morphotype	Total E2 (%)*		
Woody plant species	total	per seedling*			
Betula platyphylla var. japonica	6	4.0 ± 1.5	41.8 ± 22.3		
Populus maximowiczii	4	1.7 ± 0.5	39.5 ± 24.8		
Quercus crispula	1	0.8 ± 0.4	17.4 ± 16.7		
<i>Salix hultenii</i> var. <i>angustifolia</i>	4	1.9 ± 0.8	29.3 ± 16.2		
Salix integra	4	2.3 ± 0.5	17.3 ± 9.2		
Salix sachalinensis	3	2.0 ± 0.9	24.7 ± 12.7		

*Standard deviations are indicated

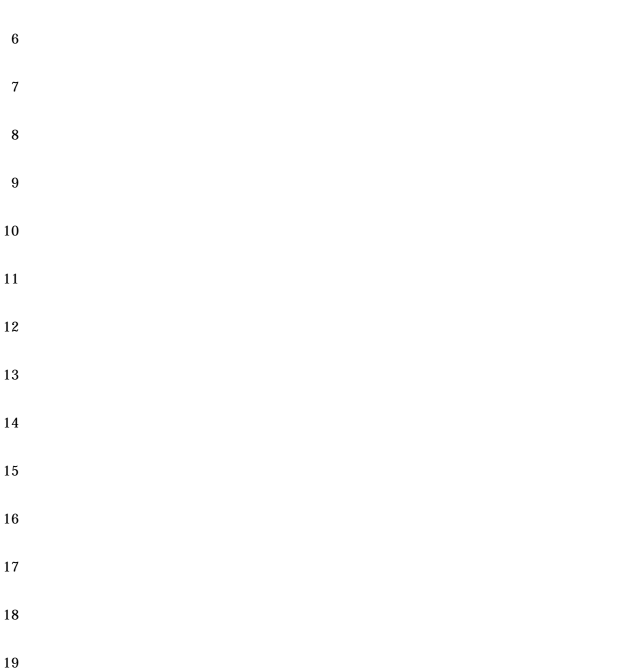


Table 3 ECM fungi detected on each ECM type with their best Blast match

Tuo o *	* ECM type**	Possible identity	Blast match	overlap (bp)	Similarity (%)	RFLP pattern (bp)***										
I ree*						<i>Hin</i> f	a F	linf	b <i>Hin</i> f	С	<i>Alu</i> a	Alu	b A	4 <i>lu</i> c	Alu	
Вр	<i>Bp</i> -1	Thelephoraceae 1	DQ195592.1	413	94	310		220	150)	300	260		200	10	0
	<i>Bp-</i> 2	Hebeloma mesophaeum	AY311521.1	313	99	n.d.					n.d.					
	Бр-2	Unidentified 5	-	410	-	n.d.					n.d.					
	<i>Bp-</i> 3	Leccinum scabrum	AF454585.1	436	99	860		440			570	400		120		
	<i>Bp-</i> 4	Thelephoraceae 5	AB211278.1	404	96	350		160	120)	500	130				
	<i>Bp-</i> 5	Unidentified 3	-	358	-	240		190	120)	+					
	<i>Bp-</i> 6	Thelephoraceae 4	AF184742.1	421	95	360		200	150)	470	130				
Pm	<i>Pm</i> -1	Scleroderma bovista	AB099901.1	216	95	n.d.					n.d.					
	Pm-2	Thelephoraceae 1	DQ195592.1	438	95	290		200	140)	310	280		210	10	0
	Pm-3	Laccaria amethystea	AB211270.1	431	99	400		350			420	380		100		
	Pm-4	Inocybe lacera	AY750157.1	405	100	380		250			330	210		180		
	<i>F 111</i> -4	Thelephora terrestris	AF272921.1	470	98	350		190	100)	440	140				
Qc	Qc-1	Thelephora terrestris	AJ549972.1	417	98	380		200	100)	450	150				
Sh	Sh-1	Hebeloma mesophaeum	AY311521.1	431	99	410		340			320	270		210		
	Sh-2	Thelephoraceae 1	DQ195592.1	460	95	320		200	140)	290	260		190	10	ð
	Sh-3	Thelephora terrestris	AY230241.1	468	98	370		210	100)	440	130				
	Sh-4	Thelephoraceae 2	U83475.1	391	97	350		180			520	210				
Si	Si-1	Laccaria amethystea	AB211270.1	405	100	390		340			420	370		120		
	Si-2	Hebeloma mesophaeum	AY311521.1	408	99	400		340			290	220		190		
	Si-3	Thelephoraceae 3	AF184742.1	400	95	210		190	160)	470					
	Si-4	Thelephora terrestris	AY230241.1	431	99	370		210	110)	450	150				
Ss	Ss-1	Hebeloma sp.	AY320395	500	98	420		350			320	280		250	21	ð
	Ss-2	Thelephoraceae 1	DQ195592.1	488	95	290		190	140)	300	260		190	10	ð
	35-2	Unidentified 1	AB096869	350	97	380		190	100)	+					
	Ss-3	Unidentified 1	AB096870	505	96	n.d.					n.d.					

* Ss S. sachalinensis, Pm P. maximowiczii, Sh S. hultenii var. angustifolia, St S. integra, Qc Q. crispula, Bp B. platyphylla var. japonica, Bm B. maximowicziana, Lk L. kaempferi.

The assigned two names for one ECM type indicates that some ECM types were misunderstood as identical but were differentiated by PCR-RFLP. * n.d. not detected, + not craved.

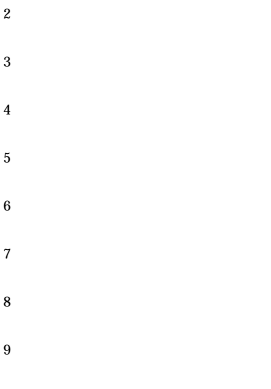


 Table 4 Percentage of colonization (Ec)** and frequencies (F) of each ECM fungi observed in ECM hosts*** established at the study site on Mt. Usu, Hokkaido, Japan.

ECM fungi	Вр		Qc		Pm		Sh		Si		Ss		
ECWIUNg	Ec (%)	F (/6)	Ec (%)	F (/6)	Ec (%)	F (/9)	Ec (%)	F (/9)	Ec (%)	F (/6)	Ec (%)	F (/12)	
Laccaria amethystea					12.1 (14.2)	6			9.6 (6.9)	6			
Inocybe lacera					18.9 (9.1)*	7*							
Hebeloma mesophaeum	2.2 (1.8)*	4*					18.3 (12.9)	8	3.4 (3.0)	5			
Hebeloma sp.											12.8 (12.9)	7	
Scleroderma bovista					74.1	1							
Leccinum scabrum	48.8	1											
Thelephora terrestris			17.4 (16.7)	5	18.9 (9.1)*	7*	30.0 (21.3)	3	1.3 (1.6)	2			
thelephoraceae 1	19.8 (23.0)	5			76.7	1	3.2 (3.7)	5			17.3 (9.0)*	12*	
thelephoraceae 2							11.4	1					
thelephoraceae 3									26.5	1			
thelephoraceae 4	1.2 (0.9)	4											
thelephoraceae 5	8.2 (9.9)	5											
Unidentified 1											17.3 (9.0)*	12*	
Unidentified 2	4.8 (3.6)	6											
Unidentified 3	2.2 (1.8)*	4*											

* Percentage of colonization and frequencies of these fungi is obscured because two fungal species were included in one ECM type. ** Average and standard deviation (in parenthesis) were presented. ***See Table 3

