1	Category of paper: Short note
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3	Microfungi associated with withering willow wood in ground contact near Syowa
4	Station, East Antarctica for 40 years
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22AbstractData are rather lacking on the diversity of microfungi associated with 23exotic plant substrates transported to continental Antarctica. We examined the 24diversity and species composition of microfungi associated with withering woody 25shoots of saplings of Salix spp. (willows) transplanted and in ground contact 26near Syowa Station, East Antarctica for more than 40 years. The willow 27saplings originated from Hokkaido, Northern Japan, and were experimentally 28transplanted in 1967-1968, but died within a few years. Dead willow shoots, 29unbranched and standing on bare ground for approximately 50 years, were used 30 for the isolation of fungi with the surface disinfection method. A total of 43 isolates were retrieved from 32 (78%) of the 41 shoots tested. The fungal isolates 3132were classified into 18 molecular operational taxonomic units (MOTUs) based on

33	the similarity of rDNA ITS sequences at the 97% criterion. Leotiomycetes was
34	the most common class in terms of the number of isolates and MOTUs, followed
35	by Dothidiomycetes, Sordariomycetes, and Eurotiomycetes. Molecular
36	phylogenetic affinities suggested that the closest relatives of the MOTUs were
37	saprobic and root-associated fungi. The result of the present study suggested
38	that Cadophora luteo-olivacea is widespread in soils throughout Antarctica and
39	likely indigenous.
40	
41	Keywords Continental Antarctica • Fungi • Root endophyte • Salix •
42	Syowa Station
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44	Introduction
45	
46	Ice-free regions of continental Antarctica, comprising only about 2% of the
47	continent, are cold and arid, and strong selection pressures are imposed on plant
48	establishment and soil development. Despite the harsh environment, previous

49	studies have reported the occurrence of free-living fungi in soils and in
50	association with bryophytes in coastal outcrops of continental Antarctica (e.g.
51	Azmi and Seppelt 1997; Tosi et al. 2002, 2005; Newsham et al. 2009). Recent
52	studies have examined fungal populations in historically-introduced exotic
53	materials and found a significant overlap of fungi isolated from these materials
54	and fungi isolated from environmental samples in pristine locations (Farrell et
55	al. 2011). A significant effect of exotic substrates on indigenous soil fungi has
56	also been found (Arenz et al. 2011). However, data are still lacking regarding the
57	diversity of microfungi associated with exotic plant substrates transported to
58 59	continental Antarctica. The purpose of the present study is to examine microfungi associated with withering woody shoots of saplings of <i>Salix</i> spp.
60	(willows) in ground contact in Syowa Station, East Antarctica for 40 years.
61	

62 Materials and methods

63

64 Study site and sample collection

66	Samples were collected near Syowa Station on East Ongul Island, Lützow-Holm
67	Bay, East Antarctica (60°00'47"S, 39°34'57"E, 16 m a.s.l.). In February 1967,
68	saplings of dwarf deciduous shrubs Salix pauciflora and S. reinii, 10-20 cm in
69	height and originating from Hokkaido, Northern Japan, were transplanted at
70	experimental sites near Syowa Station by Dr. T. Hoshiai of the 8th Japanese
71	Antarctic Research Expedition (JARE-8) to test their growth and survivorship.
72	These saplings endured through winter, sprouted, and bloomed in the next
73	summer of 1968, but not all sprouted in the summer of 1969 (Hoshiai 1970).
74	Additional saplings were transplanted by Dr. Y. Endo of JARE-9 in 1968, giving
75	a similar result of the sapling producing leaves the next year but dying within a
76	few years because of the adverse environment of Antarctica (Hoshiai 1970).
77	During JARE-51 in 2009-2010, we found dead willow shoots still standing on the
78	experimental site. In February 2010, a total of 41 withering shoots (aboveground
79	parts without leaves, soil, or belowground parts, approximately 3 cm in height,
80	and 1-3 mm in basal diameter) were collected with tweezers, preserved in paper

81 bags, stored at 2°C, and taken back to the laboratory in Japan.

82

83 Fungal isolation

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Fungi were isolated from shoots using the surface disinfection method according 85 86 to Osono et al. (2012). The surface-disinfected shoots were plated on 9-cm Petri 87 dishes containing 2% lignocellulose agar (LCA) modified as described by Miura 88 and Kudo (1970) (glucose 0.1%, KH₂PO₄ 0.1%, MgSO₄•7H₂0 0.02%, KCl 0.02%, 89 NaNO₃ 0.2%, yeast extract 0.02%, and agar 2% (w/v)), two shoots per plate. Note that the modified LCA of Miura and Kudo (1970) does not contain lignin or other 90 recalcitrant compounds. The modified LCA was used because its low glucose 91 92content suppresses the overgrowth of fast-growing fungal species (Osono and Takeda 1999). The plates were incubated in darkness at 10°C and observed for 4 93 94weeks after the disinfection. Any fungal hyphae or spores appearing on the plates were subcultured onto fresh LCA plates, incubated, and observed 95 96 micromorphologically. Isolates were then used for molecular analysis as 97 described below.

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99 Molecular methods

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101 Genomic DNA was extracted from mycelia that had been cultured on 2.5% malt 102extract agar overlaid with a cellophane membrane following the modified CTAB 103 method described by Matsuda and Hijii (1999). Polymerase chain reactions 104 (PCR) were performed using a Quick Taq HS DyeMix (Toyobo, Osaka, Japan). 105Each PCR reaction contained a 50 µl mixture (21 µl distilled water, 25 µl master 106mix, 3 μ l ca. 0.5ng/ μ l template DNA, and 0.5 μ l of each primer (final, 0.25 μ M)). 107 To PCR amplify the region including the rDNA ITS and 28S rDNA D1-D2 108domain, the primer pair ITS1f (Gardes and Bruns 1993) and LR3 (Vilgalys and 109 Hester 1990) was used. Each DNA fragment was amplified using a PCR thermal 110 cycler (DNA engine; Bio-Rad, Hercules, CA, USA) using the following thermal cycling schedule. The first cycle consisted of 5 min at 94°C, followed by 35 cycles 111 112of 30 s at 94°C, 30 s at 50°C for annealing, 1 min at 72°C, and a final cycle of 10 113 min at 72°C. The reaction mixture was then cooled at 4°C for 5 min. PCR 114products were purified with a QiAquick PCR Purification Kit (Qiagen, Germany) according to the manufacturer's instructions. 115116Purified PCR products were sequenced by FASMAC Co., Ltd. 117 (Kanagawa, Japan). Sequencing reactions were performed in a Gene Amp PCR 118 System 9700 (Applied Biosystems, USA) using a BigDye Terminator V3.1 119 (Applied Biosystems), following the protocols supplied by the manufacturer. The 120 fluorescent-labeled fragments purified from the were unincorporated 121terminators using an ethanol precipitation protocol. The samples were resuspended in formamide and subjected to electrophoresis in an ABI 3730xl 122123sequencer (Applied Biosystems). 124The sequences determined in this study were deposited in the DNA Data Bank of Japan (DDBJ) (AB752244-AB752287). The rDNA ITS sequences 125126 were compared with available rDNA sequences in the GenBank database using 127BLASTN searches (Altschul et al. 1990). For phylogenetic analysis, MAFFT ver. 1286 (Katoh and Toh 2008) was used for preliminary multiple alignments of

129	nucleotide sequences. Final alignments were manually adjusted using BioEdit
130	(Hall 1999). Alignment gaps were treated as missing data, and ambiguous
131	positions were excluded from the analysis. The phylogenetic tree was conducted
132	by maximum likelihood (ML) methods (Felsenstein 1981) with the best fit
133	nucleotide substitution model based on the lowest Bayesian Information
134	Criterion (BIC) score. To estimate clade support, the bootstrap procedure of
135	Felsenstein (1985) was employed with 1000 replicates. These analyses were
136	carried out using MEGA5 (Tamura et al. 2011).
136 137	carried out using MEGA5 (Tamura et al. 2011). The isolates were grouped into molecular operational taxonomic units
137	The isolates were grouped into molecular operational taxonomic units
137 138	The isolates were grouped into molecular operational taxonomic units (MOTUs) according to the similarity of rDNA ITS sequences at the 97% criterion.
137 138 139	The isolates were grouped into molecular operational taxonomic units (MOTUs) according to the similarity of rDNA ITS sequences at the 97% criterion. The frequency of occurrence of MOTU was calculated as a percentage of the

- **Results**

145	Fungi were isolated from 32 (78%) of the 41 shoots tested for isolation. A total of
146	43 isolates were obtained, and these were classified into 18 MOTUs (Table 1, Fig
147	1). Leotiomycetes was the most frequent class, including 29 isolates of 10
148	MOTUs, followed by Dothidiomycetes (9 isolates, 4 MOTUs), Sordariomycetes (3
149	isolates, 2 MOTUs), and Eurotiomycetes (2 isolates, 2 MOTUs) (Fig. 2). The
150	most frequent MOTUs were MOTU1 in the Leotiomycetes that had 100%
151	sequence match of the ITS region to Cadophora luteo-olivacea (7 isolates),
152	MOTU9 in Leotiomycetes (7 isolates), and MOTU18 in Dothidiomycetes (5
153	isolates) (Table 1, Fig. 2).

Discussion

Some of the microfungi associated with dead willow shoots in the present study
are classed as saprobic fungi (Table 1). For example, *Cadophora luteo-olivacea*(MOTU1) is a saprobe occurring in many habitats including wood, soil, and
plants (Gramaje et al. 2011). Several *Cadophora* species, including *C*.

161	luteo-olivacea, have also been isolated from soils and historic wood along the
162	Ross Sea region of Antarctica (Arenz et al. 2006) and have the potential to cause
163	soft rot in wood (Blanchette et al. 2004). Similarly, Phialocephala lagerbergii,
164	which had 99% sequence match of the ITS region to MOTU3, is known to be a
165	wood-inhabiting fungus (Grünig et al. 2009). Geomyces vinaceus, an anamorph
166	of Pseudogymnoascus roseus and which had 100% sequence match of the ITS
167	region to MOTU8, is associated with wood, soil, and roots (Rice and Currah
168	2006). Coniochaeta lignaria, which had 99% sequence match of the ITS region to
169	MOTU13, has been shown to have lignocellulose-degrading enzymes (Lopez et al.
170	2007), which can facilitate growth and energy acquisition in dead willow shoots
171	consisting of structural lignin and cellulose polymers.
172	We noted that root-associated microfungi were isolated frequently from
173	the dead willow shoots (Table 1). For example, Ilyonectria robusta and Phoma
174	sclerotioides, which had 99% sequence match of the ITS region to MOTU14 and
175	MOTU17, respectively, are root-rot fungi (Wunsch and Bergstrom 2011; Cabral
176	et al. 2012). <i>Phialocephala fortinii</i> , which had 99% sequence match of the ITS

177	region to MOTU5, and also possibly MOTU2 in Phialocephala, is a common
178	endophyte of plant roots and is widespread in sub-Antarctic ecosystems and also
179	present in continental Antarctica (Grünig et al. 2008; Newsham et al. 2009).
180	Jumpponen et al. (2003) detected a DNA sequence with 99% similarity to P .
181	fortinii in a rhizoid of the liverwort Cephaloziella varians on the Antarctic
182	Peninsula.
183	It is unclear whether these fungi were widespread or localized in their
184	distribution in Antarctica and whether they were indigenous to Antarctica or
185	introduced along with the saplings in soil from Japan. MOTU1, one of the most
186	frequent taxa (Table 1), had 99% to 100% sequence match (with query coverage
187	between 89% and 97%) of the ITS region to Cadophora luteo-olivacea isolated
188	from wood and soil in the Ross Sea Region (DQ317327, Arenz et al. 2006;
189	GU212374, Blanchette et al. 2010) and along the Antarctic Peninsula (FJ911899,
190	Rosa et al. 2010; HQ438025, Gonçalves et al. 2012). This result suggested that
191	this fungus is widespread in soils throughout Antarctica and likely indigenous.
192	Similarly, Geomyces vinaceus (OTU8) was isolated from moss samples in

193 Victoria Land on the west coast of the Ross Sea (Tosi et al. 2002), but the
194 distribution of this fungus in Antarctica remains unknown and deserve further
195 researches.

196 It is unclear whether the fungi isolated in the present study were active 197or dormant in dead shoots. However, the supply of exotic woody substrates, such 198as dead willow shoots, can contribute to fungal abundance, as the natural lack of 199 organic material in Antarctica limits the densities of fungal populations (Arenz 200 et al. 2011). To exist in Antarctica, fungi need to be able to tolerate the harsh 201environment, and Antarctic fungi have a variety of physiological traits that 202enable them to survive under cold and dry conditions (Robinson 2001), including 203 cold tolerance, accumulation of intercellular trehalose and polyols, secretion of 204antifreeze proteins, and enzymes active at low temperatures. Future studies will 205include physiological evaluations of these fungal isolates and measurements of 206 activity at low temperatures.

207

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213	
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1 Hirose et al. Table 1.

- $\mathbf{2}$
- 3

4	Table 1 The number of isolates and blast identity results (in percentage) of fungal molecular operational taxonomic units
5	(MOTUs) isolated from withering willow shoots and sequence accession number for the closest relative found at GenBank.

			Closest match at Genbank	
Class	MOTU	Number of isolates	(Accession number)	Sequence similarity %
Leotiomycetes	1	7	Cadophora luteo-olivacea (GU128589)	100
	9	7	Leotiomycetes sp. (JQ759481)	99
	2	4	<i>Phialocephala</i> sp. (FM999988)	99
	7	3	Leotiomycetes sp. (JQ758759)	99
	5	2	Phialocephala fortinii (EU888625)	99
	8	2	Geomyces vinaceus (AJ608972)	100
	3	1	Phialocephala lagerbergii (AB190400)	99
	4	1	Helotiales sp. (AB598096)	92
	6	1	<i>Clathrosporium intricatum</i> (EF029192)	95
	10	1	<i>Tetracladium</i> sp. (AJ890435)	99
Eurotiomycetes	11	1	Exophiala salmonis (GU586858)	99
	12	1	Penicillium turbatum (AY213679)	100
Sordariomycetes	14	2	<i>Ilyonectria robusta</i> (JF735265)	99

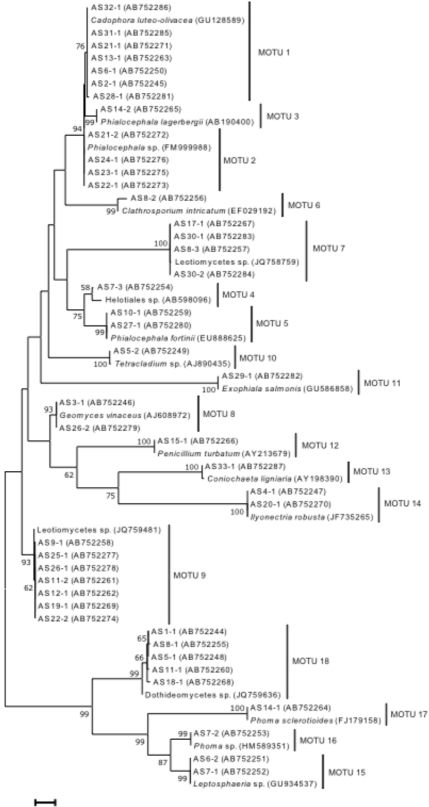
	13	1	<i>Coniochaeta ligniaria</i> (AY198390)	99
Dothidiomycetes	18	5	Dothideomycetes sp. (JQ759636)	98
	15	2	Leptosphaeria sp. (GU934537)	99
	16	1	<i>Phoma</i> sp. (HM589351)	100
	17	1	Phoma sclerotioides (FJ179158)	99

- 1 Figure legend
- $\mathbf{2}$

Fig. 1 Maximum-likelihood (ML) phylogeny inferred from rDNA ITS sequences including 18 fungal molecular taxonomic units (MOTUs) isolated from withering willow shoots. The evolutionary model used was the Kimura 2-parameter model (Kimura 1980) with a discrete Gamma distribution (+G, parameter = 0.7952) and a proportion of Invariant sites (+I, 34.3127% sites) to allow for non-uniformity of rates among sites. Bootstrap values for the ML analysis are indicated for corresponding branches.

Fig. 2 Rank-abundance relationship of fungal molecular taxonomic units
(MOTUs) isolated from withering willow shoots. Black bar, Leotiomycetes; blank
bar, Eurotiomycetes; shaded bar, Sordariomycetes; gray bar, Dothidiomycetes.

 $\mathbf{2}$



0.05

