Molecular analysis of lichen-associated bacterial communities

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Abstract

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Keywords

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Introduction

The genomic exploration of microbes from poorly studied ecological niches has become an exciting endeavour in recent years. Niches as diverse as soil, water and air are being studied by various techniques, ranging from culture techniques to metagenomic approaches (e.g. Venter et al., 2004). This is a promising perspective for the investigation of further, biologically rich habitats. Among these, diverse groups of cryptogams including lower plants and fungi could be particularly suitable as hosts for bacterial communities. A recent investigation of bryophyte-associated bacteria revealed numerous bacterial phylotypes (Opelt & Berg, 2004), some of which could not be assigned to genera. Another interesting habitat is certainly provided by fungal symbioses. Whereas the involvement of bacteria in mycorrhizal symbioses is wellstudied (Garbaye, 1994), we focus here on another important symbiotic life style of fungi, the lichen association.

Of all fungal symbioses, lichens are a rather particular case. The lichen association of fungi and algae contributes to a substantial evolutionary radiation of the mycobionts (Gargas *et al.*, 1995; Lutzoni *et al.*, 2001), and also allows their algal partners to grow well under environmental

The bacterial communities associated with 11 different lichen samples (belonging to eight different species) from different habitats were investigated. The culturable aerobic-heterotrophic fraction of the bacterial communities was isolated from nine lichen samples on protein-rich and sugar-rich/N-free media. Thirty-four bacterial isolates were purified and pooled into groups (phylotypes) by analysis of the ribosomal internal transcribed spacer polymorphism. Twenty five phylotypes were identified, each comprising between one and three isolates. One isolate of each phylotype was partially sequenced and the resulting 16S rRNA gene sequences were compared in a phylogenetic analysis. Three genera of Firmicutes, four of Actinobacteria and three of Proteobacteria were identified. Two phylotypes, belonging to the phyla Actinobacteria and Proteobacteria, respectively, were not identified at genus level. Some bacterial taxa were retrieved frequently in different lichen species sampled in the same or different sites. *Paenibacillus* and *Burkholderia* phylotypes seem to be common in lichens. Luteibactor rhizovicina was found in three different lichens of two different regions. In a cultivation-independent approach, total DNA was extracted from 11 lichen samples. Molecular fingerprints of the bacterial communities were obtained by PCR-amplification of the internal transcribed spacer region, and sequencing of selected bands indicated the presence of additional bacteria.

> situations that would usually not be favourable in biological isolation. This evolutionary successful lifestyle apparently arose early in the evolution of ascomycetes (Lutzoni et al., 2001), before the Lower Devonian (Taylor et al., 1995). The lichen symbiosis is maintained by approximately one fifth of all known extant fungi, and by more than 42% of known Ascomycota. Lichens are ubiquitous and are found from sub-polar ranges to tropical rainforests. They are also able to grow on diverse substrates and sometimes under extreme ecological conditions. Since most lichens are exceptionally drought-tolerant and slow-growing organisms, they provide an unusual and long-living ecological niche for additional microorganisms, which may include other fungi (well-studied lichen parasites, as well as less known epi- and endobionts (Petrini et al., 1990; Prillinger et al., 1997; Miadlikowska, personal communication)) and prokaryotes.

> The earliest reports about non-cyanobacterial prokaryotes in lichens were contradictory, and likely due to a misinterpretation of crystallized secondary compounds (Uphof, 1925; Suessenguth, 1926). Clear evidence for the presence of bacteria in lichens was then provided by a series of papers that appeared long before the emergence of molecular methods. Henkel & Yuzhakova (1936) and Iskina (1938) detected

nitrogen-fixing bacteria in lichens by cultivation on nitrogen-free Ashby medium, and they assigned these strains to *Azotobacter*. While Krasil'nikov (1949) could not confirm these results with the sample he studied, Scott, (1956) mentioned again the frequent finding of *Azotobacter* in lichens. Besides *Azotobacter*, other genera were previously reported from lichens, such as *Bacillus* (Henkel & Plotnikova, 1973), *Beijerinckia* (Panosyan & Nikogosyan, 1966), *Clostridium* (Iskina, 1938) and *Pseudomonas* (Henkel & Plotnikova, 1973). While an involvement in nitrogen fixation for lichens was repeatedly considered in these works, the finding of actinobacteria prompted Zook (1983) to suggest also a defensive role for bacteria in lichens. Moreover, Lenova & Blum (1983) suggested that up to millions of bacterial cells could be present per gram of a lichen thallus.

The analysis of bacteria in these works relied only on phenotypic methods using cultured isolates and therefore our knowledge about the taxonomic diversity of lichenassociated bacteria is still rather limited. However, a precise determination of strains using molecular data is required to gain further insights and to assess, for example, whether bacterial strains are selective for their host lichens or evolved resistances against the antimicrobial activity of lichen secondary metabolites (Ingólfsdóttir et al., 1985). Only one study so far has characterized the diversity of actinomycetes in lichens using DNA fingerprinting (Gonzáles et al., 2005). In this contribution we present a more general molecular approach to culturable lichen-inhabiting bacteria and study their diversity in selected lichens using DNA sequence data. Moreover, we show first results of a cultivation-independent technique suitable for studying all the bacterial communities associated with lichens.

Materials and methods

Lichen samples

We analyzed 11 different lichen samples, belonging to eight species, sampled at five different sites. For further details on

Table '	1.	Lichens	used	in	this	work
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the selected species, see Table 1. Each sample was taken with washed instruments and put in sterile bags. The samples were immediately frozen and conserved until further analysis. Sub-samples of approximately 0.3–0.5 g of the thalli were washed in sterile water and utilized for the isolation of bacteria and for direct extraction of total DNA.

Bacterial isolation

To isolate the external bacteria, nine subsamples of lichens were vortexed in 0.8% NaCl solution for 60 s and 100 μ L of the solution was plated on both Tryptone-Yeast extract (TY) and sugar-rich/N-free media (5g glucose, 5g mannitol, 0.8 g K₂PO₄, 0.2 g MgSO₄ · 7H₂O, 0.15 g CaCl₂, 0.04 g FeSO₄ · 7H₂O, 0.005 g Na₂MO₄ · 2H₂O, 15 g agar with volume made up to 1 L with distilled water and pH adjusted to 7.0). The sub-samples were then surface-sterilized by immersion for 4 min in H₂O₂ (9%) and washed in sterile water prior to isolation of the internal bacteria. Thallus fragments were subsequently crushed in 500 μ L of sterile water with a sterile scalpel and 100 μ L of the resulting suspension were then plated on the same media as above.

The plates were incubated at 25 °C for 15 days. For each plate, the colonies showing distinct phenotypes were purified by plating them separately on new plates with the same media. The pure cultures were numbered according to the lichen sample and the origin (external surface or internal parts, respectively), as well as the isolation medium (e.g. Bint1: bacterium isolated on TY medium from the internal thallus of the lichen sample 'B', GestV2N: bacterium isolated on N-free medium from the external surface of the lichen sample 'G'). All isolates are stored at -80 °C in 20% glycerol at the Dipartimento di Biologia Cellulare e dello Sviluppo, Palermo, Italy.

Molecular characterization of the isolates and identification of phylotypes

Bacterial genomic DNAs were extracted by the lysozymeproteinase K-sodium dodecyl sulphate method (modified

Lichen sample	Lichen species	Sampling site and altitude	Date of sampling
A	Cladonia digitata	Austria, Styria, Rabenwald, 1200 m	10/2004
В	Cladonia rangiferina	Austria, Styria, Rabenwald, 1200 m	10/2004
C	Cladonia coniocraea	Austria, Styria, Botanical Garden of Graz, 365 m	10/2004
D	Cladonia pyxidata	Austria, Styria, Rabenwald, 1200 m	10/2004
E	Cladonia coccifera	Austria, Styria, Rabenwald, 1200 m	10/2004
G	Cladonia pyxidata	Austria, Styria, Plankogel, 1440 m	5/2005
Н	Pseudevernia furfuracea	Austria, Styria, Plankogel, 1520 m	5/2005
J	Hypogymnia physodes	Austria, Styria, Plankogel (on moss), 1440 m	5/2005
К	Hypogymnia physodes	Austria, Styria, Plankogel (on bark) 1520 m	5/2005
R	Roccella phycopsis	France, Normandy, St. Malo, 10 m	3/2005
S	Roccella fuciformis	France, Normandy, St. Malo, 10 m	3/2005

from Sambrook et al. (1989) by increasing the reagent concentration to 2.4 mg mL^{-1} lysozyme, 0.5 mg mL^{-1} proteinase K and 0.8% sodium dodecyl sulphate). The bacterial isolates were assigned to groups with similar banding profiles of amplified ribosomal internal transcribed spacers between the 16S and 23S ribosomal genes (ITS 16S-23S). Because of the high level of polymorphism of the ribosomal spacer, the ITS 16S-23S profiles (showing one to numerous bands per profile) are regarded as specific for bacterial strains. This allowed us quickly to screen the bacterial diversity. The isolates with the same profile were considered to belong to the same bacterial phylotype. For the ITS 16S-23S amplification we used 0.3 µM of each of the optimized primers ITSF (5'-GTCGTAACAAGGTAGCCG-TA-3') and ITSR eub (5'-GCCAAGGCATCCACC-3') (Cardinale et al., 2004) in a reaction volume of 25 µL, containing 100-150 ng template DNA, 2 units of Taq DNA polymerase (Sigma-Aldrich Corp., St. Louis, MO) and 0.2 mM of each dNTP. PCR was carried out for 30 cycles under the following conditions: 90 s at 95 °C, 60 s at 55 °C and 90 s at 72 °C in a thermal cycler T-personal (Biometra, Goettingen, Germany). An initial hot start (5 min at 95 °C) and a final extension (5 min at 72 $^{\circ}$ C) were also performed. Finally, 5 μ L of PCR product was checked by agarose gel electrophoresis and ethidium bromide staining. The ITS profiles of different isolates were compared by aligning the bands using a DNA marker with bands between 100 and 10 000 nucleotides (DNA Ladder mix, FERMENTAS, St. Leon-Rot, Germany). Additionally, Amplified Ribosomal DNA Restriction Analyses (ARDRA; Urzi et al., 2001; Lagacé et al., 2004; Ntougias et al., 2004) were in some cases performed with the endonucleases Hinfl, TaqI and AluI to confirm the results of the ITS screening and to assign isolates with ambiguous ITS profiles to a particular phylotype.

Identification of the phylotypes

Representative isolates of all detected phylotypes were further characterized by 16S rRNA gene sequencing and phylogenetic analysis. The amplification of the 16S rRNA gene was performed with the primers fD1 and rD1 (Weisburg et al., 1991) using the same PCR protocol described above for the ITS 16S-23S amplification. The PCR products were visualized by agarose gel electrophoresis and quantified by comparison with the bands of the DNA Ladder Mix marker (FERMENTAS). About 60 ng of each product were used for sequencing reactions, which were performed by the BMR laboratory of the University of Padova, Italy (http:// bmr.cribi.unipd.it/). The sequence electropherograms were checked with the programs ABIVIEW (EMBOSS package, http://emboss.life.nthu.edu.tw/emboss/), EDITVIEW (http:// www.appliedbiosystems.com/support/software/) and FINCHTV (http://www.geospiza.com/finchtv/index.htm). To identify

similar sequences that are available in the NCBI Genbank, sequences were used in BLASTn searches (Altschul *et al.*, 1997, http://www.ncbi.nlm.nih.gov/blast/). The sequences from the isolates are available from Genbank/EMBL under the accession numbers AM062703 to AM062725.

Phylogenetic position of the isolates

The 16S rRNA gene sequences were aligned using CLUSTALX (Thompson *et al.*, 1994), and a distance matrix was calculated by the DNADIST software of the PHYLIP package (Felsenstein, 2004; http://evolution.genetics.washington. edu/phylip.html) using the F84 algorithm (Felsenstein & Churchill, 1996). The Neighbour-Joining (Saitou & Nei, 1987) phylogenetic tree was constructed with the NEIGHBOR software of the PHYLIP package. The topology of the tree was statistically tested by performing 1000 bootstrap resamplings of the data with the program SEQBOOT and a Majority Rule consensus tree was obtained by the program CONSENSE of the PHYLIP package. The tree of Fig. 1 was prepared with the TREEVIEW program (Page, 1996; http:// taxonomy.zoology.gla.ac.uk/rod/treeview.html).

Cultivation-independent analysis of the lichenassociated bacterial communities

After external sterilization by immersion in H₂O₂ (9%) for 4 min, total DNA was extracted from approximately 0.3-0.5 g of lichen thalli using the CTAB/chloroform-isoamyl alcohol method as previously described (Cubero et al., 1999). The regions of DNA including the 16S ribosomal gene, the internal transcribed ribosomal spacer and approximately 100 nucleotides of the 23S ribosomal gene were amplified using 5 µL of crude extracted DNA as template, 0.4 µM of each fD1 and FGPL 132-38 primers (Quatrini et al., 2002), 0.2 mM of each dNTP, 2.5 units of Tag DNA polymerase (Sigma-Aldrich) and 2 mM MgCl₂, according to the PCR-protocol previously described for 16S rRNA gene and ITS 16S-23S amplifications. A second nested amplification was performed to amplify the ITS 16S-23S region of all bacterial communities using 10 µL of the first PCR product as template and the primers ITSF and ITSReub. The concentrations of other reagents were the same as above. About 200 ng of PCR products were visualized in both ethidium bromide-stained agarose gel (2.5% NuSieve agarose, USB, Cleveland, OH) and in silver stained (Bassam et al., 1991), native acrylamide gel (5% acrylamide : bisacrylamide, Sigma-Aldrich). The banding patterns were compared by aligning the bands with the DNA size marker. The lengths of detectable bands ranged between about 150 and 1200 nucleotides. The fluorescence intensity relative to the overall intensity of each profile was optically evaluated and the data were saved in a Microsoft[©] Excel matrix. This

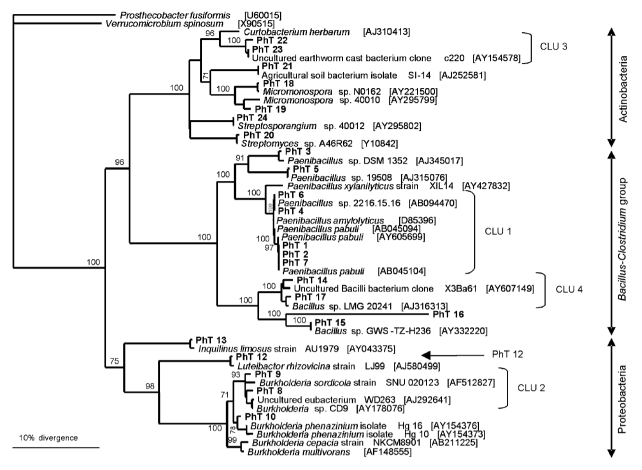


Fig. 1. Neighbour-joining tree, obtained using an alignment of 600 nucleotides of the 16S rRNA gene sequences, showing the phylogenetic relationships of the lichen-associated bacteria. Numbers on the internal nodes are bootstrap support values of 1000 re-samplings. Clusters (CLU1 – CLU4) are explained in the text.

matrix was used for the statistical analysis of the similarity between the profiles by using the NTEDIT and NTSys2.0 software (Rohlf, 1998). The tree was constructed using the Unweighted Pair Group Method with Arithmetic Mean (UPGMA) algorithm, based on the average taxonomic distances [Dist coefficient (Sneath & Sokal, 1973), Dice similarity index (Atlas & Bartha, 1993)]. The genetic diversity of the bacterial communities was evaluated by the ecological diversity index of Shannon-Wiener $(H = -\Sigma P_i)$ $\log P_i$) (Atlas & Bartha, 1993) and the dominance index of Simpson $(D = \Sigma P_i^2)$ (Hunter & Gaston, 1988), where P_i is the ratio between the intensity of fluorescence of each *i*th band and the total fluorescence intensity of the profile (normalized as percentage). Commonly present or otherwise intense bands were excised from the NuSieve gel and the DNAs were recovered by using the GFXTM Purification Kit (Amersham Biosciences, Little Chalfont, UK). The ITS region of four bands was sequenced and the sequences are available from Genbank/EMBL under the accession numbers DQ343830 to DQ343833.

Results

Bacterial isolation

Thirty four morphologically distinct bacteria were isolated from nine lichen samples (between one and eight from each lichen). Eighteen were from the external surfaces and 16 from the internal thallus parts (Table 2). Twenty-four bacteria were isolated on Tryptone-Yeast extract medium (TY) and 18 on Glucose-Mannitol-N-free medium. Generally, few colonies (between 1 and 26) of each phenotype grew on the isolation plates; only the phenotypes represented by CestV1, Dint1, Eint1b, GestV1, GestL2 (Table 2) formed more than 100 (up to thousands) colonies on each plate. Ten of the 24 isolates on TY medium were able also to grow efficiently on N-free medium.

Screening of the isolates

The analysis of the ribosomal internal spacer polymorphism was used to group the 34 isolates to 25 distinct phylotypes.

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Phylotype	lsolate	Host lichen	Most closely related sequence (Genbank accession number)	sequence sequence similarity in % (no. of nucleotides)	Taxonomical position of isolate (most related species)	Notes/reference of most related sequence
PhT 1	Bint1 [*]	Cladonia rangiferina	Paenibacillus pabuli (AB045104)	99.3 (893)	Firmicutes, Paenibacillaceae (Paenibacillus pabuli)	Chitinolytic aerobic endospore-forming bacteria
	BestV1 Aint*1	Cladonia digitata				
PhT 2	BestV3	Cladonia rangiferina	Paenibacillus pabuli (AB045104)	99.4 (933)	Firmicutes, Paenibacillaceae (Paenibacillus pabuli)	Chitinolytic aerobic endospore-forming bacteria
PhT 3	Eint3	Cladonia coccifera	Paenibacillus sp. DSM 1352 (AJ345017)	98 (978)	Firmicutes, Paenibacillaceae (Paenibacillus agarexedens)	Agarolytic aerobic endospore-forming soil bacteria Uetanabaro <i>et al.</i> (2003)
PhT 4	Eint1a	Cladonia coccifera	Bacterium H17 (AY345554)	99.9 (860)	Firmicutes, Paenibacillaceae (Paenibacillus amviolytricus)	Isolated from "microbial hotspot" in the Hawaiian Archineland
PhT 5	Hint1N	Pseudevernia furfuracea	Paenibacillus sp. 19508 (A 1315076)	99.1 (810)	Eirmicutes, Paenibacillaceae (Paenibacillus mendelii, Paenibacillus nhvllosnhaerae)	Isolated from biodeteriorated mural paintings in a nerronolis tomb Hevrman & Swince (2001)
PhT 6	Jint1	Hypogymnia physodes (on moss)		100 (742)	Firmicutes, Paenibacillaceae (Paenibacillus amvlolvticus)	Isolated from "microbial hotspot" in the Hawaiian Archinelago
PhT 7	JestL1	Hypogymnia physodes (on moss)	Paenibacillus pabuli (AB045104)	99.8 (803)	Firmicutes, Paenibacillaceae (Paenibacillus pabuli)	Chitinolytic aerobic endospore-forming bacteria
PhT 8	Dint1	<i>Cladonia pyxidata</i> (from Rabenwald)	uncultured eubacterium WD263 (AJ292641)	98 (1003)	Betaproteobacteria, Burkholderiaceae	Isolated from polychlorinated biphenyl-polluted soil Nogales et al. (2001)
	DestL 1 BestV4	Cladonia rangiferina			(Burkholderia glathei, Burkholderia	
		, , ,			sordidicola)	
PhT 9	GestV1N	<i>Cladonia pyxidata</i> (from Plankogel)	<i>Burkholderia</i> sordidicola strain SNU 020123 (AF512827)	97.9 (948)	Betaproteobacteria, Burkholderiaceae (Burkholderia sordidicola, Burkholderia glathei)	Isolated from the white-rot fungus Phanerochaete sordida Lim et al. (2003)
	HestL1N JestL1N	Pseudovernia furfuracea Hypogymnia physodes				
PhT 10	EestV2	(keun nu) Cladonia coccifera	Burkholderia phenazinium isolate	98.5 (729)	Betaproteobacteria, Burkholderiaceae (Burkholderia phenazinium)	Component of naphthalene-degrading soil guilds Wilson <i>et al.</i> (2003)
PhT 11	GestV2N	Cladonia pyxidata (from Plankogel)	(c/c+c+i+) of Br	nss	Betaproteobacteria, Burkholderiaceae	I

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PCB-degrading bacterium associated with plant roots	Isolated from clinical sample	Soil bacterium Felske et al. (1999)	1		1	Soil bacterium	Goto <i>et al.</i> (2000)	Isolated from coastal sediments Zhao <i>et al.</i>	(2004)	Herbicides-producing bacterium	I	Lukow (1999)	uncultured bacteria associated with the intestine of the earthworm <i>Lumbricus rubellus</i> Singleton <i>et al.</i> (2003)	uncultured bacteria associated with the intestine of the earthworm <i>Lumbricus rubellus</i> Singleton <i>et al.</i> (2003)
Gammaproteobacteria, Xanthomonadaceae (Luteibactor rhizovicina)	Alphaproteobacteria, Rhodospirillaceae (Inquilinus limosus)	Firmicutes, Bacillaceae (Bacillus niacini, Bacillus bataviensis)	Firmicutes, Bacillaceae Bacillus mycoides, Bacillus cereus, Bacillus weihenstephanensis)		Firmicutes, Staphylococcaceae (Staphylococcus epidermidis)	Firmicutes, Bacillaceae (Bacillus niacini)		Actinobacteria Micromonosporaceae	(Micromonospora matsumotoense)	Actinobacteria Micromonosporaceae (Micromonospora olivasterospora, Micromonospora auratinigra)	Actinobacteria Streptomycetaceae (Streptomyces sanglieri, Streptomyces yanii)	Actinobacteria, Cellulomonadaceae (Cellulomonas xylanilytica)	Actinobacteria, Microbacteriaceae (Frigoribacterium genomovar 2 and 3, Curtobacterium herbarum)	Actinobacteria, Microbacteriaceae (Frigoribacterium genomovar 2 and 3, Curtobacterium herbarum)
99.9 (718)	99.4 (712)	99 (686)	99.6 (687)		(009) 06	99.5 (778)		99.3 (730)		96.8 (619)	99.8 (813)	100 (804)	99.8 (410)	100 (720)
Luteibactor rhizovicina isolate OUCZ70 (AY785744)	Inquilinus limosus strain AU430 (AY043373)	<i>Bacillus</i> sp. isolate IDA216	Bacillus sp. (AY167860 and other 17 sequences)		<i>Staphylococcus</i> <i>epidermidis</i> (AY030342 and other 31 sequences)	Uncultured bacterium clone X20 (DQ083105)	<i>Bacillus niacini</i> (AB021194)	Micromonospora sp.	N0162 (AY 221500)	<i>Micromonospora</i> sp. 40010 (AY295799)	<i>Streptomyces</i> sp. strain A46R62 (Y 10842)	Agricultural soil bacterium isolate SI-14 (AJ252581)	Uncultured earthworm cast bacterium clone c220	(Arcultured) earthworm cast bacterium clone c220 (AY154578)
Cladonia coniocraea Cladonia rangiferina Cladonia pyxidata (from Rabenwald)	Cladonia rangiferina	Cladonia coccifera	Cladonia coccifera		Cladonia pyxidata (from Plankogel)	<i>Hypogymnia physodes</i> (on bark)		Hypogymnia physodes	(on bark)	Hypogymnia physodes (on bark)	Hypogymnia physodes (on bark)	Hypogymnia physodes (on bark)	Cladonia pyxidata (from Plankogel)	Cladonia pyxidata (from Plankogel)
CestV1 BestL2 DestL2	BestL3	EestL2	Eint 1b	EestL1	GestL2	KestL1		Kint1N		Kint2N	KestL 1N	KestL2	GestV1	<u>Gint1</u>
PhT 12	PhT 13	PhT 14	PhT 15		PhT 16	PhT 17		PhT 18		PhT 19	РҺТ 20	PhT 21	PhT 22	PhT 23

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				16S rRNA gene		
			Most closely related sequence (Genbank	sequence similarity in % (no. of		
Phylotype	Isolate	Host lichen	accession number)	nucleotides)	Taxonomical position of isolate (most related species) Notes/reference of most related sequence	Notes/reference of most related sequence
PhT 24	Gint1N	<i>Cladonia pyxidata</i> (from Plankogel)	Streptosporangium sp. 99 (511) 319E04 Streptosporangium sp. 319B09	99 (511)	Actinobacteria, Streptosporangiaceae (Streptosporangium vulgare)	1
PhT 25	GestL3N	<i>Cladonia pyxidata</i> (from Plankogel)	nss	nss	Alphaproteobacteria	1

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à 5 e E 5 dentification the Underlined isolates were used for

Statistically not significant

The ITS 16S-23S profiles showed one to seven different bands for each isolate and we were generally able to unambiguously recognize the bacteria with the same profile. The isolates with unclear differences in ITS 16S-23S profiles (i.e. isolates Aint1 and Bint1, Table 2) were assigned to phylotype based on Amplified Ribosomal DNA Restriction Analysis (ARDRA) performed with three restriction endonucleases (AluI, TaqI and HinfI). Phylotypes 9 and 12 were represented in external parts of three different lichen species, respectively (phylotype 9 in three different genera, phylotype 12 in three species of Cladonia). Both phylotype 1 and phylotype 8 were found in two different Cladonia species. Cladonia digitata and Cladonia rangiferina contained phylotype 1 internally, and phylotype 8 was found inside Cladonia pyxidata and externally in C. rangiferina. Phylotype 15 comprised isolates from both the surface and the inside of Cladonia coccifera. All other phylotypes are represented by a single isolate (Table 2).

Identification of strains and phylogenetic analysis

The 16S ribosomal gene of one isolate representing each of the phylotypes was partially sequenced. Identification by BLASTn searches for significant sequence similarity revealed that seven phylotypes (nine isolates) represent the genus Paenibacillus (Firmicutes), four phylotypes (eight isolates) belong to the genus Burkholderia (Betaproteobacteria), and three phylotypes (four isolates) can be assigned to the genus Bacillus (Firmicutes). Phylotype 12 (three isolates) is related to Luteibactor rhizovicina (Gammaproteobacteria), whereas phylotypes 22 and 23 (two isolates) belonged to the Microbacteriaceae (Actinobacteria); however, an unambiguous identification at genus level was not possible. The other phylotypes were identified as Micromonospora sp. (two phylotypes, two isolates), Streptomyces sp. (one phylotype, one isolate), Streptosporangium sp. (one phylotype, one isolate), Cellulomonas sp. (one phylotype, one isolate), Staphylococcus sp. (one phylotype, one isolate) and Inquilinus limosus (one phylotype, one isolate). Phylotype 25, containing one isolate belonging to the Alphaproteobacteria, was not identified at the genus level (Table 2). The phylogenetic analysis of the 16S ribosomal gene sequences resulted in a highly resolved Neighbour-Joining tree with moderate to high bootstrap support values for most clades (Fig. 1). A monophyletic clade of closely related phylotypes (CLU1) can be clearly identified in the Paenibacillus branch. The clade contains the phylotypes 1, 2, 4, 6 and 7, which are closely related to the species P. pabuli, and P. amylolyticus. CLU1 includes seven of the nine isolates identified as Paenibacillus sp., isolated on TY medium from both the internal thallus and the external surface of four lichen species. Phylotype 3 (from C. coccifera) and phylotype 5

(from Pseudevernia furfuracea) represent two further Paenibacillus species isolated from the internal parts of their host thalli (Table 2). A second cluster (CLU2) includes Burkholderia strains, including the phylotypes 8 and 9. These are closely related to the species Burkholderia glathei and B. sordidicola. CLU2 includes six of the eight isolates identified as Burkholderia sp., isolated on TY (phylotype 8) and N-free medium (phylotype 9) from the external surface of five lichen samples (from four different species, one of them from distant sampling sites). A third cluster (CLU3) contained phylotypes 22 and 23 in the Actinobacteria branch; both of these phylotypes were formed by a single isolate identified as Micromonospora sp., related to Micromonospora auratinigra (PhT 23) and Micromonospora matsumotoense (PhT 22) and both were isolated from the internal thallus of Cladonia pyxidata. A fourth cluster (CLU4), including the phylotypes 14 and 17, encompasses the Bacillus branch. These phylotypes are closely related to the species Bacillus bataviensis and Bacillus niacini, and were isolated from the surface of C. coccifera and Hypogymnia physodes, respectively. Distinct from this group are two other phylotypes from Cladonia species: phylotype 15 (related to Bacillus sp. according to Genbank information) and phylotype 16 (related to Staphylococcus epidermidis). Phylotype 12 from three Cladonia species was identified as Luteibactor rhizovicina, which was recently described from the rhizosphere of Hordeum vulgare (Johansen et al., 2005). All others phylotypes were isolated from the external surfaces of lichens, and grouped with other strains found on lichens.

Cultivation-independent analysis

As the DNA extracted from the 11 lichen samples by the CTAB/chloroform-isoamyl alcohol method could not directly be visualized on an ethidium bromide-stained agarose gel, we arbitrarily used 5 µL of crude extract as template for a first PCR reaction to amplify the 16S ribosomal gene and the ITS 16S-23S with conserved primers for Eubacteria. The PCR products were hardly visible on an agarose gel and we therefore used $10\,\mu\text{L}$ of the PCR product as template for a nested PCR to amplify the ITS 16S-23S. The multiple PCR products were clearly visible as distinct bands of variable intensity (Fig. 2). We repeated this approach several times and with different visualization methods to confirm the consistency of the results. The profiles - considering size and fluorescence intensity - were compared to assess the variation in banding patterns between the different communities. A matrix was constructed and used for the calculation of the ecological indices of Shannon-Wiener (diversity index) and Simpson (dominance index). The values of the Shannon-Wiener index ranged between 0.90 for the bacterial community of C. coccifera (lichen sample 'E') and 2.28 for the bacterial community of C. digitata (lichen sample 'A'). The

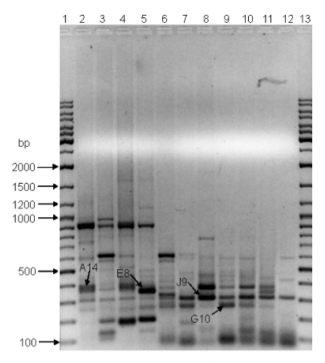
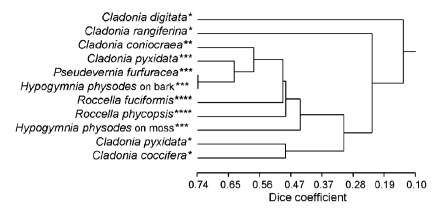
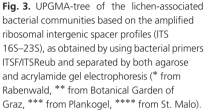


Fig. 2. Bacterial internal transcribed spacer profiles obtained by analysis of total DNA extracts from surface-sterilized thalli of 11 lichen samples (2.5% agarose gel, ethidium bromide stained). Excised and sequenced bands are indicated (see text for details). Lanes 1 and 13: DNA marker; lanes 2–12: lichen samples in the following order: A, B, D, E, R, H, J, G, K, C and S.

average values, considering all 11 lichen-associated bacterial communities, were 1.71 ± 0.38 for the Shannon-Wiener diversity index and 0.23 ± 0.097 for the Simpson dominance index. The comparison showed a high specificity of profiles from the different samples (including unique bands). On the other hand, several bands are shared among different profiles, indicating the occurrence of similar bacteria in different lichens, and from different geographical regions. Four bands were excised and sequenced (Fig. 2). BLAST searches indicated the presence of additional bacteria that were not detected by culture methods. A14 showed the same identity values in the BLAST search with Nitrosomonas and Nitrosospira ssp., which are ammonia-oxidizing Betaproteobacteria. Band E8 did not match with any ITS sequence in Genbank and represents a hitherto unknown bacterial type. Band J9 fits well with Propionibacterium ssp., an obligate anaerobic Actinobacterium, and G10 was identified as Bacillus subtilis, which has previously been found in lichens (Henkel & Plotnikova, 1973).

The comparison of the profiles showed relative similarity values of 14–74% (Fig. 3). According to the ITS fingerprints, the differences of the bacterial community seem to be rather high among the sampling sites, and there is no clear congruence with the relationships of the hosting lichen.





The greatest diversity was found among the *Cladonia* samples, whereas the bacterial communities associated with different lichens sampled in Plankogel, St. Malo and the Botanical Garden of Graz formed a cluster with similarity values of 44%. The bacterial communities from two distantly collected *C. pyxidata* are diverse, whereas the bacterial communities associated with lichens from Rabenwald did not form a unique cluster, owing to the low similarity between the bacterial communities of both *C. digitata* and *C. rangiferina* and the other communities.

Discussion

Although lichen symbioses generally comprise two obligate partners with clear functional roles, i.e. the photobionts (either green algae or cyanobacteria, or sometimes both) and the mycobionts, additional fungi and bacteria can occur as optional symbionts in lichen thalli. In this contribution we used DNA sequence data to investigate the composition of the bacterial communities associated with lichens from temperate habitats and to assess the phylogenetic position of all detected culturable strains. The characterization of the culturable bacterial community was carried out in a twostep approach. First, with the ribosomal intergenic spacer analysis we assessed the genetic variation of strains cultivated from lichens, and second, by the analysis of rRNA gene sequences of a representative of each strain group, we assigned most bacteria to their respective genera and placed the discovered strains in a phylogenetic framework. However, as we generally purified only distinct phenotypes from each lichen sample, we may still have missed other genetically different strains with the same morphology. Nonetheless, we found 25 different phylotypes representing discrete bacterial taxa. The strains were assigned to different genera, of which three belong to Firmicutes, four are found among the Actinobacteria and three among the Proteobacteria. The association of Paenibacillus and Burkholderia with lichens seems to be common, irrespective of relationship

and habitats of the host. Their presence is not surprising, as these genera are known also from associations with diverse other fungi. For example, Burkholderia was isolated from Basidiomycota, from Glomeromycota, and from Zygomycota (Bianciotto et al., 2000; Lim et al., 2003; Partida-Martinez & Hertweck, 2005), and Paenibacillus is found in Basidiomycota and in Glomeromycota (Budi et al., 1999; Poole et al., 2001; Bertaux et al., 2003). Several genetically distinct strains within these genera are associated with lichens. We also detected representatives of actinomycetes, which have previously been found in other studies (Zook, 1983; Gonzáles et al., 2005). However, we did not detect bacteria of the genus Azotobacter, repeatedly mentioned in the literature and isolated from other lichens (reviewed in Lenova & Blum, 1983), nor did we find other genera previously isolated from lichens, such as Clostridium, Bejerinckia or Pseudomonas. We assume that the composition of the bacterial communities in lichens is affected by diverse biotic and abiotic factors, which include the phylogenetic position of the lichens, but also the geographic origin, the substrate, the microhabitat conditions, and the pattern of fungal secondary metabolites.

Our direct analysis of bacterial diversity by PCR approaches using total DNA extracts of surface-sterilized material (using H_2O_2 to degrade superficial DNAs) is a first step towards assessing the entire bacterial diversity within lichen thalli. The results revealed variation among all samples analysed, especially in the mostly soil- and moss-inhabiting representatives of *Cladonia*. Some shared bands suggest the presence of similar strains in different lichens, which agrees with the data from the cultured fraction. Sequence analysis of selected bands revealed additional bacterial strains and further analysis of cloned PCR products is the focus of ongoing studies.

A clear statement cannot be made about the ecological role of lichen-inhabiting bacteria. It is, nevertheless, interesting that many strains are capable of growing readily on N-free media. In the case of *Paenibacillus*, this agrees with published data about strain-specific N-fixing capacities in this genus. Eight strains are known in this genus as nitrogen fixers (Rodriguez Coelho et al., 2003). All our lichenassociated strains of Paenibacillus grow readily on N-free medium, but their closest relative according to rRNA gene sequence data, P. pabuli, is not known as a nitrogen-fixer. Our data would be in accordance with a nitrogen-fixing role of lichen-associated bacterial strains, but further experiments are needed to assess whether fixed nitrogen is available in significant amounts to the symbionts. There is no clear evidence for a defensive role (Zook, 1983), but as antifungal properties are known from bacteria, e.g. from strains in Burkholderia (Opelt & Berg, 2004), this hypothesis should be investigated further. It also remains to be studied whether bacteria use extracellular compounds produced by lichens for their growth and to what extent they are involved in the degradation of lichens (e.g. a chitinolytic activity is known from Paenibacillus and Streptomyces). As bacteria are commonly observed in decaying lichens and especially in the soil interface (e.g. Asta et al., 2001), it is also possible that bacteria could benefit from phenolic substances leaching from lichens (Stark & Hyvärinen, 2003). On the other hand, it might also be argued that lichen-associated bacteria could have a helper effect for the establishment of fungal-algal symbioses, similar to mycorrhizal helper bacteria (Garbaye, 1994). So far, lichen-associated bacteria seem to be present on the surface of thalli or in intercellular spaces (e.g. De los Ríos et al., 2005); their endosymbiotic occurrence is not known. As this was found in diverse non-lichenized fungi, Ascomycota (Barbieri et al., 2000), Basidiomycota (Bertaux et al., 2003, 2005), Glomeromycota (Bianciotto et al., 2000), and Zygomycota (Partida-Martinez & Hertweck, 2005), further investigations might also test whether endosymbioses with bacteria also occur in lichens. The ubiquitous presence of bacteria in lichens will have to be considered in studies of certain paralogs of functional genes in lichen mycobionts, such as MSAS type I polyketide synthases (Kroken et al., 2003), which may occur both in fungi and bacteria. This is also underlined by the recent finding of an abundance of these genes in lichen-associated actinomycetes (Gonzáles et al., 2005). Moreover, the occurrence of Burkholderia or Propionibacterium raises the question whether lichens might also act as a reservoir for opportunistic pathogens, as has been shown for rhizosphere-associated bacteria (Berg et al., 2005).

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