

Characterization of Endolithic Culturable Microbial Communities in Carbonate Rocks from a Typical Karst Canyon in Guizhou (China)

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Abstract

The endolithic environment is a ubiquitous habitat for microorganisms and a critical interface between biology and geology. In this study, a culture-based method and the phylogenetic analysis based on 16S rRNA and internal transcribed spacer (ITS) sequences were used to investigate the diversity of endolithic bacteria and fungi in two main types of carbonate rocks (namely dolomite and limestone) from Nanjiang Canyon in Guizhou karst area, China. The results of bacterial diversity indicated that all bacteria isolated from dolomite and limestone rocks were divided into 4 bacterial groups, including *Proteobacteria*, *Actinobacteria*, *Firmicutes* and *Bacteroidetes*. For these two kinds of rocks, *Proteobacteria* was the first dominant group, and *Gammaproteobacteria* occupied the greatest proportion which might be closely related to *Pseudomonas* in phylogeny to be the most dominant genera after isolation. *Actinobacteria* and *Bacillus* bacteria were also widespread in these two kinds of rock environments. There were only 9 and 8 strains of fungi isolated from dolomite and limestone respectively, which all belonged to *Ascomycota*. To the best of our knowledge, this is the first report on diversity of endolithic culturable bacteria and fungi in carbonate rocks in Guizhou karst region. These microorganisms may play an important and unprecedented role in the carbonate rock weathering during the long history of geological evolution.

Key words: endolithic bacteria, endolithic fungi, carbonate rock

Introduction

The endolithic environment, the tiny pore and crack space in rocks, is a ubiquitous habitat for microorganisms and a critical interface between biology and geology (Friedmann, 1982; Walker and Pace, 2007a). Microorganisms native to the endolithic environment must possess adaptive ability and tolerance to severe stresses including rapid temperature variations, oligotrophy, desiccation, and high UV flux (Sigler *et al.*, 2003; Selbmann *et al.*, 2013). Communities of microorganisms that inhabit endolithic environments include autotrophic and heterotrophic bacteria, fungi, algae, and lichens (Sigler *et al.*, 2003). The endolithic microorganisms rarely existed by means of single colony formed by a single species, while they usually grew collectively to form microbial community with complex, various structures and functions (Gorbushina *et al.*, 2004; Gorbushina, 2007). Nowadays, endolithic microorganisms have been observed not only in a variety of extreme terrestrial ecosystems but also in temperate environments

(Gerrath *et al.*, 2000; Wierzchos *et al.*, 2006; Horath and Bachofen, 2009; Wong *et al.*, 2010; Tang *et al.*, 2012). They also have been detected inhabiting a variety of rock types such as halite, sandstone, quartz, gypsum, dolomite and limestone (Dong *et al.*, 2007; Gorbushina, 2007; Tang *et al.*, 2012; Roldan *et al.*, 2014).

Nowadays, many investigations of endolithic microbial communities used culture-independent techniques (Sigler *et al.*, 2003; Walker and Pace, 2007b; Horath and Bachofen, 2009; Wong *et al.*, 2010). Although, culture-independent methods can provide important information about the diversity of phylogenetic groups in endolithic environments, pure cultures are still indispensable for the determination of the metabolic activity of microorganisms, which can be significantly different even among closely related phylotypes (Spring *et al.*, 2015). In addition, the culturable microorganisms account for about 0.001 ~ 15% of the total microorganisms of environmental samples (Tamaki *et al.*, 2005). Thus, a considerable part identified by culture-independent method is uncultured microbial species. Pure

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culture method played and plays an irreplaceable role in exploring microbial community structures, especially in the following aspects, such as new species or special functional microbes, *etc* (De Leo *et al.*, 2012; Krakova *et al.*, 2015). However, to date, no study has examined the endolithic microbial community in Guizhou karst environment using culture-dependent method. This method has the potential to improve our understanding of microbial diversity and facilitate functional studies of endolithic microorganisms.

Guizhou, a province in southwest of China, is one of the three largest developing karst areas in the world. The carbonate rock area is 130,000 km², covering 73.8% of total land surface area of the province. The weathering of carbonate rocks influences the geochemical compositions of rocks, soils, the atmosphere, and organisms, and the transfer process of matter and energy in the karst environment (Lian *et al.*, 2010). Endolithic microorganisms have been found to be directly and indirectly involved in the weathering of carbonate rocks. Papida *et al.* (2000) found that a mixed microbial population exacerbated physical weathering of carbonate rocks. The endolithic microorganisms play an important and unprecedented role in the carbonate rock weathering during the long history of geological evolution. So, the aim of the present study was therefore to investigate the endolithic microbial diversity in dolomite and limestone rocks in karst area of Guizhou Province using culture-dependent techniques. Combined with the culture-independent method at the early stage (Tang and Lian, 2012; Tang *et al.*, 2012), endolithic microbial diversity of carbonate rocks could be reflected more fully. Meanwhile, the research laid the foundation to further explore the carbonate rock weathering by using the microbial resources in the karst endolithic environment.

Experimental

Materials and Methods

Site description and sample collection. Nanjiang canyon (26°56'N, 106°58'E) is a typical karst canyon, which is praised as "Karst Ecosystem Museum". It is located in Kaiyang County of Guizhou Province, southwest China. Details of the site are given in our previous study (Tang and Lian, 2012; Tang *et al.*, 2012). There are a lot of exposed carbonate rocks in this area. The dolomite and limestone rocks are weathered and porous. The soil developed on its weathering crust is non-zonal soil with shallow soil layers and no apparent genetic layers. In this area, three Triassic dolomite (D1-D3) and three Triassic limestone samples (L1-L3) up to a depth of ~1 cm were collected using a sterile rock chisel and

placed in sterile bags on ice. The samples were kept at 4°C until subjected to microbial isolation. All samples were collected in September 2009.

Powder X-ray diffraction. For mineralogy in the dolomite and limestone rock samples, powder X-ray diffraction patterns were obtained using CuK α radiation at 40 kV with an X-ray diffractometer (D/Max-2200/PC, Rigaku). The results revealed 2 dolomite samples to be pure dolomite rocks and one dolomite sample to be mostly dolomite (99%) with trace amounts of anorthose. The 3 limestone samples were mostly calcite (97–99%) containing trace of dolomite and quartz.

Isolation and enumeration of microorganisms. In accordance with the requirements of aseptic technique, the mineral powder of each crush rock sample was scraped. After serial dilution, they were coated on beef extract-peptone agar medium (5 g/l beef extract, 10 g/l peptone and 15 g/l agar) and Martin culture medium (1 g/l K₂HPO₄, 0.5 g/l MgSO₄·7H₂O, 5 g/l peptone, 10 g/l glucose, 15 ~ 20 g/l agar and 3.3 ml/l 1% streptomycin aqueous solution). Then they were reversely cultivated under constant temperature of 30°C. After 2 ~ 3 days for bacteria or 5 ~ 7 days for fungi, the colony counting could be obtained. The single colonies with different phenotypes, were picked up and the purified strains were transferred to the corresponding agar slants for conservation at 4°C.

DNA extraction, PCR amplification, and sequencing. DNA extraction and PCR amplification were performed using the TIANcombi DNA Lyse&Amp PCR Kit (Tiangen Biochemical technology Co., Ltd., China). The bacterial universal primers 27F (5'-AGAGTTT-GATCCTGGCTCAG-3') and 1492R (5'-GGTTAC-CTTGTTACGACTT-3') and fungal universal primers ITS1 (5'-TCCGTAGGTGAACCTGCGG-3') and ITS4 (5'-TCCTCCGCTTATTGATATGC-3'), were used to amplify the bacterial 16S rDNA and fungal internal transcribed spacer (ITS) sequences, respectively. All of the primers were synthesized by Shanghai Sangong Co., Ltd., China.

All reactions were carried out in a 50 μ l reaction mixture, containing 25 μ l of 2 \times PCR Reagent, 1.25 μ l of forward primer and reverse primer (10 μ M) respectively and 5 μ l of DNA template. PCR was run under the following conditions: initial denaturation at 94°C for 5 min; followed by 30 circles of 1 min denaturation at 94°C, 1 min annealing at 52°C, 2 min extension at 72°C, and a final extension step of 10 min at 72°C. The results of PCR amplification were confirmed using 1% agarose gel, stained with SYBR Safe 10.000 \times in DMSO (Invitrogen). PCR products were subsequently purified using PCR purification kit (E.A.N.A. Gel Extraction kit, OMEGA, USA) and subjected to sequencing in an ABI PRISM 3730 automatic sequencer (Shanghai Sangon Co., Ltd., China).

Phylogenetic analysis. DNA sequences were analyzed using Bellerophon (Huber *et al.*, 2004) and CHIMERA_CHECK (Cole *et al.*, 2003) software to remove chimeric artifacts. The sequences were considered to be of the same phylotype if they were $\geq 97\%$ similar to one another over the region of the 16S or ITS rRNA gene sequenced (Stackebrandt and Ebers, 2006). Then the effective sequences were compared with known sequences in the NCBI database (<http://www.ncbi.nih.gov/>) by Basic Local Alignment Search Tool (BLAST) (Altschul *et al.*, 1990). The sequences were aligned using Clustal X software (Thompson *et al.*, 1997), and phylogenetic trees were constructed with the Mega 4.0 program package using the neighbor-joining method (Kumar *et al.*, 2004). Bootstrap confidence values were obtained with 1000 replicates. The trees were constructed by calculating the Kimura distance (Kimura, 1980).

Nucleotide sequence Accession numbers. The nucleotide sequences reported in this study have been deposited in GenBank under accession numbers JN650544-JN650602.

Results

Isolation and enumeration of microorganisms.

Beef extract-peptone agar medium was utilized to isolate bacteria of rock samples. There were 39 strains and 26 strains of bacteria with different phenotypic characteristics in dolomite and limestone respectively. Fungi of rock samples were separated by Martin culture medium, and dolomite and limestone respectively obtained 9 and 8 strains of fungi with different colony phenotypes. It was calculated that the culturable bacteria of dolomite and limestone were about 2.3×10^5 CFU/g and 2.0×10^5 CFU/g by using dilution-plate method; while the culturable fungi were about 7.7×10^3 CFU/g and 3.0×10^3 CFU/g respectively.

Diversity of the bacterial 16S rRNA gene sequences from dolomite rocks. There were 39 effective sequences in total by sequencing 16S rDNA for the isolates of 39 strains of dolomite. The alignment results of similarity indicated that they were classified into 4 phyla, 15 genera and 26 species (Table I). These 39 strains of bacteria showed high similarity with the known 16S rDNA sequences in database, namely 98 ~ 100%.

The sequence analysis of 16S rDNA (see Table I and Fig. 1) illustrates that 39 strains of bacteria from dolomite were classified into 4 major bacteria groups. More than 53.8% of them belonged to *Proteobacteria*, which including 12 strains of *Gammaproteobacteria*, accounting for 30.8%; 5 strains of *Alphaproteobacteria*, accounting for 12.8%; and 4 strains of *Betaproteobacteria*, accounting for 10.2%. In addition, 10 strains were from *Actinobacteria*, and accounted for 25.7%, while 7 strains

were *Firmicutes*, accounting for 17.9%. Moreover, one strain belonged to *Bacteroidetes*, accounting for 2.6%.

Gammaproteobacteria, as one subphylum of *Proteobacteria*, accounted for the highest ratio of culturable bacteria in the endolithic dolomite environment so that it was regarded as the most dominant group obtained by isolation. The 10 strains of bacteria in the group, accounting for 25.6%, had close relations with *Pseudomonas* in phylogeny, and the similarity of 16S rDNA sequences was 99%. Thus, the group was the most dominant bacterial genus acquired by isolation. Therein, four strains were closely associated with *Pseudomonas putida*, 1 strain was highly homologous to *Pseudomonas nitroreducens*. In *Alphaproteobacteria*, there were 2 strains which were closely related to *Brevundimonas* and *Sphingobium* respectively and 1 strain had homology of 99% with *Sphingobium*. Besides, 4 strains of bacteria belonged to *Betaproteobacteria*, where 2 strains were *Duganella* and the other 2 strains belonged to *Marseille*.

Actinobacteria was the second dominant group of the culturable bacteria from dolomite. In this group, 4 strains of bacteria had a close genetic relationship with *Arthrobacter*; three of them were highly homologous with *Rhodococcus*; the homology of two strains and *Streptomyces zaomyeticus* (FJ792553) reached 100%; and the rest one was 99% homologous with *Microbacterium oxydans* (HQ202812).

There were 7 strains of bacteria in *Firmicutes*, including 6 *Bacillus*, which were the second dominant genera of the culturable bacteria from dolomite. In this group, three of them were 99% homologous with *Bacillus cereus* (GU969128), two showed high homology with *Bacillus sphaericus* (DQ286315), and one was 100% homologous with *Bacillus simplex* (JF496520). Moreover, DCB46 was 99% homologous with *Paenibacillus* sp. (FJ006903).

Only one strain of bacteria belonged to *Bacteroidetes*, which was 99% homologous with *Dyadobacter* sp. (HQ231938).

Diversity of the bacterial 16S rRNA gene sequences from limestone rocks. 16S rDNA of 26 bacterium isolates from limestone was sequenced to obtain 26 effective sequences in total. The relationships between 16S rDNA sequence of the bacteria isolated from limestone and the most similar sequence in GenBank database are listed in Table II. It can be seen that they were divided into 4 phyla, 13 genera and 17 species. There was a high similarity between the 26 strains of bacteria and the known 16S rDNA sequences in GenBank database. Among them, the similarity of LCB37 was 96% only, while that of the rest was higher than 97%.

As indicated by 16S rDNA phylogenetic analysis, the 26 strains of bacteria isolated from limestone were assigned to 4 major phylogenetic groups (Fig. 2). It can

Table I
Similarity analysis of 16S rDNA sequences of isolates from dolomite rocks

Representative isolate (accession no.)	No. of isolates in OUT*	Nearest type strain (accession no.)	Similarity (%)
Actinobacteria	10		
<i>Arthrobacter</i>	4		
DCB14(JN650552)	2	<i>Arthrobacter</i> sp. (HM165265)	99
DCB16(JN650554)	1	<i>Arthrobacter</i> sp. (HQ166103)	99
DCB48(JN650561)	1	<i>Arthrobacter</i> sp. (FN794213)	99
<i>Microbacterium</i>	1		
DCB18(JN650556)	1	<i>Microbacterium oxydans</i> (HQ202812)	99
<i>Rhodococcus</i>	3		
DCB15(JN650553)	1	<i>Rhodococcus</i> sp. (JF900087)	99
DCB17(JN650555)	2	<i>Rhodococcus</i> sp. (EF612316)	99
<i>Streptomyces</i>	2		
DCB61(JN650569)	2	<i>Streptomyces zaomyceticus</i> (FJ792553)	100
Bacteroidetes	1		
<i>Dyadobacter</i>	1		
DCB47(JN650560)	1	<i>Dyadobacter</i> sp. (HQ231938)	99
Firmicutes	7		
<i>Paenibacillus</i>	1		
DCB46 (JN650559)	1	<i>Paenibacillus</i> sp. (FJ006903)	99
<i>Bacillus</i>	6		
DCB1(JN650544)	3	<i>Bacillus cereus</i> (GU969128)	99
DCB23(JN650558)	1	<i>Bacillus simplex</i> (JF496520)	100
DCB8(JN650548)	2	<i>Bacillus sphaericus</i> (DQ286315)	99
Proteobacteria	21		
Alphaproteobacteria	5		
<i>Brevundimonas</i>	2		
DCB19(JN650557)	1	<i>Brevundimonas vesicularis</i> (FJ999941)	99
DCB60(JN650568)	1	<i>Brevundimonas</i> sp. (DQ778038)	99
<i>Methylobacterium</i>	1		
DCB49(JN650562)	1	<i>Methylobacterium</i> sp. (FN868959)	99
<i>Sphingobium</i>	2		
DCB53(JN650564)	2	<i>Sphingobium amiense</i> (JF459961)	100
Betaproteobacteria	4		
<i>Duganella</i>	2		
DCB58(JN650566)	1	<i>Duganella</i> sp. (JF778719)	99
DCB59(JN650567)	1	<i>Duganella zoogloeoides</i> (NR_025833)	99
<i>Massilia</i>	2		
DCB52(JN650563)	1	<i>Massilia</i> sp. (DQ257420)	99
DCB56(JN650565)	1	<i>Massilia</i> sp. (AB552860)	98
Gammaproteobacteria	12		
<i>Enterobacter</i>	1		
DCB6(JN650546)	1	<i>Enterobacter amnigenus</i> (DQ481471)	99
<i>Pseudomonas</i>	10		
DCB7(JN650547)	3	<i>Pseudomonas</i> sp. (FN547413)	99
DCB11(JN650550)	2	<i>Pseudomonas</i> sp. (AM934695)	99
DCB13(JN650551)	4	<i>Pseudomonas putida</i> (EU118779)	99
DCB74(JN650570)	1	<i>Pseudomonas nitroreducens</i> (EU500825)	99
<i>Serratia</i>	1		
DCB5(JN650545)	1	<i>Serratia fonticola</i> (HQ407251)	99

* The strains were clustered into operational taxonomic units (OTUs) at a level of sequence similarity $\geq 97\%$.

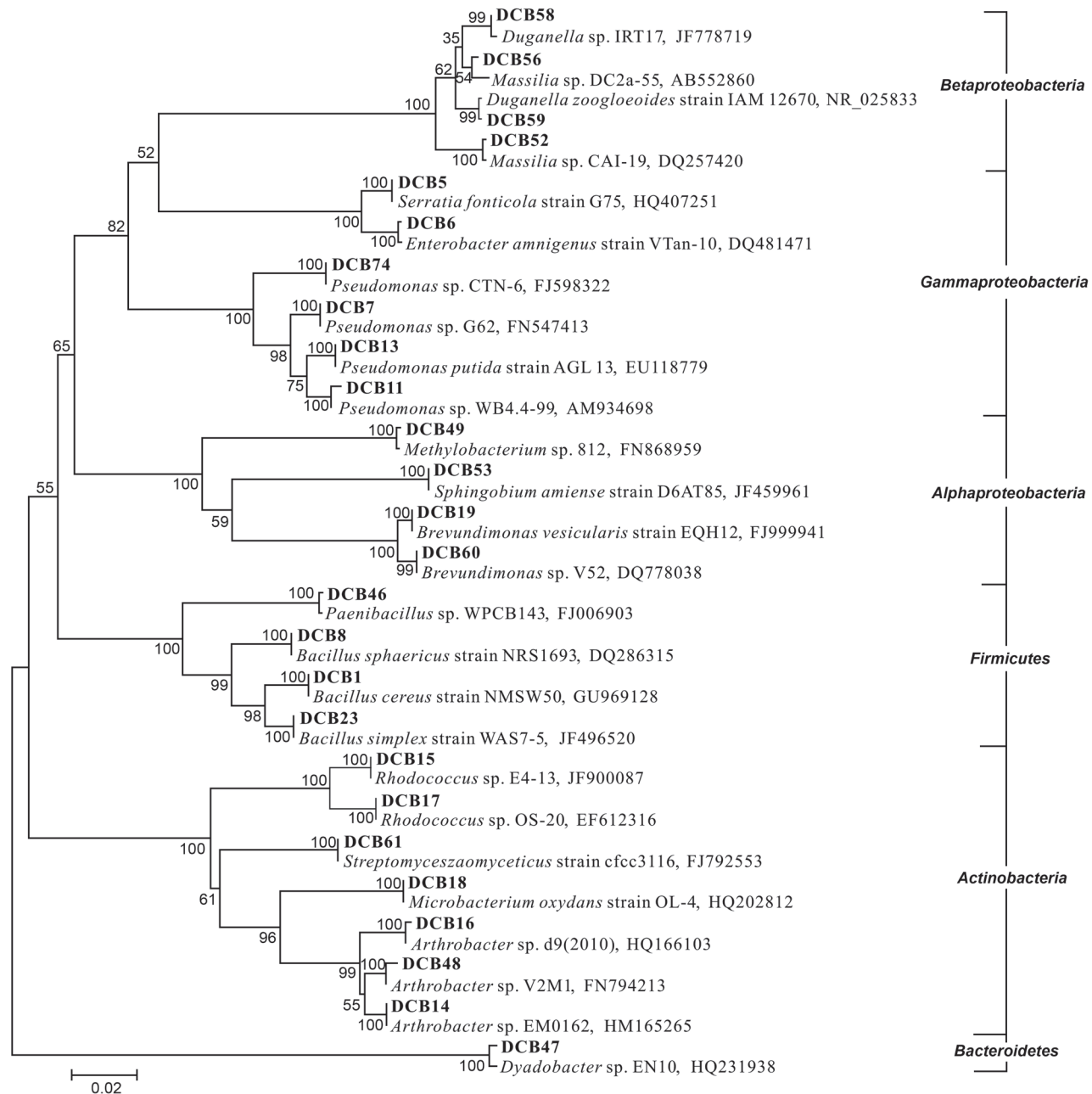


Fig. 1. Phylogenetic relationship based on 16S rRNA gene sequences of culturable bacterial clones isolated from dolomite rocks (in bold type) with closely related sequences from the GenBank database. Neighbor-joining trees; bootstrap values (1,000 replicates) are shown at the nodes

be known that *Proteobacteria* played a dominant role in bacterial diversity and had 18 strains, taking up 69.3%, including 8 *Betaproteobacteria* and *Gammaproteobacteria*, accounting for 30.8% respectively; 2 strains of *Alphaproteobacteria*, accounting for 7.7%. In addition, three strains belong to *Actinobacteria* and three of them were *Bacteroidetes*, which accounted for 11.5%. Moreover, there were 2 strains of *Firmicutes* (7.7%).

As for *Proteobacteria*, 8 of them belonged to *Gammaproteobacteria* subphylum, and all of them were *Pseudomonas*, which was regarded as the first dominant genera of culturable bacteria from limestone. In the subphylum, LCB43 had 100% similarity to dolo-

mite isolate DCB13, which means that they were the same kind of bacteria. In addition, the similarity between them and *P. putida* (EU118779) reached 99%. The other 7 strains of bacteria were 100% similar to *Pseudomonas* sp. (FJ006865), while as for the species of *Pseudomonas*, it was needed to be further identified. There were 8 strains of *Betaproteobacteria*, including 4 *Marseille bacteria*, which were sub-dominant genera of culturable bacteria from limestone. Moreover, 2 *Janthinobacterium*, a *Duganella*, and a *Methylibium* were included. Only 2 strains in *Alphaproteobacteria* showed 100% homology with *Brevundimonas intermedia* (JF915342).

Table II
Similarity analysis of 16S rDNA sequences of isolates from limestone rocks.

Representative isolate (accession no.)	No. of isolates in OUT*	Nearest type strain (accession no.)	Similarity (%)
Actinobacteria	3		
<i>Arthrobacter</i>	1		
LCB65 (JN650582)	1	<i>Arthrobacter</i> sp. (HQ231937)	99
<i>Microbacterium</i>	1		
LCB38 (JN650577)	1	<i>Microbacterium oxydans</i> (HQ316116)	99
<i>Oerskovia</i>	1		
LCB39 (JN650578)	1	<i>Oerskovia paurometabola</i> (HM244951)	100
Bacteroidetes	3		
<i>Flavobacterium</i>	2		
LCB34 (JN650574)	1	<i>Flavobacterium</i> sp. (HQ425305)	98
LCB37 (JN650576)	1	<i>Flavobacterium</i> sp. (DQ178976)	96
<i>Sphingobacterium</i>	1		
LCB40 (JN650579)	1	<i>Sphingobacterium</i> sp. (AM411998)	99
Firmicutes	2		
<i>Bacillus</i>	1		
LCB77 (JN650587)	1	<i>Bacillus cereus</i> (GU969128)	99
<i>Staphylococcus</i>	1		
LCB44 (JN650581)	1	<i>Staphylococcus</i> sp. (EU855210)	99
Proteobacteria	18		
Alphaproteobacteria	2		
<i>Brevundimonas</i>	2		
LCB32 (JN650573)	2	<i>Brevundimonas intermedia</i> (JF915342)	100
Betaproteobacteria	8		
<i>Methylibium</i>	1		
LCB69 (JN650585)	1	<i>Methylibium</i> sp. (FJ615290)	99
<i>Duganella</i>	1		
LCB66 (JN650583)	1	<i>Duganella zoogloeoides</i> (NR_025833)	99
<i>Janthinobacterium</i>	2		
LCB30 (JN650572)	2	<i>Janthinobacterium</i> sp. (FN293043)	99
<i>Massilia</i>	4		
LCB36 (JN650576)	2	<i>Massilia</i> sp. (DQ257420)	99
LCB71 (JN650586)	1	<i>Massilia</i> sp. (AB552860)	98
LCB67 (JN650584)	1	<i>Massilia</i> sp. (FR865960)	99
Gammaproteobacteria	8		
<i>Pseudomonas</i>	8		
LCB43 (JN650580)	1	<i>Pseudomonas putida</i> (EU118779)	99
LCB28 (JN650571)	7	<i>Pseudomonas</i> sp. (FJ006865)	100

* The strains were clustered into operational taxonomic units (OTUs) at a level of sequence similarity $\geq 97\%$

Three strains of bacteria in *Bacteroidetes* group were closely related to *Flavobacterium* and *Sphingobacterium*, while 3 strains of bacteria in *Actinobacteria* group were *Arthrobacter* sp., *Microbacterium* sp. and *Oerskovia* sp. respectively. Besides, the similarity between LCB77 of *Firmicutes* group and dolomite isolate DCB1 was up to 100%. They were 99% similar with 16S rDNA sequences

of *B. cereus* (EU979024), namely type strain. In addition, an isolate of *Firmicutes* had close relations with *Staphylococcus*.

Diversity of culturable fungi from dolomite and limestone rocks. ITS sequences for 9 strains of fungi isolates from dolomite and 8 strains of fungi isolates from limestone were compared with the sequence

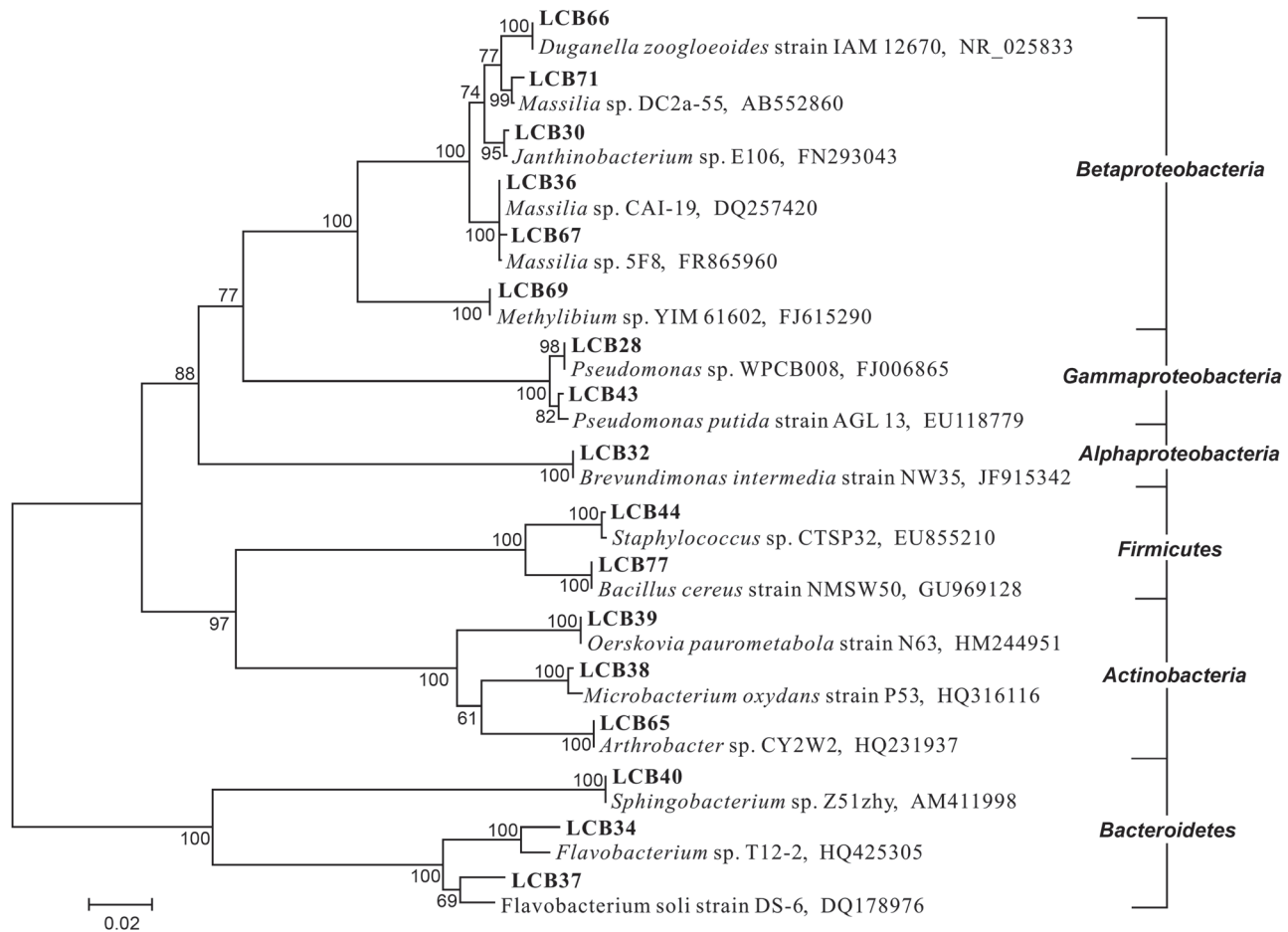


Fig. 2. Phylogenetic relationship based on 16S rRNA gene sequences of culturable bacterial clones isolated from limestone rocks (in bold type) with closely related sequences from the GenBank database. Neighbor-joining trees; bootstrap values (1,000 replicates) are shown at the nodes

of GenBank database to obtain the most similar sequences. Table III shows the relationships between ITS sequences of fungi isolates from dolomite and limestone and the most similar sequences in GenBank database. The results indicated that 9 isolates from dolomite were classified into 1 phylum, 6 genera and 7 species; while the 8 strains of fungi isolates from limestone were divided into 1 phylum, 7 genera and 8 species. They all had high similarity (98 ~ 100%) with the reported ITS sequences of GenBank database, including 12 strains with 100% similarity.

A phylogenetic tree was constructed based on the sequences with high similarity between the reference sequences in database and the strains isolated from dolomite and limestone, as shown in Fig. 3. The results manifested that 9 strains of fungal isolates from dolomite and 8 strains from limestone were *Ascomycota*. Among the fungal isolates from dolomite, 3 of them were *Aspergillus*, which was the genus with the most isolates. There were 2 strains with 100% similarity to a strain of *Aspergillus niger* (EF592173), while the other one had 100% similarity with a strain of *Aspergillus oryzae* (HQ285588). There was also an isolate belonging

to *Penicillium*, which was 100% similar with *Penicillium crustosum* (GQ340556). Besides, the other four strains were closely associated with *Aureobasidium pullulans* (JF439462) of *Aureobasidium*, *Myrothecium verrucaria* (GQ131886) of *Myrothecium*, *Purpureocillium lilacinum* (HM439952) of *Paecilomyces* and *Trichoderma* sp. (GQ497170) respectively, and the homology was as high as 99 ~ 100%.

The 8 fungi isolates from limestone belonged to 7 genera of *Ascomycota*, including *Alternaria*, *Bipolaris*, *Chaetomium*, *Cryptococcus*, *Fusarium*, *Penicillium* and *Trichoderma*. Except that there were 2 isolates in *Chaetomium*, the other ones were divided into different genera. From the phylogenetic tree, it can be seen that isolate LCF94 and the fungal isolate from dolomite, namely DCF87 were 100% similar, which suggested that they were the same kind of fungi. In addition, the similarity between them and *P. crustosum* (GQ340556) of *Penicillium* was 100%. Moreover, isolate LCF98 and fungal isolate from dolomite DCF83 showed similarity of 100%. It means that they were the same kind of bacteria and both of them were *Trichoderma*. Therefore, the fungal strains of *Penicillium* and *Trichoderma* were

Table III
Similarity analysis of ITS sequences of isolates from dolomite and limestone rocks.

Representative isolate (accession no.)	No. of isolates in OUT*	Nearest type strain (accession no.)	Similarity (%)
Dolomite samples			
<i>Ascomycota</i>	9		
<i>Aureobasidium</i>	1		
DCF84 (JN650590)	1	<i>Aureobasidium pullulans</i> (JF439462)	100
<i>Aspergillus</i>	3		
DCF86 (JN650591)	2	<i>Aspergillus niger</i> (EF592173)	100
DCF91 (JN650594)	1	<i>Aspergillus oryzae</i> (HQ285588)	100
<i>Myrothecium</i>	1		
DCF88 (JN650593)	1	<i>Myrothecium verrucaria</i> (GQ131886)	99
<i>Penicillium</i>	1		
DCF87 (JN650592)	1	<i>Penicillium crustosum</i> (GQ340556)	100
<i>Purpureocillium</i>	2		
DCF81 (JN650588)	2	<i>Purpureocillium lilacinum</i> (HM439952)	100
<i>Trichoderma</i>	1		
DCF83 (JN650589)	1	<i>Trichoderma</i> sp. (GQ497170)	100
Limestone samples			
<i>Ascomycota</i>	8		
<i>Alternaria</i>	1		
LCF97 (JN650600)	1	<i>Alternaria</i> sp. (HQ833813)	100
<i>Bipolaris</i>	1		
LCF96 (JN650599)	1	<i>Bipolaris</i> sp. (FJ770073)	98
<i>Chaetomium</i>	2		
LCF92 (JN650595)	1	<i>Chaetomium</i> sp. (HQ607819)	99
LCF93 (JN650596)	1	<i>Chaetomium nigricolor</i> (JF439467)	99
<i>Cryptococcus</i>	1		
LCF99 (JN650602)	1	<i>Cryptococcus albidus</i> (AF444355)	100
<i>Fusarium</i>	1		
LCF95 (JN650598)	1	<i>Fusarium oxysporum</i> (EF495237)	99
<i>Penicillium</i>	1		
LCF94 (JN650597)	1	<i>Penicillium crustosum</i> (GQ340556)	100
<i>Trichoderma</i>	1		
LCF98 (JN650601)	1	<i>Trichoderma</i> sp. (GQ497170)	100

* The strains were clustered into operational taxonomic units (OTUs) at a level of sequence similarity $\geq 97\%$

isolated from samples of dolomite and limestone. It can be seen that the two types of fungi existed in both kinds of rocks studied in the research.

Discussion

As a special habitat, carbonate rock provided a great challenge to the organisms living in it owing to the features of rocks, such as arid environment, nutrition deficiency, great temperature fluctuations, *etc.* However, there were various microorganisms on its surface and in its cracks, which played a crucial role in

biogeochemical cycle. Although molecular biological method has been developed during exploring and revealing microbial diversity, it is still necessary to separate and purify microorganisms to understand their ecological functions in biosphere. Thereby, culture-dependent method is still an essential approach to research microbial diversity.

A total of 65 bacterial strains and 17 fungal strains were isolated from dolomite and limestone by using pure culture method. The 39 isolated bacterial strains from dolomite samples were divided into 4 phyla, 15 genera and 26 species; while the 26 bacterial strains of limestone were classified into 4 phyla, 13 genera

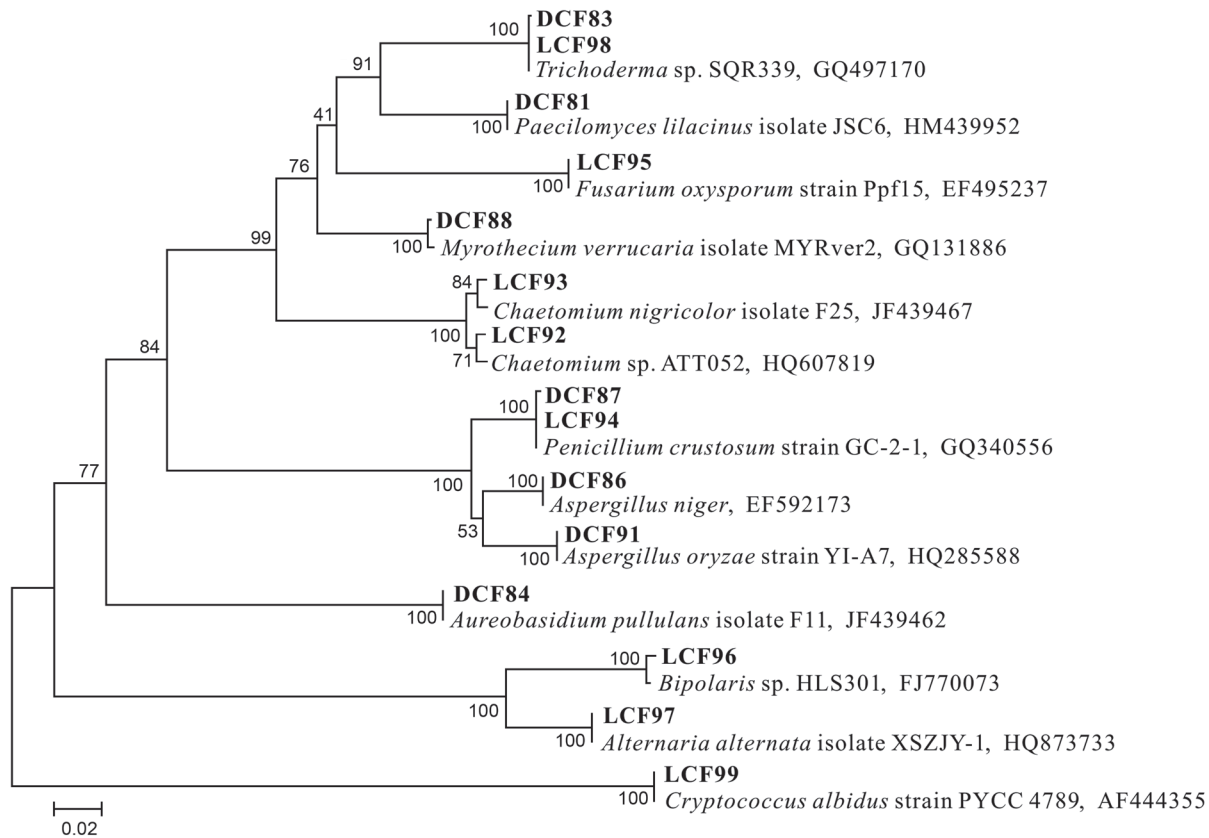


Fig. 3. Phylogenetic relationship based on ITS gene sequences of culturable fungal clones isolated from dolomite (DCF) and limestone (LCF) rocks (in bold type) with closely related sequences from the GenBank database. Neighbor-joining trees; bootstrap values (1,000 replicates) are shown at the nodes

and 17 species. The main phyla of the isolated bacteria from both kinds of rocks were consistent, but there were some differences in species. In addition, the diversity of bacteria was much higher than that of fungi. For these two kinds of rocks, *Proteobacteria* was the first dominant group, and *Gammaproteobacteria* occupied the greatest proportion. Besides, it was closely related to *Pseudomonas* in phylogeny to be the most dominant genera after isolation. *Pseudomonas* is a genus composed of ubiquitous metabolically versatile Gram-negative bacteria, capable to grow in a variety of environmental conditions (Belgini *et al.*, 2014). Some researchers found that *Pseudomonas* also existed in the soil of Antarctic area even in deep sea and it was connected with polycyclic aromatic hydrocarbons (PAHs). Moreover, some *Pseudomonas* even was directly involved in degradation (Ma *et al.*, 2006; Cui *et al.*, 2008). A lot of pseudomonads bacteria were found in the dolomite and limestone rock endolithic environments, but their roles in rock environment were still unclear. The proportion of *Actinobacteria* in two kinds of rocks was greater. Meanwhile, the group is widespread in soil. An explanation for the high fraction of *Actinobacteria* in Guizhou karst region could be due to their strong cell wall, the capability of forming spores, and their high GC-content. These characteristics would

allow their survival in harsh environments. Many bacteria belonging to *Bacillus* were isolated from the rock environment. Owing to the kind of bacteria forming the spores, which had powerful resistance to poor environment, they could survive in rock environment. Our results confirmed that members of the *Proteobacteria*, *Actinobacteria* and *Bacillus* are ecologically significant constituents of carbonate rocks. However, future work is needed to determine the functions of these important groups in karst environments.

There were only 9 and 8 strains of fungi isolated from dolomite and limestone respectively, which all belonged to *Ascomycota*. Besides, they were common fungi in environment which suggested that the culturable fungi resources were very limited on the two kinds of rocks. In view of numerous slow growing fungi colonize carbonate rocks, the cultivating time of fungi in the study might be short. With longer periods of time, it is possible that a more realistic and abundant fungal diversity would have been registered. Nevertheless, the roles of these fungi in rock environment have not been reported so far. Hence, it is necessary to study the functions of these microorganisms in karst rock environment, especially the roles in the process of weathering and pedogenesis of rocks by further utilizing these microbial resources.

Combining the research results and our previous report about the endolithic microbial diversity in the carbonate rocks in karst area of Guizhou by using culture-independent method (Tang and Lian, 2012; Tang *et al.*, 2012), it was indicated that the microbial diversity of carbonate rocks was abundant. There were autotrophic microorganisms, such as *Cyanobacteria* with nitrogen fixation and photosynthesis, anoxygenic phototrophic *green non-sulfur bacteria*, as well as heterotrophic microorganisms, for instance, *Proteobacteria*, *Acidobacteria*, *Bacteroides*, *Firmicutes* and heterotrophic fungi as well as *Archaea*. The autotrophic microorganisms could be the main primary producer in rock environment to provide nutrients for heterotrophic microorganism by fixing the carbon and nitrogen in the air. However, it was not enough for the numerous heterotrophic groups in rock environment merely depending on the nutrient substances supplied by autotrophic microorganisms (Lian *et al.*, 2010). Owing to it tended to be more serious for heterotrophic microorganisms in rock environment, they survived by intercepting the few soil particles and nutrients in air and water except for being symbiotic with autotrophic microorganisms (Sterflinger, 2000; Viles and Gorbushina, 2003; Gorbushina, 2007; Lian *et al.*, 2010; 2011). The rock microorganisms were in a relation of mutual cooperation and symbiosis (Lian *et al.*, 2010). The main purpose of cooperation between different microorganism groups was to retain water and gain limited trace nutrients to sustain life activity and population continuity (Sterflinger, 2000; Gorbushina *et al.*, 2003; Lian *et al.*, 2011). Once the environmental conditions, such as temperature and humidity fitted for growth, they would quickly breed, spread and accelerate the weathering of minerals (Chen *et al.*, 2014). The rock microorganisms in karst area played an irreplaceable role in the long geological evolution process of weathering and pedogenesis of carbonate rocks. Therefore, the future study is expected to further investigate the microbial community in the carbonate rock environments and deeply recognize it in special habitat so as to profoundly research its biogeochemistry function, such as the weathering and pedogenesis of carbonate rocks by using these microbial resources.

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