



34 Abstract

35 This study investigated the long-term effect of environmental physical factors on the
36 relative abundance of bacteria and the consequential landscape evolution in karst topography,
37 focusing mainly on the effects of limestone weathering and calcite precipitation. The Narrow-
38 Sky located in the upper part of Takangshan is a small gulch of Pleistocene coralline limestone
39 formation in southern Taiwan. The landscapes were different in the karst walls between the
40 opening and the inner of gulch due to the variation of physical parameters such as sunlight
41 penetration, humidity, and temperature. A metagenomic approach was used out to determine
42 the relationship of microbial community structures on the landscapes in various habitats around
43 the gulch, namely on the inner and outer limestone wall, the water collected from speleothems
44 surface, and the ground soil at the outer wall. The total organic carbon content was measured
45 in solid samples to evaluate the biomass of the habitats. Our results showed that the biomass
46 of habitats in the opening of the gulch was two times higher than the that inside where light
47 penetration was lower. We also found that speleothems only occurred at the inner wall inside
48 the gulch, where the environment exhibited water drips running through the surface of
49 speleothems and less light penetration. The metagenomics in each habitat were surveyed to
50 measure the sequence similarity of operational taxonomic units relative to urease-producing
51 bacteria and weathering-associated bacteria available in the National Center for Biotechnology
52 Information database. Our data revealed that the metagenomics of the inner wall and water
53 samples exhibited more sequences that were similar to those of urease-producing bacteria,
54 whereas the outer wall showed more sequences that were similar to those of weathering-
55 associated bacteria, suggesting that bacteria facilitated the formation of limestone weathering
56 and calcite precipitation for various habitats. This study revealed the pivotal roles of
57 microorganisms in governing the geological evolution on the limestone landscape.

58



59 **Introduction**

60 Weathering and calcite precipitation are two opposite activities that affect the dynamic
61 changes of the karst landscape. Although weathering and calcite precipitation can occur in
62 abiotic conditions, several lines of evidence from cave studies or laboratory data have shown
63 that microorganisms can accelerate the reactions that promote the formation of calcium
64 carbonate and the breakdown of calcite in situ (Gat et al., 2014;Sulu-Gambari, 2011;Castanier
65 et al., 1999;Lian et al., 2008;Jones, 2017). During the breakdown of carbonate rocks, microbial
66 colonies build up on rock surfaces, resulting in rock decomposition by acidification and
67 moisturization onto the surfaces (Wu et al., 2017;Hutchens, 2009;Uroz et al., 2009). The
68 obtainment of nutrients from the rock surface further promotes the release of organic ligands,
69 which in turn facilitate the release of mineral elements, thus creating a positive feedback loop
70 (Lian et al., 2008;Uroz et al., 2007). Many studies have documented that the mineral
71 dissolution of rocks in a flow-through system was higher in the presence of microorganisms,
72 whereas the dissolution was enhanced in the groups of surface-attached microorganisms,
73 especially when compared with the unattached ones (Ahmed and Holmström, 2015;Seiffert et
74 al., 2014;Jacobson and Wu, 2009). Many bacterial strains have been reported to have the ability
75 to adhere to rock surfaces and establish the weatherability (Sulu-Gambari, 2011). For example,
76 *Shewanella oneidensis* can recognize silicate and oxide mineral surfaces and cause further
77 weathering associated reactions (Lower et al., 2001). To date, many studies have categorized
78 the bacteria of weatherability (Uroz et al., 2009;Sulu-Gambari, 2011;Lian et al., 2008).

79

80 Many bacteria can induce the biomineralization processes of calcium carbonate
81 precipitation that render the formation of stalactite. The microbial-induced reaction is mainly
82 carried out by urease-producing bacteria in the presence of ammonium ions in the alkaline
83 environment. The identified urease-producing bacteria have been investigated extensively
84 (Anbu et al., 2016;Wei et al., 2015;Ercole et al., 2001;Jones, 2017;Animesh and Ramkrishnan,
85 2016;Abo-El-Enein et al., 2012). The microbial communities of karst habitats are diverse, and
86 their components largely depend on the locations and composition of limestone (Barton and
87 Northup, 2007;Ortiz et al., 2014;Tomczyk-Żak and Zielenkiewicz, 2015;Engel, 2010).
88 Temperature, light intensity, and light penetration are important parameters that control the
89 developing orientation of microbial communities. Researchers have shown that
90 microorganisms, operating together with the local environmental conditions, play important
91 roles in remodeling the landscapes of karst (Castanier et al., 1999;Mortensen et al.,



92 2011;Qabany et al., 2012;Anbu et al., 2016). However, how physical factors affect microbial
93 communities and the consequent geological changes remains unknown. Determining how
94 microbial relative abundance shifted in response to changes in environmental factors and the
95 consequent geological evolution can enable us to better understand the effect of
96 microorganisms on the dynamic alterations of karst landscapes.

97

98 The most abundant limestone is found in warm and humid regions. Because of its porous
99 and loose property, limestone can be easily infiltrated by rainfall or groundwater to form
100 trenches, shallow concavity, or clefts. Limestone landscapes in Taiwan are scattered all over
101 the island and can be found in the Hengchun Peninsula, east coastal areas, central range, and
102 southwest of Taiwan. The tectonic studies of Takangshan revealed that the upthrow consists of
103 large lenses of Pleistocene anticline, 4 km in length and 2 km in width (Lacombe et al., 1997;
104 M.L. Hsieh and Knuepfer, 2001). The crest of Takangshan is covered with coral reef limestones
105 (with an average thickness of 40 m), which are interbedded in clastic layers. On top of the hills,
106 expanding vegetation coverages, coupled with erosion soil, are commonly observed in most of
107 these limestone landscapes. The Narrow-Sky is a nickname for a mountain crack located at a
108 limestone hill in the Tainliao district of Taiwan. The dimensions (length, width, and height) of
109 the gulch is approximately 100 m, 2 m, and 12 m, respectively. Because of vegetation
110 coverages and its topographic features, the exposure of sunlight at different spots inside the
111 gulch is different. For example, sunlight can penetrate the limestone wall of the opening
112 through the vegetation coverage, while it is relatively dim inside the path of the gulch.
113 Moreover, moisture and temperature are also different between the opening and the center of
114 the gulch. The most tangible difference between the inner and outer wall of the gulch is the
115 formation of speleothems, which are plentiful in the inner gulch and are nonexistent in the outer
116 section. Because microbial communities are sensitive to changes in environmental physical
117 factors, the microbial composition in different locations may have adapted to the environment
118 according to physical factors, which may play a role in reshaping the gulch scenery.

119

120 In this study, we investigated the effect of physical factors on microbial communities in the
121 limestone landscape. With the recent advent of next-generation sequencing (NGS) platform
122 and computational methods, we could conduct genome studies on microbes to determine the
123 relationship between environmental factors in their habitats, such as sunlight penetration during
124 daytime, humidity, and pH and the relative abundance of microbes. We collected samples from
125 limestone walls at the opening and inside the gulch, from water dripping at the inner limestone



126 wall, and from the soil of the outer weathered limestone at the gulch opening: we collected
127 these samples to extract DNA. Genomic DNA extracted from these samples was further
128 subjected to the PCR amplification of 16S rRNA gene sequences by using the Illumina's MiSeq
129 system. Bioinformatics tools were employed to explore DNA reads in operational taxonomic
130 units (OTUs). The DNA sequence in each OTU was blasted with the sequences which is current
131 weathering bacteria and urease-producing bacteria available in the National Center for
132 Biotechnology Information (NCBI) database.



133 **2. Materials and methods**

134

135 **2.1 Sample Site Description and the Collection of Samples and Physical Parameters**

136 We collected samples from the limestone gulch of Tainliao (120° 21' 19.1" E, 22° 51'
137 00.7" N): the location is illustrated in Figure 1. For collecting the microorganisms in the
138 surface of limestone walls, sterile cotton swabs were used to wipe the surface areas of
139 sampled spots. The samples were collected in a tube and sent to the laboratory to measure
140 total organic carbon and extract DNA for subsequent metagenomics studies. The physical
141 factors in sampling spots including illumination, temperature of the air or soil, humidity, and
142 pH of soil were recorded.

143

144 **2.2 DNA Extraction and PCR for Metagenomics Analysis**

145 The procedure modified from kit of Genomic DNA from soil (Macherey-Nagel) was used
146 to extract bacterial DNA from limestone samples. The detailed procedure was described in
147 detail in our previous study (Huang et al., 2018). In short, DNA in a bulk of soil fraction was
148 isolated and eluted for the PCR amplification of 16S rRNA gene sequences at V3-V4 regions
149 by using Illumina's MiSeq system to create paired-end sequencing data. The target sequence
150 was amplified through PCR by using mixed forward and reverse primers. After separation
151 through electrophoresis in agarose gel, PCR products with expected sizes were harvested.

152

153 **2.3 Metagenomics Library Construction and Analysis**

154 The Illumina Nextera XT index kit was used in the second-stage PCR for the addition of
155 the index. The raw data of forward and reverse reads were aligned using CLC bio's analysis
156 platform (Genomic Workbench v.8.5) with Q20 as a threshold to generate output fasta files.
157 Fasta files were further processed using the sequence analysis tool USEARCH. All sequence
158 fills were merged together, followed by removing duplicates and clustering sequences into
159 OTUs at 97% pairwise identity with the minimum cluster size being set at 2 to construct an
160 OTU-reference library. A comparison between samples and the reference library at a level of
161 97% sequence identity was made to yield an OTU table, and the number of reads in each
162 OTU was revealed. A 16s UTAX reference database was employed for the assignment of
163 taxonomy for query sequences in the OTU-reference library. We analyzed each habitat by
164 aligning the data, relative abundance, and biodiversity with a heatmap and principal
165 coordinate analysis (PCoA).

166



167 **2.4 Functional Bacteria Analysis**

168 To investigate urease-producing bacteria and weathering-associated bacteria in each
169 habitat, a bioinformatics approach was used to find functional bacteria based on the similarity
170 of DNA sequences. In this method, tables of urease-producing bacteria and weathering
171 bacteria—including bacteria for surface recognition, surface attachment, and mineral
172 dissolutions—were selected from previously published papers in which their corresponding
173 16S DNA sequences were downloaded from the NCBI database, as shown in Supplementary
174 Tables I and II. The DNA sequence tables were used as references to construct phylogenetic
175 trees by employing the Molecular Evolutionary Genetics Analysis 7 (MEGA 7) program with
176 the setting of parsimony, neighbor-joining, and maximum likelihood analyses. The similarity
177 between adjacent pairs of OTU sequences and reference sequences was tested using the
178 NCBI nucleotide BLAST program. DNA sequences with a similarity of more than 95% were
179 defined as urease-producing bacterial lineages or weathering-associated bacterial lineages of
180 corresponding bacteria, and their read numbers were manually selected to calculate their
181 populations.

182

183



184 **3. Results**

185

186 **3.1 General Description of Environmental Factors**

187 The location of the limestone gulch in Tainliao and the path to the mountain gulch are
188 demonstrated in Figure 1. The left panel of the figure shows the locations of Tainliao, and the
189 right panel describes in detail where soil and karst samples were collected. The water samples
190 were collected from the drippings of stalactites in the gulch. Various environmental
191 parameters were assessed, namely, illumination, temperature, humidity in air, humidity in
192 soil, and pH in soil. The illumination in the gulch was relatively low all year around, ranging
193 from approximately 20 to 600 Lux in a location where reflected light is available, and
194 ranging from approximately 5 to 70 Lux on the wall when measured from 9 AM to 6 PM on a
195 shiny summer day. The illumination at the opening of the gulch ranged from approximately
196 100 to 800 Lux at a brighter location, but it ranged from approximately 60 to 650 (Lux) on
197 the limestone wall. On the same day, the illumination at an open space around the gulch was
198 approximately 8,000 Lux, 150,000 Lux, 85,000 Lux, and 4,000 Lux at 9 AM, 12 noon, 3 PM,
199 and 6 PM, respectively. The temperature in the inner of the gulch was 2°C – 4°C lower than
200 the temperature at the opening of the gulch. The humidity in the soil versus air was 100%
201 versus 70% ± 5% at the inner gulch and 37.5% ± 22% versus 60% ± 5% at the opening. The
202 pH of the soil was approximately 4.4 – 5 at the inner gulch and approximately 6.2 – 6.6 at the
203 opening. The total organic carbon content in the inner karst wall, the outer karst wall, and the
204 soil of the outer ground was 3.9% ± 0.2%, 7.7% ± 1%, and 9.1% ± 0.5%, respectively. In
205 short, the inner gulch was a zone of relatively lower light penetration compared with the
206 opening of the gulch. The humidity in the air was similar in the inner and outer gulch. The
207 relative light penetration in the outer gulch may affect the level of local biosynthesis,
208 resulting in a higher total organic carbon content in the areas.

209

210 **3.2 The Microbial Community Structure in Various Karst Habitats Based on Results**
211 **of the NGS Platform**

212 The metagenomic sequence data from different habitats, namely from the outer soil, the
213 outer karst, the inner karst, and from dripping water, contained a similar level of assembled
214 reads that were clustered into OTUs, revealing a high variety in numbers, as shown in Table
215 1. The average read number was more than 400,000. The sample from the soil of the outer
216 gulch had the highest OTUs, whereas the water sample had the lowest OTUs. Because the



217 Shannon index of the sample from the outer gulch was the highest, the effective number of
218 species was also the highest.

219

220 Figure 2 shows the relative abundance of OTUs in different habitats, which contained 22
221 phyla in total. Our results revealed that the soil sample from the outer gulch had the highest
222 alpha diversity, whereas the water sample had the lowest alpha diversity. Four major phyla,
223 namely Proteobacteria, Acidobacteria, Actinobacteria, and Cyanobacteria, accounted for 80%
224 of total microbial species in all the groups. Moreover, Cyanobacteria was present in both
225 limestone walls and was absent in water and soil habitats. Although Actinobacteria can be
226 found in freshwater habitats, our results revealed that they accounted for only <0.4% of the
227 relative abundance in the karst dripping water. The right panel of Figure 3 shows the heatmap
228 of OTUs in various habitats. Our data revealed that microorganisms around the gulch were
229 considerably diverse, and the OTU pattern of the water sample was markedly different from
230 those of other samples. The habitats of the outer gulch, outer soil, and outer karst wall more
231 closely resembled each other than the outer and inner karst wall. These findings suggest that
232 the effect of light penetration and moisture overwhelmed the effects of chemical
233 compositions in the karst walls. The right panel of Figure 3 shows the PCoA distribution of
234 dominated OTUs in the environment, indicating that the distribution of bacteria in the karst
235 gulch was considerably diverse. Many unique OTUs were present in the water habitat (blue
236 square). Although the number of each OTU between the samples of the outer karst wall and
237 soil more closely resembled (Figure 3, right panel), the PCoA showed that the distribution of
238 many dominant OTUs in inner (orange cross) and outer (green cross) karst walls was
239 adjacent to each other, suggesting that the sequences of dominant species in these two
240 habitats were similar to each other.

241

242 **3.3 Distribution of Weathering Bacteria in the Karst Gulch**

243 Table 2 shows the OTUs of habitats with sequences that had >95% similarity to reference
244 sequences of weathering bacteria. Supplementary Table I shows bacteria collected in previous
245 studies, indicating that they were capable of promoting the functions of weathering in rocks.
246 Most of the weathering-associated bacteria in habitats belonged to the phyla Proteobacteria.
247 Although 220 species of weathering-associated bacteria were used as references, only <10%
248 of them showed similarity to OTU sequences in the karst habitats in this study. The relative
249 abundance of weathering-associated bacteria in each habitat is shown in Figure 4. The sample
250 from the inner karst gulch contains the last portions of bacteria relative to weathering-



251 associated bacteria. The dominant genus in the rock and soil of weathering-associated bacteria
252 in the karst gulch was *Sphingomonas* whereas *Noviherbaspirillum* was the unique genus in the
253 water sample. The existence of weathering-associated bacteria in water indicated that water
254 plays a role in mediating the propagation of weathering bacteria (Ahmed and Holmström,
255 2015). Because studies have revealed that microorganisms suspended in liquid can still lead to
256 the dissolution of elements from rocks, *Noviherbaspirillum* can facilitate the weathering
257 process of the inner karst wall. However, the relative impact of these bacteria on the weathering
258 process remains unclear.

259

260 **3.4 Distribution of Urease Producing Bacteria in the Karst Gulch**

261 Table 3 shows the OTUs of habitats with sequences that had >95% similarity to reference
262 sequences of urease-producing bacteria, which by definition are microbes that can synthesize
263 enzymes for urea hydrolysis, resulting in subsequent biocalcification in the presence of
264 calcium ions. The reference table of urease-producing bacteria is shown in supplementary
265 Table II. The relative abundance of urease-producing bacteria in each habitat is shown in
266 Figure 5. Urease-producing bacterial lineages in the inner karst wall were related to *Bacillus*
267 *megaterium*, *B. subtilis*, and *B. mycoides*. In water samples, urease-producing bacterial
268 lineages were closely related to *B. megaterium* and *Halomonas denitrificans*. Urease-
269 producing bacteria in both the inner karst wall and dripping water can contribute to the
270 progress of biocalcification on the inner karst wall. Although the total organic carbon content
271 on the outer karst wall was two times higher than on the inner karst wall, the numbers of
272 urease-producing bacterial lineages might only exist in marginal amounts on the outer karst
273 wall due to the extremely low portion of relative abundance of the bacteria (0.003%, Figure
274 5). The relative abundance of urease-producing bacterial lineages on the inner wall (both in
275 water and the karst wall) was approximately 200 times higher than that on the outer karst
276 wall. The high portion in relative abundance of urease-producing bacteria on the inner wall
277 indicated that a persistent stalactite formation occurs on the habitat, which is consistent with
278 its ecological landscape.

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283 **4. Discussions**

284

285 Understanding the microbial diversity in the karst landscape provides insights into how
286 bacteria survive in extreme environments and the consequence of geological evolution after
287 their interaction. Many studies focusing on the abundance of microorganisms in karst caves
288 have showed a large microbial diversity in limestone landscapes (Engel, 2010; Ortiz et al.,
289 2014; Ortiz et al., 2013; Tomczyk-Żak and Zielenkiewicz, 2015; Zepeda Mendoza et al., 2016).
290 Most of these studies have confirmed that Actinobacteria and Proteobacteria were the dominant
291 species in karst samples. Our study results revealed that total OTUs distributed in the phyla of
292 Actinobacteria, Proteobacteria, Cyanobacteria, and Acidobacteria in karst habitats were
293 approximately 3500, suggesting the extreme diversity of microorganisms in karst landscapes
294 in our studied site. The bacterial communities from different geological areas exhibited
295 regional difference. For example, the majority of bacterial phyla in karst soil in Guizhou China
296 were Proteobacteria, Actinobacteria, Acidobacteria, and Planctomycetes (Zhou et al., 2009).
297 Our data of karst soil revealed that this habitat exhibited the highest microbial diversity. We
298 posit that weathering bacteria present in the outer karst wall and soil contribute to the nutrient
299 level of the soil, causing a higher total organic carbon content and Shannon index of the soil
300 habitat. Our study indicated that light penetration, together with other physical parameters,
301 specify the development of particular microbial communities as showed in Figure 3. In the
302 long run, the subtle changes of the composition of microbial communities alter the geochemical
303 reactions, rendering the variation of karst landscapes.

304

305 With the application of the NGS platform for acquiring metagenomic data in various karst
306 habitats, we could examine the effects of physical parameters on the evolution of microbial
307 communities and the consequential changes in the microenvironment. To make the most of the
308 metagenomic data, we used the sequence similarity tool, BLAST, to determine the likeness of
309 representative DNA sequences of OTUs compared with the functional bacteria available in the
310 NCBI database. Although the relative abundance in the phylum levels of karst habitats was
311 similar, the compositions of functional bacteria tested in each habitat were substantially
312 different. We set the cutting point of similarity at 95% to compare functional bacteria in various
313 habitats, which is approximately the level of the genus. However, it is still under debate
314 whether the 95% cutoff in the DNA sequence similarity is a proper setpoint to cluster a category
315 of functional bacteria. Based on this calculus, our data revealed a large difference in the final
316 results, as shown in Figures 4 and 5, suggesting that a considerable difference exists in the



317 relative abundance of functional bacteria in different habitats. Further confirmation of specific
318 functional bacteria in various habitats can be achieved through molecular cytogenetic
319 techniques, such as fluorescence in situ hybridization.

320

321 We hypothesized that the two primary activities of karst landscapes, namely weathering
322 and stalactite formation, might affect the dynamic changes and geographic evolution of karst
323 walls. Functional bacteria associated with these activities were analyzed based on the NGS
324 platform. Our data revealed that a drastic shift in key microorganisms, weathering bacteria and
325 urease-producing bacteria, occurred in the habitats of various physical parameters, suggesting
326 that these parameters play a role in the initiation of different paths in geological evolution. The
327 differences in functional bacterial compositions in various habitats supported the fact that the
328 speleothem formation occurred primarily in the inner karst wall in the gulch, suggesting
329 physical conditions in the inner karst wall favor the growth of urease-producing bacteria and
330 promote calcite precipitation. Studies on Cyanobacteria and calcium precipitation have shown
331 that microorganisms may highly enhance the precipitation of CaCO₃ minerals in hot spring
332 water (J. Kaźmierczak et al., 1996; Kawano and Obokata, 2007). In this study, the relative
333 abundance of Cyanobacteria in the inner karst wall was twice as large as the relative abundance
334 of Cyanobacteria in the outer karst wall, suggesting that the environment of the inner karst is
335 favorable for the development of Cyanobacteria and the consequential mineral precipitation.

336

337 Biocalcification has been widely applied in the ecosystem for many purposes, including
338 land consolidation, groundwater control, crack remediation, and immobilization of toxic metals
339 (Anbu et al., 2016; Kumari et al., 2016; S. Animesh and Ramkrishnan, 2016; S.A. Abo-El-
340 Enein et al., 2012; Uroz et al., 2007). Various bacteria, shown in supplementary Table II,
341 effectively produce urease, resulting in the precipitation of calcite. Although many
342 environmental factors that affect the growth conditions of urease-producing bacteria have been
343 tested, none of the previous studies have investigated the effects of sunlight penetration on the
344 natural selection of bacterial development. In the study of calcifying bacteria in the Stiffe cave,
345 *Bacillus* and *Arthrobacter* were isolated from natural habitats, which might have contributed
346 to speleothem formation. In this study, several distinct features were found from the data of the
347 NGS platform and the analysis of total organic carbon. First, we found that *B. megaterium* and
348 *H. denitrificans* were the predominant species among calcifying bacteria. Second, urease-
349 producing bacteria were dominant in the inner karst wall. Finally, urease-producing bacterial
350 lineages were also present in the dripping water of the inner wall, which possessed different



351 species of urease-producing bacteria. Most importantly, the interface between water dripping
352 and the inner karst wall was subjected to the biocalcification effects of both urease-producing
353 bacteria. Our data suggests that bacteria in the water drips of the inner karst wall play an
354 important role in facilitating speleothem formation.

355

356 Remarkably, habitats with a lower relative abundance of urease-producing bacteria
357 showed a higher value in relative abundance of weathering bacteria. Meanwhile, the TOC was
358 higher in samples at the gulch opening compared with the sample in the inner wall. We
359 concluded that sunlight and nutrient levels may be two factors affecting TOC in these habitats.
360 Sunlight is an important source providing energy for the accumulation of biomass. Light
361 penetration provided a discriminatory growth condition to heterotrophic microorganisms on
362 habitats in inner and outer walls. In the gulch, more than 90% of the luminance from sunlight
363 was filtered out by the vegetation coverage at the opening of the gulch, and the karst structure
364 of the steep wall further filtered off 0% to 85% of light penetration inside the gulch, depending
365 on the angle of the sun and the horizon, which affects the photosynthesis reaction in these areas.
366 We also noticed that the effective number of species in the soil increased drastically, suggesting
367 that an elevation of mineral nutrients, one important consequence of weathering effect on rocks,
368 caused by the weathering process could provide a favorable growth condition for many other
369 bacteria in the soil. Previous studies have documented how the dissolution of calcite can be
370 enhanced in the presence of heterotrophic microorganisms (Jacobson and Wu, 2009). Our data
371 revealed that the composition of Acidobacteria increased in the habitat of soil, which is
372 consistent with that of a previous study (Zhou et al., 2009). We propose that light penetration
373 plays a pivotal role in natural selection to promote the growth of weathering-associated bacteria,
374 which in turn increase the nutrient level in situ and favor the development of microorganisms.

375

376 **Conclusion**

377 Given an example of the karst landscape, we provided evidence regarding how physical
378 parameters change the microbial community and the consequential landscape evolution.
379 Furthermore, we showed that light penetration regulates the microbial population, leading to
380 the breakdown of calcite, whereas the chemical composition of limestone might deliver certain
381 conditions that limit the growth of bacterial species. These factors, namely light penetration,
382 water dripping, moisture, the chemical composition of karst, and selected bacteria that are
383 intertwined, shape the weathering process and stalactite formation. The natural selections of
384 bacteria were achieved by the preferential growth of two bacterial groups: urease-producing



385 bacteria in the inner karst wall and weathering bacteria in the outer karst wall. Our data reveals
386 a causal relationship between environmental factors that contribute to the remodeling of the
387 topography and are mediated by microorganisms. To the best of our knowledge, this study is
388 the first to address the distinct role of bacteria in the water dripping of karst in biocalcification
389 and the effect of light penetration in the microenvironment on the colony selection of microbial
390 communities.
391



392 Figures and tables



393

394 Figure 1. The Location of the Limestone Gulch

395 Figure (A) shows a series maps of increased scale pointing to the limestone gulch. Figure (B)

396 shows the view of the gulch. Both sides of the path opening have stairs leading to the center

397 of the gulch. We took samples from outer limestone wall of the gulch (OL, Figure b1), the

398 soil on the outer ground (OS, Figure b2), the inner limestone wall of the gulch (IL, Figure

399 c3), and water dripping from the wall (WA). The inner and outer limestone walls exhibit

400 distinct landscapes in the stalactite formation.

401



402

	OS	OL	IL	WA
TOC	9.0±0.4 (%)	7.7±0.9 (%)	3.9±0.1 (%)	-
Reads	296,726	433,210	477,281	448,446
OTUs	2470	831	1899	467
Phyla	20	15	19	18
Shannon - Wiener index	6.2	4.1	3.9	3.6
Effective number of species	511	61	51	36

403

404

405

406

407 Table 1. The results of Total Organic Carbon, the Basic Information from the NGS Platform,

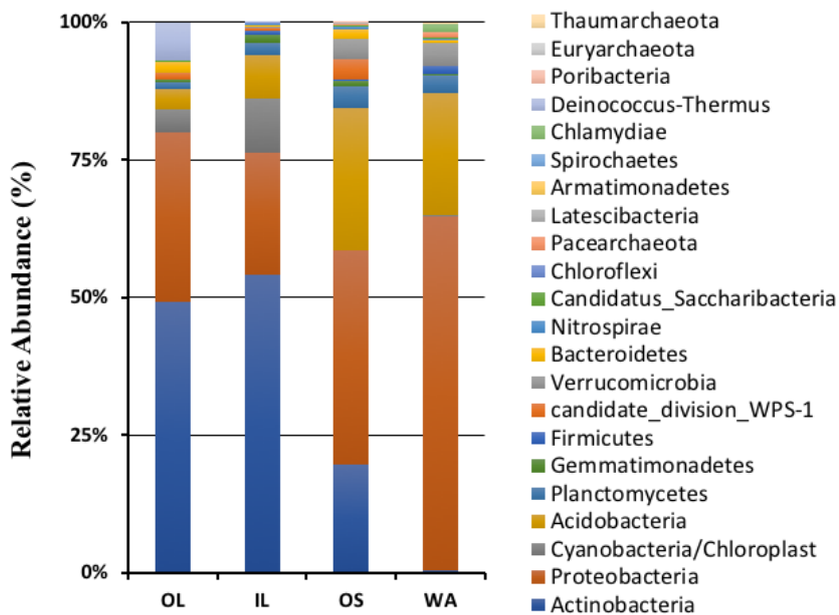
408 and the Bacterial Biodiversity in 4 Different Habitats

409 The symbols of OS, OL, IL, and WA represent sample sites of outer ground, outer limestone

410 wall, inner limestone wall and water, respectively.



411

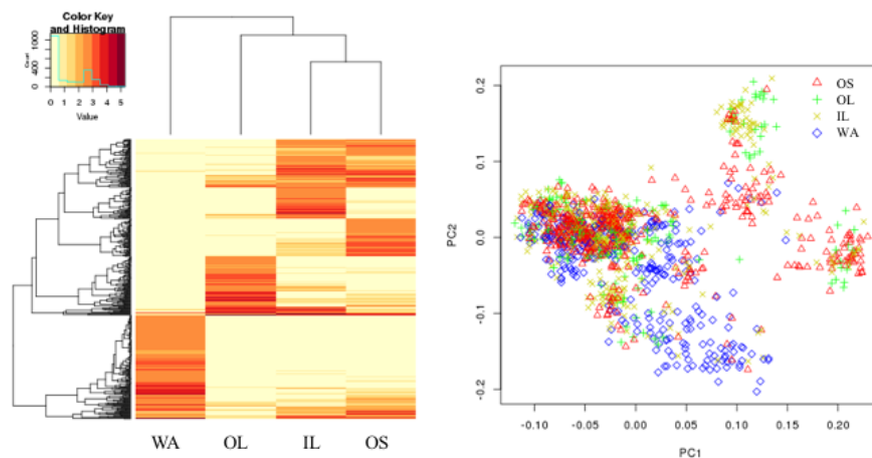


412

413 Figure 2. The Relative Abundance of 4 Various Habitats from the Karst Landscape in Tainliao

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418 Figure 3. (A) The Heatmap of OTUs Based on the Read Number in Different Habitats in

419 Tainliao. (B) The PCoA Distribution Based on the Distance Calibration from DNA

420 Sequences of OTUs in 4 Habitats in Tainliao.



421
 422 Table 2. The Taxonomy of OTUs and the Bacterial References with the Sequencing
 423 Similarity Higher than 95% to Weathering-Associated Bacteria

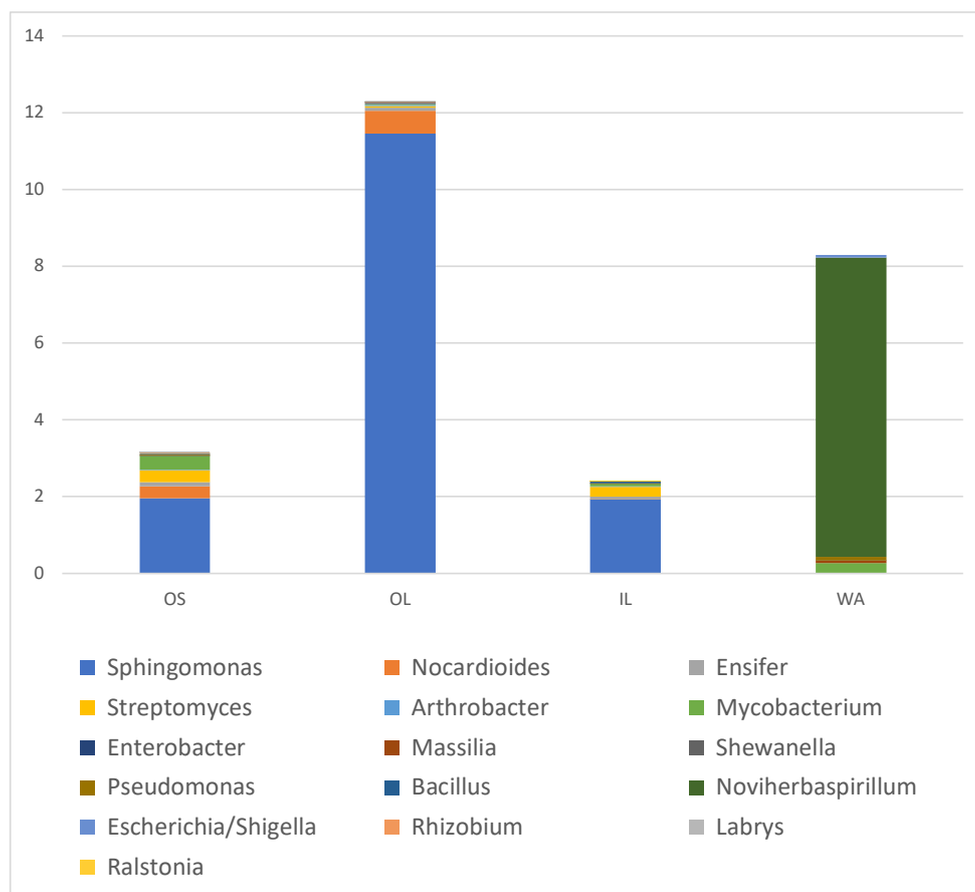
Classification	OTU	Reference of bacteria	Sequence ID	Identities	Taxonomy
Alphaproteobacteria	karst949	Labrys sp.	LC372609.1	398/407(98%)	Labrys
	karst918	Sphingomonas anadarae	AB261013.1	394/405(97%)	Sphingomonas
	karst12	Sphingomonas sp.	AF385529.1	400/406(99%)	Sphingomonas
	karst1757	Sphingomonas sp.	AF385529.1	393/406(97%)	Sphingomonas
	karst2759	Sphingomonas sanguinis	D13726.1	385/403(96%)	Asticcacaulis
	karst1653	Aminobacter sp.	AB905480.1	391/408(96%)	Ensifer
	karst341	Aminobacter sp.	FM886907.1	401/407(99%)	Ensifer
	karst1961	Rhizobium leguminosarum	D14513.1	401/407(99%)	Rhizobium
Betaproteobacteria	karst578	Janthinobacterium sp.	AM071372.1	424/433(98%)	Massilia
	karst3282	Janthinobacterium sp.	AB252072.1	417/429(97%)	Massilia
	karst5	Collimonas sp.	FR729923.1	419/432(97%)	Noviherbaspirillum
	karst1267	Collimonas sp.	FR729923.1	413/437(95%)	Ralstonia
Gammaproteobacteria	karst1216	Enterobacter	AB616140.1	431/433(99%)	Enterobacter
	karst397	Citrobacter rodentium	AB682287.1	415/432(96%)	Escherichia/Shigella
	karst3185	Shewanella morhuae	AB205576.1	421/433(97%)	Shewanella
	karst595	Pseudomonas stutzeri	AJ006107.2	431/435(99%)	Pseudomonas
	karst2678	Pseudomonas fluorescens	FJ972536.1	422/433(97%)	Pseudomonas
	karst464	Pseudomonas sp.	AJ417069.1	425/434(98%)	Pseudomonas
Gram-positive	karst1814	Pimelobacter simplex	AY509240.1	411/423(97%)	Nocardioides
	karst2873	Arthrobacter oxydans	LN774480.1	395/413(96%)	Arthrobacter
	karst175	Streptomyces lividans	AB184695.1	403/411(98%)	Streptomyces
	karst372	Mycobacterium colombiense	AM062764.1	401/423(95%)	Mycobacterium
	karst3247	Mycobacterium colombiense	AM062764.1	407/427(95%)	Mycobacterium
	karst1108	Mycobacterium colombiense	AM062764.1	401/424(95%)	Mycobacterium
	karst1032	Mycobacterium colombiense	AM062764.1	406/425(96%)	Mycobacterium
	karst2144	Mycobacterium ratisbonense	AJ271863.1	393/413(95%)	Mycobacterium
	karst101	Mycobacterium sp.	X84978.1	408/411(99%)	Mycobacterium
	karst1188	Bacillus subtilis	AB018487.1	417/434(96%)	Bacillus
	karst1300	Bacillus mycoides	AB547222.1	432/432(100%)	Bacillus
	karst47	Pimelobacter simplex	AY509240.1	399/411(97%)	Nocardioides
	karst886	Streptomyces lividans	AB184826.1	399/409(98%)	Streptomyces
	karst2729	Streptomyces lividans	AB184695.1	398/413(96%)	Streptomyces
	karst2914	Kocuria polaris	AJ278868.1	403/426(95%)	Arthrobacter



424 Table 3: The Taxonomy of OTUs and the Bacterial References with the Sequencing
425 Similarity Higher than 95% to Urease-Producing Bacteria

Classification	OTU	Reference of bacteria	Sequence ID	Identities	Taxonomy
Actinobacteria	karst1300	Bacillus mycoides	AB547222.1	432/432(100%)	Bacillus
Firmicutes	karst791	Bacillus megaterium	JX893034.1	419/433(97%)	Bacillus
	karst260	Bacillus megaterium	JX893034.1	416/431(97%)	Bacillus
	karst2293	Bacillus megaterium	JX893034.1	411/430(96%)	Bacillus
	karst1188	Bacillus subtilis	AB018487.1	417/434(96%)	Bacillus
Gammaproteobacteria	karst189	Halomonas denitrificans	AM229317.1	418/432(97%)	Halomonas
	karst2279	Halomonas denitrificans	AM229317.1	414/432(96%)	Halomonas

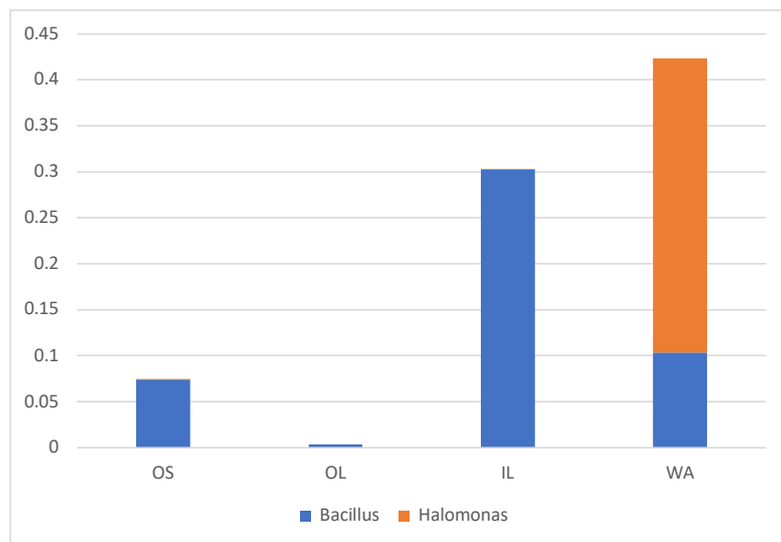
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428 Figure 4: The Relative Abundance of Weathering-Associated Bacterial Lineages in Each

429 Habitat



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431 Figure 5: The Relative Abundance of Urease-Producing Bacterial Lineages in Each Habitat



432 References

433

434 Abo-El-Enain, S. A., Ali, A. H., Talkhan, F. N., and Abdel-Gawwad, H. A.: Utilization of
435 microbial induced calcite precipitation for sand consolidation and mortar crack remediation,
436 HBRC Journal, 8, 185 – 192, 10.1016/j.hbrj.2013.02.001, 2012.

437 Ahmed, E., and Holmström, S. J. M.: Microbe – mineral interactions: The impact of surface
438 attachment on mineral weathering and element selectivity by microorganisms, Chemical Geology,
439 403, 13-23, 10.1016/j.chemgeo.2015.03.009, 2015.

440 Anbu, P., Kang, C.-H., Shin, Y.-J., and So, J.-S.: Formations of calcium carbonate minerals by
441 bacteria and its multiple applications, SpringerPlus, 5, 10.1186/s40064-016-1869-2, 2016.

442 Animesh, S., and Ramkrishnan, R.: Study on effect of microbial induced calcite precipitates on
443 strength of fine grained soils, Perspectives in Science 8, 198—202, 10.1016/j.pisc.2016.03.017,
444 2016.

445 Barton, H. A., and Northup, D. E.: Geomicrobiology in cave environments: past, current and
446 future perspectives, Journal of Cave and Karst Studies, 69, 163-178, 2007.

447 Castanier, S., Métyer-Levrel, G. L., and Perthuisot, J. P.: Ca-carbonates precipitation and
448 limestone genesis — the microbiogeologist point of view, Sedimentary Geology, 126, 9 – 23,
449 1999.

450 Engel, A. S.: Microbial Diversity of Cave Ecosystems, in: Geomicrobiology: Molecular and
451 Environmental Perspective, edited by: Barton, L. L., Mandl, M., and Loy, A., Springer, 219-238,
452 2010.

453 Ercole, C., Cacchio, P., Cappuccio, G., and Lepidi, A.: Deposition of calcium carbonate in karst
454 caves: role of bacteria in Stiffe's cave, International Journal of Speleology, 30A 69-79, 2001.

455 Gat, D., Tsesarsky, M., Shamir, D., and Ronen, Z.: Accelerated microbial-induced CaCO₃
456 precipitation in a defined coculture of ureolytic and non-ureolytic bacteria, Biogeosciences, 11,
457 2561-2569, 10.5194/bg-11-2561-2014, 2014.

458 Hutchens, E.: Microbial selectivity on mineral surfaces: possible implications for weathering
459 processes, Fungal Biology Reviews, 23, 115-121, 10.1016/j.fbr.2009.10.002, 2009.

460 Jacobson, A. D., and Wu, L.: Microbial dissolution of calcite at T=28°C and ambient pCO₂,
461 Geochimica et Cosmochimica Acta, 73, 2314-2331, 10.1016/j.gca.2009.01.020, 2009.

462 Jones, B.: Review of calcium carbonate polymorph precipitation in spring systems, Sedimentary
463 Geology, 353, 64-75, 10.1016/j.sedgeo.2017.03.006, 2017.

464 Lian, B., Chen, Y., Zhu, L., and Yang, R.: Effect of Microbial Weathering on Carbonate Rocks,
465 Earth Science Frontiers, 15, 90-99, 2008.

466 Lower, S. K., Hochella, M. F. J., and Beveridge, T. J.: Bacterial recognition of mineral surfaces:
467 nanoscale interactions between *Shewanella* and α -FeOOH, Science, 292, 1360-1363, 2001.

468 Mortensen, B. M., Haber, M. J., DeJong, J. T., Caslake, L. F., and Nelson, D. C.: Effects of
469 environmental factors on microbial induced calcium carbonate precipitation, J Appl Microbiol,
470 111, 338-349, 10.1111/j.1365-2672.2011.05065.x, 2011.

471 Ortiz, M., Legatzki, A., Neilson, J. W., Fryslie, B., Nelson, W. M., Wing, R. A., Soderlund, C. A.,
472 Pryor, B. M., and Maier, R. M.: Making a living while starving in the dark: metagenomic insights
473 into the energy dynamics of a carbonate cave, ISME J, 8, 478-491, 10.1038/ismej.2013.159, 2014.



- 474 Qabany, A., Soga, K., and Santamarina, C.: Factors affecting efficiency of microbially induced
475 calcite precipitation, *Journal of Geotechnical and Geoenvironmental Engineering*, 138, 992-1001,
476 10.1061/(ASCE)GT.1943-5606.0000666, 2012.
- 477 Seiffert, F., Bandow, N., Bouchez, J., von Blanckenburg, F., and Gorbushina, A. A.: Microbial
478 Colonization of Bare Rocks: Laboratory Biofilm Enhances Mineral Weathering, *Procedia Earth
479 and Planetary Science*, 10, 123-129, 10.1016/j.proeps.2014.08.042, 2014.
- 480 Sulu-Gambari, F.: Bacterially-induced dissolution of calcite: The role of bacteria in limestone
481 weathering, Master of Science, Department of Earth and Planetary Science, McGill University, 96
482 pp., 2011.
- 483 Tomczyk-Żak, K., and Zielenkiewicz, U.: Microbial Diversity in Caves, *Geomicrobiology Journal*,
484 33, 20-38, 10.1080/01490451.2014.1003341, 2015.
- 485 Uroz, S., Calvaruso, C., Turpault, M. P., Pierrat, J. C., Mustin, C., and Frey-Klett, P.: Effect of the
486 mycorrhizosphere on the genotypic and metabolic diversity of the bacterial communities involved
487 in mineral weathering in a forest soil, *Appl Environ Microbiol*, 73, 3019-3027,
488 10.1128/AEM.00121-07, 2007.
- 489 Uroz, S., Calvaruso, C., Turpault, M. P., and Frey-Klett, P.: Mineral weathering by bacteria:
490 ecology, actors and mechanisms, *Trends Microbiol*, 17, 378-387, 10.1016/j.tim.2009.05.004, 2009.
- 491 Wei, S., Cui, H., Jiang, Z., Liu, H., He, H., and Fang, N.: Biomineralization processes of calcite
492 induced by bacteria isolated from marine sediments, *Braz J Microbiol*, 46, 455-464,
493 10.1590/S1517-838246220140533, 2015.
- 494 Wu, Y.-w., Zhang, J.-c., Wang, L.-j., and Wang, Y.-x.: A rock-weathering bacterium isolated from
495 rock surface and its role in ecological restoration on exposed carbonate rocks, *Ecological
496 Engineering*, 101, 162-169, 10.1016/j.ecoleng.2017.01.023, 2017.
- 497 Zhou, J. P., Huang, Y., and Mo, M. H.: Phylogenetic analysis on the soil bacteria distributed in
498 karst forest *Brazilian Journal of Microbiology*, 40, 827-837, 2009.
- 499