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Light attenuation as a control for microbiogeomorphic features: Implications for coastal cave speleogenesis

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1 **Light attenuation as a control for microbiogeomorphic features: implications for coastal cave**
2 **speleogenesis**

3
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13
14 **Abstract**

15 San Salvador (Bahamas) is a carbonate island with dozens of flank margin caves formed in the
16 phreatic zone by fresh seawater mixing within the freshwater lens. These caves have no direct
17 connection with the sea, and form at or close to the tidally influenced fluctuating water table. After
18 sea-level fall, in their subaerial parts caves are enlarged mainly by rock dissolution and by erosion
19 close to the water level, condensation-corrosion and breakdown processes. For understanding the
20 geomorphological features observed in these caves and how they are related to light attenuation, we
21 investigated three sampling sites in the tidally influenced zone of Lighthouse Cave, which has been
22 re-invaded by seawater during the Holocene sea-level highstand. A freshwater lens no longer exists
23 within or adjacent to the cave. Rock samples were collected above and below the internal lake
24 shores close to the entrance, and in the twilight and dark zones of this cave. Light and electron
25 microscopy examinations were conducted for detecting microbial cells, as well as bioconstruction
26 and bioweathering features. In addition, a high precision laser scanner was used for characterising

27 sample microtopography. Our data showed that the microtopography and geomorphology of the
28 lake shore samples (cave entrance) are dominated by bioweathering, whereas the samples of the
29 twilight and dark zones are controlled by a combination of both bioweathering and bioconstructive
30 processes depending on light availability. Bioconstructive structures, such as semi-planar
31 lamination, at the fluctuating water level of the Lighthouse Cave show that dissolution due to water
32 mixing of sea and freshwater in the Holocene is no longer the most important speleogenetic process.
33 We propose that the geomorphological evolution is strongly influenced by the degree of rock
34 diagenesis more than the initial mechanism of speleogenesis.

35

36 **Keywords:** Flank margin caves; mixing dissolution; tides; bioconstruction-bioweathering processes

37

38 **1. Introduction**

39 San Salvador Island is located in the eastern part of the Bahamas within the Bahamian Archipelago
40 in the Atlantic Ocean (Fig.1A). This island is about 11 km wide and 19 km long and lies in a
41 tectonically stable area, which has been influenced by eustatic sea-level change during the
42 Quaternary. The island is characterised by a sequence of Pleistocene shallow-water carbonate
43 deposits covering the oceanic crust basement (Meyerhoff and Hatten, 1974; Supko, 1977; Carew
44 and Mylroie, 1985). The dissolution of Bahamian carbonates produced karstic features such as
45 karren, shallow depressions, blue holes, and the well-known *flank margin caves* (Roth et al., 2006;
46 Labourdette et al., 2007). *Flank margin caves* (FMC) (Mylroie and Carew, 1990; Harris et al., 1995;
47 Gulley et al., 2016) generally present subhorizontal branches that develop at the edge of the
48 freshwater lens where the area of vadose/phreatic water mixing and fresh-seawater mixing zones
49 are superimposed. The water mixing produces a renewed aggressive solution that further dissolves
50 carbonate, thus forming caves (James and Choquette, 1984; Mylroie and Carew, 1995). FMC
51 developed in eogenetic limestone (diagenetically immature carbonate rocks with high primary
52 porosity), such as the case of Lighthouse Cave, have been described from many carbonate islands

53 (e.g., Mylroie et al., 2001; Vacher and Mylroie, 2002; Mylroie and Mylroie, 2007; Kourampas et
54 al., 2015) and also on carbonate coasts on large islands or continental margins (D'Angeli et al.,
55 2015b; White and Webb, 2015; Bontognali et al., 2016; De Waele et al., 2017, 2018). Nevertheless,
56 FMC can also develop in more highly lithified carbonate rocks (telogenetic limestones) (Mylroie et
57 al., 2008; Otoničar et al., 2010; Ruggieri and De Waele, 2014; D'Angeli et al., 2015a), during past
58 high sea levels.

59 FMC in eogenetic rocks are mainly characterised by spongy morphologies, maze areas, dead-end
60 passages, cusped walls and irregular chambers that narrow inland. Presence of phreatic dissolution
61 pockets are common, and the absence of high-velocity structures, turbulent-flow wall sculptures
62 (i.e., scallops) and stream sediments (Waterstrat et al., 2010) indicates a diffuse or laminar flow
63 speleogenetic environment. Wave processes are not required in FMC speleogenesis; exclusively sea
64 level change (mainly due to coastal uplift or eustatic fluctuation) can influence their position
65 (Mylroie and Carew, 1988). These caves form without entrances; access to them and/or the invasion
66 of daylight into them, only occurs post-speleogenesis, when denudational processes breach their
67 tops and/or sides.

68 It is well known that caves are low energy environments that can preserve fragile speleothems,
69 sediments, and archaeological remains over long time spans (e.g., Van Hengstum et al., 2011;
70 Winkler et al., 2016). In addition, they can host unique microbial communities adapted to
71 subsurface environmental conditions, such as absence of light and low organic matter input
72 (Tomczyk-Żak and Zielenkiewicz, 2016). In general, the cave entrance (strongly influenced by
73 surface conditions) and the twilight zone (with limited light penetration but still important
74 temperature and relative humidity variations) are dominated by phototrophs, whereas the dark zone
75 (characterised by absence of light and stable environmental conditions the year round) is dominated
76 by chemotrophs (Northup and Lavoie 2001; Mejía-Ortíz et al., 2018; Popović et al., 2019).
77 Microorganisms interact with minerals and promote bioweathering (Naylor and Viles, 2002) and
78 biomineralisation processes (Barton and Northup, 2007; Riquelme et al., 2015). They can also have

79 an important, albeit poorly understood, role in ecosystem engineering (Phillips, 2016). The key
80 biogenic and biochemical processes that create distinctive morphological features in caves are: i)
81 microbially-mediated mineral dissolution, and ii) microbially-mediated mineral precipitation
82 (Riquelme et al., 2015). Sulphur, iron and/or manganese oxidising bacteria in contact with
83 carbonate rocks, increasing local acidity through redox reactions and secretion of organic acids or
84 exoenzymes (Sand, 1997), produce intense mineral dissolution (Northup and Lavoie, 2001; Miller
85 et al., 2014) and secondary mineral deposition such as manganese oxides and moonmilk deposits
86 (Hill and Forti, 1997; Gradziński et al., 1997; Miller et al., 2012, 2018). Microbially-mediated
87 precipitation of minerals, especially carbonates, has been frequently observed (Tisato et al., 2015;
88 Bontognali et al., 2016), but more detailed research is needed to better understand the main
89 processes involved. For instance, microalgae and cyanobacteria, growing close to cave entrances or
90 in the twilight zone, can precipitate CaCO_3 by fixing carbon dioxide or can trap and bind particles
91 transported by flowing water or wind (Contos et al., 2001). Most interestingly, “crayback” or
92 “lobster” biomediated mineral growth known as “cyanobacterial subaerial stromatolites” (in the
93 twilight zone) have been found in caves in New South Wales, Australia (Cox et al., 1989a, 1989b),
94 and in Borneo (Lundberg and McFarlane, 2011, 2012). Crusts and coatings of iron-oxyhydroxides
95 (Peck, 1986; Provencio and Polyak, 2001; Frierdich and Catalano, 2012) and manganese-oxide
96 crusts have been found on cave walls (Onac et al., 1997; Northup et al., 2000; Lozano and Rossi,
97 2012). These authors demonstrated that rock weathering is influenced by microbial processes
98 (Dotson et al., 1999; Northup et al., 2003). In addition, Onac et al. (2001) and Audra et al. (2019)
99 reported bat guano to be an important source for mineral growth.

100 To date there have been few investigations on the microbially-mediated processes that operate in
101 each cave zone, and how they may affect the type and spatial patterns of morphological features in
102 cave systems (an exception is Coombes et al., 2015). Here we have sought to address some of these
103 knowledge gaps by investigating the microbiogeomorphic processes developing at the entrance,
104 twilight and dark zones of a flank margin cave system (Lighthouse Cave) located in the Bahamas.

105 In addition, we attempted to identify consistent biogeomorphic features and/or processes that can be
106 associated with ecosystem engineering and likely to construction niches sensu Phillips (2016).
107 Hence, the aim of this study was to understand how biological processes can influence eogenetic
108 carbonate rocks in the development of peculiar micromorphological features (e.g., mineral
109 precipitation, pitting/etching, boreholes) in zones of the cave system with different natural light
110 conditions, and assess how present (post-mixing) flank margin cave evolution may be influenced by
111 secondary bioweathering or bioconstruction processes.

112

113 **2. Materials and Methods**

114 **2.1. Site description and sampling**

115 San Salvador Island has a tropical climate, with daily average temperatures of 25-28°C (Gulley et
116 al., 2015), generating a very high potential evaporation rate of > 1300 mm/yr (Crump and Gamble,
117 2004). The Lighthouse Cave is a limestone cave system located on the NE coast of San Salvador
118 Island (The Bahamas) (Fig.1B), and has no direct connection to the sea.

119 The cave is 402 m-long, and half of the passages show water bodies still influenced by tidal
120 fluctuation (1 m range). The water bodies inside the cave have an overall salinity of 33 PSU,
121 conductivity of 52 mS/cm, 26.6°C, and 7.21 pH (McGee et al., 2010). From the geomorphological
122 point of view, the cave is composed of one large central chamber, adjacent smaller halls, and
123 ramifying dead-end branches developed between 2 to 11 m a.s.l., following the flank of a dune, and
124 mainly formed during 5e high stand. The main entrance and two minor entrances are vadose pits
125 created after the flank margin cave speleogenesis. The main morphologies are characterised by a 3D
126 maze and a tubular branch ending abruptly, domes, arches; big halls alternate with small and narrow
127 passages, and bell holes are clearly visible on the ceilings. Bat guano deposits are abundant in the
128 twilight zone, close to the cave entrance, and represent the most important source of organic matter
129 and phosphate minerals in Lighthouse Cave (Onac et al., 2001).

130 Cave walls are generally white, but dark brown crusts are also visible along the tidal zone (Mylroie,
131 2014). Three replicates from the three different sampling sites (Fig. 2) were collected in Lighthouse
132 Cave in February 2014 during low tide (local tidal vertical range of ~0.80 m). Three cm-long rock
133 samples were taken using a geologist hammer and stored in sterile plastic bags. All the rock
134 samples were collected from calcarenitic eolianite limestone belonging to Owl's Hole Formation
135 (Middle Pleistocene age) (Panuska et al., 1999; Kindler et al., 2010). The sampling sites were: i) a
136 limestone rock exposed in the intertidal zone of a hypersaline lake located in a mangrove forest, 20
137 m W of the main cave entrance (sampling site: SS1); ii) a limestone wall in the twilight zone of the
138 cave, 20 m from the entrance, close to the water table, approximately 10-15 cm below mean high
139 tide level (sampling site: SS2), and iii) a limestone wall near the water table, approximately 25 cm
140 below high tide level and located ~60 m from the cave entrance (sampling site: SS3).

141

142 **2.2. Microtopographic characterisation**

143 To identify the processes involved in the present-day cave evolution, the millimetre-scale
144 microtopographic irregularities of each rock sample were examined using a high-precision laser
145 scanner. This instrument minimises measurement errors and resolution problems associated with
146 conventional roughness meters (Bourke et al., 2008), and creates digital terrain models (DTMs). We
147 used a micro Epsilon high-precision laser scanner at the University of Glasgow, with a maximum
148 distance between the laser and sample of ~35 mm. Contour map analyses were performed using the
149 ArcGIS system. Six profiles (NNE-SSW) were drawn across each rock sample (using a systematic
150 random sampling design) to measure roughness values (Giaccio et al., 2002; Gomez-Pujol et al.,
151 2006; Naylor et al., 2012; Moses et al., 2014). Roughness values were obtained using the following
152 ratio, after Whitehouse (2012):

$$\frac{\text{Surface length (mm)}}{\text{Profile length (mm)}}$$

153

154 **2.3. Stereomicroscopy and ESEM analysis**

155 Stereomicroscopy and Environmental Scanning Electron Microscopy (ESEM) investigations were
156 performed to observe newly formed crystals and recognise microbial communities that contributed
157 to microtopographic changes on the rock surfaces (Moses et al., 2014; Coombes et al., 2015). These
158 observations were conducted at the University of Glasgow, based on the methods of Naylor and
159 Viles (2002) and Coombes et al. (2011, 2015). Three rock chips, $\sim 4 \text{ cm}^2$, were studied at each
160 sampling site (Table 1) to generate semi-quantitative data on biogeomorphological processes
161 operating in each zone of the cave system. The top surface of each chip was observed using
162 Olympus SZ x7 and Olympus Bx41 microscopes equipped with an Olympus DP25 camera for the
163 recognition of biological features, such as photosynthetic-based biofilms, shiny filaments, fossils,
164 white creamy deposits, and black coatings (after Naylor and Viles, 2002, and Coombes et al., 2011,
165 2015). Ten random replicate points (without overlap) were chosen on each chip and studied at three
166 magnifications ($\times 2$, $\times 3.2$ and $\times 5.6$), which were the most suitable for capturing the spatial variation
167 in organisms across all three sampling zones.

168 The top surface and a cross-section of each chip were subsequently studied using a FEI Quanta
169 200F Environmental SEM, operated at 20 kV, and equipped with a EDAX Energy Dispersive X-ray
170 spectrometer (EDS). The chips were uncoated as they were observed in low vacuum mode.
171 Secondary electron (SE) and backscattered electron (BSE) images were acquired. Ten replicate
172 points on the top surface of each rock chip were selected randomly (without overlap) to study the
173 occurrence of bioconstructions (e.g., biochemical encrustations, extracellular polymeric substances
174 (EPS), microbial filaments, algae, foraminifera, minerals) and bioweathering features (e.g.,
175 microboring, dissolution features, microcracks, biological pitting/etching). Likewise, six replicate
176 points (without overlap) were chosen from each chip to analyse the same features in cross-section.

177 The occurrence of features observed by stereomicroscopy and ESEM were classified using
178 “SACFOR” abundant scale (Hiscock, 1996) with the following classification: superabundant (80-
179 100%), abundant (40-79%), common (20-39%), frequent (10-19%), occasional (5-9%), rare (1-4%)
180 and absent (0%). This method is used to describe and quantify the abundance of marine benthic

181 flora and fauna in biological surveys (Jones and Pinn, 2006; Howarth et al., 2011). It has been also
182 employed in the analysis of micro-scale biological growths involved in weathering and erosive
183 processes of engineering materials (Coombes et al., 2011).

184

185 **2.4. Statistical analysis**

186 Statistical analyses were conducted for the recognition of microbial patterns related to different
187 natural light conditions throughout the cave system and for understanding the intensity of
188 bioweathering and bioconstructive processes. The occurrence of biological features in the three
189 sampling sites (SS1, SS2, SS3), as observed by stereomicroscopy, were first tested for normality
190 and subsequently analysed using the ANOVA method to check for homogeneity of variance (Table
191 1) (Underwood, 1997; Coombes et al., 2011). Each biological feature observed with
192 stereomicroscopy (i.e., algae, shiny filaments, fossils, white creamy deposits, black coatings) was
193 treated as a fixed factor and analysed for all “sampling sites” (SS1, SS2, SS3) and magnifications
194 ($\times 2$, $\times 3.2$, $\times 5.6$) using an Excel data sheet. Sampling sites and magnifications are the two sources of
195 variation. The comparison of combined “Sampling site observations” x “Magnification
196 observations” was made using the analysis of variance: single factor.

197 Biological features observed by ESEM were also analysed (Table 1). We treated each sampling site
198 individually, separating bioconstructive from bioweathering features and analysed them using the *t*-
199 Student test (two tailed test: two-samples assuming unequal variance) in two situations (on the top
200 surface and in cross-section) to understand if their presence is influenced by a specific location and
201 factor.

202

203 **3. Results**

204 **3.1. Rock microtopography**

205 The microtopographic analysis performed for all rock samples collected in Lighthouse Cave
206 revealed differences in the surface microreliefs according to their location. The contour map

207 analysis showed how microtopography changed within the different cave zones. For each rock
208 sample (one chip per sampling site; see Table 1), five distinctive classes of microrelief were
209 established (Table 2): i) sample SS1, located on a karren stone in the intertidal zone of a hypersaline
210 lake, depicted three classes of relief, ranging from 0 to 4-8 mm; ii) in SS2, collected in the twilight
211 zone of the cave, we observed the whole range of relief classes (from 0 to >12 mm), and iii) sample
212 SS3, from the dark cave interior and with the smallest range of roughness values (mean roughness
213 1.08), four classes of relief (from 0 to 8-12 mm), were noticed (Table 2).

214

215 **3.2. Microscopy observations**

216 Different biological features were observed by stereomicroscopy at three different magnifications
217 (Fig. 3), comprising filaments of staghorn-like algae (Fig. 3A), biofilms of green algae (Fig. 3B)
218 and biological-like filamentous structures (Fig. 3C). The occurrence of biological features tended to
219 decrease from site SS1 towards SS3 (Fig. 4), whereas the shiny filaments (Fig. 3D), fossils (Fig.
220 3E), and black coatings (Fig. 3F) tended to increase towards the cave darkness (Fig. 4). Brown
221 coatings were reported in all samples. The occurrence of these biological structures was measured
222 as shown in Figure 4, based on the SACFOR scale. Magnification of x5.6 revealed the most
223 representative results across all the studied sampling sites. Brown and black coatings showed the
224 highest occurrence for all the studied samples, contrasting with generic red algae and fossils.
225 *Cladophora* algae, white creamy deposits and black coatings were solely observed in SS1, SS2 and
226 SS3, respectively (Fig. 4).

227 White creamy deposits (Fig.5A, 5B) were exclusively observed on the chips collected from site
228 SS2, and are preferentially located along the depressions on the exposed rock surface (Fig.5A). The
229 EDS microanalysis showed that the creamy white deposits contain the following elements: C, O, Cl,
230 P, Mg, Ca, Na, S, Al, Si, K, Fe, and Mn (see the green spot in Figure 8).

231

232 ESEM examinations were performed on three chips per sampling site, and the occurrences (%) of
233 bioconstructive and bioweathering features both on the top surface and in cross-section are reported
234 respectively in Figure 6 and 7. Generally, bioweathering features were more abundant in the chip
235 cross-section, probably due to the biological coating observed on the top surface hiding the
236 dissolution structures.

237 ESEM images of bioconstructive and bioweathering features are reported in Figure 8. EDS
238 microanalyses showed that sample SS1 is composed of microcrystalline calcite (Fig. 8B and
239 corresponding EDS spectra), with abundant salt crystals due to the presence of seawater in the
240 sample location (Fig. 9A). It was noticed that when biological crusts are less abundant on the
241 sample surface, salt can easily penetrate up to a depth of ~3.5 mm (Fig. 9). In contrast, when
242 extensive microbial mats are present on the sample surface, biological pitting and etching solely
243 extend to a depth of 1-1.5 mm (Fig. 9A). Sample SS2 is characterised by an overall porosity of 12%
244 and shows the presence of several layers (Fig. 5C), the yellowish layers seem to be more compact
245 (made of microcrystalline calcite crystals), whereas the white layer is composed of calcite with
246 copious borings, likely contributing to enhance its porosity (Fig. 5D). Fine layers and aggregates of
247 Ca-phosphate minerals are present both along the cross-section and on the sample surface, and are
248 likely related to bat guano deposits. Borings of approximately 0.5 mm diameter are clearly visible
249 along the cross-section, increasing mineral porosity (Fig. 9B).

250 The top surface of SS3 is characterised by cubic Ca-phosphate minerals (Fig. 8K and Fig. 9C) and
251 black coatings of ferromanganese oxides, whereas the inner part is made of carbonate minerals
252 enriched in Mg and acicular Ca-phosphates (Fig. 8J and Fig. 9C). Endolithic microorganisms able
253 to pit and etch rocks were also observed (Fig. 8I).

254 Table 3 summarizes and compares all the results obtained for the rock chips from each sampling
255 site.

256 The results of the ANOVA analysis for the biological features observed by stereomicroscopy are
257 reported in Table 4. The null hypothesis affirms: “The biological features are not influenced by light

258 attenuation". The obtained results show $F_{\text{values}} > F_{\text{critic}}$ in almost all biological features of the three
259 sites. This result means that the null hypothesis can be rejected because the distribution of almost all
260 the observed biological features is influenced by light attenuation. Only the brown coatings show an
261 $F_{\text{value}} < F_{\text{critic}}$, suggesting that their distribution is not influenced by light attenuation.

262 The results of Student's *t*-test analysis of ESEM observations are reported in Tables 5 and 6
263 (bioconstructive and bioweathering features, respectively). We used Student's *t*-test to understand
264 whether bioconstructive-dissolution features related to the same process. Analysing bioconstructive
265 features on the top surface and cross-sections from the three sampling sites showed that the P
266 (probability) values for SS1 and SS2 are higher (Table 5) than the critical value (0.05). This result
267 means that the *t*-test analysis is not significant and so it is difficult to reject null hypothesis.

268 Bioconstructive features in SS1 and SS2 could be related to the same process (i.e., the presence of
269 blue filaments). Conversely, the P value of SS3 is lower (Table 4) than the critical value (0.05), so
270 null hypothesis can be rejected because the bioconstructive features might be related to a different
271 process (i.e., the presence of shiny filaments or Mn-Fe oxide-hydroxide precipitation). Analysing
272 bioweathering features, we saw that P values are much higher (see Table 6) than P critical value
273 (0.05), demonstrating the test analysis not to be significant. The same process (i.e., the presence of
274 microorganisms able to pit and etch the rock) can be responsible for the observed phenomena.

275

276 **4. Discussion**

277 Contrasts in light attenuation inside the cave have an important influence on biological colonisation.
278 Based on visual inspections and microscopy observations, sampling sites SS1 and SS2 are
279 dominated by phototrophic organisms. Their behaviour affects microtopography, mineralogy and
280 geochemistry of the rock substrate.

281 We sought to assess how the micromorphology of karst environments changes across an
282 environmental and process gradient. Biological features occur in all samples, and three different
283 associations were recognised using light and scanning electron microscopy, one for each sampling

284 site. We observed that blue filaments, *Cladophora* and red algae (including staghorn algae) tend to
285 decrease in abundance from the hypersaline lake toward the cave interior owing to light attenuation,
286 whereas shiny filaments, fossils and black coatings tend to increase along the same profile. This
287 result agrees with Coombes et al. (2015), who studied the Puerto Princesa Underground River in the
288 Philippines, suggesting that the sensitivity of microbial communities to light strongly influences the
289 nature and types of biogeomorphological processes that operate in cave systems.

290 The increase of fossils (in SS3) is related to a marine inflow (through fractures), whereas shiny
291 filaments and black Mn-coatings might be controlled by microorganisms able to thrive in nutrient-
292 poor dark locations.

293 Light attenuation plays an important role in influencing the behaviour of biological communities
294 involved in landform processes, bioweathering and bioconstruction within the underground
295 environment. As suggested by Phillips (2016), light attenuation is an interesting candidate for
296 “niche construction”. Niche construction means that biogeomorphic ecosystem engineering
297 influences natural selection (such as stromatolite formation and/or Ca-nitrate precipitation in dry
298 cave deposits). As a matter of fact, it is well-known that geomorphic processes (such as
299 cementation-precipitation, weathering, erosion and deposition) can be both microbially-controlled, -
300 induced, –influenced, and/or abiotic (Viles, 2012).

301 To provide a clear understanding, the data obtained in this study are separately discussed for each
302 sampling site, focusing on the main biogeomorphological processes.

303

304 **4.1. Sampling site SS1**

305 The microtopography and high surface roughness of the limestone rock from sampling site 1,
306 located in a hypersaline lake near the cave entrance, were promoted by biological activity as
307 revealed by microscopy observations (boreholes and pittings). On the top surface of SS1 we also
308 observed a salt penetration band (~6 mm), as well as changes in rock porosity associated with
309 endolithic growth. It is well known that salt crystallisation, similar to microgelivation, can actively

310 contribute to the weathering of rocks, especially through its penetration into pore spaces and rock
311 fractures (Williams and Robinson, 1998; Matsuoka, 2001; Moses et al., 2014). The depth of
312 subsurface deterioration depends on parameters such as porosity, permeability, lithology and
313 moisture, as well as climatic conditions and biological activity (Matsuoka, 2001). Epilithic
314 organisms may weaken the top surface of rocks (e.g., by boring, pitting and etching), whilst the
315 endolithic ones (as observed in SS3 samples) may affect the rock just beneath the surface by
316 enlarging porosity or fractures, through chemical-physical reactions that change their
317 microenvironment (Friedmann, 1982; Bell, 1993; Viles, 2000; 2012). Phototrophs were the
318 dominant organisms colonising the rock surface in SS1, and, likely helped by salt penetration and
319 gastropod grazing activities, contributed in weakening and disaggregating particles, which were
320 subsequently removed by seawater (wave and tide fluctuations) and/or by wind. These processes
321 together would have created the observed profile characterised by microvalleys and -ridges.

322

323 **4.2. Sampling site SS2**

324 The SS2 rock chips showed greater roughness (Table 2), and their profile is more pronounced than
325 SS1 (Table 3). The white creamy deposits observed along the depressions of the chip surface, with a
326 complex chemistry (O, Cl, C, P, Mg, Ca, Na, S, Al, Si, K, Fe, and Mn), are likely related to bat
327 guano deterioration. The fine layers and the presence of aggregates of Ca-phosphate minerals
328 associated with the white deposits support this hypothesis. The rock samples from SS2 are
329 characterised by laminae with different colours and porosity/permeability as previously described.
330 The yellowish layers (first and third) (Fig. 9C) are relatively more compact (made of
331 microcrystalline calcite) and less porous than the second whitish layer in which boreholes occur
332 extensively (diameter of $10 \pm 4 \mu\text{m}$). These microcrystalline layers may be the product of
333 subsequent weathering processes that also changed the primary porosity (Winkler, 1997; Nicholson,
334 2001; Tuğrul, 2004).

335 Black (1933) described deposits in Andros Island (Bahamas) whereby trapping and binding
336 processes involved the presence of cyanobacteria. Thus, the laminations are likely related to
337 trapping and binding of detrital grains and microfossils supplied by tidal fluctuation and recall
338 microbialites (Burne and Moore, 1987). Similar dark-brown and reddish crusts were observed along
339 the intertidal zone in several flank margin caves in Croatia (e.g., Otoničar et al., 2010) and are likely
340 related to microorganisms such as bacteria and red algae. These laminated deposits can be defined
341 as a “biological boundstone” according to Black (1933). The organisms involved in its formation
342 live in dark, quiet, shallow water, and in the tide-influenced twilight zone of the cave environment,
343 where nutrients are delivered by guano that also behaves as source of acids and organic matter
344 input. The extensive borings in the whitish layer of the SS2 chip cross-section suggests that
345 endolithic microorganisms were involved in the formation of these layers (Fig.5D).

346

347 **4.3. Sampling site SS3**

348 The sampling site SS3 is located in the deepest and darkest part of Lighthouse Cave. The top
349 surfaces of SS3 chips showed smoother texture than samples SS1 and SS2 as revealed by high
350 precision laser scanner measurements (Table 4). Black coatings were observed on the sample
351 surfaces, mainly composed of cubic Ca-phosphate minerals and Mn-Fe oxides-hydroxides. In
352 addition, shiny filaments and microfossils were visible. The internal structure of SS3 chips was
353 characterised by porous carbonate minerals enriched in Mg, and acicular Ca-phosphates with pitted
354 and etched crystal surfaces, likely caused by endolithic microorganisms. Similar deposits have been
355 described by Spilde et al. (2009) and were defined as “Speleosols” (i.e., “soil-like materials formed
356 in caves”). They are made of ferromanganese deposits related to two different processes involving
357 the activity of Mn-Fe oxidising and acid-producing microbiological communities (Spilde et al.,
358 2005; Miller et al., 2012): 1) alteration of the cave wall, leaching of soluble elements and
359 subsequent enrichment in Al, Fe, Mn and trace elements, and 2) deposition of secondary minerals
360 (mainly Mn-Fe oxides-hydroxides). Usually these structures have three components: an external

361 dark-coating/crust or speleosol, a punk rock (a porous and altered portion of host rock; Hill, 1987)
362 and bedrock.

363

364 **5. Conclusions**

365 Microtopography of the exposed rock surfaces within and close to Lighthouse Cave varies with
366 location (e.g., entrance, twilight zone or deep into the cave). Going from outside (SS1) to deep
367 inside (SS3) there is a general flattening of microrelief.

368 In general, light attenuation, together with organic matter supply, rock type, age, and diagenetic
369 maturity play an important role in influencing the behaviour of biological communities involved in
370 rock surface processes. We found that bioweathering is more intense on samples collected outside
371 the cave (SS1), likely due to the presence of phototrophs that, dissolving and weakening the rock,
372 disaggregate particles that are subsequently removed by wind erosion, creating typical ridges and
373 valleys in the rock surface microtopography. Nevertheless, episodic gastropod grazing actions
374 might “reset and shape” the overall microtopography. Conversely, within the cave environments,
375 chemotrophs facilitate both bioweathering (endolithic boreholes) and bioconstructive processes by
376 dissolving mineral grains and/or inducing secondary mineral precipitation (e.g., Mn-Fe oxides-
377 hydroxides), respectively.

378 The above described biogeomorphological structures, especially the ones found along the cave
379 walls (SS2-SS3) at the fluctuating water level, testify that, nowadays in this peculiar flank margin
380 cave, dissolution/corrosion processes due to fresh seawater mixing are less active.

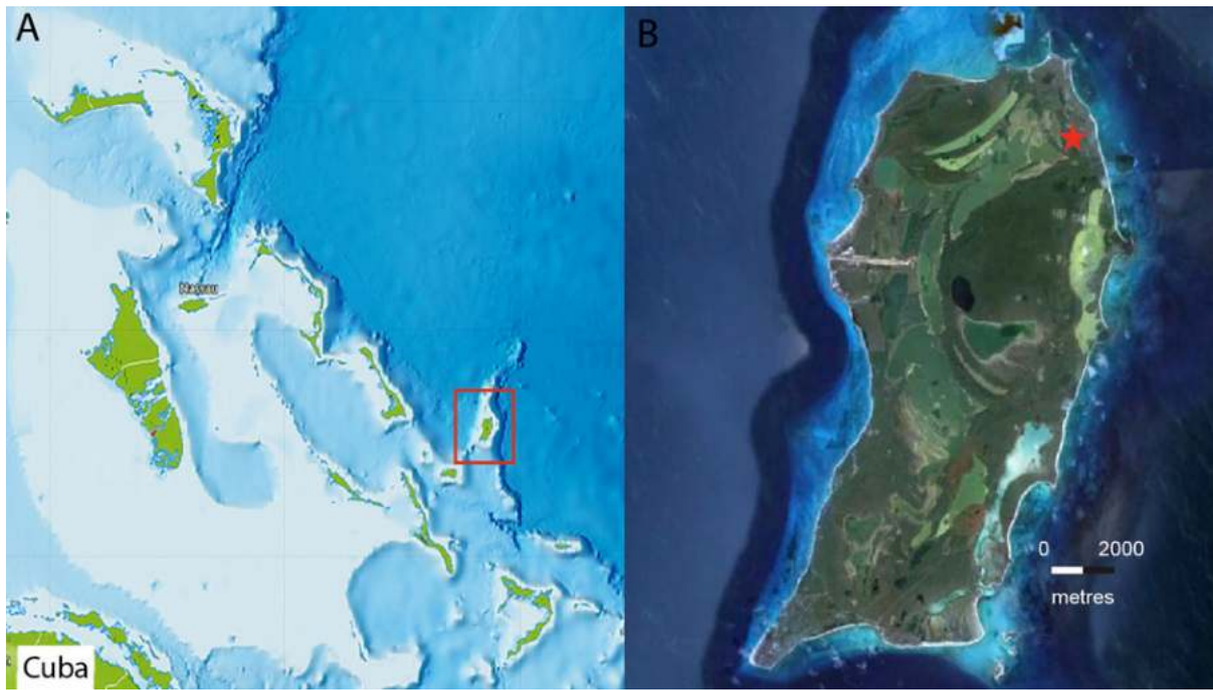
381 In addition, we propose that the geomorphological evolution is strongly influenced by the degree of
382 rock diagenesis (eogenetic (immature) limestones in Lighthouse Cave vs. telogenetic (mature)
383 limestones of Puerto Princesa Underground River) more than the initial mechanism of
384 speleogenesis.

385

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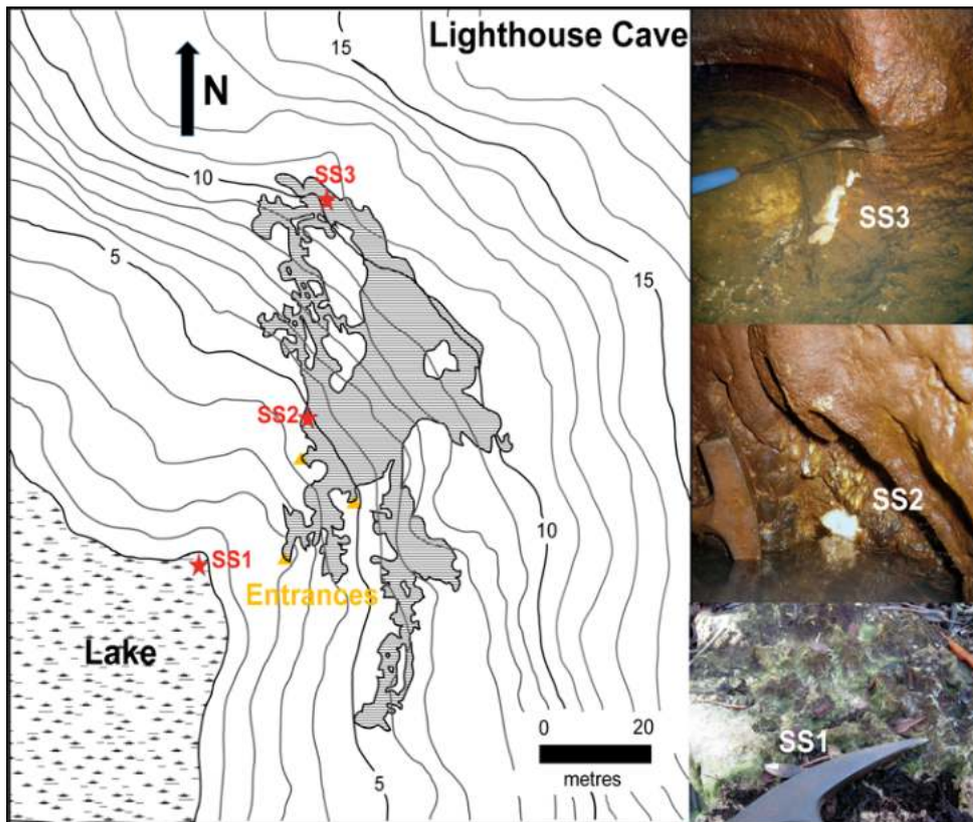
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400

401 Figure 1.

402



403

404 Figure 2.

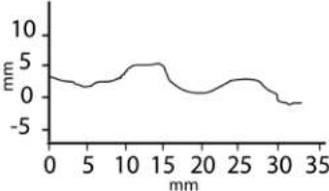
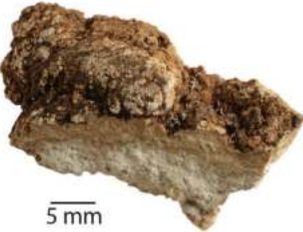
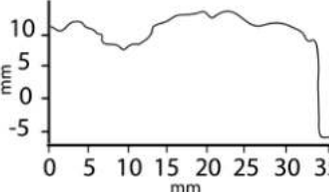
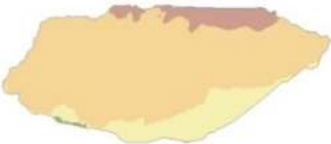
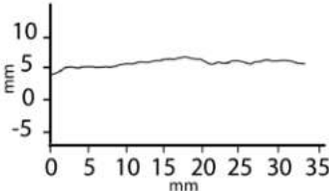
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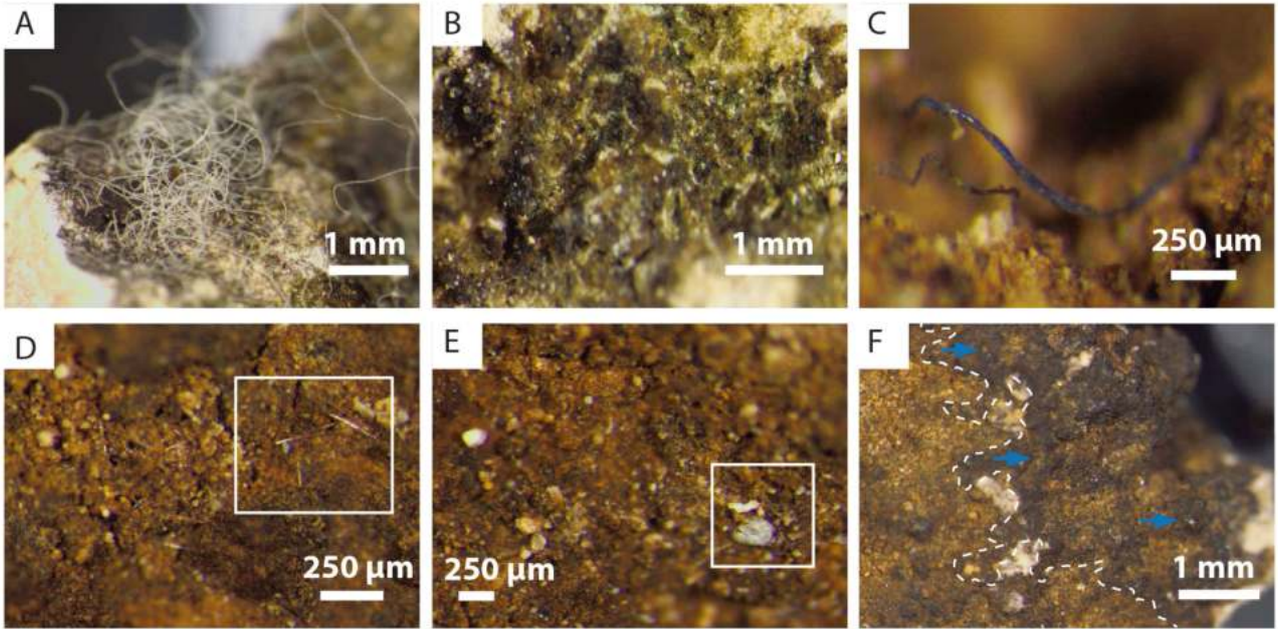
406 Table 1.

Method	Number of chips studied from each sampling site	Number of observations per chip
Laser scanner	1	1
Roughness measurements	1	6
Light microscopy	3	10
ESEM (surface)	3	10
ESEM (cross-section)	3	6

407

408 Table 2.

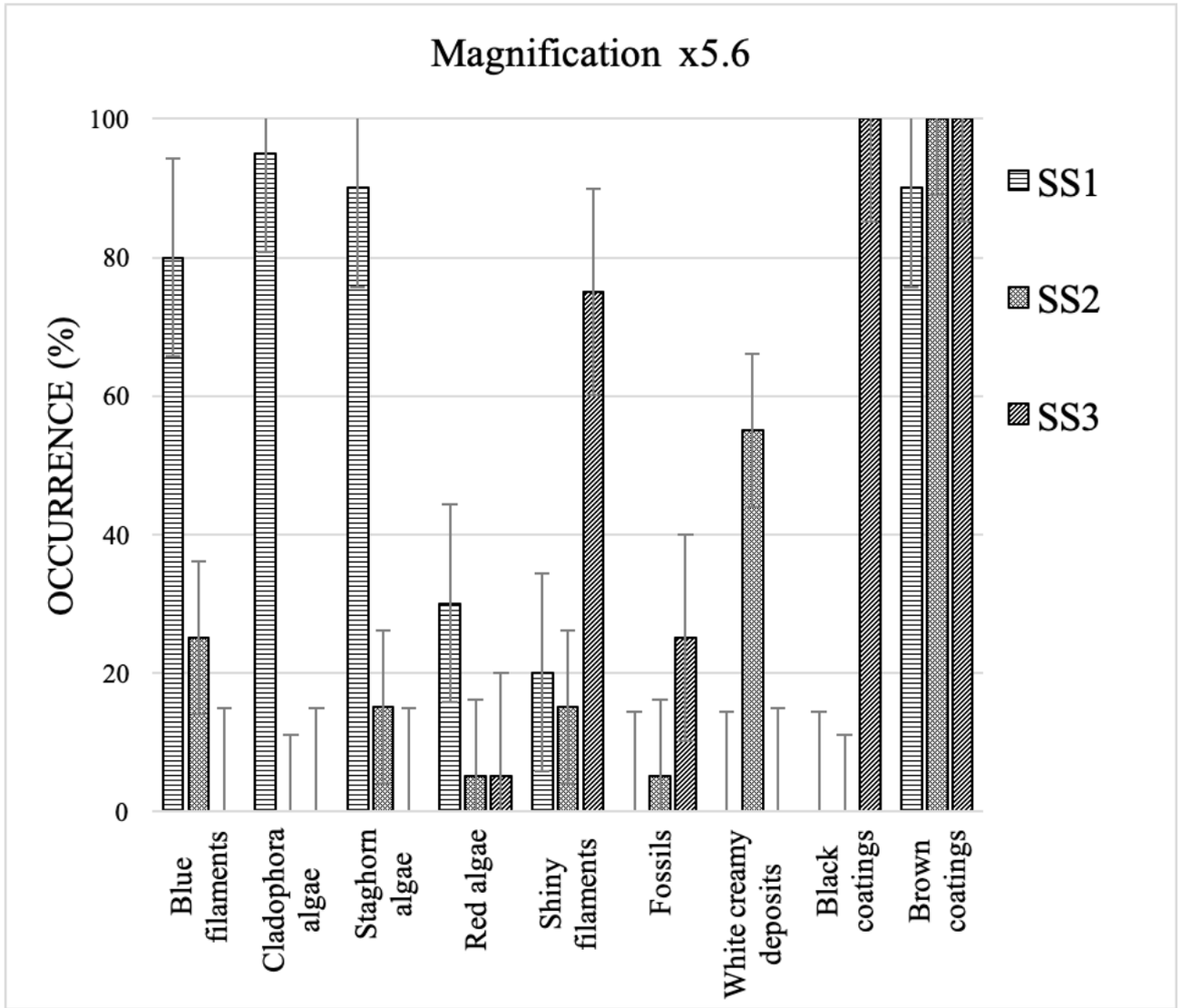
Sampling site	Stereomicroscopy image	DTM Contour map analysis	Microrelief classes scale	GIS Contour Map Analysis		Surface roughness value and representative profile
				Total area [mm ²]	Total area (%) per microrelief class	
SS1			1 < 0 mm	407 ±3	1) 36.0 ±0.5	1.15 – 1.33 
			2 0-4 mm		2) 53.0 ±0.3	
			3 4-8 mm		3) 11.0 ±0.2	
			4 8-12 mm		4) 0	
			5 > 12 mm		5) 0	
SS2			1 < 0 mm	369 ±6	1) 13 ±0.8	1.43 – 2.17 
			2 0-4 mm		2) 1.5 ±0.8	
			3 4-8 mm		3) 15.0 ±2	
			4 8-12 mm		4) 51.5 ±0.6	
			5 > 12 mm		5) 19.0 ±0.07	
SS3			1 < 0 mm	382 ±2	1) 0.4 ±1	1.03 – 1.13 
			2 0-4 mm		2) 15.0 ±0.2	
			3 4-8 mm		3) 77.0 ±0.3	
			4 8-12 mm		4) 7.6 ±0.09	
			5 > 12 mm		5) 0	



409
410

Figure 3.

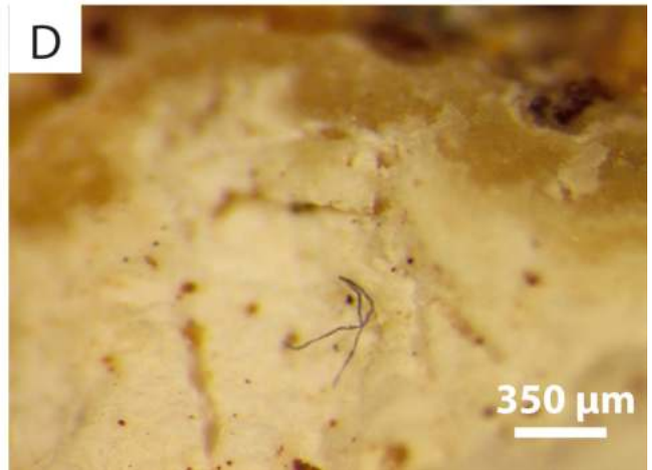
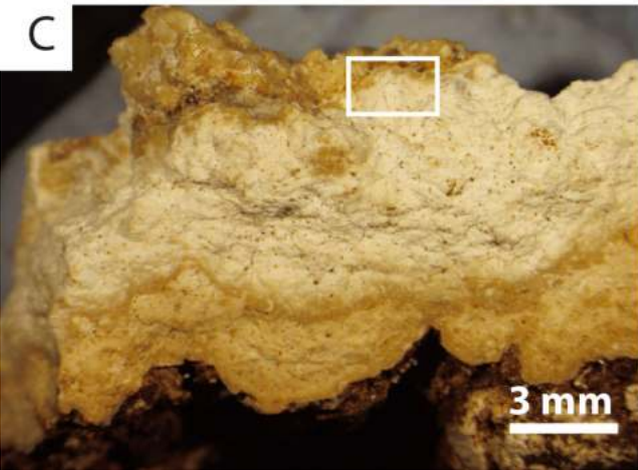
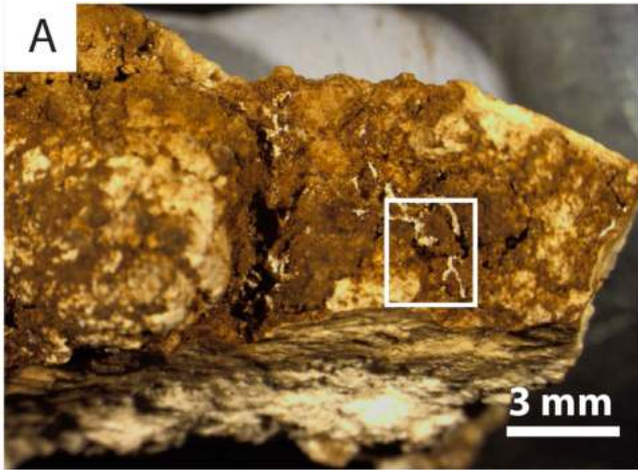
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413 Figure 4.

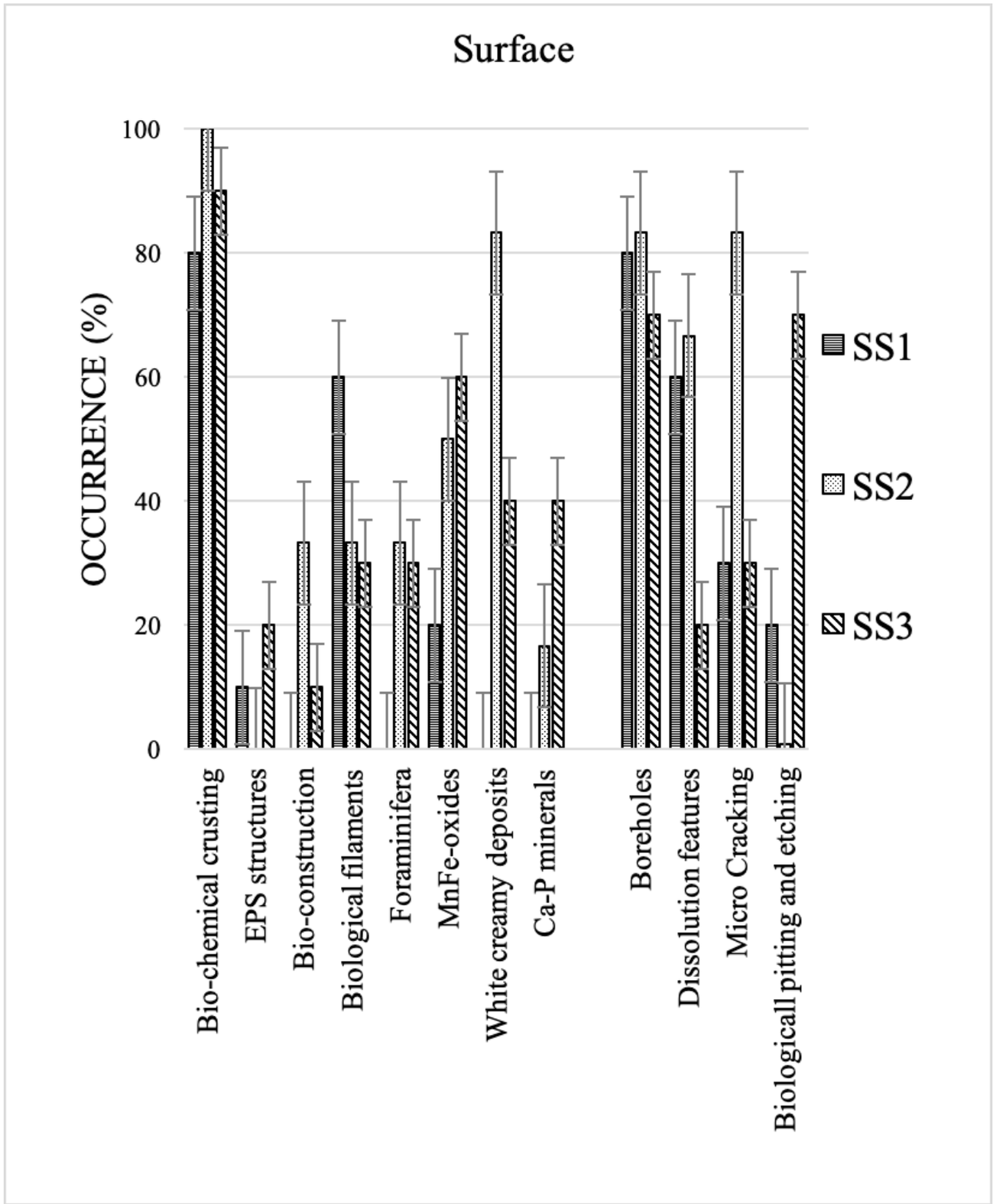
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416 Figure 5.

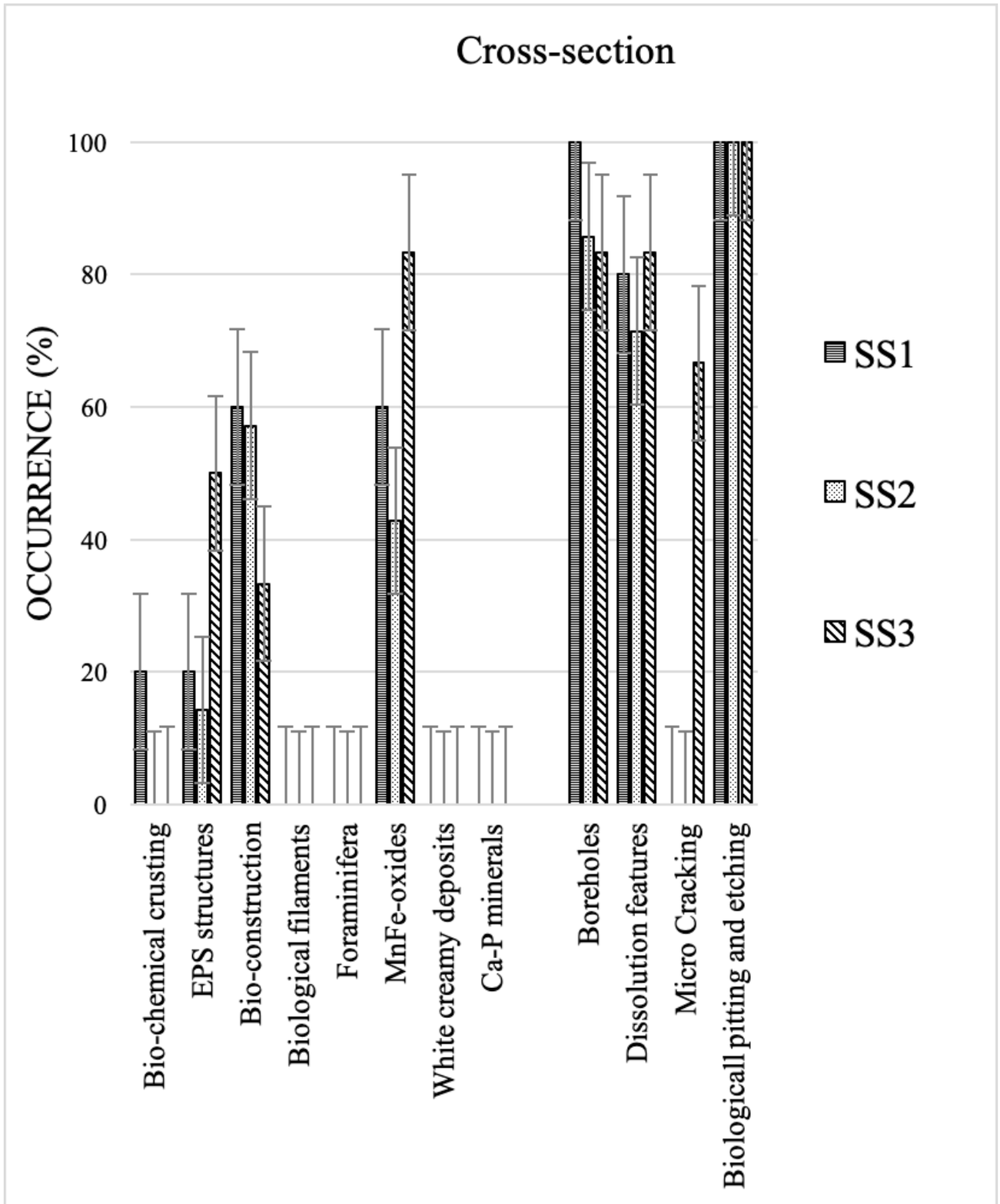
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419 Figure 6.

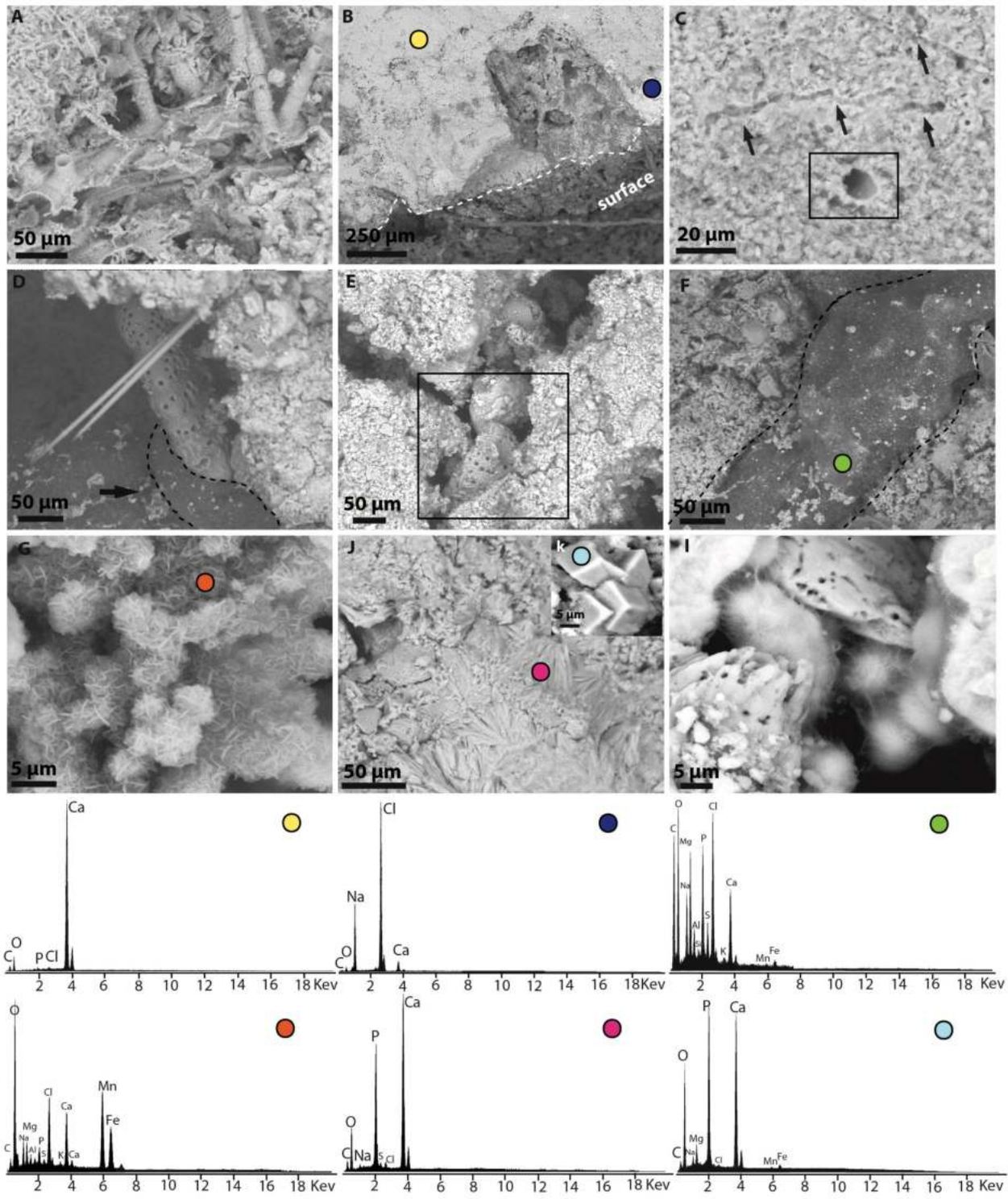
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422 Figure 7.

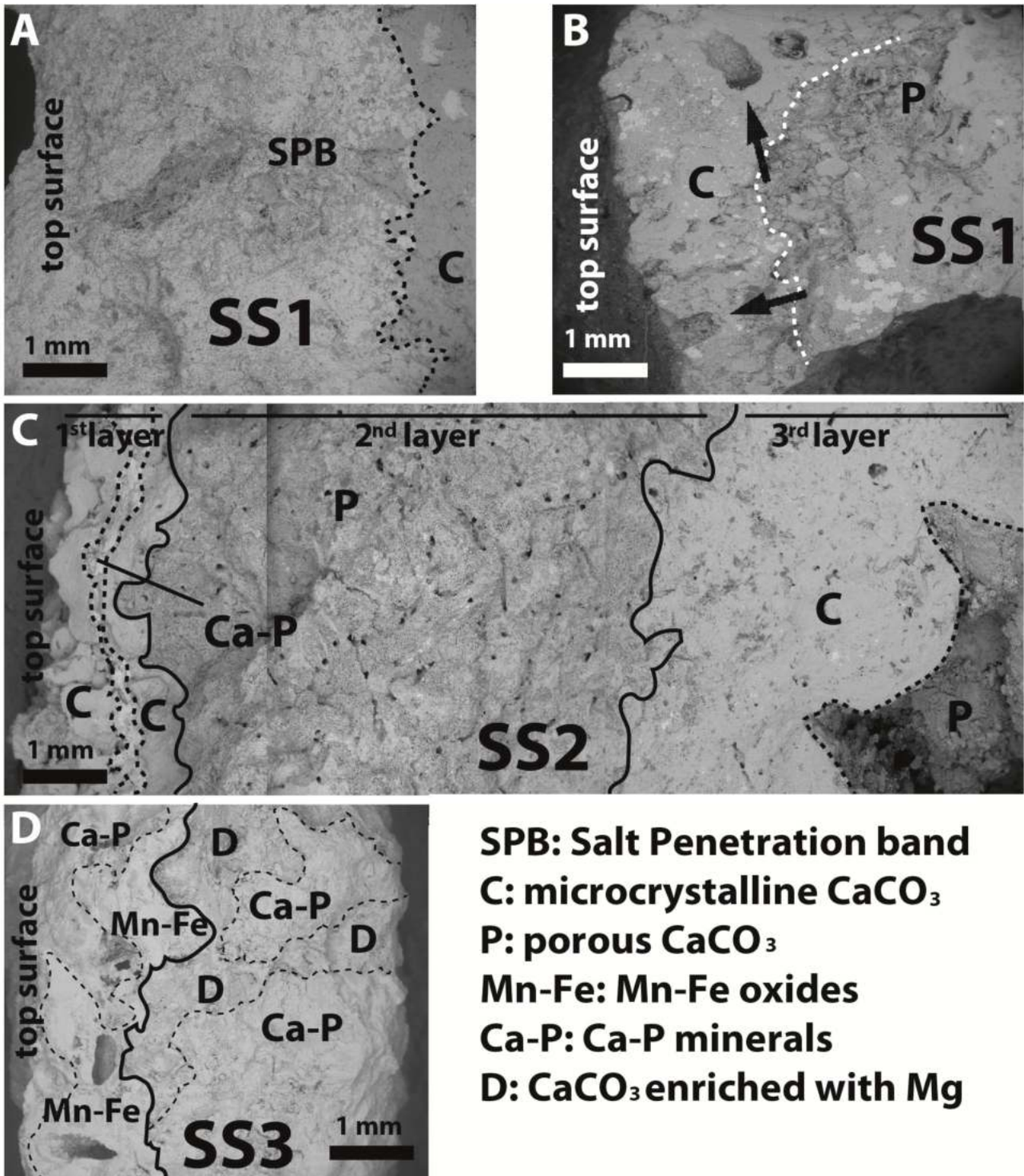
423



424

425 Figure 8.

426



427

428 Figure 9.

429

	SS1	SS2	SS3
Sampling Site	<ul style="list-style-type: none"> hypersaline lake located in a mangrove forest 20 m W of the main cave entrance 	<ul style="list-style-type: none"> limestone wall in the twilight zone of the cave, 20 m from the entrance, close to the water table, approximately 10-15 cm below mean high tide level 	<ul style="list-style-type: none"> limestone wall near the water table, approximately 25 cm below high tide level and located ~60 m from the cave entrance
Rock Characteristics	<ul style="list-style-type: none"> Salt penetration band Pitted and etched band 	<ul style="list-style-type: none"> Semi-parallel laminae (yellowish and whitish) Endolithic microboring 	<ul style="list-style-type: none"> Black coatings of Mn-Fe oxides-hydroxides covering Mg-enriched carbonate rocks
Rock Properties	<ul style="list-style-type: none"> Salt minerals promote rock weathering and changes in porosity and permeability 	<ul style="list-style-type: none"> Creamy deposits (O, Cl, C, P, Mg, Ca, Na, S, Al, K, Fe, Mn) Ca-phosphate minerals 	<ul style="list-style-type: none"> Ca-phosphate minerals Mn-Fe oxides-hydroxides Fossils
Weathered Rind	<ul style="list-style-type: none"> Green brownish crust 	<ul style="list-style-type: none"> Brown-reddish crust and depression filled by creamy white deposits 	<ul style="list-style-type: none"> Dark-brownish crust and black coatings
Dominant Microbial features	<ul style="list-style-type: none"> <i>Cladophora</i> algae Staghorn algae Red algae Blue filaments Brown coatings 	<ul style="list-style-type: none"> Red algae Brown coatings 	<ul style="list-style-type: none"> Brown coatings Likely microbial organisms associated with shiny filaments
Microtopography (Roughness value = RV)	<ul style="list-style-type: none"> Valleys and Ridges RV 1.15-1.33 	<ul style="list-style-type: none"> Well-developed ridges and troughs RV 1.43-2.17 	<ul style="list-style-type: none"> Flat surface RV 1.03-1.13

433 Table 4.

Biological features	df W	df B	ms W	ms B	F crit	F	P
Blue filaments	6	2	8	416	5.14	156	<0.000
<i>Cladophora</i> algae	6	2	5	624	5.14	401.28	<0.000
Staghorn algae	6	2	3	644	5.14	724.75	< 0.000
Red algae	6	2	19	150	5.14	23.31	0.0015
Shiny filaments	6	2	33	202	5.14	18.24	0.0028
Fossils	6	2	8	16	5.14	5.61	0.042
White creamy deposits	6	2	<1	213	5.14	961	<0.000
Brown coatings	6	2	2	2	5.14	3	0.125

434

435

436 Table 5.

Bioconstructive fts.	df	Ms	Mc	Vt	t-crit	t-value	P
SS1	16	2.2	0.9	5.2	2.12	1.25	0.23
SS2	16	2.4	0.9	3.1	2.12	1.88	0.08
SS3	16	3.5	1.1	5.3	2.12	2.25	0.04

437 df = Degrees of freedom; Ms = mean bioconstructive features observed on the top surface; Mc = mean bioconstructive
438 features observed along the cross-section; Vt = total variance; t-crit = *t* value critic; *t*- value = *t*- value obtained from
439 statistical analysis; P = probability.

440

441

442 Table 6.

Bioweathering features	df	Ms	Mc	Vt	t-crit	t-value	P
SS1	6	4.7	3.5	6.6	2.44	0.68	0.52
SS2	6	4.7	4.5	4.9	2.44	0.16	0.87
SS3	6	4.7	5	3.8	2.44	-0.18	0.86

443

444

445 **Figure and table captions**

446 Figure 1. A) Location of San Salvador Island, Bahamas, and B) Lighthouse Cave on the NE coast of
447 San Salvador (the red star shows the position of the cave).

448

449 Figure 2. Lighthouse Cave plan (modified from Roth, 2004) and location of sampling sites above
450 present sea level (red stars). The images on the right show the sampling sites SS1, SS2, SS3.

451 Hammer represents the scale of the pictures.

452

453 Table 1. Number of rock chips and observations performed for each analysis.

454

455 Table 2. Contour map analysis of rock samples from each sampling site. The most representative
456 rock chip is shown using stereomicroscopy and DTM images. In addition, microrelief classes scale
457 (1. green, 2. yellow, 3. orange, 4. brown, 5. white), the respective area for each microrelief class, the
458 surface roughness, and representative profile are also reported.

459

460 Figure 3. Biological features observed by stereomicroscopy. A) Site SS1: staghorn algae (red algae)
461 and *Cladophora* algae are clearly visible; B) Site SS1: *Cladophora* algae; C) Site SS2: blueish
462 biological-like filament; D) Site SS3: several tiny shiny filaments are clearly visible in the white
463 square; E) Site SS3: a small fossil (juvenile stage foraminifera) is visible against the brownish
464 background; F) Site SS3: black coatings (blue arrows).

465

466 Figure 4. Occurrence of micro-scale biological features observed by stereomicroscopy (x5.6). Three
467 chips were analysed per sampling site (SS1, SS2 and SS3). Ten points were observed on each chip
468 (without overlap), giving thirty points for each sampling site (at x5.6 the area of an analysed spot is
469 6 mm^2). An occurrence of 100% means that a biological feature is observed in all ten points of each

470 chip. These measurements were based on the SACFOR scale: superabundant (80-100%), abundant
471 (40-79%), common (20-39%), frequent (10-19%), occasional (5-9%), rare (1-4%) and absent (0%).

472

473 Figure 5. Stereomicroscopy images of site SS2; A) White creamy deposits located along the
474 depressions on exposed rock surfaces. B) Detail of A; C) Cross-section of SS2 showing several
475 layers with different colours; the whitish layer is extensively bored; D) Detail of C showing
476 microboring.

477

478 Figure 6. Histogram showing the occurrence of bioconstructive and bioweathering features
479 observed on the chip top surfaces from each sampling site (SS1, SS2 and SS3). We analysed three
480 chips from each sampling site, using ESEM.

481

482 Figure 7. Histogram showing the occurrence of bioconstructive and bioweathering features along
483 the chip cross-sections from each sampling site (SS1, SS2 and SS3) using ESEM.

484

485 Figure 8. ESEM images of bioconstructive and bioweathering features on chip top surfaces and
486 cross-sections. All are BSE images unless stated otherwise. The coloured dots represent the position
487 where EDS spectra were obtained. A) Bioconstructions on the top surface of SS1, particularly
488 mineralised filaments; B) Biological pitting and etching in the cross-section of SS1; C) Dissolution
489 features (black arrows) and boring (black square) in cross-section of SS1; D) Foraminifera on the
490 top-surface of SS2. The black arrow indicates white creamy deposits (that in the BSE image have a
491 dark grey colour); E) Two foraminifera on the top surface of SS2; F) White creamy deposits on the
492 top surface of SS2 (these white deposits are dark in BSE); G) Manganese oxides in the cross-section
493 of SS3 (SE image); J) Acicular crystals of Ca-phosphate on the top surface of SS3; K) Cubic
494 crystals of Ca-phosphate on the top surface of SS3; I) Microboring caused by endolithic organisms
495 on mineral grains in the cross-section of SS3.

496 Figure 9. ESEM-BSE images of representative cross-sections from the three sampling sites (SS1,
497 SS2 and SS3). A) salt penetration band (SPB) is observed on the surface of SS1; B) biological
498 pitting (due to microalgal growth) is visible (black arrows); C) SS2 is characterised by several
499 layers of calcite with different porosity and permeability. Borings are clearly visible along the
500 cross-section in the second layer; D) the top surface of SS3 is characterised by cubic Ca-phosphates
501 and black coating of Mn-Fe oxides, whereas the inner part is made of carbonates enriched with Mg
502 and acicular Ca-phosphates.

503

504 Table 3. Short description of the main results for the three sampling sites from Lighthouse Cave,
505 regarding rock characteristics and properties, weathered rind, dominant microbial features, and
506 microtopography values.

507

508 Table 4. Statistical results obtained using the ANOVA test for the occurrence of biological features
509 as observed by stereomicroscopy. $df W$ = degrees of freedom within group; $df B$ = degrees of
510 freedom between groups; $ms W$ = mean square variance within group; $ms B$ = mean square
511 variance between groups; F = ratio of variance; P = significance.

512

513 Table 5. Student's t -test (two tailed test: two-samples assuming unequal variance) analysis of
514 bioconstructive features observed using ESEM in each sampling site.

515

516 Table 6. Student's t -test (two tailed test: two-samples assuming unequal variance) analysis of
517 bioweathering features observed using ESEM in each sample.

518

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