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Biofilms Confer Resistance to Simulated Extra-terrestrial Geochemical Extremes — Source link

Adam R. H. Stevens, Delma S. Childers, Mark Fox-Powell, Charles S. Cockell Institutions: University of Aberdeen, University of St Andrews Published on: 10 Jan 2017 - bioRxiv (Cold Spring Harbor Labs Journals)

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¹ Biofilms Confer Resistance to

² Simulated Extra-terrestrial

Geochemical Extremes

- 4 Adam H. Stevens^{*1}, Delma Childers^{1,2}, Mark Fox-Powell^{1,3}, Charles S. Cockell¹
- 5 1. UK Centre for Astrobiology, School of Physics and Astronomy, University of Edinburgh, UK
- 6 2. MRC Centre for Medical Mycology at the University of Aberdeen, Aberdeen Fungal Group,
- 7 Institute of Medical Sciences, Foresterhill, Aberdeen, UK
- 8 3. Department of Earth and Environmental Sciences, University of St. Andrews, UK
- 9 *Corresponding author: adam.stevens@ed.ac.uk ORCIRD 0000-0001-8335-4143

10 **1 ABSTRACT**

11 Biofilms improve microbes' resistance to a variety of extreme physical and chemical conditions on 12 Earth. The discovery of putative aqueous environments on other planetary bodies such as Mars 13 motivates an interest in understanding the viability of life, and the potential role of biofilms, in previously unexplored geochemical extremes. We investigated the loss of viability of planktonic cells 14 15 and biofilms of Sphingomonas desiccabilis (a Gram-negative, desiccation resistant, soil crust-forming 16 organism) to simulated Martian brines. These brines were produced from geochemical modelling of 17 past aqueous environments on Mars, and their high sulfate concentrations make them different to 18 most terrestrial brines, although similar briny environments have been found in locations such as the Basque Lakes in Canada or in deep subsurface groundwater systems. Biofilms grown on basaltic 19 20 scoria were subjected to the simulated martian brines and the viability of cells was measured over 21 time and compared to equivalent planktonic cultures. Crystal violet assay was used to measure how 22 the biomass of the biofilms changed over time in response to the brines. While certain brines were 23 highly hostile to microbial viability, we found that biofilms that were desiccated prior to being 24 treated with brines maintained viability over a longer treatment period when compared to 25 planktonic cells. Our results show that biofilms confer short-term protection to the harsh osmotic, ionic, and acidic conditions of Mars-relevant brines. However, in the most extreme simulated brines, 26 27 even biofilms eventually lost viability. By demonstrating that biofilms confer protection to conditions 28 that are potentially analogous to current day recurrent slope lineae (thought to be produced by the 29 flow of briny fluids) on Mars, our results show that contaminant biofilm-forming microorganisms 30 may have a greater chance of surviving in so-called 'Special Regions' on Mars, with implications for planetary protection in missions that aim to explore these regions. 31

2 INTRODUCTION

33	Biofilms are produced by a variety of organisms and generally consist of cells attached to a surface
34	(and each other) in an exuded matrix of extracellular polymeric substance (EPS) [1]. Biofilms can be
35	formed by single species or complex microbial communities and are generally produced in response
36	to the presence of a surface or external cues and stressors. Microbial biofilms increase the resistance
37	of the inhabitants to a number of extreme conditions including immunological aggression [2],
38	antibiotics [3], bactericidal chemicals [4], rapid changes in temperature and pH [5], UV radiation [6]
39	and salinity [7,8]. Biofilm-producing microorganisms are therefore present and thrive in a range of
40	environments on Earth and may have been involved at an early stage as one of the steps in the
41	development of the first cells [9].
42	Although biofilms confer resistance to extremes found on Earth, their ability to provide protection
43	against entirely novel chemical conditions found on other planetary bodies remains unknown. The
44	search for extra-terrestrial habitable environments centres on the availability of liquid water, and
45	aqueous environments are now known to exist on a number of planetary bodies including Mars,
46	Europa, Enceladus and Titan [10-14]. Many of these aqueous environments contain combinations of
47	anions that are rare on Earth. For example, the high concentrations of sulfate and other divalent ions
48	in Martian brines, a consequence of the specific geological history of that planet, have consequences
49	for habitability, making ionic strength an important determinant of whether the brines will support
50	life [15]. Recent observations suggest that recurrent slope lineae (RSL) features on Mars are created
51	by the action of flowing water which is saturated with dissolved perchlorate salts [16]. Such brine
52	compositions have not been found on Earth. In this study, we investigated the resistance of biofilms
53	to aqueous environments analogous to environments on Mars such as RSL currently potentially
54	active [16] or in quasi-tidal areas that existed during the phase of the planet's history when it lost its
55	surface water to atmospheric escape or to retreat into subsurface ice [17,18], but also have

- 56 implications for the icy moons in the outer solar system, which are thought to have high levels of
- 57 dissolved salts in their liquid water interiors [19].
- 58 We hypothesise that production of biofilms might protect microorganisms and allow them to
- 59 maintain viability following exposure to compositionally diverse brines with high levels of dissolved
- 60 salts. Our results have implications for the adaptations that would be required by any putative life in
- 61 extra-terrestrial environments and for planetary protection, *i.e.* whether terrestrial organisms
- 62 accidently introduced into extra-terrestrial brines could persist.

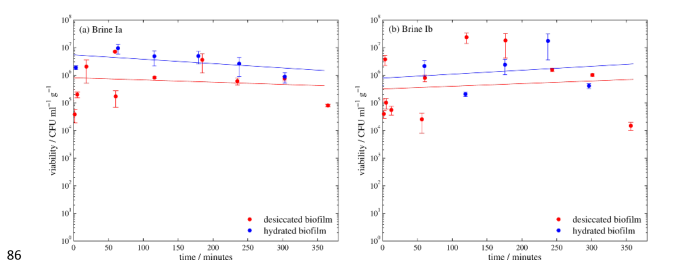
63 **3 RESULTS**

64 3.1 VIABILITY AFTER BRINING

Figures 1-4 show the viability of *S. desiccabilis* biofilms over time when immersed in Mars-analogue
brines. Each time point was an independent culture (triplicate pieces of scoria with biofilm on them)
exposed to brine for a given length of time. Experimental replicates are collected on the same axes.
Error bars are the standard error of the biological triplicates from each experiment. Best-fit lines
were calculated using a linear regression of all experimental replicates.

70 In Type I brines (Fig. 1) there was only minor decreases in viability over more than 300 minutes when 71 dried or when hydrated. In some cases there appears to be growth during brine immersion (Ib). In 72 brine IIa (Fig. 2a), viability was maintained for over 300 minutes whether the biofilms had been dried 73 or not. In brine IIb (Fig 2b) the dried biofilms showed a decrease in viability, decreasing to below 74 detection levels after around 240 minutes. Hydrated biofilms showed a more rapid decrease in 75 viability, as viable cells were unrecoverable after less than 30 minutes of brine treatment. In type III 76 brines (Fig. 3), viability of cells in the hydrated biofilms reduced to below detection limits after 77 around 30 minutes, which was the shortest timepoint measured. There was more variation in the 78 maintenance of viability of dessicated biofilms treated with the type III brines. For brine IIIa there

was a reduction in cell viability over the time course, but not to below detection limits, and not in a
linear fashion. In brine IIIb there was a rapid loss of viability to below detection levels within
approximately the first 60 minutes, though individual cultures in some experimental replicates
showed some measured viability at longer timescales. Brine IIIb has the lowest water activity, the
lowest pH and highest ionic strength of all the brines tested. There was a rapid and complete loss of
viability in around 100 minutes with both desiccated and hydrated biofilms in type IV brine (Figure
4).



87 Figure 1 – Viability of S. desiccabilis cells in desiccated and hydrated biofilms after periods of immersion in Type I brines,

88 measured by counting colony forming units (CFU). Each data point is a biological triplicate result, with standard error bars,

89 and trend lines are plotted as linear regressions of data from several systematic replicates.

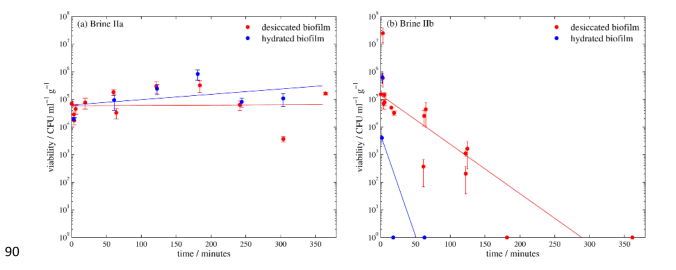


Figure 2 - Viability of S. desiccabilis cells in desiccated and hydrated biofilms after periods of immersion in Type II brines,
 measured by counting colony forming units (CFU). Where no CFUs were countable, results are plotted at 10⁰. Each data
 point is a biological triplicate result, with standard error bars, and trend lines are plotted as linear regressions of data from
 several systematic replicates.

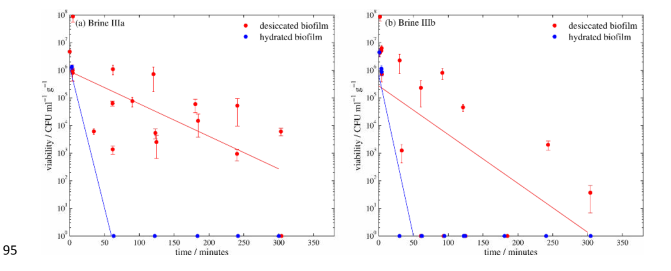
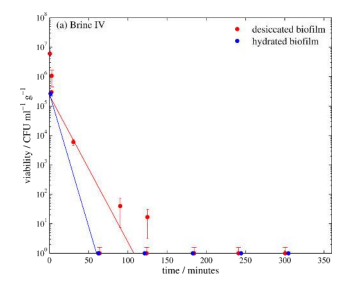


Figure 3 - Viability of S. desiccabilis cells in desiccated and hydrated biofilms after periods of immersion in Type III brines,
 measured by counting colony forming units (CFU). Where no CFUs were countable, results are plotted at 10⁰. Each data
 point is a biological triplicate result, with standard error bars, and trend lines are plotted as linear regressions of data from
 several systematic replicates.



100

Figure 4 - Viability of S. desiccabilis cells in desiccated and hydrated biofilms after periods of immersion in Type IV brine,
 measured by counting colony forming units (CFU). Where no CFUs were countable, results are plotted at 10⁰. Each data
 point is a biological triplicate result, with standard error bars, and trend lines are plotted as linear regressions of data from
 several systematic replicates.

Planktonic cultures of *S. desiccabilis* showed broadly similar responses to the 'weaker' brines but far stronger responses to the 'harsher' brines compared to biofilm cultures. There was no decrease in viability over several hours of immersion in brines Ia, Ib and IIa. In brines IIb, IIIa, and IV there was a total loss in viability within less than 10 minutes (the shortest timepoint measured). In brine IIIb it was not possible to recover any planktonic cells into a pellet, suggesting they had lysed within 10 minutes.

111 To provide a more quantitative comparison of viability in the different brines, we calculated a linear regression for the results of each experiment and measured the gradient. Table 2 shows the gradient 112 of these regression fits and the corresponding R² value to assess the quality of the fits. A larger 113 negative gradient implies a more rapid loss of viability, and a smaller R² value suggests more 114 115 variation in biofilm structure and growth between experimental replicates. The two instances of an R2 value of 1 are for experiments with only one measureable time point after the initial controls and 116 117 so the fit is only for two points. These values are graphed in Figure 5. Brines IIb, IIIa, IIIb and IV show 118 the most rapid loss of viability and there is a clear difference between desiccated and hydrated

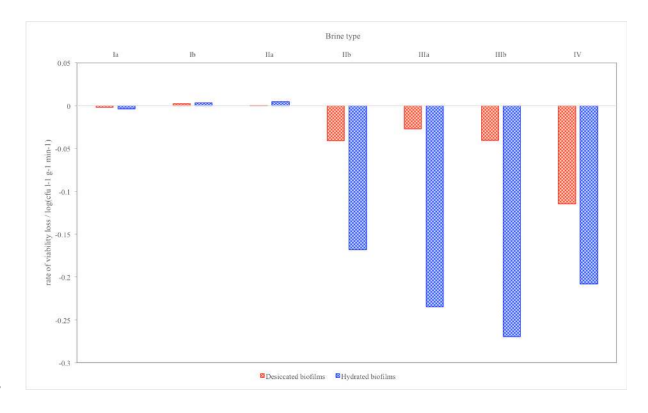
119 biofilms, with the latter losing viability ~2-10 orders of magnitude more rapidly when subjected to

these brines.

Brine	Rate of viability loss / log(cfu l ⁻¹ g ⁻¹ min ⁻¹)					
	Desid	ccated	Hydrated			
	Fit gradient	R ²	Fit gradient	R ²		
la	-0.0018	0.098	-0.0036	0.362		
Ib	0.0022	0.095	0.0033	0.210		
lla	0.0002	0.083	0.0045	0.301		
llb	-0.0411	0.770	-0.1679	0.643		
Illa	-0.0271	0.497	-0.2346	1.000		
IIIb	-0.0407	0.358	-0.2697	0.869		
IV	-0.1147	0.709	-0.2081	1.000		

121 Table 1 - Rate of viability loss of S. desiccabilis cells in desiccated and hydrated biofilms when immersed in Brines Ia to IV,

122 measured from linear regressions of the curves shown in Figures 2-5.





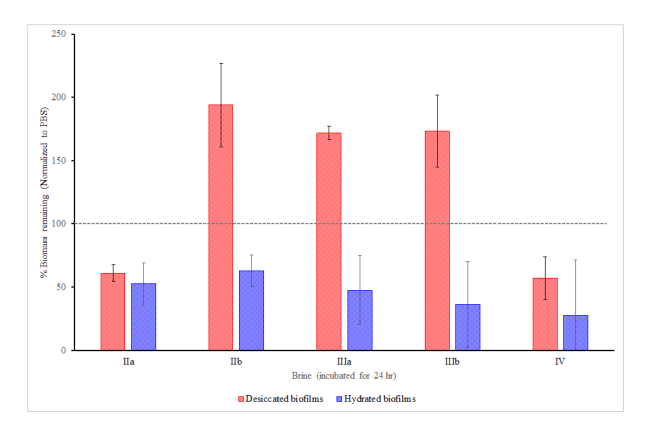
125 measured from linear regressions of the curves shown in Figures 1-4.

126 **3.2 Repeated Brining Cycles**

- 127 We also tested the viability of biofilms on scoria after repeated cycles of brining and desiccation,
- 128 repeating the process used in Section 3.1. Viability was tested after two to five cycles of brine
- immersion. CFU counts after the extraction procedure were negative for all cycles tested.

130 **3.3 BIOMASS ASSAY**

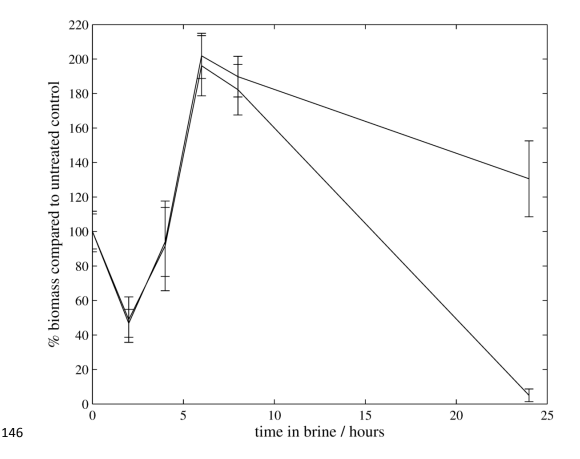
- 131 As Type I brines had little effect on the viability of cells in biofilms, these were not included in CV
- assays. Figure 6 shows the relative biomass present after a 24 hour incubation in the Type II, III and
- 133 IV brines for both desiccated and hydrated biofilms. All of the hydrated biofilms showed a reduction
- in biomass after 24 hours, suggesting the biofilms were broken down by the brines. In the case of
- the desiccated biofilms, there was also a reduction in biomass after 24 hours in brines IIa and IV, but
- in brines IIb, IIIa and IIIb, the desiccated biofilms showed an increase in presumptive biomass over 24
- 137 hours, by up to 250%.



138

139 Figure 6 – Percentage biomass remaining after immersion in brines IIa to IV for 24 hours for desiccated and hydrated

- 140 *biofilms. Error bars are calculated from biological and systematic triplicates.*
- 141 To investigate this increase in biomass, we also performed a time series using the same
- 142 experimental procedure, with brine IIb over another 24 hour period (Fig. 7). During incubation in the
- brine there was an initial reduction in biomass followed by an increase over several hours, which
- 144 peaked at around 6 hours after exposure to brine. After this increase, the biomass began to reduce,
- 145 from a peak of 200% to an average of around 100% after 24 hours.



147 Figure 7 – Biomass remaining over time for desiccated biofilms in Brine IIb. Two experiments were performed.

148 **4 DISCUSSION**

149 Other planetary bodies in the solar system harbour aqueous environments that differ markedly in 150 geochemical composition from aqueous environments on the Earth. These differences are a consequence of contrasting geological histories, which have produced the rocks and atmospheres 151 152 that ultimately determine water chemistry. 153 In this study we sought to address the hypothesis that as on Earth, specific adaptations of microbial 154 growth would influence the extent to which organisms could persist in extreme extraterrestrial 155 aqueous environments. In particular, biofilms are known to provide resistance to chemical extremes [e.g. 4,7]. These data have implications for planetary protection that is concerned with whether 156 terrestrial microorganisms could grow in extraterrestrial aqueous environments into which they are 157

inadvertently introduced and, more speculatively, the adaptations that would be required for

159 extraterrestrial life, if it exists, to grow in these alien geochemical extremes.

160 We found that combined physicochemical extremities of the brines had a significant effect on

161 microbial survival. Generally, those brines with high water activity, low ionic strength and moderate

162 pH show little effect on the viability of *S. desiccabilis* cells, whether grown in a planktonic state or in

a biofilm.

164 Brines with low pH and high ionic strength generally had a large impact on viability, making

planktonic cells unrecoverable in less than 10 minutes. Our data suggest that water activity is one

166 factor that influences the ability of biofilms to survive brine exposure. For example, brine IIIa has a

167 high water activity, low pH and high ionic strength and had a lesser impact on viability than brine IIb,

168 which has a similar similar pH and ionic strength, but lower water activity. This observation is

supported by the rapid loss of cell viability for planktonic cells and biofilms immersed in brine IIIb,

170 which has the lowest water activity of the brines tested.

171 However, factors other than water activity also appear to play a role in loss of viability in biofilms.

Brine IV caused a more rapid loss in viability than Brine IIb, although it has a higher water activity,

173 more moderate pH and lower ionic strength. Brine IV is the only solution with perchlorate ions,

174 which may suggest that these ions may be deleterious to cells, perhaps because of the oxidizing

175 capability of perchlorate [20].

176 The resistance to loss of viability caused by the extreme brines is likely to be caused by a

177 combination of physical protection and metabolic adaptation. Biofilms are known to provide a

barrier between cells and the external hostile environments [3,21], in this case providing protection

against the physicochemical extremes of the brine.

As shown by the more rapid reduction in viability measured with hydrated biofilms compared to
desiccated biofilms, the physical protection offered by the biofilm is not the only factor that allows

the cells to remain viable for longer in the brine. Desiccation adds yet another level of protectionagainst extreme brine exposure.

184 The crystal violet assay suggested that when desiccated, biofilms showed an increase in presumptive 185 biomass after several hours in the brine, a result not seen with hydrated biofilms and repeated 186 consistently in desiccated biofilms. One biological explanation for these observations might be that 187 desiccated biofilms are able to begin production of metabolites (such as membrane-stabilising 188 disaccharides) or additional EPS during initial exposure to osmotic stress. Production of EPS is a 189 known response to osmotic stress/desiccation [22]. Since matric (drying) water stress is equivalent 190 to osmotic water stress from high concentration solutes [22], by drying the biofilms before brining, 191 this process might act to 'prepare' the organisms by signalling the production of osmotic-resistance 192 metabolites (potentially changing the composition of the biofilm), which might continue during 193 exposure to brine prior to loss of viability. These observations merit further study. 194 Whilst our experiments show that under the most extreme brines, biofilms confer resistance to loss 195 of viability, we were unable to recover viable cells after 5 hours in the harsher brines (Brines IIb, IIIa, 196 IIIb and IV). We cannot rule out that other organisms or communities might be able to adapt to

these extreme conditions, but our results do not show long-term survival. Furthermore, we tested the hypothesis that in the desiccated state, biofilms would enhance survival in transient aqueous systems, such as those proposed to occur in Martian RSLs. Under repeated cycles of desiccation and exposure to brine, which simulated the conditions that might occur in RSL, we found that viability was completely lost after the first cycle in all cases.

We have shown that biofilms formed by microorganisms offer protection against the extreme conditions of Mars-analogue brine solutions. Cells in biofilms desiccated before being immersed in brines remained viable for orders of magnitude longer than planktonic cells or hydrated biofilms under the same conditions. Some of these brines had low water activity, low pH and high ionic content that would otherwise make them uninhabitable [15], but *S. desiccabilis* biofilms retained

207 viability in them for short periods of time, despite them quickly destroying S. desiccabilis cells in the 208 planktonic state. Even though S. desiccabilis is not an extreme halophile, nor is it known to possess 209 exceptional tolerance to osmotic shock, its biofilm forming capability provides resistance to high 210 levels of dissolved salts. It was selected here for this ability to form biofilms and known desiccation 211 tolerance. Halophilic biofilm-forming organisms that are exposed to geochemical regimes similar to 212 those investigated here would likely possess even greater resistance. 213 These results have implications for the designation of RSL as 'special regions' for planetary protection purposes [23]. Given that terrestrial spacecraft almost certainly carry a bioload, some 214

215 portion of which may be in biofilm form, even the transient aqueous environment suggested by RSL

216 could present an environment that hardy terrestrial microbes could exploit or leave contaminating

217 material in, especially if protected by a biofilm.

218 Finally, our results add to the expanding body of literature showing the protective properties of

219 biofilms in extreme environments. Prior studies have shown biofilms' capabilities to protect resistant

220 cells against osmotic stress, but not with brines with properties as different or as extreme as those

used in our study, such as concentrated sulfate brines. Although these brines are more relevant to

222 extra-terrestrial environments, they are also relevant to some terrestrial locations such as the

223 Basque lakes of British Columbia, which are sulfate-rich water bodies [24], or deep subsurface

groundwater systems [25].

225 **5 MATERIALS AND METHODS**

226 **5.1 STRAINS AND GROWTH CONDITIONS**

227 This study examined viability and cell biomass in the organism Sphingomonas desiccabilis (S.

228 desiccabilis), a Gram negative, rod-shaped, chemoheterotrophic, strictly aerobic bacterium originally

isolated from a soil crust in the Colorado Plateau, USA [26]. Biofilms were cultured on pieces of

- basaltic scoria (highly vessiculated Icelandic Iava (*Bar-Be-Quick Barbeque Products, Burnley, UK*) see
 Figure 8), chosen as an analogue for martian regolith and to provide a surface for the biofilm to form
 on. Overnight cultures of *S. desiccabilis* grown in R2A medium at 20°C [27], were normalised by cell
 density using OD₆₀₀ measurements and inoculated onto ~5 mm pieces of scoria at a concentration of
 50 µl in 4 ml of R2A medium and grown for one week at 20°C. Biofilm formation during this period
 was confirmed by visual inspection.
- 236



- 237
- Figure 8 Basaltic scoria used as a Mars analogue growth substrate a) larger piece and b) broken down to ~5 mm piece for
 use in well plates.
- After a week of growth, the supernatant was removed from the scoria. The biofilm-covered scoria pieces were then either a) dried in a laminar flow hood overnight or b) had their media replaced with PBS in order to halt growth but prevent desiccation. This allowed us to compare the response of desiccated biofilms to hydrated biofilms. $1-2 \times 10^7$ cells planktonic cells were collected from 500 µl of the same normalised overnight cultures (incubated with shaking to reduce biofilm growth) for planktonic brine exposure experiments.

5.2 Exposure to brines

247	The biofilm inoculated scoria was then treated with Mars-analogue brines over a time course. The
248	brines were synthesised based on computational reconstructions of putative martian brines by Tosca
249	et al. [28], as used by Fox-Powell et al. [15], and include four types at two stages in their evaporation
250	sequence (a and b). Type Ia and Ib are alkaline carbonate-chloride brines, which are analogous to the
251	conditions in Gale Crater approximately 3.7 billion years ago [29] and the fluids that interacted with
252	the Nakhla martian meteorite [30]. Type IIa and IIb are Mg-SO ₄ -Cl dominated brines, characteristic of
253	widespread Hesperian evaporite deposits such as those at Meridiani Planum [31]. Type IIIa and IIIb
254	are similar in composition to Type II but contain higher levels of dissolved iron, and are extremely
255	acidic.
256	In addition to the six Mars analogue brines, a magnesium perchlorate solution was also included
256 257	In addition to the six Mars analogue brines, a magnesium perchlorate solution was also included (Type IV), given observations of magnesium perchlorate in the martian polar regolith by the Phoenix
257	(Type IV), given observations of magnesium perchlorate in the martian polar regolith by the Phoenix
257 258	(Type IV), given observations of magnesium perchlorate in the martian polar regolith by the Phoenix lander [32] and more recently associated with RSL following spectral observations [16]. This brine
257 258 259	(Type IV), given observations of magnesium perchlorate in the martian polar regolith by the Phoenix lander [32] and more recently associated with RSL following spectral observations [16]. This brine was a solution of magnesium perchlorate with a concentration to give a water activity matching that

Brine type	la	lla	Illa	lb	llb	IIIb	IV
NaHCO ₃	0.126						
KHCO ₃	0.028	0.041		2.237			
КСІ	0.022	0.020	0.075	3.776	1.033	1.142	
MgCl ₂ .6H ₂ O	0.001	0.056			1.154	3.007	
NaCl		0.154	0.189	1.266	2.265	1.036	
MgSO ₄ .7H ₂ O		2.068	3.066		2.550		
FeSO ₄ .7H ₂ O			1.225			2.313	
FeCl ₂ .4H ₂ O			0.208			0.985	
HCI					0.038	0.113	
Mg(ClO4) ₂							3.097
Water activity	0.97	0.92	0.90	0.80	0.63	0.56	0.67
рН	8.2	6.2	1.8	8.3	2	0.6	5.5
Ionic Strength	0.18	8.5	12	5	11.6	13	9.29

	mol/liter								I
263	Table 2 - Salts added to mix	« Mars analo	gue brines. C	Concentration	ns are in mole	es litre ⁻¹ . Add	apted from F	ox-Powell et	al. [15

Table 2 - Salts added to mix Mars analogue brines. Concentrations are in moles litre ⁻¹. Adapted from Fox-Powell et al. [15]

and originally calculated from Tosca et al. [28].

265 Repeated brining was tested by removing the brines from the scoria after an hour. They were then

dried in a sterile flow hood until there was no visible surface moisture on the scoria, and the process

267 repeated.

268 **5.3 VIABILITY AFTER BRINING**

269 After being immersed in the brine, the scoria was washed twice in 1X PBS (Phosphate Buffered

270 Saline) and then suspended in PBS. The suspension was vortexed briefly and sonicated for two

271 minutes to extract cells from the biofilms. Planktonic cells that had been brined in the same manner

were centrifuged at 8 g for 10 minutes, the brine removed and the cells re-suspended in 1 ml of PBS.

273 Cell suspensions were plated onto nutrient agar plates in serial dilution and the CFUs quantified

after two days of growth at 20°C to give a measure of viability after immersion after a given amount

of time in the different brines. Results were collected for each brine for both dried and hydrated

biofilms in biological triplicate and systematic replicate at different time points.

277 5.4 DETERMINATION OF BIOMASS

We performed a crystal violet (CV) staining to investigate how the brines affect the overall biomass of the biofilms. Biofilms were grown using the same method as described above and stained using a protocol adapted from Childers et al. [33]. The biofilm-coated scoria were incubated in the different brines, or PBS as a control for 24 hours, and then washed three times in PBS. Each scoria piece was then stained with 0.4% crystal violet solution for 15 minutes at room temperature, washed three times with PBS and destained for 15 minutes in 1 mL of 33% acetic acid solution. The optical density of the destain solution was used to quantify the total biomass of biofilm left on each piece of scoria

- after brining, with the PBS incubation forming the 'zero brine' control. The OD₅₇₀ was measured by
- spectrophotometer with 33% acetic acid as the blank.

287

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