Report from Anonymous Referee #1

Manuscript BG-2023-51 manuscript: "Properties of exopolymeric substances (EPS) produced during cyanobacterial growth: potential role in whiting events."

1. Anonymous Referee #1

The authors conducted a culture-based study and used a complementary arsenal of carefully done measurements to verify their results and observations and infer how the precipitation of CaCO3 in pelagic cyanobacterial blooms may occur (during whiting events). FTIR, pH drift assays, EPS compositional analyses, etc. They were able to show that the precipitation of CaCO3 (calcite, and to a lesser extent vaterite) coincided with the magnitude of EPS production and the available functional groups on the EPS occurring in early stationary phase cultures. Larger precips were formed during the exponential phase, and smaller, more abundant precips were formed during the stationary phase.

The larger precipitates early on (Exponential phase cultures) and smaller precipitates observed in later stationary phase is somewhat puzzling. But interpretations were made that help to explain these outcomes, especially when considering natural bloom systems.

The "pH of cultures was around 10, and remained steady". Given that cultures were grown under 12/12 light/dark cycles, was pH measured in darkness? It should be clarified if pH was measured during light conditions or dark, or both. Please clarify?

Author's response: *Synechococcus* cultures were grown under a light/dark cycle of 12 hours each. However, pH measurements were exclusively carries out during the light cycle. The pH values were measured approximately 3-4 hours after the completion of the dark cycle. This explains the consistently high pH values depicted in Figure 1B. We have now incorporated this information into the main text (Line 113).

Figure 1. What may have caused the dip in both pH (B), and numbers of cells (A) during stationary phase (near day 40) in Experiment 1? Any suggestions?

Author's response: Indeed, in Experiment 1, we observed an atypical phenomenon where the cultures appeared to undergo a collapse around day 40. Both pH and cell density values decreased but started to increase again within approximately 4 days. One possible explanation for this unusual pattern could be a disruption or change in nutrient availability. This could include phosphorous (PO_4^{3-}) deficiencies or a transition to a different nitrogen (N) source (e.g., from NH_4^+ to NO_3^-). Another probable factor is the depletion or insufficiency of CO_2 during this specific phase of cultivation. The shift in the carbon source from CO_2 to HCO_3^- might trigger a metabolic response in the cells, leading to their re-adaptation, which in turn could influence cell growth and account for the observed pattern in Experiment 1. It is important to note that this explanation is merely a supposition based on information gathered from the literature (e.g. Rückert et al., 2004) and that further measurements would be necessary to confirm this hypothesis. Nevertheless, when comparing pH values and cell numbers between Experiment 1 and Experiment 2, no significant differences were found (p-value > 0.05, as shown in Figure AA and AB).

Anova: Single Factor										
SUMMARY										
Groups	Count	Sum	Average	Variance					р	Н
Column 1	6	58,48292	9,747153	0,923947					Column 1	Column 2
Column 2	6	58,845	9,8075	0,735155				Time (day	Experiment 1	Experiment 2
								0	8,24	8,58
								3	8,94	8,87
ANOVA								14	10,28	10,10
Source of Variation	SS	df	MS	F	P-value	F crit		28	10,45	10,28
Between Groups	0,010925	1	0,010925	0,01317	0,910906	4,964603		41	9,89	10,50
Within Groups	8,29551	10	0,829551					56	10,70	10,52
Total	8.306435	11								
	-,- ,- ,- ,- ,- ,- ,- ,- ,- ,- ,- ,- ,-									

Figure AA. Anova Single-factor statistical test comparing pH values obtained from growth experiments 1 and 2.

Anova: Single Factor										
SUMMARY							 		cell density	y (cells.L-1)
Groups	Count	Sum	Average	Variance					Column 1	Column 2
Column 1	6	58,48292	9,747153	0,923947				Time (day	Experiment 1	Experiment 2
Column 2	6	58,845	9,8075	0,735155				0	9,52E+10	7,12E+10
								3	2,58E+11	2,03E+11
								14	1,74E+12	5,65E+11
ANOVA								28	2,10E+12	1,48E+12
Source of Variation	SS	df	MS	F	P-value	F crit		41	1,81E+12	1,24E+12
Between Groups	0,010925	1	0,010925	0,01317	0,910906	4,964603		56	2,69E+12	1,44E+12
Within Groups	8,29551	10	0,829551							
Total	8,306435	11								

Figure AB. Anova Single-factor statistical test comparing cell numbers obtained from growth experiments 1 and 2.

Using FTIR, highest protein levels (line 256, 257) were indicated, and later using colorimetric assays of protein (line 271) it is stated that highest protein occurred in EPS also during early stationary phase (also shown in Table 3) – good verification!

Author's response: Thank you.

The FTIR results for EPS are especially informative and helpful. The authors should consider summarizing these in a separate Table for easier reference by the reader.

Author's response: We conducted an FTIR analysis of the EPS extracted at various *Synechococcus* growth stages and included the corresponding results in the supplementary materials document as Table 1S.

Although this is a laboratory-based study, it sheds light on a longer standing issue of how whiting events occur during blooms in natural systems. The authors are to be commended on the nice, careful work examining this whiting-related process. The ms was well written, and only minor changes are suggested.

Author's response: We thank the reviewer for the positive feedback.

Report from Referee #2

Manuscript BG-2023-51 manuscript: "Properties of exopolymeric substances (EPS) produced during cyanobacterial growth: potential role in whiting events."

2. <u>Referee #2: Dr. Tobias-Hunefeld, Sven</u> sven.hunefeldt@igb-berlin.de

Summary:

The authors investigated the role of EPS in carbonate precipitation during a Synechococcus bloom experiment. EPS characteristics were assessed with FT-IR spectroscopy, SDS-PAGE and forced carbonate mineral precipitation experiments. The authors separated the bloom into exponential, and early and late exponential stages. Stationary phases contained higher abundances of negatively charged groups than during the exponential phase, leading to smaller (< 50 um) but more abundant carbonate minerals in the stationary phases, whereas the exponential phase contained larger (> 50 um) but less abundant carbonate percipitates.

I believe this manuscript gives a small but important contribution to the field. I especially applaud the many different approaches taken to measure the EPS and associated factors. My comments mainly center around cutting down on the amount of references, only using the key ones, and ensuring that the different sections (introduction/results/discussion) don't intersect with each other. Other comments include ways to convey data reliability and acknowledging limitations.

Major comments:

The abstract should make clearer what techniques were used for measurements, at least the ones that the major findings were drawn upon.

Author's response: Thank you for this remark. We have followed your suggestion and this information is now added to the abstract (Lines 21-23).

The graphical abstract is well made and explains the stated findings well. However, I am unsure of what the 'Negatively charged groups' are, is it the connecting lines of the different phases?

Author's response: The negatively charged groups are represented by the "connecting lines" and resemble a honeycomb-like structure.

In that case I would remove the Exponential/Early stationary/Late stationary illustrations and explanations from the Legend.

Author's response: We acknowledge the reviewer's observation regarding the repetition of the terms "Exponential/Early stationary/Late stationary phase." In response, we have taken their suggestion and eliminated them from the legend and decided to keep it solely as subtitles in the Graphical abstract.

There are many references throughout the introduction and discussion, to the degree that the text in the intro is 50% references and 50% informative text. I would focus more on relevant and landmark references and use these key references to make your point. Rather than having 3-5 references for every statement. For example: L83-84: There are 3 references that say that cyanobacteria are known EPS producers, and then another 2 that state this is especially true during blooms. It would be best to have only those last two as they are the most relevant to your point.

Author's response: After thorough examination, we carefully reviewed the entire manuscript once again. As a result, we have made the decision to retain only the three most relevant references for each statement. We acknowledge the reviewer's point that certain subsections contained an excessive number of citations. However, it is important to emphasize that is a lab-based study in which we present a [theoretical] model that we have extrapolated to the natural system. To effectively integrate our findings with the natural system scenario, it is imperative to access wide range of disciplines and explore relevant literature.

Methods seem to match the stated goals of the study. Many measurements however were done in duplicate, with no justification as to why. Such as protein, sugar and glycosaminoglycan quantifications. Unless measurements were extremely similar, triplicates would convey a greater degree of measurement precision. I would recommend either measuring an additional sample (if available), stating the reason for why duplicates were chosen for this, or acknowledging this as a study limitation.

Author's response: Protein measurements were conducted in quadruplicate, sugar measurements in triplicate and GAGs (glycosaminoglycans) measurements in duplicate. The reported values in the study represent the mean of these measurements taken from two replicates of EPS samples. Therefore, the reported averages were obtained from a total of eight protein measurements, six sugar measurements and four GAG measurements. We have now included this information in the main text at lines 265-266.

As there were two experimental runs that were temporally displaced, I would have liked to see a brief comparison of their measurements to show that these two test runs are not significantly different.

Author's response: Comparison of pH and cell number measurements of Experiment 1 and Experiment 2 showed that they are not significantly different (p-value > 0.05, see Figure AA and Figure AB). This information is now included in the main text (Lines 192, 198).

Anova: Single Factor										
SUMMARY										
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Figure AA. Anova Single-factor statistical test comparing pH values obtained from growth experiments 1 and 2.

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SUMMARY									cell densit	y (cells.L-1)
Groups	Count	Sum	Average	Variance					Column 1	Column 2
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Within Groups	8,29551	10	0,829551							
Total	8,306435	11								

Figure AB. Anova Single-factor statistical test comparing cell numbers obtained from growth experiments 1 and 2.

Additionally, is there an explanation why experiment 1 had a 7 day and experiment 2 had a 14 day exponential phase?

Author's response: Although the inoculum was consistent between Experiment I and II (we calculated the same final optical density), the initial cell count on day 0 was ~ 1.3 times higher in Experiment I (9.52 × 10^{10} cells. L^{-1}) than in Experiment II (7.12 × 10^{10} cells. L^{-1}). This difference in cell density could potentially account for the faster cell growth and shorter exponential phase observed in Experiment I.

With the difference in exponential phase, why were the chosen sample days the same for both experiments? Specifically: D0, D14, D8, D56.

Author's response: In our previous publication (Martinho de Brito et al., 2022), we demonstrated that the exponential growth phase of *Synechococcus* spp. lasts for a minimum of 18 days. Based on this finding, we selected day 14 as the first sampling point in the current study, representing the mid of the exponential phase. The choice of the second sampling point was based on the observed stability in cell numbers and pH values in Experiment 1, suggesting the transition from the exponential phase to the stationary phase. Thus, we selected day 28 for the second sampling point. Initially, we anticipated sampling during the death phase for the final time point. However, as stated in the main text, we did not observe a distinct death phase during the 56-day experiment. Consequently, we collected the last sample during the late stationary phase. For Experiment II, we established the same sampling time to correlate the obtained data with EPS production. Although the number of cells may not be the same in both experiments, the growth stages of the cultures overlap significantly (p-value = 0.91).

By picking only crystals above 10 µm, could this be introducing some bias? Or was the 10 µm limit chosen due to methodological limitations? If yes, should be stated. Otherwise, the 10 µm size cut off should be justified.

Author's response: Based on the observations made during the precipitation experiment, we identified two primary size classes. To analyze these size classes, we used an image analysis software and selected two crystal size classes: crystals with a size $< 50 \mu m$ and crystals with a size $> 50 \mu m$. It is important to note that the $< 50 \mu m$ size class encompasses all crystals ranging from $0 \mu m$ to $< 50 \mu m$, which includes crystals smaller and larger than 10 μm . Therefore, crystals above 10 μm were counted and their counts are included in the reported results. Please refer to the "Crystal count and size distribution" (Subsection 2.6.2) in the Materials and Methods section for a more detailed explanation of this counting process.

Please add confidence intervals/standard deviations to Figure 5.

Author's response: Each one of the curves shown in Figure 5 represents pH values that were measured at a frequency of every 2 seconds, resulting in a minimum of 400 measurements per curve. Due to the large number of measurements and to maintain the clarity of the graph, we opted not to include the standard deviation (SD) for each curve. Including the SD in the graph would render it unreadable and overcrowded. However, it is important to note that the general trend is preserved. We have included a supplementary Figure (Figure 2S) that displays the replication of the *in vitro* inhibition of calcium carbonate precipitation, demonstrating the consistent pattern observed in the main figure.

I would also like to see the statistics represented on all relevant figures. Between negative controls and measured values, as well as between time points.

Author's response: After a careful examination of all the tables and figures of the manuscript, we confirmed that only Table 1 was lacking the addition of standard deviations (SD). We have now corrected this error and added the necessary SD values to Table 1. Please refer to Table 1 (Lines 213-215) in the manuscript for this updated information.

Please increase the readability of the introduction and discussion, lots of results were included and some points are made and justified either in the next discussion section or 20 lines below. Based on this there were some other points to make:

As EPS characteristics are phytoplankton/organism specific it should be noted that these findings need to be expanded on with additional studies and other phytoplankton.

Author's response: We acknowledge the reviewer's valuable input regarding this significant aspect. As a result, we have included the relevant information in the Discussion section of the manuscript (Line 476).

Tables and Figures that were already presented in the results section should not be mentioned again in the discussion.

Author's response: We have implemented changes throughout the entire text to address this concern. Specifically, we have removed several references to tables and figures from the discussion section, as suggested by the reviewer.

The same is true for the discussion section, do not reiterate results in this section, but rather focus only on the implications and what this means for the field.

Author's response: We have made revisions and modifications to the Discussion section as required.

Ensure that conclusion drawn from the data are found only in the discussion, and not in the results. Such as L276, L278-279.

Author's response: We have removed these sentences from the Results section and that information is now included in the Discussion section.

Please ensure that you do not discuss or provide justification for the chosen methods in your results, but rather in the introduction, methods section, or discussion (as relevant).

Author's response: We have changed the text accordingly.

Please make clear how discussion L381-399 relates to whiting events. The text is very information dense, however how it relates to phytoplankton associated whiting events is not clearly stated.

Author's response: We agree that there is significant repetition within this paragraph. As some of this information was already presented in the Introduction section, we have condensed the paragraph into three lines for brevity and clarity. Please refer to Lines 451-454 for the summarized version.

I believe it would help to have L400-417 before L381-399 or remove L381-399 from the manuscript.

Author's response: After careful consideration, we have decided to retain the original version of Lines 381-399. We believe that this section presents the narrative in a logical and comprehensive manner. Our aim was to establish a connection between cell growth/metabolic activity (high photosynthetic activity and high pH levels and alkalinity, as well as CO₂ limitation) with EPS production, exploring its impact on carbonate precipitation. The same structured approach was consistently applied throughout the discussion of all *Synechococcus* growth stages, as outlined in Sections 4.1, 4.2, and 4.3.

As EPS characteristics are phytoplankton/organism specific it should be noted that these findings need to be expanded on with additional studies and other phytoplankton.

Author's response: We acknowledge the reviewer's valuable input regarding this significant aspect. As a result, we have included the relevant information in the Discussion section of the manuscript (Line 476).

Minor comments (please also see attached pdf with comments):

The limit of detection was stated in µg.L⁻¹ in text, however Table 1 values are in µM. The same units should be used throughout the manuscript.

Author's response: The limit of detection was modified and is now expressed in µM (Line 209).

I would also be interested in day 0 values throughout the text, as it was stated that they were measured but are not shown in the manuscript.

Author's response: We agree that this was confusing as stated and we have rectified it by amending the caption of Table 1 (Line 212). The measurements were done in the medium before the inoculation (t0). Therefore, in Table 1, t0 corresponds to the column entitled "Initial concentrations in the medium".

Only Figure 2 is required. Table 2 should then be relegated to the supplementary.

Author's response: We prefer to keep Table 2 and Figure 2 in the main text as they complement each other. The calculations of cell-specific EPS production described in Figure 2 were derived from the data presented in Table 2, which includes information on cell yield and EPS production at the three different growth stages. By including both Table 2 and Figure 2, we aim to enhance the clarity and comprehension of the study's findings.

FT-IR mentions a lot of different spectra, it would be best to only focus on the ones from the discussion or those relevant to the findings.

Author's response: We strongly believe that providing a comprehensive description of the Infra-Red spectra is highly useful for a deeper comprehension of the EPS, including its composition and structure. To our knowledge, there is still limited availability of this type of information for exopolymeric substances. Therefore, we emphasize the importance of highlighting and describing all the peaks in the spectra. Additionally, we would like to emphasize that the other reviewer of this paper suggested moving Table 1S "Attribution of main infrared absorption bands of EPS samples" from the supplementary materials to the main text due to its relevance. The reviewer found this information highly pertinent, further reinforcing the significance of including it in the main text.

Can Figure 4 be redone with the inclusion of a negative control?

Author's response: Usually the use of Alcian Blue (AB) does not require a negative control since it is not expected to stain non-polyanionic substances such as proteins and sugars. This is a commonly accepted practice, and several studies examining polyanionic matrices on gels have been published without including a negative control (Marie et al., 2007; Gaspard et al., 2008). There are instances where non-polyanionic matrices exhibit no staining at all with AB, as demonstrated by Pavat et al. (2012). Moreover, in our gel, the lane corresponding to the molecular weight (MW) ladder can be considered as an appropriate negative control.

Properties of exopolymeric substances (EPS) produced during cyanobacterial growth: potential role 1

in whiting events 2

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- 9 ⁴Department of Marine Sciences, The University of Connecticut, Groton, CT 06340, USA
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- 12 13

14 Abstract 15

- Extracellular polymeric substances (EPS) are an important organic carbon reservoir in many pelagic and benthic environments. The production of EPS is intimately 16
- 17 associated with the growth of phyto- and picoplankton. EPS plays a critical role in carbonate precipitation through the binding of cations and by acting as a nucleation
- 18 site for minerals. Large-scale episodes of fine-grained calcium carbonate precipitation in the water column (whiting events) have been linked to cyanobacterial blooms,
- 19 including of Synechococcus spp.. The mechanisms that trigger these precipitation events are still debated. We pose that the cyanobacterial EPS, produced during
- 20 exponential and stationary growth phases plays a critical role in the formation of whitings. The aim of this study was to investigate the production of EPS during a
- two-month cyanobacterial growth, mimicking a bloom. The production and characteristics of EPS were examined in different growth stages of Synechococcus spp. 21
- 22 using various techniques such as FT-IR spectroscopy, colorimetric and SDS-PAGE assays. We further evaluated the potential role of EPS in carbonate precipitation
- 23 through in vitro forced precipitation experiments. EPS produced during the early and late stationary phase contained a larger amount of negatively charged groups
- than present in EPS produced during the exponential phase. Consequently, a higher Ca²⁺ binding affinity of the stationary phase-EPS led to the formation of a larger 24
- 25 amount of smaller carbonate minerals ($\leq 50 \mu m$) compared to crystals formed in exponential phase-EPS, which were less abundant and larger ($\geq 50 \mu m$). These findings
- were used to establish a conceptual model for picoplankton bloom-mediated CaCO₃ precipitation that can explain the role of EPS in whitings (see graphical abstract). 26
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1. Introduction 32

33 1.1 Significance of this study

- 34 Massive carbonate precipitation episodes in the water column, also referred to as 'whiting events' are a well-known phenomenon of modern freshwater (Schultze-35 Lam et al., 1997; Hodell et al., 1998; Stanton et al., 2021) and marine environment (Shinn et al., 1989; Robbins and Blackwelder, 1992; Larson and Mylroie, 2014).
- 36 Whitings are caused by large-scale precipitation of micron-sized calcium carbonate particles (visible from space) and represent a major sink in the carbon cycle. The
- 37 particles associated with whitings can make up a major sedimentary constituent of the modern-day and ancient carbonate rock records (Pomar and Hallock, 2008).
- 38 Whiting events can be triggered by a combination of biological and physicochemical processes. Among the biological mechanisms that have been studied in this
- 39 context, picocyanobacterial proliferations have often been invoked in the initiation of whitings (Hodell et al., 1998; Thompson, 2000; Obst et al., 2009). Photosynthesis
- 40 increases pH levels and alkalinity during cyanobacterial blooms, ultimately causing the saturation state of calcium carbonate to rise, thereby leading to its potential
- precipitation. The role of Synechococcus spp. bloom-forming cyanobacteria in CaCO₃ precipitation has been demonstrated in laboratory experiments (Yates and 41
- 42 Robbins, 1998; Dittrich et al., 2003; Obst et al., 2009; Bundeleva et al., 2014; Martinho de Brito et al., 2022) and observed in field investigations (Wells and Iling,
- 1964; Thompson et al., 1990; Dittrich and Obst., 2004). Change in temperature, salinity, CO2 pressure as well as turbulence are some of the physicochemical factors 43
- 44 that can lead to the formation of supersaturated solutions and subsequent precipitation of CaCO₃ thus initiating the whiting. Even though several possible biogenic
- and abiotic mechanisms have been identified, the formation of whitings is still poorly understood. 45
- 1.2 Overview of phytoplankton blooms 46

Phytoplankton blooms, including those of picoplankton, are dense accumulations of cells resulting in a visible discoloration of the surface water layers (Reynolds and 47 Walsby, 1975; Huisman et al., 2018). Their occurrence has been recorded worldwide in marine and freshwater bodies (Paerl et al., 2001; Paerl and Huisman, 2008; 48 Ploug, 2008). Light intensity, water temperature, nutrient availability, weather conditions and hydrodynamics are key factors that determine the onset and persistence 49 of a bloom. Blooms are typically seasonal, frequently observed during late spring or summer, and can be dominated by picoplankton (Huisman et al., 2018). Some 50 phytoplankton organisms, notably cyanobacteria, may produce toxins and form large-scale harmful algal blooms (Paerl et al., 2001). The intensity and frequency of 51 52 cyanobacterial blooms have been increasing due to anthropogenic eutrophication (Heisler et al., 2008; O'Neil et al., 2012), a trend expected to exacerbate due to 53 climate change (Lürling et al., 2018). Cyanobacteria comprise a diverse group of photoautotrophic organisms that play a pivotal role in global primary production and 54 are key players in the biogeochemical cycles of carbon, nitrogen and oxygen (Callieri and Stockner, 2000; Raven et al., 2017). The unicellular cyanobacterium Synechococcus is one of the most abundant photosynthetic microorganisms on Earth (Whitton and Potts, 2012), which contribute substantially to the picoplankton 55 community in marine (Murphy and Haugen, 1985; Coello-Camba and Agustí, 2021) and freshwater environments (Weisse, 1993) that can form dense blooms 56 57 (Schultze-Lam et al., 1992; Phlips et al., 1999; Dittrich and Obst, 2004).

1.3 Phytoplankton blooms and CaCO₃ precipitation 58

- During the occurrence of dense phytoplankton blooms, high rates of photosynthetic activity lead to a rapid depletion of CO₂ in the surface waters, increasing alkalinity. 59
- Depending on the buffering capacity of the water, this could result in pH values ≥ 9 to as high as 11 (Ibelings and Maberly, 1998; Zepernick et al., 2021). Consequently, 60

61 the inorganic carbonate equilibrium shifts towards carbonate (CO_3^{2-}) . Some cyanobacteria possess a carbon concentrating mechanism (CCM) that converts HCO_3^{-1} to

62 CO₂ through the action of carbonic anhydrase enzymes (Price et al., 1998; Badger et al., 2002) and produce hydroxide ions (Kupriyanova and Pronina, 2011). The

63 activity of extracellular carbonic anhydrase (eCA) may contribute to the create an alkaline microenvironment in the extracellular polymeric substances (EPS)

64 surrounding the cyanobacterial cells (Price et al., 2002; Dupraz et al., 2009). When OH⁻ ions are released during photosynthesis it causes the pH to rise, which favors

65 carbonate mineral precipitation, assuming there are enough calcium ions available (Kamennaya et al., 2012). Consequently, during blooms, carbonate minerals can

66 form on EPS or precipitated in the microenvironment surrounding cyanobacterial cells.

67 **1.4 The role of EPS**

Cyanobacteria are known producers of EPS, especially during blooms (Pannard et al., 2016; Liu et al., 2018). EPS serve as a boundary between cells and their 68 69 immediate environment (Whitton and Potts, 2012) and may act as a template for CaCO₃ nucleation (Dupraz and Visscher, 2005; Dupraz et al., 2009; Kamennaya et 70 al., 2012). EPS are high molecular weight organic molecules composed of polysaccharides, proteins, nucleic acids and lipids (Pereira et al., 2009; Marvasi et al., 2010; 71 Decho and Gutierrez, 2017). This complex mixture of molecules may contain specific monomer components, such as uronic or sialic acids (monosaccharides), aspartic or glutamic acids (amino acids) or functions (sulfate, phosphate), which carry negative charges in physiological conditions and can therefore bind cations, such as 72 73 Ca²⁺, and promote the nucleation of CaCO₃ crystals (Trichet and Defarge, 1995; Dupraz et al., 2009; Walker et al., 2019). Conversely, polyanionic EPS in solution 74 can inhibit crystal growth by poisoning the faces of growing nuclei by an adsorption mechanism, according to a classical and accepted view prevailing for other macromolecules of similar charge properties: synthetic peptides (Wheeler et al., 1991), skeletal proteins (Wheeler et al., 1981; Addadi and Weiner, 1985), coccolith-75 76 associated polysaccharides (Borman et al., 1982) or natural organic matter dissolved in seawater (Mitterer and Cunningham, 1985). The production and composition 77 of EPS differ among different species of microorganisms and their type of metabolism and depend on environment in which they live, stressors (e.g., nutrient availability, pH, temperature, light, salinity) and the stage of their growth (Pereira et al., 2009; Pannard et al., 2016; Martinho de Brito et al., 2022). The deprotonation 78 of functional groups at elevated pH enhances the binding capacity of cations such as Ca^{2+} and controls crystal nucleation and growth by reducing the interfacial energy 79 barrier between the crystal and the EPS substrate (Dupraz et al., 2009; Dittrich and Sibler, 2010). EPS play a two-fold role in carbonate formation by initially inhibiting 80

81 (through Ca²⁺ binding) and subsequently promoting carbonate precipitation by releasing calcium ions during EPS alteration and degradation (Dupraz and Visscher,

82 2005. Furthermore, through specific functional group composition and structural architecture, EPS may also exert control over the mineralogy, morphology and/or

83 abundance of the minerals that are formed (Trichet and Defarge, 1995; Dupraz et al., 2009).

84 **1.5 The goal of this study**

We have previously reported that the pH of Synechococcus cultures increased when grown in a non-buffered medium (Martinho de Brito et al., 2022). In these growth conditions, the production of EPS was enhanced compared to growth in a buffered medium. Furthermore, the EPS from cells grown in non-buffered conditions contained more negatively-charged functional groups that impacted the properties of the carbonate minerals that precipitated. The current study further investigates

the properties of EPS produced during different growth phases of Synechococcus spp. Over an extended incubation time (mimicking a prolonged natural bloom). We

89 aim to better understand the role of cyanobacterial blooms in carbonate precipitation through EPS production and develop a conceptual model of picoplankton-

90 mediated organomineralization to explain the biological origin of whiting events.

91 **2. Materials and Methods**

92 2.1 Synechococcus PCC7942 strain and culture growth conditions

93 Synechococcus PCC7942 was obtained from the Centre de Ressources Biologiques de l'Institut Pasteur (Paris). Cultures were grown in a one-third-strength, non-

 $buffered \ liquid \ BG-11 \ medium \ (Allen, 1968; Rippka \ et al., 1979). \ The \ medium \ consists \ of \ (per \ liter): 1.5 \ g \ of \ NaNO_3; 0.04 \ g \ of \ K_2 HPO_4 2H_2 O; 0.075 \ g \ of \ MgSO_4 7H_2 O; 0.075 \$

95 0.036 g of CaCl₂2H₂O; 6 mg of citric acid combined with 6 mg of ferric citrate; 0.001 g of Na₂EDTA2H₂O and 0.02 g of Na₂CO₃. Trace metal solutions contained

96 (per liter) 2.86 mg of H₃BO₃; 1.81 mg of MnCl₂4H₂O; 0.222 mg of ZnSO₄7H₂O; 0.39 mg of Na₂MoO₄2H₂O; 0.079 mg of CuSO₄5H₂O and 0.0494 mg of

97 $Co(NO_3)_26H_2O$. Cultures were incubated at room temperature (21°C±2), in a light/dark cycle of 12h/12h under 36.8 μ E m⁻² s⁻¹ of photon irradiance while shaken at

98 200 rpm in a Cimarec i Multipoint Stirrer, 6 Position, 2000 rpm, 3L per Multipoint, 100-240 VAC rotary shaker.

99 2.2 Experimental design of Synechococcus-bloom formation

- 100 Two independent growth experiments were performed in 1L glass serum bottles containing 800 mL of 1/3 BG-11 medium adjusted to pH 7.5, sealed with silicone caps
- 101 to allow gas exchange. Cells used for the inoculum (pH = 9.2) were pre-cultured in a full-strength BG-11. Immediately after inoculation (30 mL/bottle), the pH
- 102 increased to approximately 8.2.

103 2.2.1 Experiment I

- In the first growth experiment, six bottles were inoculated with Synechococcus PCC7942. Cell growth and EPS production were examined. Optical density (OD_{750nm}),
 pH and cell counts were monitored weekly (2-3 times by week). EPS was extracted on days 14, 28 and 56 of cultivation (two bottles were harvested at each sampling
 time).
- 107 **2.2.2 Experiment II**
- 108 The second growth experiment was performed in quadruplicate. Chlorophyll a (Chla), extracellular carbonic anhydrase activity (eCA), nutrients (NO₃⁻ and PO₄³⁻) and
- calcium concentration were analysed at 0, 14, 28 and 56 days of cultivation. pH values, OD and cell counts were also assessed at longer intervals (once per week) than
 in Experiment I.

111 2.3 Growth assessment

2.3.1 pH values, optical density (OD) and cell counts 112

- 113 The pH value was measured about 3-4 h after the light cycle started with a CRISON GLP 21 pH meter (Crison Instruments SA, Alella, Spain). Cell growth was
- 114 monitored through cell counts and OD₇₅₀ measurements. Cell counts were performed using a counting chamber (Neubauer, Mariangela, Germany) by randomly
- 115 selecting five fields of view and counting approximately 100-200 cells. The OD at 750 nm of a 1-ml sample of the culture was measured in a Bio-Rad SmartSpec Plus
- Spectrophotometer (Bio-Rad, Hercules, CA, USA). 116

2.3.2 Chlorophyll-a extraction 117

- Chla was extracted from 2 ml culture aliquots using a methanol extraction method (Stal et al., 1984). Following the extraction in the dark at 4 °C, samples were 118
- 119 centrifuged. The Chla absorbance was measured in the supernatant at 665 nm using a Bio-Rad SmartSpec Plus Spectrophotometer (Bio-Rad, Hercules, CA, USA).

120 2.3.3 Extracellular carbonic anhydrase activity

- 121 The extracellular carbonic anhydrase (eCA) activity was measured using a BioVision Carbonic Anhydrase Activity Assay Kit Kit (BioVision, Ref. K472-100, Abcam,
- 122 Waltham, MA, USA) according to the manufacturer's specifications. Aliquots of ~ 5 ml were analysed immediately after the collection. To avoid cell lysis and
- intracellular CA contamination, samples were not centrifuged. The cells were separated from the supernatant by using a 1 mL syringe and a 0.20 µm NALGENE® 123
- 124 syringe filter. The absorbance was measured in a Bio-Rad Model 680 Microplate Reader at 405 nm.

2.3.4 Nitrogen, phosphorus and calcium measurements 125

- Phosphate, nitrate and calcium concentrations were determined in the growth medium at 0, 14, 28 and 56 days of cultivation. Cells were removed by centrifugation 126
- 127 and filtration through a 0.20 µm Millipore filter under a mild vacuum. The samples were stored at 4°C in the dark until measured by ion chromatography. Analyses
- were realized within the PEA²t technical platform of the Chrono-Environment Laboratory UMR6249 (Université de Franche-Comté, Besançon, France) and the Ca²⁺ 128
- 129 concentration was determined by ICP-AES (dual axial and radial view iCAP Pro XP model with fast loop, Thermofisher Scientific, Courtaboeuf, France) available at
- the University of Franche-Comté, Besançon, France. 130

131 2.4 EPS extraction and purification

- EPS were extracted from the Synechococcus cultures as previously described by Martinho de Brito et al. (2022). EPS were harvested after 14, 28 and 56 days of 132
- cultivation. Cyanobacterial cells were inspected by microscopy to ensure that no cell lysis had occurred during the extraction process. The pure EPS fractions were 133
- 134 obtained by ultrafiltration (>10 kDa = retentate) for volume reduction and the weight of the material was determined following by dialysis (using a 1 kDa Membrane)
- 135 lyophilization on a high-precision analytical balance (Quintix 35-1S, Sartorius, Gottingen, Germany).

2.5 EPS characterization 136

2.5.1 Fourier Transform-Infrared Spectroscopy 137

- FT-IR spectra were obtained from freeze-dried EPS on an FT-IR Bruker Alpha spectrometer (Bruker Optics SARL, Marne la Vallée, France) fitted with an Attenuated 138
- 139 Total Reflectance (ATR) ALPHA-P device equipped with a mono-reflection diamond crystal. A total of 24 scans were performed on each sample at a spectral
- 140 resolution of 4 cm⁻¹ in the 4000–375 cm⁻¹ wavenumber range. The qualitative assignment of absorption bands was performed by comparison with spectra available in
- 141 the literature (Coates, 2000).

142 2.5.2 Protein, sugar and glycosaminoglycan [quantification]

- The total protein content of EPS was determined using the Bicinchoninic acid assay (Pierce® BCA Protein Assay Kit) and bovine serum albumin as the standard. The 143
- 144 total sugar content was determined by a modified phenol-sulfuric acid method (Dubois et al., 1956) and xanthan and dextran were used as standards (Sigma-Aldrich,
- St. Louis, MO, USA). The total glycosaminoglycan (GAGs) content was quantified using the Blyscan Assay according to the manufacturer's protocol (Blyscan Kit 145
- 146 B1000, Biocolor Ltd., Antrim, UK) with chondroitin sulphate as the standard. All assays were carried out in duplicated EPS samples.

2.5.3 Visualization of polyanionic macromolecules on Alcian Blue stained gels 147

Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) followed by Alcian Blue staining (Wall and Gyi, 1988) were used to separate and to stain 148 negatively charged macromolecules (10- > 170 kDa), respectively. Alcian Blue is a dye that specifically binds to glycoconjugates with an acidic character (e.g., 149 150 containing carboxylated or sulfated functional groups). Samples were analysed on one-dimensional precast gradient protein gels (TGX Gel 4-15%, 90 mm x 70 mm) 151 on a Mini-Protean 3 cell (Bio-Rad, Hercules, CA, USA), according to the method previously described by Martinho de Brito et al. (Martinho de Brito et al., 2022). Prior to migration, samples were heat-denatured in standard 2x Laemmli sample buffer (5 min., 99°C, ref. 1610737, Bio-Rad). A pre-stained protein ladder 152 (Euromedex, #06P-0111; MW: 10 kDa to > 170 kDa) was used as a reference. 153

154 2.5.4 Inhibitory effect of EPS using pH-drift assay

155 The capacity of negatively charged functional groups in EPS to inhibit the in vitro precipitation of calcium carbonate was tested with the pH-drift assay (Wheeler et al., 1981; Marin et al., 2000; Kawaguchi and Decho, 2002). This assay was performed as previously described by Martinho de Brito et al. (2022). Briefly, the pH was 156 recorded by a pH meter (Laboratory Research Grade Benchtop pH/mV Meter with 0.001 pH Resolution-HI5221) connected to a PC via a USB cable. Data were 157 recorded by the HANNA HI92000 software. The pH was measured every two seconds for ~15 min. The shape of the curve (after reaching its maximum, about one 158 minute after T0) reflects directly the inhibitory capacity of the tested EPS: a fast decrease in pH (decreasing exponential) indicates ongoing precipitation i.e. the 159 absence of inhibition. A delayed decrease in pH, resulting in a plateau around pH 8, indicates an inhibitory effect, proportional to the length of the plateau. Between 160 161 each experiment, the electrode was refreshed with dilute acid and blank tests (without EPS) were performed.

164 **2.6 Interaction of EPS with the in vitro precipitation of CaCO₃**

165 The potential of the EPS matrix to interact with the precipitation of calcium carbonate was tested via the diffusion method in the presence of a closed ammonia-CO₂

saturated atmosphere (Albeck et al., 1993). 200 μ L of the mixture containing pre-filtered (0.22 μ m) CaCl₂ solution (10 mM) and EPS at increasing concentrations (3, 18, and 36 μ g.mL⁻¹) were incubated in duplicate in 16-well plates (Lab-Tek, Nunc/Thermo Scientific, Rochester, NY, USA). The EPS concentrations were selected

168 to match the EPS yields at the extraction times (14, 28 and 56 days of cultivation). The plastic covers of the well plates were perforated to allow the reaction between

169 CaCl₂ solutions containing EPS and ammonium bicarbonate. The well plates were placed in a desiccator that was incubated at 4°C in the dark for 72 hrs. At the

170 completion of the incubation period, the pH value was measured in each well, the overlying solutions were carefully removed to dryness and CaCO₃ crystals analysed.

171 Blank experiments were performed without any EPS. The experiment was carried out in duplicates.

172 **2.6.1 Morphology and mineralogy of the crystals**

173 The 16-well plates containing crystals were used in two manners: first, the morphology of the CaCO₃ crystals was checked with a tabletop scanning electron microscope

174 (Hitachi TM 1000, Ibariki, Japan) in back-scattered electron mode. To this end, the glass plate base was unsealed from its plastic well part and directly observed

175 without carbon or gold sputtering. Secondly, the polymorph of the calcium carbonate minerals was determined by FT-IR spectroscopy using an FT-IR Bruker Alpha

176 (Bruker Optics, SARL, Champs-sur-Marne, France). Mineral phases were determined by comparison of the spectra with the reference spectra available in the RRUFF

177 Project database (<u>https://rruff.info</u>, accessed on January1st, 2022).

178 **2.6.2 Crystal counts and size distribution**

179 CaCO₃ crystals were counted directly in the 16-well plates using an inverted microscope (Nachet, Paris, France) equipped with Mosaic 2.2.1 image analysis software.

Images were processed to obtain crystal sizes (average width and length of size classes $< 50 \,\mu\text{m}$ and $> 50 \,\mu\text{m}$) and the total count of crystals in each well. A total of ten fields of view (10 squares) accounting for 15.5 mm² were analysed. The results are reported as the mean \pm standard deviation.

182 **2.7 Statistical analysis**

183 All the data concerning Synechococcus growth and EPS production are representative of two independent experiments with two technical replicates (four replicates

184 for EPS extracted at 56 days of culture). The results are reported as the mean ± standard error of the mean. Statistical significance was assessed by performing single-

185 factor ANOVA tests; p-values < 0.05 were statistically different.

186 **3. Results**

187 **3.1 Trends in Synechococcus PCC7942 growth experiments and pH evolution**

188 Cell density and pH values increased over the Synechococcus cultivation period (Figure 1A and 1B). The growth of Synechococcus cells showed a typical pattern

including a brief lag phase (~6-7 days) followed by a 7-day (experiment I) and 14-day (experiment II) exponential phase and finally a stationary phase. The stationary

190 phase (early stationary phase) was reached after 14 and 21 days of growth in experiment I and II, respectively, and lasted until day 56 of cultivation in both experiments

191 (late stationary phase) (Figure 1A). Growth experiments I and II started with a similar cell density of approximately 10¹⁰ cells.L⁻¹ and demonstrated reproducible

- 192 growth patterns (p-value = 0.91). At the time of inoculation, cell density was 9.5×10^{10} in experiment I and 7.1×10^{10} cells.L⁻¹ in experiment II (Figure 1A).
- 193 Synechococcus grew exponentially until reaching a maximum of 1.7×10^{12} in experiment I at 14-day of growth and 1.5×10^{12} cells.L⁻¹ after 21 days of growth in
- 194 experiment II. At the end of the exponential growth phase, the cell numbers levelled off and achieved a stable growth stage (stationary phase). Typical evolutions of

195 pH values in culture media during the Synechococcus growth experiments are presented in Figure 1B. As a general trend, pH is linked to the photosynthetic activity

of cyanobacteria. The pH levels rose rapidly during the exponential phase in both experiments, reaching around 10, and stayed steady during the stationary phase.

- 197 While experiment I experienced significant pH fluctuations during the latter part of the stationary phase, overall, the pH evolution trends for both experiments are
- 198 comparable (p-value = 0.91; Figure 1B). The p-values for pH and cell numbers showed that the two independent growth experiments are not significantly different.







199

Figure 1. Evolution of biomass of Synechococcus PCC 7942 culture (A) and pH evolution (B) during exponential, early and late stationary phases. The vertical dotted lines (B)

201 represent the stage transition between lag, exponential and stationary phases. Each value is the mean \pm SD of all replicate values.

202 **3.2 Extracellular carbonic anhydrase**

The activity of extracellular carbonic anhydrase (eCA) in solution changed slightly over the growth experiment (Figure S1). The highest eCA activity (~1600) was detected after 14 days of culture, during the exponential phase. The lowest activity was measured after 56 days of growth, in the late stationary phase.

205 **3.3 Nutrient concentrations during growth**

High nitrate concentrations supported exponential growth and high cell density (Table 1). The results show that a major decrease in nitrate and phosphate concentrations occurred during the exponential growth phase and remained slowed down progressively over the stationary phase. At the end of the stationary phase, the phosphate concentration had decreased to approximately 30% of its initial level. On the other hand, the nitrate concentration was still high, with approximately 67% of its initial concentration remaining. Ammonium concentration was below the limit of detection (2.22μ M). Calcium concentrations decreased gradually and accounted for the total calcium concentration of 81% in the late stationary phase. Other medium constituents should be present in excess and were thus not measured.

211

Table 1. Concentrations of NO₃⁻, PO₄³⁻ and Ca²⁺ (µM) in the culture medium before inoculation (initial concentrations in the medium) and during exponential, early and late stationary

213 of Synechococcus growth phases are given as mean concentrations of four replicates (n=4).

		Synechococcus growth phases						
Major anions and cations (µM)	Initial concentrations in the medium	Exponential	Early stationary	Late stationary				
NO ₃ -	7082±58.7	5731±328.9	5544±57.9	4716±250.1				
PO ₄ ³⁻	68±0.6	39±4.7	41±2.2	21±6.7				
Ca ²⁺	102±0.5	91±2.1	88±1.7	83±4.8				

214

215 **3.3 Abundance of EPS**

The recovery yields of the EPS produced (mean \pm SD) resulting from the applied extraction method are listed in Table 2. The EPS yields varied from 2.9 \pm 0.5 to 18.6 \pm 2.1 mg.L⁻¹ during exponential and early stationary phases and reached the highest yield of 35.4 \pm 4.2 mg.L⁻¹ at 56 days of culture, in the late stationary phase (Table 2). When the values were normalized per cell yield, results showed that the EPS concentration increased significantly between the exponential and late stationary phases (p-value < 0.05) (Figure 2). Synechococcus continuously produced EPS during the 56-day experiment. In the first 14 days of growth, cells grew exponentially and EPS production was deficient. Between exponential and early stationary phases, EPS production increased by a factor of five to seven, reaching a maximum in the late stationary phase, after the 56-day growth experiment.

222

223 Table 2. Cell yield, total EPS production and cell-specific EPS production in Synechococcus PCC7942 cultures during exponential, early and late stationary growth phases. Data

224 represent the means of two independent experiments.

	Time of harvest (growth phase)						
	Exponential	Early stationary	Late stationary				
Cell yield (cells.L ⁻¹)	$(161.6\pm21.6)10^{10}$	$(211.2\pm6.0)10^{10}$	$(268.8 \pm 14.4)10^{10}$				
EPS yield (mg.L ⁻¹)	2.9±0.5	18.6±2.1	35.4±4.2				
Cell-specific EPS production (mg. cells ⁻¹)	(1.9±0.6)10 ⁻¹²	(8.8±0.8)10 ⁻¹²	(13.1±0.9)10 ⁻¹²				





26

227

- Figure 2. Cell-specific EPS production during the exponential, early and late stationary phases. MEAN±SD replicates from (n=2).
- **3.4 Chemical properties of EPS**
- 231 **3.4.1 FT-IR spectroscopy of EPS**
- 232 FT-IR spectroscopy was used to check the overall EPS properties and composition. The IR spectra of EPS harvested during the exponential, early and late stationary
- 233 phases of the growth experiment are depicted in Figure 3. The three spectra show strong similarities, exhibiting characteristic absorption bands for polysaccharides

- 234 and protein moieties (highlighted in Figure 3 by vertical-coloured areas). However, differences in sample composition were also revealed by the presence of additional 235 absorptions indicated by arrows in Figure 3. Interestingly, the spectrum of the exponential phase EPS exhibits a strong band, isolated at 582 cm⁻¹, which according to the literature on EPS could be assigned to a C-X stretch of alkyl halides (Kavita et al., 2011). Bands at 811-868 cm⁻¹, most likely representing the glycosidic linkage 236 237 between sugar monomers, were only present in EPS extracts in the early and late stationary phases. Bands at 1039 – 1128 cm⁻¹ (C–O and C–O–C stretching vibrations) 238 could be assigned to polysaccharides and polysaccharide-like structures (Wang et al., 2012) and were observed in all EPS samples (Figure 3, blue area). In contrast, the small shoulders observed in the early and late stationary phase EPS, at \sim 1242 and 1244 cm⁻¹ correspond to sulfate groups (nS=O stretching vibrations). Low-239 intensity bands observed in the range of 1370-1450 cm⁻¹ are assigned to CH₃ and CH₂ deformations (bends) of proteins (Kansiz et al., 1999). These absorption bands 240 241 were more evident in EPS obtained during the early stationary phase. The bands present in the range of 1660 and 1540 cm⁻¹ are attributed to C=O and C-N stretching vibrations and are characteristic of Amide I and II functions (Figure 3, orange and green areas, respectively), which are typically associated with proteins (Coates, 242 243 2000). Spectra of the early stationary phase EPS showed higher peaks of protein than those observed in EPS from exponential and late stationary phases. The medium bands at 1730 and 1727 cm⁻¹, present in samples extracted from exponential and late stationary phases, can be attributed to C=O stretching vibrations resulting from 244 lipids and fatty acids (Kansiz et al., 1999). Absorptions in the range of 2960–2850 cm⁻¹ corresponding to C-H stretching vibrations of aliphatic hydrocarbons and 245 possibly indicative of long-chain polymers (e.g., sugars or proteins), were observable in all EPS extracts. The amide A band (3345 cm⁻¹), characteristic of the N–H 246 247 vibration of peptide groups in proteins, is present in all spectra (Figure 3, yellow area), but is particularly visible on the early stationary phase EPS spectrum. In the
- samples at 14 and 56 days of growth, this band is included in shoulders due to the presence of OH absorptions centred at 3342 and 3351 cm⁻¹, respectively. The list
- of band assignments is summarized in supplementary material (Table S1).



Figure 3. FT-IR spectra of EPS produced during the exponential (black line), early (green line) and late (blue line) stationary phases. Amide A absorbs in the range of 3342-3351

252 cm⁻¹ (yellow area), amides I-II at 1542–1650 cm⁻¹ (orange and green areas), sulfate groups at ~1242-1244 cm⁻¹, polysaccharides at ~1040–1070 cm⁻¹ (blue area), and the β -glycosidic 253 linkages are visible as a shoulder at ~867cm⁻¹.

254 3.4.2 Protein, sugar and glycosaminoglycan (GAGs) contents

The EPS produced during the exponential growth phase revealed the lowest concentration of protein ($79\pm9\ \mu$ g. mg⁻¹ EPS) (Table 3). The highest protein concentration was measured in EPS produced during the early stationary phase ($253\pm42\ \mu$ g.mg⁻¹ EPS), whereas during the late stationary phase EPS, the protein concentration decreased by ~ two-fold. When accounting for the cell yield at times of EPS extraction, cells produced EPS with ca 11-15 times more protein in the stationary phase than in the exponential phase. The sugar content in the EPS harvested during the three different growth stages did not vary significantly. The EPS produced during the exponential phase contained a slightly higher sugar content ($584\pm9\ \mu$ g of xanthan and $504\pm78\ \mu$ g of dextran equivalents. mg⁻¹ EPS) than that measured in EPS produced during the early and late stationary phases (1.8 times and 1.3 times lower, respectively). Our results show that, over the cultivation time, cells enhanced the

- 261 production of larger amounts of glycosaminoglycans (GAGs) which can be associated with amino sugars and glycoproteins. The highest fraction of sulfated groups
- 262 (GAGs) to total EPS ($217\pm143 \ \mu g \text{ GAGs.mg mg}^{-1} \text{ EPS}$) was found in the late stationary phase EPS.
- 263
- Table 3. Protein, sugar and glycosaminoglycan content of the harvested EPS at times 14, 28 and 56 days of Synechococcus PCC7942 culture. Values represent the average of four,
- three and two measurements of protein, sugar and GAGs, respectively, in two EPS replicated samples (n=2).

	Time of EPS harvesting (days/growth phase)						
Components of FDS wield	14 days	28 days	56 days				
Components of ErS yield	Exponential	Early stationary	Late stationary				
Protein (μg·mg ⁻¹ EPS)	79 <u>+</u> 9	253±42	128±13				
Cell-specific protein production (µg protein.cell ⁻¹)	$(1.5\pm0.6) \times 10^{-10}$	(2.2±0.1)× 10 ⁻⁹	(1.7±0.0)×10 ⁻⁹				
Sugar (µg xanthan equivalents·mg ⁻¹ EPS)	584 <u>+</u> 95	326 <u>±</u> 26	434 <u>±</u> 11				
Cell-specific sugar production (µg xanthan equivalent.cell ⁻¹)	(1.0±0.2)×10 ⁻⁹	$(2.8\pm0.1)\times10^{-9}$	(5.7±0.2)×10 ⁻⁹				
Sugar (µg dextran equivalents∙mg ⁻¹ EPS)	504 <u>+</u> 78	292 <u>+</u> 22	381±90				
Cell-specific sugar production (µg dextran equivalent.cell ⁻¹)	(8.9±1.4)×10 ⁻¹⁰	(2.6 ±0.1)×10 ⁻⁹	(5.0±0.2)×10 ⁻⁹				

Glycosaminoglycans (μg GAGs·mg ⁻¹ EPS)	4 ± 0	31±13	217±143
Cell-specific GAGs production (µg GAGs.cell ⁻¹)	(5.5±5.5)×10 ⁻¹²	$(2.6\pm0.8)\times10^{-10}$	(3.0±2.0)×10 ⁻⁹
GAGs/Sugar (xanthan) ratio	0.01 ± 00	0.09 ± 00	0.51±0.3
GAGs/Sugar (xanthan) ratio	0.01 ± 00	0.10 ± 00	0.58 ± 0.4

267 **3.4.3 SDS-PAGE**

The results of gel electrophoresis after the migration of exponential, early and late stationary phase EPS samples are illustrated in Figure 4. Replicates showed similar 268 band patterns that are distributed between 10 and > 170 kDa. A sharp greenish band in the migration front is strongly stained in late stationary phase EPS (Figure 4) 269 270 and may correspond to chlorophyll. A less pronounced smear is visible in extracts obtained from the early stationary phase (Lanes 4 and 5). Bands of ≤ 10 kDa were 271 not detected in the EPS produced during the exponential phase (Lanes 2 and 3). A marked smear pattern is evidenced in all EPS extracted between 10-26 kDa: one 272 prominent band was individualized at 17 kDa. A discrete blue smear (> 17-43 or 55 kDa) is evidenced in exponential phase EPS samples (Lanes 2 and 3) and is less 273 obvious in EPS samples from the early and late stationary phase (lanes 4-5 and 6-7, respectively). No specific bands were individualized in the > 17-43 kDa molecular mass range, for the three growth phases. A band at about 45-47 kDa was strongly stained in exponential phase only. An area between 43 and 170 kDa was noted in all 274 275 EPS extracts, accounting for 5-6 individualized bands that may correspond to the consecutive addition of an identical 'module', because the progression is logarithmic: is clearly seen in the early and late stationary phase lanes (lanes 4-7). The individualized bands were densely stained in EPS from the late stationary phase, including 276 a smear at ~43-55 or 72 kDa (Lanes 6 and 7) and a prominent band at > 170 kDa (Lanes 6 and 7). 277

278



279

Figure 4. SDS-PAGE of EPS harvested during exponential (lanes 2-3), early (lanes 4-5) and late (lanes 6-7) stationary phases. Alcian blue staining was applied. The molecular ladder

281 (MW) reference is shown in lane 1.

282 **3.4.4 pH-drift assay**

Recordings of the pH-drift assay are shown in Figure 5. The pH-drift assay determined the inhibitory effect of the EPS matrix (e.g., negatively charged functional groups) on the rate of CaCO₃ precipitation. Negatively charged groups of EPS can bind calcium ions from the solution and inhibit the nucleation of carbonates. When CaCO₃ minerals start to nucleate, the pH of the solution decreases. Results show that the inhibitory effect was concentration-dependent and clear differences were visible between EPS extracted in the exponential (Figure 5A), early (Figure 5B) and late (Figure 5C) growth phases. EPS matrices from the stationary phase of culture growth (Figures 5B and 5C) exhibited a stronger inhibitory effect on CaCO₃ precipitation than the EPS extracted during the exponential phase (Figure 5A). Complete inhibition was only reached in EPS from early and late stationary phases when 50 μ g of EPS.mL⁻¹ was tested. In this case, a drop in pH was not observed and nucleation of crystals did not occur (Figure 5B and 5C), which means that the inhibition was total. Conversely, the exponential phase EPS exhibited less inhibition of CaCO₃

290 precipitation (Figure 5A). The shorter plateau shows that the mineral-binding capacity of the matrix delayed CaCO₃ precipitation but that consequently the pH dropped

and visible precipitates formed, showing a less powerful inhibitory effect of the EPS compared to stationary phases EPS matrices.



- Figure 5. In vitro inhibition of calcium carbonate precipitation by using EPS extracted during exponential (A), early (B) and late (C) stationary phases. Each panel shows the effect of six different EPS concentrations (0, 100, 200, 300, 400 and 500 μ g EPS. mL⁻¹) on CaCO₃ precipitation, using the pH-drift assay method. The drop in pH indicates nucleation of CaCO₃ (= precipitation) and a plateau indicates inhibition of precipitation. A larger plateau indicates a higher Ca-binding capacity of the matrix and thus stronger inhibition. Complete inhibition was observed when 50 μ g of EPS solution from early and late stationary phases were used (e.g., see arrows). The results in each panel represent single experiments. Replication showed identical results (see Supplementary Figure S2).
- 298

3.5 Calcium carbonate crystallization in the presence of EPS

300 Forced CaCO₃ experiments were performed using a control solution (without EPS) and EPS solutions, at same pH, with concentrations of 3, 18 and 36 μg.mL⁻¹. Each

301 concentration corresponds to the EPS yield at different growth stages: exponential phase (= $3 \mu g EPS.mL^{-1}$), early ($18 \mu g EPS.mL^{-1}$) and late ($36 \mu g EPS.mL^{-1}$) stationary

302 phases. The crystals formed in the various EPS solutions showed different morphological (Figure 6) and mineralogical (Figure S3) features as well as distinct crystal

303 sizes and distributions compared to those formed in control solution (Figure 7).

304 3.5.1 Mineral morphology

A preliminary light microscopic analysis was carried out in order to identify the most significant samples to analyse by SEM (Figure 6). The morphology of crystals precipitated in the negative controls was very homogeneous and predominantly composed of calcite rhombohedrons that sometimes formed polycrystalline aggregates of size > 50µm (Figure 6A). All control solutions tested for the various EPS harvested during exponential and stationary phases showed similar crystal characteristics. In the EPS solutions, CaCO₃ crystals showed both rhombohedral and spheroidal morphologies (Figure 6B-D). The morphology of crystals appears to change with increasing EPS concentrations. Spherical minerals formation was observed in the exponential phase-EPS solution (Figure 6B) and were less frequent in the EPS solution from early stationary phase (Figure6C). In the late stationary phase-EPS solution, rhombohedrons represented the prevalent crystal morphology while spherical minerals were absent (Figure 6D).

312



313

- Figure 6. In vitro forced CaCO₃ precipitation assay in (A) the absence of the EPS (control solution) and in the presence of EPS extracted during the (B) exponential, (C) early and (D) late stationary phases under increasing EPS concentrations of 3, 18 and 36 μ g.mL⁻¹, respectively. The images show two different CaCO₃ morphologies: rhombohedral (white squares)
- 316 and spheroidal (white circles), in some cases shown as polycrystalline crystals. The scale bar (black) at the bottom right of the images is 500 µm.

317 **3.5.2 Crystal mineralogy**

318 The crystals' mineralogy was assessed by FT-IR microscopy performed on selected individual crystals of $> 10 \mu m$ (Figure S3). The results revealed that calcite was

319 the only CaCO₃ polymorph formed in the control solution. Calcite and vaterite formed in all EPS solutions tested. The FT-IR spectra revealed that all rhombohedrons

320 and polycrystalline aggregates with "sharp edges" represent calcite polymorphs. In contrast, spheroidal crystals revealed a vaterite signature (Figure S3).

321 **3.5.3 Crystal size and distribution**

322 The results from image analysis showed that a larger quantity of crystals precipitated in the stationary phase (early and late) EPS solutions (Figure 7) and that major

differences were also observed in crystal size distribution (Figure 7). A comparison of the class of small crystal sizes (< 50 μ m) with the large crystal size class

- 324 (>50μm) showed a clear trend of an increasing total number of small crystals in the stationary phase EPS solutions compared to those formed in the EPS solutions
- from the exponential phase. The size reduction of the crystals at 18 and 36 mg/L (Figure 7, early and late stationary phases) suggests a partial inhibitory effect of the
- 326 EPS on the formation of calcium carbonate.



328 Figure 7. Total numbers of small (< 50 µm) and large (> 50 µm) crystal size classes of precipitated CaCO₃ in EPS solutions obtained from exponential and stationary phases, in EPS concentrations of 3, 18 and 36 µg.mL⁻¹, respectively. 329

330 4. Discussion

331 Our study demonstrates that the amount and properties of EPS change significantly (p-value < 0.05) at the three different stages of Synechococcus growth in an 332 artificial bloom experiment. Cells continuously produce EPS that increases in concentration and become more negatively charged in the stationary phase. We sampled 333 this EPS over the exponential, early and late stationary phases and studied its role in carbonate mineral precipitation. Based on this, a conceptual model was developed 334 to correlate the findings of this investigation with the potential EPS production of the naturally occurring picoplankton blooms and its possible involvement in whiting 335 events. Though natural blooms experience a variety of factors that are not represented in the experiments described in this paper, the first part of the discussion is focussing only on the experimental data, whereas the interaction of these basic processes with other biotic and abiotic factors acting in the environment is discussed 336

337 afterwards.

352

338 4.1 Exponential growth phase

339 Macronutrients, such as nitrogen (N) and phosphorus (P) promote the initiation of cyanobacterial blooms (Reynolds and Walsby, 1975; Paerl, 2008; Xu et al., 2015). 340 In our growth experiment, the beginning of the exponential phase (and the persistence of bloom) (Figure 1A) was positively correlated with the high initial nutrient 341 concentration in the medium (Table 1). Environmental factors such as water temperature, light intensity. hydrodynamics and availability of dissolved inorganic carbon 342 (DIC) are also important determinants of cyanobacteria bloom development (Clark and Flynn, 2000; Dokulil and Teubner, 2000; Havens, 2008). Blooms can 343 dramatically alter the supply of inorganic carbon for photosynthesis, which causes the pH to increase (Ibelings and Maberly, 1998). In the early exponential phase of 344 our batch cultures, the high photosynthetic activity of cyanobacteria cultures resulted in fast pH increase thereby reducing the total inorganic carbon of the grown medium. Light and CO2 are the sources of energy and carbon for cyanobacteria, and are of critical importance for their growth (Takahashi et al., 2004). At pH 9 345 (Figure 1B), the concentration of CO₂ predicted is close to zero (< 1μ M) and the HCO₃⁻ concentration is 475 μ M (PhreeqC data). A similar scenario was observed in 346 natural blooms occurrence: the population of cyanobacteria draws down the partial pressure of CO₂ (pCO₂) in the photic zone, increasing the surface water pH up to 347 9-10 (Ibelings and Maberly, 1998; Verspagen et al., 2014) and CO₂ concentration can become completely depleted or reach values close to zero (Maberly, 1996). 348 349 Under extreme conditions, the concentration of HCO₃⁻ can also become markedly reduced (Talling, 1976; Maberly, 1996). When the rate of photosynthesis is greater 350 than the combined rate of resupply of CO₂ from the atmosphere and DIC in the hypolimnion, deviation from the air equilibrium occurs, favouring CaCO₃ precipitation. The pH of most aquatic systems ranges from 7.5-8.1 and keeps inorganic carbon primarily in the form of bicarbonate (O'Neil et al., 2012). In poorly buffered systems, 351 such as highly productive lakes, the pH and speciation of DIC experience large fluctuations which vary widely on a scale from daily (diel) to episodic, to seasonal

(Maberly, 1996) with diel variations as high as two pH units and 60 µmol DIC.L⁻¹ (Maberly, 1996). Because CO₂ favors the C₃ photosynthesis (C₃ cycle operation of 353 Calvin-Beson cycle), the high pH of ~ 10 in our growth medium could be associated with carbon limitation (Ibelings and Maberly, 1998; Verspagen et al., 2014). 354 355

356 To alleviate CO₂ limitation, cyanobacteria have developed an efficient CO₂-concentrating mechanism (CCM) (Aizawa and Miyachi, 1986; Badger and Price, 1992; 357 Badger et al., 2002; Burnap et al., 2015) and can use bicarbonate as an inorganic carbon source for photosynthesis (Price et al., 1998; Giordano et al., 2005; Sandrini 358 et al., 2016). By activating CCM, cyanobacteria concentrate CO₂ by a factor of up to a thousand (Badger and Andrews, 1982; Badger et al., 2002; Price, 2011). CO₂-359 deficient conditions experienced during the exponential phase of our growth experiment, coupled with the continuous cellular demand for inorganic carbon to support 360 photosynthetic carbon fixation likely led the cells to activate CCM. The predicted concentrations of CO₂ and HCO₃⁻ in the growth medium (PhreeqC data) in the early 361 and late exponential phase infer that Synechococcus cells actively transported across the membrane and accumulated DIC into the cell, where the HCO₃⁻ pool was utilized to generate elevated CO2 levels around Rubisco (Badger et al., 2002; Price et al., 2008). The CCM of cyanobacteria accomplishes very high carbon 362 concentrating factors (Cextrenal : Cinternal) at deficient specificity factors of RuBisCo (Tortell, 2000; Tortell et al., 2000). CCM involves bicarbonate transporters in the 363 cell membrane, intracellular (iCA) and extracellular (eCA) carbonic anhydrase enzymes and concentrated RuBisCO activity located in carboxysomes (Badger et al., 364 2006; Price et al., 2008; Rae et al., 2013). CA converts HCO₃⁻ to CO₂ (Badger and Price, 1994), which increases the external pH in close proximity to the cells. In our 365 study, eCA activity was \sim 1.6-2.0 times higher during the exponential growth phase and reduced gradually through the stationary phase (Supplementary Figure S1). 366 The strongly stained band only present in the exponential phase-EPS at around 45-47 kDa (Figure 4, lanes 2, 3) may be indicative of eCA, as reported by Kupriyanova 367 et al., 2018, but this requires further investigation. Another plausible explanation for the 45-47 kDa band could be the presence of chlorophyll f synthase, which 368

typically migrates at around 46 kDa (Shen et al., 2019). Similarly, Yang et al. (2023) measured the CA anhydrase in solution over a 30-day growth experiment with

370 Synechococcus PCC 7942 and reported an increase over the lag phase and large fluctuations over the exponential phase. During the stationary phase, CA did not vary

371 greatly but a minor decrease was recorded in the late stationary phase (Yang et al., 2023). In our study, the higher eCA activity recorded could explain the strongly

stained ~45-47-kDa band that was only identified in our SDS-PAGE gels of EPS produced during the exponential phase (Figure 4, lanes 2-3). The molecular weight

(MW) of this band is similar to a 42-43 kDa eCA previously identified by Kupriyanova et al. (2018) and discussed by Martinho de Brito et al. (2022). As explained
 in the Results section 3.4.3, we cannot exclude that the band is chlorophyll f synthase, which seems to show up around 46 kDa. A more substantiated demonstration

of the identity of the SDS-PAGE band will require other approaches (beyond the scope of the present study), such as micro-sequencing of the purified 43 kDa band

376 or the use of a CA-specific antibody.

377

378 Active uptake of HCO₃⁻ and accumulation of Ci species requires the input of metabolic energy e.g., ATP (BCT1 HCO₃⁻ transporter), NADPH or reduced ferredoxin (CO₂ uptake) or coupling to an electrochemical Na⁺ gradient (SbtA or BicA HCO₃⁻ transport) (Badger et al., 2002; Price et al., 2008). This energetic cost may therefore 379 reflect on the growth rates achieved. Synechococcus PCC 7942 grows at > 80% of its maximum growth rate when provided with HCO₃ as its main inorganic carbon 380 381 source (Miller et al., 1984). During the exponential phase, the carbon production from photosynthesis is mainly allocated for biomass production, not for EPS synthesis. 382 During this phase (Figure 2 and Table 2), the small amount of EPS produced comprises a higher proportion of sugars and lower amounts of protein and GAG compared 383 to EPS produced during the stationary phase (Figure 3 and Table 3). Our study indicates that rather than proteins, sugars are the major component in all EPS extracts. This finding is supported by the data obtained from FTIR analysis (Figure 3). The smaller amount of negatively charged groups of the EPS during the exponential 384 385 phase (Figure 4, lanes 2-3) compared to those of EPS from the early and late stationary phases (Figure 4, lanes 4-7) resulted in weak to moderate inhibitory capacity (Figure 5A). The main phenomenon observed in the pH-drift assay (Figure 5) is the initial Ca binding to negatively charged groups in EPS prior to carbonate addition, 386 which initiates CaCO₃ precipitation. This results in a decrease of pH. The pH drift assay showed that EPS from exponential phase (Figure 5C) having a larger plateau, 387 and thus a lower a calcium binding capacity then the EPS from the stationary phase. This observation was further corroborated by the forced precipitation experiments, 388 389 which showed that EPS from the exponential phase induced small amount of mostly large-sized carbonate crystals (>50 µm), very similarly to the negative control experiment (Figure 7) (Martinho de Brito et al., 2022). The high concentration of Ca^{2+} in the medium (83 μ M) compared to the initial [Ca²⁺] at the beginning of the 390 391 experiment (103 μ M), indicates that a small amount of calcium ions was bound to negatively charged functional groups of EPS (Table 1, see [Ca²⁺]). 392 In our batch experiment, cells continue to grow exponentially for ~20 days of cultivation. At this point, cultures reached the maximum cell density (Figure 1A) and pH values ranged between 10-11 (Figure 1B). Based on our calculations, under these alkaline conditions, CO_2 was completely depleted ($1.7 \times 10^{-3} \,\mu\text{M}$) in the growth 393 medium, whereas HCO₃⁻ was extremely low (~ 79 μ M). Thus, the dominant inorganic carbon speciation was CO₃²⁻ (421 μ M). Because cells cannot take up CO₃²⁻ and 394 395 HCO₃⁻ concentration seems to be insufficient to cover the carbon demands of cyanobacterial growth, we assume that this may have been the cause of cell numbers

396 starting to level off (Figure 1A, early stationary phase). Consequently, cultures entered a stationary state due to a lack of inorganic carbon availability required to

397 increase cell population (Miller et al., 1984; Mayo et al., 1989; Verspagen et al., 2014). The excess of nutrients measured in the medium in the late exponential phase

398 (Table 1) suggested that the specific growth rate was not limited by nutrient availability but by a rather low level of CO₂ carbon content.

399400

401 **4.2 Early stationary phase**

402 Insufficient CO₂ availability is considered to be the external stress factor constraining the growth rate of cyanobacteria (Maberly, 1996; Hein, 1997; Ibelings and Maberly, 1998) and low [HCO₃-] could sustain a constant population density for at least ~ 40 days (See Figure 1A, stationary phase). Our results suggest that at this 403 point, carbon fixation was allocated to EPS synthesis, not to biomass production (Miller et al., 1984). Increased EPS production is usually associated with external 404 stress factors (Rossi and De Philippis, 2015), including high pH conditions (Martinho de Brito et al., 2022). Moreover, metabolic stress may also alter the composition 405 of EPS (Babele et al., 2019; Martinho de Brito et al., 2022). In the present study, the negative functional group abundance increased, resulting in a higher acidity of 406 407 EPS (Figure 4, lanes 4-5) due to an increase in protein and sulfated glycan (GAG) (Table 3). In the pH conditions of the early stationary phase, all the functional 408 groups of the EPS matrix are deprotonated and are able to bind calcium ions (Figure 5B) (Dupraz and Visscher, 2005; Braissant et al., 2007; Dittrich and Sibler, 2010) and bind calcium more efficiently nanometric nuclei in formation (if their formation is thermodynamically favoured). We suggest that the increased calcium-binding 409 capacity of the EPS probably accounts for lower the Ca²⁺ concentration measured in the medium (Table 2, see [Ca²⁺]). In our in vitro forced precipitation assay, we 410

411 measure the second effect, the inhibitory one (mineral-binding effect), which results in the production of small-sized calcium carbonate crystals (< 50 µm), in

412 comparison to what happens in the exponential phase (Figure 7).

413

414 **4.3 Late stationary phase**

As mentioned above, we assume that the continuous increase in EPS production over the late stationary phase, including an overall augmentation of negatively charged 415 functional groups (Figure 4, lanes 6-7), including GAG content (Table 2), might be a specific response to a stress scenario As expected, the present study shows that 416 the greater amount of negatively charged functional groups of EPS from the late stationary phase (Figure 4, lanes 6-7) resulted in a higher Ca-binding capacity than 417 exponential and early stationary phase-EPS (Figure 5C). Our forced precipitation experiments showed that minerals produced in the late stationary-EPS solutions are 418 smaller and more abundant than those formed in EPS solutions from the early stationary phase (Figure 7). Under natural conditions, when the Ca²⁺ supply is continuous, 419 the crystals may or may not continue to grow, depending on the physical space within the EPS matrix (Dupraz et al., 2009). Based on the high concentration of nitrate 420 (4720 µM) measured in the late stationary phase (Table 1), we assume that the abundance of this nutrient supported the persistence of the stationary phase, i.e., similar 421 422 to a prolonged bloom in natural conditions. The death phase was not observed in our 56-day-long experiment. Given that our cultures were continuously stirred, we 423 can assume that light was not limiting cyanobacterial growth. Furthermore, in natural blooms, the increase in population density may affect cells at greater depth 424 through self-shading by decreasing the light available for photosynthesis (Townsend et al., 1994). Yet, cyanobacteria (including Synechococcus) are known to be well-425 adapted to low-light conditions (Campbell and Carpenter, 1986; Palenik, 2001; Callieri et al., 2011). Additionally, the presence of sulfated constituents on late stationary phase-EPS contributes to a higher negative charge of the matrix and higher Ca-binding potential (Decho and Kawaguchi, 2003; Skoog et al., 2022), compared

to EPS extracted in the exponential phase which contained significantly lower GAG (Table 2). The present study shows that the greater amount of negatively charged

functional groups of EPS from the late stationary phase (Figure 4, lanes 6-7) resulted in a higher Ca-binding capacity than exponential and early stationary phase-EPS

- 429 (Figure 5C). Our forced precipitation experiments showed that minerals produced in the late stationary-EPS solutions are smaller and more abundant than those
- 430 formed in EPS solutions from the early stationary phase (Figure 7), suggesting an increased inhibitory ability of the late stationary-EPS.
- 431

432 **4.4 Natural bloom and formation of whitings – Conceptual model**

433 Our observations made during exponential and stationary phases can be applied to generate a conceptual model of EPS properties during a Synechococcus bloom 434 event (Figure 8A-C). The onset of a bloom starts with an increase in cell numbers, with high values in spring-summer (exceeding 10⁵-10⁷ cells.mL⁻¹) and lower values 435 in winter months (< 10⁵ cells.mL⁻¹) in both marine (Agawin et al., 1998; Phlips et al., 1999) and freshwater (Maeda et al., 1992; Tai and Palenik, 2009) environments. 436 This resembles the exponential growth phase in our study (Figure 1, exponential phase). We predict that during the initial phase of a natural bloom, there is little EPS production: cells grow relatively quickly and the carbon fixed during photosynthesis is predominantly allocated to biomass production (Figure 8A). The fast growth 437 is followed by a phase during which cell numbers level off, typically due to stress conditions, which is represented by the early stationary phase in our study. Under 438 439 certain conditions, blooms can be sustained for weeks and possibly longer (Anderson et al., 2002; Havens, 2008; Zhao et al., 2013), similar to what we observed in our growth experiments (Figure 1A, early stationary phase). The maintenance of a bloom requires continuous input of nutrients, which is also the case in our experiment 440 (Table 1) or in the case of natural systems, a turnover from lysing cells recycled by other microbes. During this phase, we did not observe a significant increase in cell 441 442 density but the production of EPS continued at a disproportionately high rate (Figure 8B-C). Our findings are in agreement with the lab studies using diatom cultures 443 which show that EPS production is low during exponential growth and increases in the stationary phase (Myklestad and Haug, 1972; Myklestad et al., 1989; Bhosle 444 et al., 1995). These authors reported that nutrient-deficient conditions enhanced the production of EPS over the growth phases. If carbon fixation continues and some critically required nutrient is lacking from the growth medium, most likely the phototrophic organisms produce carbohydrate reservoirs (Ciebiada et al., 2020). These 445 include storage polymers like glycogen and the production of other carbohydrate-rich compounds, including EPS (De Philippis et al., 1996, 2001; Decho and Gutierrez, 446 447 2017). The decline of blooms in natural environments is typically associated with nutrient, low or high light intensity, grazing or viral infection. Under these stressful

- 448 conditions, an increase in EPS production by the phyto/picoplankton community may be expected.
- 449

450 Synechococcus spp. blooms can cause whiting events (Thompson, 2000), characterized by the presence of large amounts of CaCO₃ minerals in surface water. Various mechanisms have been proposed for this phenomenon, including chemical and physical processes (Shinn and Steinen, 1989: Larson and Mylroie, 2014) as well as 451 452 biologically mediated-precipitation (Thompson and Ferris, 1990; Robbins and Blackwelder, 1992; Stanton et al., 2021). However, no consensus has been reached on 453 the precise cause of these events. Carefully transporting the results from forced precipitation experiments to a whiting event, we suggest that early in the bloom (Figure 454 8A), relatively large CaCO₃ crystals form, provided sufficient Ca²⁺ is available (Figure 8D). As the bloom continues to grow, progressively the larger quantity of 455 negatively charged functional groups in the EPS provides more cation-binding sites and thus inhibits calcium carbonate precipitation largely. Depending on the three-456 dimensional structure of the EPS and surface properties (Wang et al., 2012), nucleation may yield smaller CaCO₃ crystals (Figure 8). If this occurs, then the production 457 of a more negatively charged matrix (largely contributed by the enrichment in sulfated polysaccharides) may offer some selective advantage to the cyanobacteria 458 population, by inhibiting and/or delaying mineral precipitation and by reducing crystal size formed around the cells. This might result in slow sinking rates, extending 459 the residence time of the cyanobacterial community in the photic zone. If the bloom occurrence is short (e.g., similar to 14-28 days in our growth experiment), minerals 460 making up the whiting will be relatively larger. Consequently, the aggregates of cyanobacteria, EPS and CaCO₃ minerals may sink faster because mineral precipitation in EPS increases the cyanobacterial-specific density several-fold. The Synechococcus specific density (ρ) is 1.040 g·cm⁻³ (Reynolds, 1987), near-neutrally buoyant, 461 whereas ρ_{calcite} is 2.710 g·cm⁻³ (Lange, 1999). The production of larger amounts of more negatively charged EPS may act as a protection mechanism against carbonate 462 463 formation in the vicinity of the cell wall (Martinez et al., 2010; Bundeleva et al., 2012), thus allowing the organisms to reside longer in the photic zone. Interestingly, the production of EPS that contained sulfated groups among bacteria seems to be exclusive to cyanobacteria (Pereira et al., 2009; Maeda et al., 2021). Maeda et al. 464 465 (2021) reported that the cyanobacterium Synechocystis 6803 produced large amounts of GAG compounds during an experimental bloom formation. The authors suggested that these constituents can be advantageous for the development of surface bloom as it may increase the buoyancy, permitting cells to migrate upward 466 467 rapidly when the water column is stable (Walsby et al., 1995). Thus, GAG production may be considered as an alternative for organisms that lack gas vesicles to remain longer in the photic zone (Maeda et al., 2021). The negative charge of EPS produced containing high sulfated content also protects the community against viral 468 469 infection (Matsunaga et al., 1996). Therefore, the production of GAG by pelagic cyanobacteria contributes to stress tolerance and viral infectivity, helping in the 470 persistence of bloom. In our growth experiments, a decline in cell numbers was not observed, which would represent the end of the bloom. In the natural environment, nutrient depletion, grazing or viral lysis/infection are the most likely causes of terminating a bloom (Gons et al., 2002). The cell lysis releases organic matter, which 471 supports the growth of heterotrophic bacteria (Kjelleberg et al., 1987; Hagström et al., 1988; Kieft et al., 2021). Photosynthetically derived organic carbon is one of the 472 major carbon and energy sources for heterotrophic bacteria (Allgaier et al., 2008). These heterotrophs can degrade EPS and liberate bound Ca²⁺ (Visscher et al. 1998; 473 Ionescu et al., 2015; Diaz et al., 2017). In addition, microbial respiration will produce HCO₃-/CO₂, increasing the saturation index of CaCO₃, and may enhance the 474 whiting (Figure 8). Although our model is [somewhat] largely theoretical at this stage and explore the role of one picoplankton species, its merit is to focus on an 475 overlooked actor of whiting events, the EPS. Furthermore, it provides a conceptual framework to work with, for designing novel experiments and measurements both 476 in natural systems and at the lab bench, to validate the molecular mechanisms involved in microbial bloom associated CaCO₃ formation in marine and lacustrine 477 478 models.

479



- 481
- 482 **Figure 8.** Conceptual diagram of proposed EPS-supported carbonate precipitation mechanism explaining the origin of whiting events.

483 Data availability

484 All raw data can be provided by the corresponding authors upon request.

485 Author contributions

- 486 M.M.d.B., I.B. and P.T.V. designed the study in a project directed by P.T.V., I.B. and E.V.; M.M.d.B., I.B., P.T.V., F.M., A.W. and L.P. developed the methodology;
- 487 M.M.d.B. and I.B. carried out the laboratory measurements; M.M.d.B., P.T.V. and I.B. analysed the data; M.M.d.B. wrote the manuscript draft with significant 488 contributions of P.T.V. and I.B. M.M.d.B., P.T.V., I.B., E.V., F.M., A.W and L.P. reviewed and edited the manuscript. All authors have read and agreed to the
- 488 contributions of F.1.V. and I.D. WI.WI.d.D., F.1.V., I.D., E.V., F.WI., A.W and L.F. Teviewed and edited the manuscript. An
- 489 published version of the manuscript.

490 **Competing interests**

491 The authors declare that they have no conflict of interest.

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EPS (Exponential growth phase)	EPS (Early stationary phase)	EPS (Late stationary phase)	Band assignment
	Wavenum	ber (cm ⁻¹)	
3342	/	3351	пОН
included in OH absorption band	3281	included in OH absorption band	Amide A (<i>n</i> N–H)
/	3077	/	Amide B (<i>n</i> N–H)
2927	2959, 2922, 2851	2938	пс-н
1730	/	1727	n _{C=O}
1658	1648	1650	Amide I $(n_{C=O})$
1543	1542	1549	Amide II (n_{C-N})
1376	1448, 1401	1375	$d_{ m C-H}$
/	1305	/	Amide III (n_{C-N})
/	1242	1244	v _{S=0}
1136	1127	/	n _{C-O-C}
1043	1070	1038	n _{C-O}
867	/	811	<i>g</i> с-н
582	/	/	$d_{ m C-X}$



Figure S1. Extracellular carbonic anhydrase (eCA) activity measured at days 14, 28 and 56 of Synechococcus PCC 7942 growth experiment.



Figure S2. Replication of the *in vitro* inhibition of calcium carbonate precipitation experiment by using EPS extracted during exponential (A), early (B) and late (C) stationary phases of *Synechococcus* growth experiment. A negative control (no EPS) and EPS extracts of concentrations of 10, 20, 30, 40, and 50 μ g.mL⁻¹ were used in the CaCO₃ inhibition assay. The decrease of pH indicates precipitation and a plateau inhibition of carbonate mineral precipitation. A larger plateau (> 50 μ g) indicates stronger inhibition of calcium carbonate precipitation (e.g., see black arrows in panel B-C).



Figure S3. FT-IR spectra of (A) vaterite and (B) calcite. The two calcium carbonate polymorphs precipitated in EPS solutions produced during two *Synechococcus* growth phases. Data shown EPS produced (A) during exponential growth phase a concentration of 3 μ g·mL⁻¹ and (B) during the late stationary phase with a concentration of 36 μ g·mL⁻¹.