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Revision 1

| 2 | Precipitation of low-temperature disordered dolomite induced by |
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| 3 | extracellular polymeric substances of methanogenic Archaea Methanosarcina |
| 4 | barkeri: Implications for sedimentary dolomite formation |
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22 ABSTRACT

A correlation between methanogenesis and dolomite formation has been reported; 23 however, the mechanism underlying of this association is not fully understood. In this study, we 24 conducted forced carbonate precipitation experiments at room temperature in calcite-seeded 25 Ca/Mg carbonate solutions containing either purified non-living biomass or bound extracellular 26 27 polymeric substances (EPS) of the methanogen *Methanosarcina barkeri*. Purified non-living biomass and bound EPS was used so as to avoid the possible influence of the complex 28 components of the growing microbial culture on carbonate crystallization. Our results 29 demonstrated that non-living biomass of M. Barkeri can enhance the Mg incorporation into 30 calcitic structure and induce the crystallization of disordered dolomite. In the presence of ~ 113 31 mg/L of non-living biomass, disordered dolomite with ~41 and 45 mol% of MgCO₃ was 32 precipitated in solutions with initial Mg:Ca ratios of 5:1 and 8:1, respectively. A systematic 33 increase in the MgCO₃ contents of the precipitated Ca-Mg carbonates was also observed with the 34 increased non-living biomass concentration. Bound EPS was shown to be the component of non-35 living biomass that catalyzed the precipitation of disordered dolomite. At only $\sim 25 \text{ mg L}^{-1}$ of 36 bound EPS, disordered dolomite with ~47 and 48 mol% of MgCO₃ was precipitated in solutions 37 38 with initial Mg:Ca ratios of 5:1 and 8:1, respectively. We propose that adsorption of bound EPS to growing carbonate surfaces through hydrogen bonding is the key to catalyzing disordered 39 dolomite crystallization, and that this mechanism is also applicable to natural EPS-induced 40 41 dolomite formation. This study provides significant insight into the formation mechanism of microbial-induced dolomite with heavy δ^{13} C values. 42

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Key words: sedimentary dolomite, methanogen, EPS, catalysis, microbial-induced dolomite,
heavy δ¹³C value

46 INTRODUCTION

Although abundant in ancient rocks, dolomite is uncommon in modern sedimentary 47 environments. Present-day low-temperature dolomite formation is usually observed in 48 association with marine and other saline environments (Jones 1961; Zenger et al. 1980; Machel 49 and Mountjoy 1986; Hardie 1987; Mazzullo 2000; Warren 2000). Freshwater dolomite has also 50 been documented but its occurrence is rare (El-Sayed et al. 1991; Colson and Cojan 1996; Capo 51 et al. 2000; Whipkey et al. 2002; Roberts et al. 2004; Kenward et al. 2009). The rarity of modern 52 dolomite is largely consistent with the notorious difficulty in reproducing dolomite 53 crystallization under ambient conditions (Lippmann 1973; Oomori and Kitano 1987; Land 1998; 54 Higgins and Hu 2005), contributing to the long-existing controversy over the formation 55 mechanism of sedimentary dolomite, i.e. the "dolomite problem" (Zenger et al. 1980; Machel 56 and Mountjoy 1986; Hardie 1987; Burns et al. 2000; Mazzullo 2000; Warren 2000). 57

While there is no simple abiotic recipe for dolomite precipitation, recent studies suggest 58 that microbes are paramount to overcoming kinetic barriers to dolomite crystallization. A 59 number of metabolic pathways have been implicated in catalyzing dolomite precipitation, 60 including both bacterial sulfate reduction (BSR) and methanogenesis (Baker and Kastner 1981; 61 Baker and Burns 1985; Hardie 1987; Compton 1988; Vasconcelos and McKenzie 1997; Wright 62 1999; Burdige et al. 2000; Mazzullo 2000; Warren 2000; Van Lith et al. 2003b; Roberts et al. 63 2004; Kenward et al. 2009; Deng et al. 2010). Many carbonate precipitation studies have been 64 65 performed exploring the poorly constrained role of sulfate-reducing bacteria (SRB) in promoting dolomite precipitation (Vasconcelos et al. 1995; Nielsen and Jahn 1999; Warthmann et al. 2000; 66 Van Lith et al. 2003b; Wright and Wacey 2005; Kenward et al. 2009; Deng et al. 2010; Krause et 67 68 al.; Zhang et al. 2012a; Xu et al. 2013; Zhang et al. 2013). For example, Zhang et al. (2012a)

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demonstrated the catalytic role of dissolved sulfide, one of the major products of BSR, in 69 dolomite precipitation. However, fewer studies have been devoted to methanogens, although 70 there may exist some physiochemical rules which are common to dolomite induced by SRB and 71 methanogens. Roberts et al. (2004) and Kenward et al. (2009) conducted Ca-Mg carbonate 72 precipitation experiments in natural environment and culture media respectively, with the 73 74 involvement of methanogens and showed dolomite precipitation in natural environment. Recent molecular dynamics modeling also proves that polysaccharides (main components in EPS) can 75 lower the dehydration energy barrier (Shen et al. 2015). 76

77 In natural environments, the vast majorities of microorganisms live and grow in aggregated forms such as biofilms and flocs. The common feature of all these phenomena is that 78 microorganisms are embedded in an EPS matrix. The production of EPS matrix has been shown 79 to occur both in prokaryotic and eukaryotic microorganisms (Nielsen and Jahn 1999; Wingender 80 et al. 1999). EPS is composed of organic macromolecules including polysaccharides, proteins, 81 nucleic acids, (phospho)lipids, and other polymeric compounds. Their composition may be 82 controlled by different processes, such as active secretion, shedding of cell surface material, cell 83 lysis, and adsorption from the environment (Wingender et al. 1999). Reported functions of EPS 84 85 matrix include mediating cell adhesion to surfaces and metabolic interactions between cells and minerals, templating mineral crystallization, aggregation of cells in flocs and biofilms, 86 stabilization of the biofilm structure, formation of a protective barrier that provides resistance to 87 88 biocides, protection from UV radiation, toxic metals, the toxicity of mineral surfaces or other harmful effects, retention of water, and sorption of exogenous organic compounds for the 89 90 accumulation of nutrients from the environment (Geesey et al. 1988; Costerton et al. 1995; 91 Laspidou and Rittmann 2002; Chan et al. 2004; Harrison et al. 2007; Xu et al. 2012). EPS can

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92 also influence carbonate precipitation in multiple ways (Reid et al. 2000; Dupraz et al. 2009). Negatively-charged acidic groups within the EPS matrix can effectively bind metal cations (Li et 93 al. 2001; Perry et al. 2005; Ortega-Morales et al. 2006; Braissant et al. 2007), which can 94 therefore remove free Ca²⁺ ions from solution, inhibiting carbonate precipitation when limited 95 Ca^{2+} ions are available from the proximal surrounding environment (Kawaguchi and Decho 2002; 96 Dupraz et al. 2004; Dupraz and Visscher 2005; Gautret and Trichet 2005). Subsequently, the 97 degradation of the labile fraction of EPS, abiotically or biotically, can liberate Ca^{2+} bound to the 98 polymer to promote carbonate precipitation (Dupraz and Visscher 2005; Dupraz et al. 2009). 99 EPS can also provide nucleation sites for carbonates (Fortin et al. 1997; Nielsen and Jahn 1999; 100 Dupraz and Visscher 2005; Bontognali et al. 2008; Dupraz et al. 2009; Bontognali et al. 2010; 101 Paulo and Dittrich 2013). 102

In laboratory pure cultures, however, EPS matrix is not essential structure of 103 microorganisms, since loss of EPS matrix does not impair growth and viability of the cells as it 104 does in naturals systems (Nielsen and Jahn 1999; Wingender et al. 1999). Also, the definition of 105 EPS in pure cultures is often slightly different from that used in natural systems. EPS in pure 106 cultures is often divided into two categories: bound and soluble EPS (Hsieh et al. 1994; Nielsen 107 et al. 1997; Nielsen and Jahn 1999; Laspidou and Rittmann 2002). Bound EPS includes sheaths, 108 capsular polymers, condensed gel, loosely bound polymers, and attached organic materials. 109 Soluble EPS includes soluble macromolecules, colloids, and slimes. Bound EPS is associated 110 111 with the cell surface and is presumably crucial for biofilm formation, whereas soluble EPS is loosely associated with the cells and predominantly generated by sloughing off from bound EPS 112 113 (Xu et al. 2012).

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In previous laboratory dolomite precipitation experiments within live culture of SRB and 114 methanogen, the involvement of bound EPS has been shown in promoting dolomite nucleation 115 and growth (Van Lith et al. 2003b, a; Roberts et al. 2004; Bontognali et al. 2008; Kenward et al. 116 2009; Bontognali et al. 2014). However, within live cultures, a definitive elucidation of the effect 117 of bound EPS on dolomite precipitation can be hard to reach, since microbes, microbial 118 metabolic products, and the complex ingredients of typical culture medium may all affect 119 carbonate precipitation. For example, phosphate in SRB culture media can lead to the 120 precipitation of Ca/Mg-phosphate minerals. It also has a pronounced impact on carbonate 121 122 precipitation and can potentially obscure or alter more subtle effects on mineral precipitation, which might lead to misinterpretation of culture studies meant to simulate natural systems 123 (Gallagher et al. 2013). Some claimed dolomite precipitates are actually aragonite (Wright & 124 Wacey (2005), see their Figure 14A) and Ca-Mg-phosphates. Recently studies show that addition 125 of low dipole moment substances, such as H₂S, carboxylic acid, and methane, can change the 126 behavior of the solution, disrupt surface Mg²⁺-water complex, and promote Mg incorporation 127 into Ca-Mg-carbonates (Xu 2010; Zhang et al. 2010). Substrates like (001) surfaces of clay 128 minerals and hematite can promote heterogeneous nucleation of calcite and high magnesian 129 130 calcite, and inhibit aragonite formation due to their pseudo-hexagonal (001) surfaces (Xu et al. 2018). Low temperature abiotic synthesis with addition of smectite show precipitation of 131 disordered dolomite (Liu et al. 2019) 132

In this study, we characterized the effect of bound EPS from the methanogen *Methanosarcina barkeri* on Ca-Mg carbonate precipitation. *M. barkeri* is a anaerobic methanogenic archaea commonly isolated from mud samples in lakes and bogs and sewage samples (Stadtman and Barker 1951; Balch et al. 1979; Hippe et al. 1979; Bock et al. 1994;

Maeder et al. 2006). M. barkeri is metabolically versatile and can utilize a variety of 137 methanogenic substrates including H₂, CO₂, methanol, methylamines, and acetate. This species 138 can also adapt to one of the widest ranges of habitats for an individual methanogenic species, 139 from freshwater to high salinity water with three times the solute concentration in seawater. 140 Laboratory culture studies showed that it exhibits a dichotomous morphology, growing in 141 freshwater as large multicellular aggregates embedded in an EPS matrix, or in high extracellular 142 solute concentrations as individual cells without EPS (Stadtman and Barker 1951; Sowers et al. 143 1993; Anderson et al. 2012). In this study, we cultured *M. barkeri* in freshwater medium so that 144 we could (1) collect sufficient amount of bound EPS for carbonate precipitation experiments and 145 (2) gain a broader understanding of the potential role of methanogens in freshwater dolomite 146 formation. To the best of our knowledge, no evidence has been shown to demonstrate the 147 presence of *M. barkeri* in natural dolomite formation. However, this does not necessarily suggest 148 that M. barkeri was irrelevant to dolomite formation. Instead, a study with M. barkeri may 149 provide a chance to evaluate if dolomite formation is tied to certain microorganism species and 150 explore the geochemical and thermodynamic/kinetic "rules" which are common to all dolomite 151 (Pursher et al. 1994). 152

Instead of live cultures, we used purified bound EPS, non-living biomass and dead cell pellets (DCP) after bound EPS extraction for forced carbonate precipitation experiments at room temperature in calcite-seeded solutions. This procedure avoided the possible influence of complex components of typical cultures, which therefore makes it possible to clearly assess the contribution of bound EPS/non-living biomass/DCP to Ca-Mg carbonate precipitation. Our data demonstrated that bound EPS of *M. barkeri* can promote the incorporation of Mg into precipitating Ca-Mg carbonates and induce disordered dolomite precipitation. We propose a

- 160 plausible mechanism by which surface adsorbed bound EPS catalyzes Mg incorporation into
- 161 anhydrous Ca-Mg-carbonate. We also discussed the implications of this study to the long-lasting
- 162 "dolomite problem".

163 MATERIALS AND METHODS

164 Microorganisms and culture medium

The culture of *M. barkeri* strain MS (neotype strain) (DSM 800) was obtained from the 165 German Collection of Microorganisms (DSMZ). M. barkeri was cultivated in a near neutral pH 166 medium (pH 6.5-6.8) with the following composition (per liter): K₂HPO₄ 0.348 g, KH₂PO₄ 0.227 167 168 g, NH₄Cl 0.5 g, MgSO₄•7H₂O 0.5 g, CaCl₂•2H₂O 0.25 g, NaCl 2.25 g, FeSO₄•7H₂O 0.002 g, vitamin solution (Wolin et al. 1963) 10 ml, trace element solution (Whitman et al. 1982; Bock et 169 al. 1994) 1 ml, yeast extract (Difco) 2 g, NaHCO₃ 0.85 g, methanol 10 ml, cysteine-HCl•H₂O 0.3 170 g, and Na₂S•9H₂O 0.3 g. The medium was prepared anoxically under a N₂:CO₂ (80:20 v/v) 171 atmosphere. Methanol (50% v/v), NaHCO₃, Na₂S•9H₂O, MgSO₄•7H₂O, and CaCl₂•2H₂O were 172 added separately from sterile stock solutions after the medium was autoclaved. The stock 173 solution of NaHCO₃ was prepared under a N_2 :CO₂ (80:20 v/v) atmosphere, whereas those of 174 methanol, Na₂S•9H₂O, MgSO₄•7H₂O, and CaCl₂•2H₂O were under 100% N₂. 175

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177 Biomass collection

Biomass from *M. barkeri* was collected in the early stationary growth phase. The pH of 178 the culture when collecting biomass was \sim 5.2. No precipitates were observed in either cell-free 179 medium or live culture. Cultures were first centrifuged at 15,000 rpm for 30 min with a 180 Beckman-Coulter Avanti® J-E centrifuge. The supernatant was discarded and the biomass was 181 then washed with a N₂-sparged washing buffer containing all the inorganic ingredients in the 182 medium but not the organic ones and Na₂S. The purpose of such wash was to remove the 183 possible residue organics from the medium and other soluble microbial metabolites. The washing 184 buffer carried an ionic strength and composition close to that of the medium; otherwise some 185 bound EPS components might desorb and thus be washed away from the EPS matrix (Nielsen 186

and Jahn 1999). The washing buffer with the biomass was centrifuged at 15,000 rpm for 20 min and the supernatant was discarded. After that, ~40 ml of washing buffer was added to the washed biomass (~10-26 mg), which was then dialyzed against distilled de-ionized (DI) water for 24 h and collected for carbonate precipitation experiments. The biomass was non-metabolizing after dialysis since it was exposed to air during dialysis. To measure the dry weight of biomass in solution, a portion of the biomass solution was freeze dried at -50°C for 48 h.

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194 Extraction and quantification of bound EPS

195 The bound EPS of *M. barkeri* was extracted in the stationary growth phase. Biomass of M. barkeri was concentrated and washed following the same procedure above. Next, washing 196 buffer was added to the washed biomass to obtain a biomass concentration of 1.0 mg mL⁻¹ and 197 bound EPS was extracted from this biomass solution following a previously established 198 procedure developed for methanogenic sludges that utilizes formaldehyde and NaOH (Liu and 199 Fang 2002). It has been shown that this procedure can prevent the extracted bound EPS from the 200 contamination by intracellular substances (Nielsen and Jahn 1999; Liu and Fang 2002). The 201 detailed procedure is shown in **Fig. 1**. To obtain the concentration of bound EPS in solution, a 202 portion of dialyzed bound EPS solution was freeze dried at -50°C for 48 h for measuring the dry 203 weight. The residue DCP was also collected by adding ~40 ml of washing buffer to the residue 204 pellets and dialysis against DI water for 24 h. 205

The total carbohydrate content of bound EPS was measured using a modified phenolsulfuric acid method with glucose standards (Dubois et al. 1956). Polysaccharides (or other monomeric sugars in EPS) were first hydrolyzed to individual monosaccharides with H₂SO₄ (Pakulski and Benner 1992). To do this 1 mg of dry EPS was added into 1 mL of 12 M H₂SO₄ at

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210 room temperature for 2 h. Then 9 mL DI water was added to the slurry. Samples were briefly (3-5 s) ultrasonicated to promote the dissolution of the residue. A 5 mL aliquot of the solution was 211 pipetted into a 50 mL serum vial, crimp-sealed with Teflon liners and hydrolyzed at 100°C for 3 212 h. Then 1 mL aliquot was added in a test tube followed by 1 mL of phenol solution (5%) and 5 213 mL of 98% sulfuric acid. The tube was shaken well on a shaker. After 10 min, it was placed in a 214 water bath at 30°C for 20 min. The mixture was cooled and measured for absorbance at 490 nm 215 using an UV-Vis spectrophotometer (UV-mini 1240, Shimadzu Corp, Kyoto, Japan). The final 216 results were normalized by the dry weight of bound EPS. Bound EPS collected from three 217 218 batches of the culture was analyzed and duplicate aliquots were analyzed for each bound EPS sample. All experimental glassware used in these analyses was acid washed, rinsed with DI 219 water, and combusted at 550°C for 6 h to prevent the possible organic contamination. 220

Sugar monomer analyses were performed to investigate the monosaccharide composition 221 of the bound EPS of *M. barkeri*. This monomer composition of intact EPS was measured through 222 glycosyl analyses using gas chromatography combined mass spectrometry (GC/MS) of the per-223 O-trimethylsilyl (TMS) derivatives of the monosaccharide methyl glycosides produced from the 224 225 sample by acidic methanolysis. 400 µg of the sample was used for the analysis. 20 µg of inositol was added to the sample as an internal standard. Methyl glycosides were then prepared from the 226 dry sample by methanolysis in 1 M HCl in methanol at 80°C (18 h), followed by re-N-227 228 acetylation with pyridine and acetic anhydride in methanol (for detection of amino sugars). The sample was then per-O-trimethylsilylated by treatment with Tri-Sil (Pierce) at 80°C (0.5 h). 229 These procedures were carried out as previously described (York et al. 1986; Merkle and Poppe 230 1994). GC/MS analysis of the TMS methyl glycosides was performed on an Agilent 6890N GC 231

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interfaced to a 5975B MSD, using an Agilent DB-1 fused silica capillary column (30 m × 0.25
mm ID).

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Forced carbonate precipitation experiments with non-metabolizing biomass, bound EPSand DCP

All carbonate precipitation experiments were carried out at room temperature (22 °C) and 237 238 at least two duplicates were performed. All the glassware used in the synthesis was acid-washed, 239 rinsed with DI water and baked at 550°C for 6 h to prevent the possible organic contamination. 240 Solutions containing non-metabolizing *M. barkeri* biomass/bound EPS/DCP were diluted with DI water to obtain a range of bulk concentrations (see Table 1). Reagent grade CaCl₂•2H₂O and 241 242 MgCl₂•6H₂O powders were then added to the solutions. The concentration of CaCl₂ was fixed at 5 mM, whereas different concentrations of MgCl₂ were used (15 20 25 and 40 mM). The calcite 243 seeds were synthesized by mixing equal volumes of 500 mM CaCl₂ and 500 mM NaHCO₃. X-244 ray diffraction (XRD) analysis showed that calcite was the only phase in the synthetic seeds. 245 Scanning electron microscopy (SEM) examinations showed that the size of synthetic seeds was 246 usually several microns. The specific surface area of the seed crystals, as determined by multi-247 point N₂ BET method (Brunauer et al. 1938), was 0.2 m² g⁻¹. Synthetic calcite crystals with size 248 range of $10 \sim 20$ micrometers (0.2 g/L) were used as seeds for heterogeneous nucleation. 249 Previous studies showed that presence of calcite seed can promote the incorporation of Mg into 250 calcitic structure and inhibit aragonite precipitation (Berner 1975; Zhang et al. 2012a). 251 Experimental solutions containing biomass/bound EPS/DCP were ultrasonicated for 10 min to 252 suspend synthetic seeds and then left still for overnight so that solutions can be equilibrated with 253 atmospheric CO₂ and calcite seeds. After that, the pH of experimental solutions was measured as 254

the initial pH (**Table 1**). A geochemical program (PHREEQC) was utilized to calculate the starting chemical compositions of the control solutions (Parkhurst and Appelo 1999). The starting pH of the control solutions and PHREEQC calculations suggested that approximately 0.03-0.04 g/L out of the 0.2 g/L calcite seeds were dissolved and the control solutions were equilibrated with atmospheric CO₂.

Forced carbonate precipitation experiments were conducted with a NH₄HCO₃ drift-free 260 method (Lian et al. 2006). Crystallization reactions took place in a desiccator (dimensions $36 \times$ 261 36×41 cm). A number of Petri dishes containing experimental solutions were placed in the 262 desiccator, along with some NH₄HCO₃ powders (5 g for a total experimental solution volume of 263 500 mL) contained in separate Petri dishes. NH₃ and CO₂ produced from the decomposition of 264 NH₄HCO₃ diffused into experimental solutions where carbonate precipitation occurred. After 14 265 days, precipitates were collected by filtering solutions through a 0.22 µm membrane, rinsed with 266 DI water for several times, and air-dried. The concentrations of Ca^{2+} and Mg^{2+} in solutions were 267 also measured with inductively coupled plasma optical emission spectroscopy (ICP-OES, Varian 268 Vista-MPX, Australia) both before and after experiments at least in duplicates for each 269 270 experimental condition. Parallel control experiments were carried out with organic-free solutions. Detailed chemical conditions in carbonate precipitation experiments are listed in **Table 1**. 271

PHREEQC was also used to calculate the saturation index (SI) of control solutions with respect to disordered dolomite with ideal dolomite composition (50 mol% of MgCO₃). SI is defined as SI = Log(IAP/Ksp), whereas the IAP is the ion activity product of the dissolved mineral constituents and *K*sp is the equilibrium constant, that is $10^{-16.52}$ for disordered dolomite (Carpenter 1980). We did not calculate SI for experimental solutions containing nonmetabolizing biomass/EPS/DCP since they can bind metal cations as discussed above and we did

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not find data on their binding capacity. However, the SI of control solutions should be higherthan that of experimental solutions due to the presence of more available metal cations.

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281 XRD, SEM, transmission electron microscopy (TEM) and selected-area electron diffraction

282 (SAED) examinations

283 XRD analyses were carried out using a Rigaku Rapid II X-ray diffraction system (Mo $K\alpha$ 284 radiation). Samples were contained in thin-wall glass capillaries. Diffraction data were collected 285 on a 2-D image-plate detector. The two-dimensional images were then integrated to produce 286 conventional 20 vs. intensity patterns using Rigaku's 2DP software.

SEM samples were prepared by dispersing powders on carbon tapes and lightly carbon 287 288 coated (50-100 Å coating). SEM observations were performed using a LEO 1530 SEM equipped with energy-dispersive spectroscopy (EDS) capabilities to determine the solid-phase composition. 289 TEM and SAED measurements were done with an aberration-corrected FEG-(S)TEM 290 (Titan 80-200) which is capable of sub-Å-resolution structural and chemical imaging. Several 291 milligrams of sample were crushed between two glass slides with a few drops of ethanol. A drop 292 of the resulting suspension was placed on a holey carbon film supported by a TEM Cu grid and 293 air-dried. 294

Bulk average MgCO₃ content of synthetic Ca-Mg carbonates was measured based on the empirical curve correlating the shift of calcite (104) peak toward dolomite and MgCO₃ contents (Zhang et al. 2010). TEM-based X-ray EDS was also used to measure MgCO₃ contents of typical disordered dolomite samples (for method details, see Zhang *et al.* (2010)).

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300 Terminology of Ca-Mg carbonates

Ideal dolomite (CaMg(CO₃)₂, space group: $R\overline{3}$) has a crystal lattice consisting of alternating layers of Ca and Mg, separated by layers of CO₃, where Ca and Mg are present in equal proportions. However, very few, if any, sedimentary dolomites are truly stoichiometric CaMg(CO₃)₂ and are better represented as: Ca_(1+x)Mg_(1-x)(CO₃)₂. Most ancient dolomites are calcium-rich (Warren 2000). Mg²⁺ incorporation into calcitic structure results in the formation of various phases,

including: low-Mg calcite (LMC, space group: $R\overline{3}c$) with less than 4 mol% of MgCO₃, high Mg-307 calcite (HMC, space group: $R\overline{3}c$) with more than 4 mol% and less than 36 mol% of MgCO₃ 308 according to the proposed solvus between calcite and dolomite (Anovitz and Essene 1987), 309 disordered dolomite (with more than 36 mol% of MgCO₃ and typically Ca-rich with disordered 310 cations, i.e., instead of occurring in alternating cation layers, Ca^{2+} and Mg^{2+} ions are randomly 311 distributed; therefore, it has the same space group as calcite: $R\overline{3}c$), and dolomite (space group: 312 $R\overline{3}$) (Zhang et al. 2012a). Proto-dolomite is a poorly ordered dolomite, and generally Ca-rich 313 314 (Fang and Xu 2018).

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315 **RESULTS**

316 Saturation state of experimental solutions

After NH_4HCO_3 powder and the petri dishes containing solutions were put into the sealed 317 dessicator, the decomposition of NH₄HCO₃ started shortly and carbonate precipitation was 318 observed in 6-8 hours as indicated by the visual cloudiness in the solution. NH_4HCO_3 powders 319 320 were totally decomposed in ~ 12 hours. The pH measured one day after experiments started was fairly close to final pH measured after 14 days (Table 1). We used PHREEQC to calculate the SI 321 of control solutions with respect to disordered dolomite (SI_{dd}). However, we encountered several 322 323 problems during calculation, which prevented us from obtaining specific numbers for SI_{dd}. First, based on calculation, all the NH₃ and \sim 90% of CO₂ produced by NH₄HCO₃ decomposition 324 should be dissolved into the solution, which would produce a final pH of ~8.2. This calculation 325 is not supported by our observations that (1) the final pH was $\sim 9.2-9.3$ in all controls; (2) there 326 was still a strong smell of NH₃ gas after 14 days. Second, carbonate precipitation started before 327 all the NH_4HCO_3 was decomposed. To overcome this, we first used the final Ca^{2+} and Mg^{2+} 328 concentrations in control solutions after precipitation and a trial-and-error procedure to adjust the 329 dissolved NH₃ and CO₂ concentration to match the final pH. We found that calculated pH will 330 331 best fit the final measured pH if half of the NH_3 (31 mmol) and one quarter of CO_2 (15.8 mmol) produced by the decomposition of 5 g NH₄HCO₃ are dissolved into 500 mL solution. A SI_{dd} 332 calculated under these conditions was noted as the final SI_{dd} or the lowest SI_{dd} . Then since (1) the 333 334 pH of the solution was buffered by the dissolved NH₃ and CO₂ and (2) the pH measured one day after experiments started was fairly close to final pH, we assumed that when massive carbonate 335 precipitation started, the amounts of dissolved NH₃ and CO₂ were also 31 mmol and 15.8 mmol 336 for 500 mL solution, respectively. We used such numbers along with the measured initial Ca²⁺ 337

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and Mg^{2+} concentrations to calculate another SI_{dd} , noted as the initial SI_{dd} or the highest SI_{dd} . Calculated SI_{dd} is listed in **Table 1**. We want to emphasize that the SI_{dd} of control solutions should be higher than that of experimental solutions containing non-metabolizing biomass or bound EPS due to the presence of more available metal cations.

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343 Ca-Mg carbonates induced by non-metabolizing biomass, bound EPS, and DCP

XRD analyses of precipitated carbonates showed that non-metabolizing biomass can 344 promote Mg incorporation and Ca-Mg carbonate precipitation, especially in solutions with high 345 346 initial Mg:Ca ratios. For example, when the initial Mg:Ca ratio in solution was 3:1 or 4:1, the MgCO₃ contents of HMC precipitated in experimental solutions containing ~ 113 mg L⁻¹ of non-347 metabolizing biomass were not significantly higher than the carbonate precipitated in control 348 solutions (Fig. 2a, b; Fig. S1a, b). However, the catalytic effect of non-metabolizing biomass 349 became much more obvious with increased initial Mg:Ca ratios. Ca-Mg carbonates with ~41 and 350 ~45 mol% of MgCO₃ was precipitated in experimental solutions with an initial Mg:Ca ratio of 351 5:1 and 8:1, respectively (Fig. 2c, d; Table 1), while control solutions produced HMC with only 352 ~ 12 and ~ 18 mol% of MgCO₃, respectively (Fig. S1c, d). In addition, the non-metabolizing 353 biomass also inhibited the precipitation of aragonite (compare Fig. 2 and S1) which is generally 354 believed to compete with the crystallization of Ca-Mg carbonates (Lippmann 1973; Berner 1975). 355 The catalytic effect of non-metabolizing biomass was also supported by experiments with 356 357 different non-metabolizing biomass concentrations. The MgCO₃ contents in the synthetic Ca-Mg carbonates increased with non-metabolizing biomass concentration (Fig. 3, 4a, S2; Table 1). 358 359 Increased non-metabolizing biomass concentration also reduced the amount of aragonite in the precipitates (compare Fig. 2 and 3). With the highest concentration of biomass (~161 mg L^{-1}), 360

monohydrocalcite (CaCO₃•H₂O) also appeared (**Fig. 3**). This trend is consistent with previous studies on Ca-Mg carbonate precipitation in solutions with dissolved sulfide as catalyst, which showed that while dissolved sulfide can promote dolomite precipitation, over-dosed dissolved sulfide triggered monohydrocalcite crystallization (Zhang et al. 2012a).

Monohydrocalcite was the only crystalline phase identified in the carbonates precipitated in experimental solutions with 161mg L⁻¹ of non-metabolizing biomass and an initial Mg:Ca ratio of 8:1 (**Fig. 3d**). We suggest that an amorphous Ca-Mg carbonate phase was precipitated in addition to monohydrocalcite as indicated by the Mg removed from the solution (**Table 1**). The high concentrations of Ca^{2+} left in the solution may also reflect the formation of amorphous Ca-Mg carbonates since the amorphous Ca-Mg carbonates have higher solubility than the crystalline counterparts.

Precipitation experiments were carried out to assess the effect of bound EPS excreted by 372 M. barkeri on carbonate precipitation. Approximately 25 mg bound EPS could be extracted from 373 ~113 mg biomass. Our results clearly demonstrated the catalytic role of bound EPS in Ca-Mg 374 carbonate precipitation. The MgCO₃ contents of Ca-Mg carbonates induced by bound EPS were 375 generally higher than those by non-metabolizing biomass at same initial Mg:Ca ratio (Fig. 4b, 5; 376 Table 1). Ca-Mg carbonates with ~47 and ~48 mol% of MgCO₃ was precipitated in 377 experimental solutions with bound and an initial Mg:Ca ratio of 5:1 and 8:1, respectively (Fig. 378 5c, d; Table 1). 379

380 DCP slightly enhanced Mg incorporation compared to control experiments (Fig. 6; Table 381 1). However, the MgCO₃ contents of carbonates precipitated in DCP-bearing solutions were 382 generally much lower than those induced by bound EPS, except in experiments with an initial 383 Mg:Ca ratio of 8:1 which precipitated both HMC and Ca-Mg carbonates close to dolomite (Fig.

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6d). Precipitates in DCP-bearing solutions also contained more aragonite than those induced by
bound EPS. These data indicate that bound EPS was the active component in the nonmetabolizing biomass that promoted the precipitation of Ca-Mg carbonates close to dolomite
composition.

High-resolution SEM and TEM observations showed Ca-Mg carbonates close to 388 dolomite composition occurred as nano-crystals (~10-20 nm) overgrowing euhedral calcite seeds 389 (Fig. 7, 8a, b). On some calcite seeds, the overgrowing carbonates were not massive, which 390 therefore preserved the overall rhombohedral shape of calcite seed (Fig. 7a). However, in the 391 392 case of massive overgrowth, nano-crystals of Ca-Mg carbonate close to dolomite composition enclosed calcite seeds and formed clusters with different shapes and sizes (Fig. 7c). SAED 393 patterns showed that nano-crystals of Ca-Mg carbonate close to dolomite were not randomly 394 oriented, but rather followed the orientations of seed crystals and displayed low-angle grain 395 boundaries between neighboring nano-crystals (Fig. 8c). The [010]-zone axis SAED and fast 396 Fourier transform (FFT) patterns did not show supper-lattice reflections like (003) and $(\overline{1}05)$ 397 indicating the Ca-Mg cation order in dolomitic structure (Fig. 8c, d). Thus our synthetic product 398 was disordered dolomite. 399

400 Characterization of bound EPS

Our analyses showed that the total polysaccharide content of the bound EPS was 8.4 ± 0.5 wt%, that is, ~2.1 mg L⁻¹ out of ~25 mg L⁻¹ of bound EPS. The saccharide monomer analyses showed that mannose (~36 mol%), ribose (~30 mol%), rhamnose (~15 mol%), xylose (~10 mol%), and glucose (~7 mol%) were the dominant saccharide monomers of the polysaccharides in bound EPS (**Table 2**).

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407 **DISCUSSION**

408 EPS catalyzed crystallization of disordered dolomite

In this study, the bound EPS was extracted with a procedure which utilized formaldehyde 409 and NaOH. This procedure can minimize the contamination by intracellular substances since 410 formaldehyde can fix the cell, and thus prevent cell lysis, by reacting with the amino, hydroxyl, 411 carboxyl and sulfhydryl function groups of proteins and nucleic acids of the cell membrane 412 (Nielsen and Jahn 1999; Liu and Fang 2002). Furthermore, the addition of NaOH can cause 413 many charged groups, such as carboxylic groups in proteins and polysaccharides to be 414 dissociated, which results in a strong repulsion between the negatively charged EPS and thus 415 provides a higher water solubility of the compounds. One question, however, is whether the use 416 of NaOH and the possible reaction between bound EPS and formaldehyde played a critical role 417 in disordered dolomite precipitation. The effect of NaOH should be easily excluded because the 418 NaOH was removed from the bound EPS solution during dialysis. Regarding the effect of 419 formaldehyde, since non-metabolizing consortium biomass which was not processed with 420 formaldehyde also catalyzed similar disordered dolomite precipitation as bound PES did, the 421 422 contribution from formaldehydeshould not be significant.

Our experimental data showed that a higher initial Mg:Ca ratio in solutions generally results in higher MgCO₃ contents in precipitated Ca-Mg carbonates (**Fig. 4**), which is consistent with previous works on carbonate synthesis (Devery and Ehlmann 1981; Rushdi et al. 1992; Zhang et al. 2012b). Therefore, we may speculate that the catalytic effect of bound EPS was a result of increased Mg:Ca ratio in solution due to the binding of cations to bound EPS. That is, if fewer Mg²⁺ is bound to bound EPS than Ca²⁺, the initial Mg:Ca ratio in solution can be sharply

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429 increased. To address this question, knowledge of the binding capacity of bound EPS of M. barkeri is essential. Unfortunately, we did not find such published data. Nevertheless, previous 430 studies on the cation-binding capacity of soluble EPS extracted from SRB cultures may provide 431 some clue (Braissant et al. 2007). A Ca-binding capacity of 0.12-0.15 g_{Ca}/g_{EPS} has been 432 determined while no data for Mg is available. Although it is unlikely that the binding capacity of 433 434 the bound EPS we collected is the same as that of Braissant et al. (2007), these numbers can still offer some first-order estimations. For example, if we assume a binding capacity of 0.15 $g_{Ca} g_{EPS}$ 435 1 and 0 $g_{Mg} \ g_{EPS}{}^{-1}$, the ~25 mg L^{-1} of bound EPS will bind 0.094 mM Ca^{2+} , resulting in an 436 increase of Mg:Ca ratio from 3:1 to 3.06:1, from 4:1 to 4.08:1, from 5:1 to 5.1:1, from 8:1 to 437 8.15:1, respectively in our experiments. Such a small increase, which would be even smaller if 438 bound EPS binds less Ca²⁺ and also binds Mg²⁺, obviously cannot account for the huge 439 enhancement of Mg incorporation by bound EPS. 440

In addition, although a definitive calculation was not available in this case, the saturation index of our experimental solutions with respect to disordered dolomite was presumably high. However, control solutions with even higher saturation index still did not produce disordered dolomite. Therefore a high saturation index cannot explain the bound EPS-induced crystallization of disordered dolomite.

Here, we suggest that polysaccharides are likely to be one of the catalytic components of bound EPS since polysaccharides have been shown to mediate calcite and disordered dolomite precipitation (Braissant et al. 2003; Bosak and Newman 2005; Kawano and Hwang 2011; Zhang et al. 2012b). For example, Mg-rich disordered dolomite crystallized in solutions containing $\sim 200 \text{ mg L}^{-1}$ of agar (Zhang et al. 2012b). Kawano and Hwang (2011) showed that polysaccharides can promote the precipitation of calcite while inhibiting aragonite crystallization.

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452 Braissant et al. (2003) found that purified exopolysaccharide (xanthan EPS) exerted a strong influence on the morphology of precipitated calcite. Our analyses showed that the total 453 polysaccharide content of the bound EPS was 8.4 wt%, that is, ~2.1 mg L^{-1} out of ~25 mg L^{-1} of 454 bound EPS. This concentration is much lower than the amount of agar required to trigger 455 dolomite precipitation (Zhang et al. 2012b). This difference may be due simply to differences in 456 the properties of polysaccharides in bound EPS compared to agar, given that the Mg-457 incorporation capacities can vary significantly among different polysaccharides. For comparison, 458 disordered dolomite containing ~52 mol% of MgCO₃ crystallized in solutions containing ~200 459 mg L⁻¹ of agar and an initial Mg:Ca ratio of 8:1, whereas ~ 5 g L⁻¹ of carboxymethyl cellulose 460 was required to produce such disordered dolomite under the same solution phase conditions 461 (Zhang et al. 2012b). In this case, however, with only 25 mg L^{-1} of bound EPS, disordered 462 dolomite containing ~48 mol% was precipitated in solution with an initial Mg:Ca ratio of 8:1. 463 Therefore, it is not inconceivable that the small amounts of polysaccharides in the collected 464 bound EPS of *M. barkeri* can catalyze disordered dolomite precipitation. 465

In addition to polysaccharides, the existence of other catalytic components in bound EPS 466 should also be emphasized (Raz et al. 2000; Braissant et al. 2003; Gautret and Trichet 2005; 467 Stephenson et al. 2008; Wang et al. 2009). For example, Raz et al. (2000) found that polyacrylic 468 and polyaspartic acids can catalyze the crystallization of Ca-Mg carbonate with up to 34 mol% 469 of MgCO₃. Stephenson et al (2008) showed that a small amount of peptides in solution can 470 471 enhance the step velocity on the Ca-Mg carbonate growth hillock and slightly enhance Mg incorporation. Along analogous lines, carboxylated organic acids with a strong affinity for 472 binding Ca²⁺ compared to Mg²⁺ were shown to promote the formation of Mg-enriched 473 474 amorphous calcium carbonates (Wang et al. 2009). These possible catalytic components, along

with polysaccharides, may function synergistically to catalyze disordered dolomite nucleationand growth.

In previous studies, the role of EPS in calcium carbonate and dolomite formation was 477 mostly attributed to their ability to provide nucleation sites (Warthmann et al. 2000; Van Lith et 478 al. 2003b, a; Roberts et al. 2004; Aloisi et al. 2006; Bontognali et al. 2008; Kenward et al. 2009; 479 Deng et al. 2010; Bontognali et al. 2014). For example, Aloisi et al. (2006) found that bacterial 480 nucleation of calcium carbonate in a SRB culture was initiated at the bound EPS as nano-481 globules and these nano-globules calcified significantly only when released to the culture 482 483 medium. Aloisi et al. (2006) suggested that carbonate nucleation as nano-globules could be an important step in microbial carbonate precipitation. Here we approach the concept of "nucleation 484 sites" from a different point of view. As our SEM and TEM data showed, it was the calcite seeds 485 in our experiments that provided sites for the heterogeneous nucleation of disordered dolomite. 486 However, this does not suggest that bound EPS was not involved. Instead, as in previous studies 487 (Zhang et al. 2012a; Zhang et al. 2012b), we propose that the EPS adsorbed onto Ca-Mg 488 carbonate surfaces through hydrogen bonding between the H in the OH group of bound EPS and 489 the O in the CO_3^{2-} on carbonate surfaces. This in turn displaced surface water molecules which 490 would otherwise be associated with the hydration shell of Mg^{2+} , thereby facilitating Mg 491 incorporation and disordered dolomite crystallization. In other words, the nucleation sites for 492 disordered dolomite provided by calcite seeds will be functional only with the adsorbed bound 493 494 EPS. This hypothesis is supported by the saccharide monomer analyses of the polysaccharides in bound EPS. Our data showed that mannose (~36 mol%), ribose (~30 mol%), rhamnose (~15 495 mol%), xylose (~10 mol%), and glucose (~7 mol%) were the dominant saccharide monomers of 496 497 the polysaccharides in bound EPS. While there is no data for ribose and glucose, molecular

dynamic simulations showed that xylose, rhamnose and mannose have a stronger adsorption onto
calcite (104) surfaces than water_ (Yang et al. 2008).

Our experiments with non-metabolizing biomass also succeeded in precipitating disordered dolomite. We suggest that bound EPS of the non-metabolizing biomass may have easily desorbed from the cell surface into the solution during the dialysis and the precipitation experiments due to the possible lysis of cells. For example, ultrasonication, which is also a common procedure used for EPS extraction (Nielsen and Jahn 1999), was used to suspend added calcite seed crystals in experimental solutions. The released bound EPS then can be adsorbed onto carbonate surfaces to promote the dehydration of surface Mg^{2+} .

DCP was also shown to slightly enhance Mg incorporation compared to control 507 experiments (Fig. 6; Table 1). One possibility of the effect of DCP is that the bound EPS was 508 not totally striped from cell surface during extraction based on synthesized Ca-Mg-carbonates 509 (Fig. 10). In addition, the thick cell wall of *M. barkeri* was probably another source where this 510 catalytic effect was originated. Ultrathin sections showed a very thick (500 nm), amorphous 511 outer layer of *M. barkeri*'s cell wall which often appeared laminated (Zeikus and Bowen 1975). 512 Results of chemical analyses of isolated cell walls indicated that they consist of an acid 513 heteropolysaccharide that contains galactosamine, neutral sugars, and uronic acids (Kandler and 514 Hippe 1977; Balch et al. 1979). Therefore it is possible that the polysaccharides and some other 515 components of the cell wall also exerted catalytic effect during carbonate precipitation. 516

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518 Implications for the "dolomite problem"

519 As described above, our synthetic dolomite was finely crystalline, Ca-rich, and cation-disordered.

520 It is most likely that the inhibitory effect of Mg^{2+} ions on dolomite growth resulted in the

| 521 | extremely small size of dolomite crystals. Precipitation experiments with SRB also precipitated |
|-----|---|
| 522 | similar disordered dolomite (Bontognali et al. 2014). Modern dolomites associated with |
| 523 | methanogenesis are also generally finely crystalline, Ca-rich and poorly ordered (Pisciotto and |
| 524 | Mahoney 1981; Baker and Burns 1985; Compton and Siever 1986; Thornburg and Suess 1990; |
| 525 | Mazzullo 2000; Roberts et al. 2004). With deposition, poorly-crystallized disordered dolomite |
| 526 | can undergo maturation and recrystallization accompanied by increased cation ordering and |
| 527 | crystallinity, which can produce partially ordered proto-dolomite and eventually, fully ordered |
| 528 | dolomite (Lippmann 1973; Hardie 1987; Gregg et al. 1992; Vasconcelos and McKenzie 1997; |
| 529 | Warren 2000). Thus disordered dolomite induced by the bound EPS of methanogens can be |
| 530 | considered as a precursor to some sedimentary ordered dolomite. It was reported that the |
| 531 | dolomite or protodolomite formed in the methanogenesis zone will have high $\delta^{13}C$ values |
| 532 | (Mazzullo 2000; Greinert et al. 2001). The carbon isotope fractionation will enrich ¹³ C in |
| 533 | bicarbonate and deplete ¹³ C in methane. The dolomite formed in this environment will have high |
| 534 | δ^{13} C values, i.e., group A carbonate (Claypool and Kaplan 1974; Hennessy and Knauth 1985; |
| 535 | Burns and Baker 1987; Greinert et al. 2001; Blattler et al. 2015). The dolomite or calcian |
| 536 | dolomite formed in these areas generally contains small amounts of Fe(II) or FeCO ₃ . In extreme |
| 537 | cases, even siderite can precipitate. The existence of Fe^{2+} in dolomite also indicates a reducing |
| 538 | dolomitizing fluid that contains dissolved methane and other organics that are responsible for the |
| 539 | dolomite precipitation. It has been shown that fermenting bacteria can transfer electrons to Fe(III) |
| 540 | in sediments (Lovley 2000). The cooperative metabolism between methanogens and fermenting |
| 541 | bacteria results in CO_2 or dissolved HCO_3^- as the common product of all these reactions which |
| 542 | can result in Fe ²⁺ -bearing carbonate (Coleman and Raiswell 1993). |

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Although our results suggest a potential link between methanogens and dolomite 543 formation, caution must be exercised in extrapolating laboratory data to explaining natural 544 dolomite formation. First, no evidence has been shown to demonstrate the presence of *M. barkeri* 545 or any other methanogens in natural dolomite formation. While there is a possibility that the 546 correlation between natural dolomite formation and *M. barkeri* and other methanogens simply 547 548 has not yet been recognized, there may be some other factors that are noteworthy. First, bound EPS excreted by microorganisms in laboratory cultures is unlikely to be the same with the EPS 549 matrix produced by the same microorganisms in natural environments (Nielsen and Jahn 1999). 550 551 The production of EPS likely follows an ecophysiological response and the chemical composition and 3-D architecture of the EPS can be greatly influenced by the growth conditions 552 (Nielsen et al. 1997; Nielsen and Jahn 1999; Dupraz et al. 2009). In other words, considering the 553 different Mg-incorporation capacities of various polysaccharides, amino acids, proteins, 554 polycarboxylic acids, etc (see discussion above), the growth conditions in current experiments 555 might have a significant impact on the strength of the bound EPS in catalysis. In fact, as 556 mentioned above, previous laboratory studies showed that in high extracellular solute 557 concentrations *M barkeri* will not produce EPS (Stadtman and Barker 1951; Sowers et al. 1993; 558 Anderson et al. 2012). Therefore it is possible that in some natural environments, *M. barkeri* may 559 produce EPS matrix that might not have sufficient Mg-incorporation capacities to induce 560 (disordered) dolomite precipitation. 561

In addition, the pH and saturation index with respect to disordered dolomite in our forced carbonate precipitation experiments were presumably high compared to that of nature environments. While high pH and supersaturation conditions can be found in some evaporitic settings and saline or hypersaline environments (e.g., Wright 1999), dolomite precipitation was

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566 found in a methanogenic freshwater aquifer with neutral pH and close to equilibrium conditions (Roberts et al. 2004). Although a high pH and superstation cannot necessarily lead to disordered 567 dolomite precipitation (see discussion above), reasonable pH and supersaturation levels are still 568 required to carbonate precipitation (Dupraz et al. 2009; Gallagher et al. 2012). In fact, the pH of 569 our live *M. barkeri* culture dropped from an initial pH of 6.5-6.8 to \sim 5.2, which explains why no 570 carbonate precipitation was found in the live culture. Gallagher et al. (2012) found that the 571 utilization of different substrates by SRB resulted in different pH, alkalinity, and thereby 572 supersaturation. Since *M. barkeri* is versatile and can utilize a variety of methanogenic substrates, 573 the possibility exists that the metabolism of *M. barkeri* may result in different pH and 574 supersaturation conditions to induce the precipitation of different carbonate phases or even limit 575 carbonate precipitation. For example, in natural sediments, the major substrates for 576 methanogenesis are acetate (CH₃COO⁻) and H_2 (Lovley and Klug 1982) whose conversion to 577 methane, unlike methanol (CH₃OH), would be expected to elevate (e.g. through consumption of 578 H^+ or CO₂) rather than decrease solution pH: 579

580 $CH_3COO^- + H^+ \rightarrow CH_4 + CO_2$

581 $H_2 + CO_2 \rightarrow CH_4 + 2H_2O$

 $582 \qquad 4CH_3OH \rightarrow 3CH_4 + CO_2 + 2H_2O$

Regardless of these dissimilarities between our experimental conditions and natural environments, we propose that the physiochemical mechanism by which bound EPS of various SRB and methanogen species in laboratory cultures and EPS matrix in natural environments catalyze (disordered) dolomite precipitation should be similar. However, as discussed above, the strength in catalysis or the Mg-incorporation capacities of EPS can vary among different microorganism species or even in the same species when the growth conditions are different.

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591 IMPLICATIONS

In this study, we characterized the effect of bound EPS of the methanogen *M. barkeri* on 592 Ca-Mg carbonate precipitation. Forced carbonate precipitation experiments at room temperature 593 in calcite-seeded Ca/Mg carbonate solutions showed that non-metabolizing biomass of M. 594 Barkeri can enhance the Mg incorporation into calcitic structure and induce the crystallization of 595 disordered dolomite. Bound EPS was shown to be the component of non-metabolizing biomass 596 597 that catalyzed the precipitation of disordered dolomite. We propose a mechanism to explain the catalytic effect of bound EPS based on the adsorption of bound EPS to growing carbonate 598 surfaces through hydrogen bonding. This mechanism is also applicable to natural EPS-induced 599 dolomite formation. While our experimental conditions cannot completely mimic the natural 600 environments, this study contributes new insights into to the long-standing "dolomite problem", 601 especially the sedimentary dolomite with heavy δ^{13} C values. The formation of methane will 602 result in light carbon in methane phase and heavy carbon in aqueous carbonate that may be 603 incorporated into the carbonate mineral like dolomite in presence of the methanogens. 604

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Table 1 Chemical conditions employed in carbonate precipitation experiments with the non-metabolizing biomass, bound EPS, and 895

| 896 | DCP and | compositions | of synthetic | carbonates. | Errors represent | t standard deviation. |
|-----|---------|--------------|--------------|-------------|------------------|-----------------------|
| | | | | | | |

| | Initial Ca ²⁺ | Initial Mg ²⁺ | Initial | Final | Initial | Final Ca ²⁺ | Final Mg ²⁺ | Change in Mg ²⁺ | Final | MgCO ₃ content based on |
|---|--------------------------|--------------------------|---------|-------|--------------------|------------------------|------------------------|----------------------------|--------------------|------------------------------------|
| Experiments | (mM) | (mM) | pН | pН | SI _{dd} * | (mM) | (mM) | (mM) † | SI _{dd} * | $d_{104} \text{ (mol\%)}$ ‡ |
| | 5.2±0.2 | 15.0±0.2 | 7.8 | 9.3 | 5.29 | 0.05 ± 0.01 | 14.6±0.2 | 0.4 | 3.34 | 7±1 |
| Control | 5.2±0.2 | 20.6±0.3 | 7.8 | 9.2 | 5.36 | 0.09 ± 0.03 | 20.1±0.3 | 0.5 | 3.65 | 9.1±0.8 |
| Control | 5.2±0.2 | 24.8 ± 0.1 | 7.8 | 9.2 | 5.39 | $0.14{\pm}0.01$ | 24.2 ± 0.2 | 0.6 | 3.87 | 11.9±0.7 |
| | 5.2±0.2 | 40.5±0.3 | 7.9 | 9.2 | 5.44 | $0.19{\pm}0.01$ | 39.5±0.3 | 1.0 | 4.04 | 18±1 |
| Non- | 5.19±0.08 | 15.3±0.3 | 8.1 | 9.1 | - | 0.07 ± 0.03 | 15.0±0.4 | 0.3 | - | 5.1±0.6 |
| metabolizing | 5.19±0.08 | 21.7±0.4 | 8.2 | 9.1 | - | 0.09 ± 0.02 | 21.0±0.5 | 0.7 | - | 12±1.3 |
| Biomass | 5.19±0.08 | 26.7±0.7 | 8.2 | 9.1 | - | 0.13 ± 0.03 | 24.7±0.6 | 2.0 | - | 30±1.7 |
| $(65\pm7 \text{ mg L}^{-1})$ | 5.19±0.08 | 43.0±1.2 | 8.2 | 9.2 | - | 1.05 ± 0.08 | 40.6±0.6 | 2.4 | - | 32±2.1 |
| Non- | 5.1±0.1 | 15.7±0.3 | 8.0 | 9.1 | - | 0.14±0.04 | 15.3±0.3 | 0.4 | - | 5.7±0.1 |
| metabolizing | 5.1±0.1 | 20.9±0.4 | 8.1 | 9.1 | - | 0.17 ± 0.04 | 20.0±0.3 | 0.9 | - | 14 ± 1.1 |
| Biomass | 5.1±0.1 | 26.5±0.6 | 8.0 | 9.0 | - | 0.20 ± 0.02 | 23.3±0.5 | 3.2 | - | 41±1.3 |
| $(113\pm12 \text{ mg L}^{-1})$ | 5.1±0.1 | 43.0±1.0 | 8.1 | 9.1 | - | 0.88 ± 0.03 | 39.9±0.6 | 3.1 | - | 45±2.3 |
| Non- | 5.0±0.2 | 16.0 ± 0.2 | 7.2 | 9.0 | - | $0.14{\pm}0.01$ | 14.4±0.3 | 1.6 | - | 26±1.1 |
| metabolizing | 5.0±0.2 | 22.1±0.6 | 7.4 | 9.0 | - | 0.18±0.03 | 18.1±0.7 | 4.0 | - | 42.3±0.3 |
| Biomass | 5.0±0.2 | 27.2±0.6 | 7.5 | 9.0 | - | $0.24{\pm}0.04$ | 24.5±0.8 | 2.7 | - | 48.9±0.6 |
| $(161\pm17 \text{ mg L}^{-1})$ | 5.0±0.2 | 42.9±0.2 | 7.5 | 9.1 | - | 1.4±0.2 | 41.1±0.2 | 1.8 | - | - |
| | 5.0±0.1 | 15.2±0.5 | 6.7 | 9.1 | - | 0.05±0.01 | 14.9±0.4 | 0.3 | - | 5.4±0.6 |
| Bound EPS | 5.0±0.1 | 20.5±0.4 | 6.8 | 9.1 | - | 0.13 ± 0.01 | 18.1±0.7 | 2.4 | - | 30±2.9 |
| $(25\pm7 \text{ mg L}^{-1})$ | 5.0±0.1 | 26.1±0.7 | 7.0 | 9.0 | - | 0.16±0.02 | 22.7±0.7 | 3.4 | - | 47±2.1 |
| | 5.0±0.1 | 42.1±0.5 | 7.1 | 9.0 | - | 0.28 ± 0.01 | 38.1±0.6 | 4.0 | - | 48 ± 0.8 |
| DCD | 4.9±0.1 | 14.8±0.2 | 6.8 | 8.9 | - | 0.23±0.01 | 14.5±0.4 | 0.3 | - | 3.0±1.1 |
| $\frac{\text{DCP}}{(05+0 \text{ mg } \text{L}^{-1})}$ | 4.9±0.1 | 20.4 ± 0.1 | 6.8 | 8.9 | - | 0.12 ± 0.04 | 19.7±0.4 | 0.7 | - | 10±1.6 |
| $(95\pm9 \text{ mg L}^{-1})$ | 4.9±0.1 | 26.7±0.5 | 7.1 | 8.9 | - | 0.08 ± 0.01 | 25.9±0.5 | 0.8 | - | 15±2 |
| | 4.9±0.1 | 42.4±0.6 | 7.1 | 8.9 | - | 0.06 ± 0.01 | 39.1±0.3 | 3.3 | - | 25±2 and 46±2¶ |

*Initial and final SI with respect to disordered dolomite. See text for calculation details. †The difference between initial and final Mg²⁺ concentration.

[±]Molar content of MgCO₃ in synthetic Ca-Mg carbonates based on the Zhang et al. (2010) curve.

Two phases of Ca-Mg carbonate were identified in these precipitates.

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Table 2 Analysis of the monosaccharide composition in the polysaccharide fraction of bound 897

EPS. 898

| | mol% |
|-----------------|------|
| Mannose | 36 |
| Ribose | 30 |
| Rhamnose | 15 |
| Xylose | 10 |
| Glucose | 7 |
| Fructose | 1 |
| Glucuronic acid | 1 |

- 900 Figures' captions
 - 901 Fig. 1. XRD patterns of synthetic Ca–Mg carbonates induced by inactive biomass of *M*.
 - 902 *barkeri* (113±12 mg/L). Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond
 - to: A: aragonite; C: calcite seeds; D: Ca-dolomite; H: HMC.
 - 904 (a): HMC ($d_{104} = 3.0192$ Å, 5.6 mol% of MgCO₃) synthesized in inactive biomass-bearing
 - solutions (Mg:Ca = 3:1). A small amount of aragonite was identified in precipitates.
 - 906 (b): HMC ($d_{104} = 2.9926$ Å, 15.0 mol% of MgCO₃) synthesized in inactive biomass-bearing
 - solutions (Mg:Ca = 4:1). A small amount of aragonite was identified in precipitates.
 - 908 (c): Ca-dolomite ($d_{104} = 2.9388$ Å, 41.8 mol% of MgCO₃) synthesized in inactive biomass-
 - 909 bearing solutions (Mg:Ca = 5:1).
 - 910 (d): Ca-dolomite ($d_{104} = 2.9305$ Å, 46.7 mol% of MgCO₃) synthesized in inactive biomass-
 - 911 bearing solutions (Mg:Ca = 8:1).
 - 912
 - 913 Fig. 2. XRD patterns of synthetic HMC from control experiments with synthetic calcite
 - seeds (0.2 g/L). Peaks correspond to: A: aragonite; C: synthetic calcite; H: HMC.
 - 915 (a): Aragonite and HMC ($d_{104} = 3.0128$ Å, 8.4 mol% MgCO₃) synthesized in control solutions 916 (Mg:Ca = 3:1).
 - 917 (b): Aragonite and HMC (d_{104} = 3.0078 Å, 9.5 mol% MgCO₃) synthesized in control solutions 918 (Mg:Ca = 4:1).
 - 919 (c): Aragonite and HMC (d_{104} = 3.0027 Å, 11.5 mol% MgCO₃) synthesized in control solutions 920 (Mg:Ca = 5:1).
 - 921 (d): Aragonite and HMC (d_{104} = 2.9831 Å, 18.5 mol% MgCO₃) synthesized in control solutions
 - 922 (Mg:Ca = 8:1).

41

| 923 | Fig. 3. XRD patterns | s of svnthetic | Ca–Mg carbonates | s induced by inactiv | e biomass of <i>M</i> . |
|-----|----------------------|----------------|------------------|----------------------|-------------------------|
| 525 | 1 Stor man parton in | , or symmetric | Cu ng cu sonaces | , maacea og macei, | |

- 924 *barkeri* (65±7 mg/L). Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks
 - correspond to: A: aragonite; C: calcite seeds; H: HMC.
- 926 (a): HMC ($d_{104} = 3.0215$ Å, 4.7 mol% of MgCO₃) synthesized in inactive biomass-bearing
- solutions (Mg:Ca = 3:1). A small amount of aragonite was identified in precipitates.
- 928 (b): HMC ($d_{104} = 3.0004$ Å, 12.5 mol% of MgCO₃) synthesized in inactive biomass-bearing
- solutions (Mg:Ca = 4:1). A small amount of aragonite was identified in precipitates.
- 930 (c): HMC ($d_{104} = 2.9590$ Å, 28.6 mol% of MgCO₃) synthesized in inactive biomass-bearing
- 931 solutions (Mg:Ca = 5:1).
- 932 (d): HMC ($d_{104} = 2.9568$ Å, 30.0 mol% of MgCO₃) synthesized in inactive biomass-bearing
- 933 solutions (Mg:Ca = 8:1).
- 934

925

Fig. 4. XRD patterns of synthetic Ca–Mg carbonates induced by inactive biomass of *M*.

- *barkeri* (161±17 mg/L). Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond
 to: A: aragonite; C: calcite seeds; D: Ca-dolomite; H: HMC; M: monohydrocalcite.
- 938 (a): HMC ($d_{104} = 2.9611$ Å, 26.7 mol% of MgCO₃) synthesized in inactive biomass-bearing
- solutions (Mg:Ca = 3:1). A small amount of monohydrocalcite was observed in precipitates.
- 940 (b): Ca-dolomite ($d_{104} = 2.9373$ Å, 42.7 mol% of MgCO₃) synthesized in inactive biomass-
- bearing solutions (Mg:Ca = 4:1). Monohydrocalcite was identified in precipitates.
- 942 (c): Ca-dolomite ($d_{104} = 2.9256$ Å, 49.3 mol% of MgCO₃) synthesized in inactive biomass-
- bearing solutions (Mg:Ca = 5:1). Monohydrocalcite was identified in precipitates.
- 944 (d): No crystalline Ca–Mg carbonate precipitation was observed in inactive biomass-bearing
- solutions (Mg:Ca = 8:1). Monohydrocalcite was identified in precipitates.
- 946

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- 947 Fig. 5. The MgCO₃ contents in synthetic Ca–Mg carbonates as a function of inactive
- 948 biomass concentration and initial Mg:Ca ratio in experimental solutions.
- 949

950 Fig. 6. XRD patterns of synthetic Ca–Mg carbonates induced by EPS (27±5 mg/L) of *M*.

- **barkeri.** Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond to: A: aragonite;
- 952 C: calcite seeds; D: Ca-dolomite; H: HMC; M: Monohydrocalcite.
- (a): HMC (d_{104} = 3.0186 Å, 5.8 mol% of MgCO₃) synthesized in EPS-bearing solutions (Mg:Ca
- = 3:1). A small amount of aragonite was identified in precipitates.
- 955 (b): HMC ($d_{104} = 2.9537$ Å, 32.1 mol% of MgCO₃) synthesized in EPS-bearing solutions
- 956 (Mg:Ca = 4:1).
- 957 (c): Ca-dolomite ($d_{104} = 2.9284$ Å, 48.0 mol% of MgCO₃) synthesized in EPS-bearing solutions
- 958 (Mg:Ca = 5:1). A small amount of monohydrocalcite was identified in precipitates.
- (d): Ca-dolomite (d_{104} = 2.9281 Å, 48.2 mol% of MgCO₃) synthesized in EPS-bearing solutions
- 960 (Mg:Ca = 8:1). A small amount of monohydrocalcite was identified in precipitates.
- 961

962 Fig. 7. Comparison of the catalytic strength of inactive biomass (113±12 mg/L) and EPS

- 963 (25±7 mg/L). Approximately 25 mg EPS can be extracted from 113 mg biomass
- 964 following our extraction procedure.
- 965
- 966
- 967
- 968
- 969
- 970

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971 Fig. 8. SEM images of synthetic dolomite. "C" and "D" stand for calcite seed and

972 precipitated dolomite, respectively.

973 (a): SEM image of dolomite nano-crystals synthesized in EPS-bearing solutions growing on the

- surface of a euhedral calcite seed. Arrows indicate precipitated dolomite.
- 975 (b): A close up of the image in (a) showing that dolomite occurred as extremely small nano-
- 976 crystals.
- 977 (c): SEM image showing a calcite seed enclosed by dolomite nano-crystals.
- 978
- 979 Fig. 9. TEM examinations of synthetic dolomite.
- 980 (a): TEM image of dolomite nano-crystals synthesized in EPS-bearing solutions growing on the981 surface of a calcite seed. Arrows indicate precipitated dolomite.
- 982 (b): TEM image of dolomite synthesized in EPS-bearing solutions. Dolomite occurred as nano-
- crystals with a size of ~10-20 nm. Inset is an X-ray EDS spectrum of the synthetic dolomite that
 contained ~48 mol% of MgCO₃.
- 985 (c): SAED pattern of the dolomite in (b). The diffraction arcs suggested that there were low-
- angle grain boundaries among dolomite nano-crystals. No super-lattice reflections like (003) and
- 987 $(\bar{1}05)$ were observed on the SAED pattern. Therefore, the synthetic dolomite was disordered.
- 988 (d): High-resolution TEM image from synthetic dolomite. No supper-lattice fringes like (003)
- and $(\bar{1}05)$ were observed. Inset is a [010] zone axis FFT pattern of the image. No super-lattice
- 990 reflections were shown on the FFT indicating that the synthetic dolomite was fully disordered.

991

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993 Fig. 10. XRD patterns of synthetic Ca–Mg carbonates induced by DCP after EPS

- extraction. Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond to: A:
 aragonite; C: calcite seeds; D: Ca-dolomite; H: HMC; M: monohydrocalcite.
- 996 (a): Mg-calcite ($d_{104} = 3.0301$ Å, 2.2 mol% of MgCO₃) synthesized in DCP-bearing solutions
- 997 $(87\pm9 \text{ mg/L}, \text{Mg:Ca} = 3:1)$. Aragonite was identified in precipitates.
- 998 (b): Mg-calcite ($d_{104} = 30035$ Å, 11.3 mol% of MgCO₃) synthesized in DCP-bearing solutions
- 999 $(87\pm9 \text{ mg/L}, \text{Mg:Ca} = 4:1)$. Aragonite was identified in precipitates.
- 1000 (c): Mg-calcite ($d_{104} = 2.9980$ Å, 13.2 mol% of MgCO₃) synthesized in DCP-bearing solutions
- 1001 (87 ± 9 mg/L, Mg:Ca = 5:1). Aragonite was identified in precipitates.
- 1002 (d): Two phases of Ca–Mg carbonates were precipitated in DCP-bearing solutions (87±9 mg/L,
- 1003 Mg:Ca = 8:1). One is HMC (d_{104} = 2.9657 Å, 25.4 mol% of MgCO₃); the other Ca-dolomite (d_{104}
- $1004 = 2.9303 \text{ Å}, 46.9 \text{ mol}\% \text{ of MgCO}_3$). Monohydrocalcite was identified in precipitates.

1005

- 1 Table 1 Chemical conditions employed in carbonate precipitation experiments with the non-
- 2 metabolizing biomass, bound EPS, and DCP and compositions of synthetic carbonates. Errors
- 3 represent standard deviation.

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| Even oniver out o | Initial Ca ²⁺ | Initial Mg ²⁺ | Initial | Final | Initial | Final Ca ²⁺ | Final Mg ²⁺ | Change in Mg ²⁺ | Final | MgCO ₃ content based on |
|--------------------------------|--------------------------|--------------------------|---------|-------|--------------------|------------------------|------------------------|----------------------------|--------------------|------------------------------------|
| Experiments | (mM) | (mM) | pН | pН | SI _{dd} * | (mM) | (mM) | (mM)† | SI _{dd} * | $d_{104} \text{ (mol\%)}$ |
| | 5.2±0.2 | 15.0±0.2 | 7.8 | 9.3 | 5.29 | 0.05 ± 0.01 | 14.6±0.2 | 0.4 | 3.34 | 7±1 |
| Control | 5.2±0.2 | 20.6±0.3 | 7.8 | 9.2 | 5.36 | 0.09 ± 0.03 | 20.1±0.3 | 0.5 | 3.65 | 9.1±0.8 |
| Control | 5.2±0.2 | 24.8 ± 0.1 | 7.8 | 9.2 | 5.39 | $0.14{\pm}0.01$ | 24.2 ± 0.2 | 0.6 | 3.87 | 11.9±0.7 |
| | 5.2±0.2 | 40.5±0.3 | 7.9 | 9.2 | 5.44 | $0.19{\pm}0.01$ | 39.5±0.3 | 1.0 | 4.04 | 18±1 |
| Non- | 5.19±0.08 | 15.3±0.3 | 8.1 | 9.1 | - | 0.07 ± 0.03 | 15.0±0.4 | 0.3 | - | 5.1±0.6 |
| metabolizing | 5.19 ± 0.08 | 21.7±0.4 | 8.2 | 9.1 | - | 0.09 ± 0.02 | 21.0±0.5 | 0.7 | - | 12±1.3 |
| Biomass | 5.19 ± 0.08 | 26.7±0.7 | 8.2 | 9.1 | - | 0.13 ± 0.03 | 24.7±0.6 | 2.0 | - | 30±1.7 |
| $(65\pm7 \text{ mg L}^{-1})$ | 5.19±0.08 | 43.0±1.2 | 8.2 | 9.2 | - | 1.05 ± 0.08 | 40.6±0.6 | 2.4 | - | 32±2.1 |
| Non- | 5.1±0.1 | 15.7±0.3 | 8.0 | 9.1 | - | 0.14±0.04 | 15.3±0.3 | 0.4 | - | 5.7±0.1 |
| metabolizing | 5.1±0.1 | 20.9±0.4 | 8.1 | 9.1 | - | 0.17±0.04 | 20.0±0.3 | 0.9 | - | 14±1.1 |
| Biomass | 5.1±0.1 | 26.5±0.6 | 8.0 | 9.0 | - | 0.20 ± 0.02 | 23.3±0.5 | 3.2 | - | 41±1.3 |
| $(113\pm12 \text{ mg L}^{-1})$ | 5.1±0.1 | 43.0±1.0 | 8.1 | 9.1 | - | 0.88 ± 0.03 | 39.9±0.6 | 3.1 | - | 45±2.3 |
| Non- | 5.0±0.2 | 16.0±0.2 | 7.2 | 9.0 | - | 0.14±0.01 | 14.4±0.3 | 1.6 | - | 26±1.1 |
| metabolizing | 5.0±0.2 | 22.1±0.6 | 7.4 | 9.0 | - | 0.18 ± 0.03 | 18.1±0.7 | 4.0 | - | 42.3±0.3 |
| Biomass | 5.0±0.2 | 27.2±0.6 | 7.5 | 9.0 | - | $0.24{\pm}0.04$ | 24.5±0.8 | 2.7 | - | 48.9±0.6 |
| $(161\pm17 \text{ mg L}^{-1})$ | 5.0±0.2 | 42.9±0.2 | 7.5 | 9.1 | - | 1.4±0.2 | 41.1±0.2 | 1.8 | - | - |
| | 5.0±0.1 | 15.2±0.5 | 6.7 | 9.1 | - | 0.05±0.01 | 14.9±0.4 | 0.3 | - | 5.4±0.6 |
| Bound EPS | 5.0±0.1 | 20.5±0.4 | 6.8 | 9.1 | - | 0.13±0.01 | 18.1±0.7 | 2.4 | - | 30±2.9 |
| $(25\pm7 \text{ mg L}^{-1})$ | 5.0±0.1 | 26.1±0.7 | 7.0 | 9.0 | - | 0.16±0.02 | 22.7±0.7 | 3.4 | - | 47±2.1 |
| | 5.0±0.1 | 42.1±0.5 | 7.1 | 9.0 | - | 0.28±0.01 | 38.1±0.6 | 4.0 | - | $48{\pm}0.8$ |
| DCD | 4.9±0.1 | 14.8±0.2 | 6.8 | 8.9 | - | 0.23±0.01 | 14.5±0.4 | 0.3 | - | 3.0±1.1 |
| DCP | 4.9±0.1 | 20.4±0.1 | 6.8 | 8.9 | - | 0.12 ± 0.04 | 19.7±0.4 | 0.7 | - | 10±1.6 |
| $(95\pm9 \text{ mg L}^{-1})$ | 4.9±0.1 | 26.7±0.5 | 7.1 | 8.9 | - | 0.08 ± 0.01 | 25.9±0.5 | 0.8 | - | 15±2 |
| | 4.9±0.1 | 42.4±0.6 | 7.1 | 8.9 | - | 0.06 ± 0.01 | 39.1±0.3 | 3.3 | - | 25 ± 2 and 46 ± 2 ¶ |

*Initial and final SI with respect to disordered dolomite. See text for calculation details.
†The difference between initial and final Mg²⁺ concentration.
‡Molar content of MgCO₃ in synthetic Ca-Mg carbonates based on the Zhang et al. (2010) curve.

¶Two phases of Ca-Mg carbonate were identified in these precipitates.

3

5 Table 2 Analysis of the monosaccharide composition in the polysaccharide fraction of bound

6 EPS.

| | mol% |
|-----------------|------|
| Mannose | 36 |
| Ribose | 30 |
| Rhamnose | 15 |
| Xylose | 10 |
| Glucose | 7 |
| Fructose | 1 |
| Glucuronic acid | 1 |

Figures

Fig. 1. XRD patterns of synthetic Ca–Mg carbonates induced by inactive biomass of *M. barkeri* (113±12 mg/L). Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond to: A: aragonite; C: calcite seeds; D: Ca-dolomite; H: HMC.

(a): HMC ($d_{104} = 3.0192$ Å, 5.6 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 3:1). A small amount of aragonite was identified in precipitates.

(b): HMC ($d_{104} = 2.9926$ Å, 15.0 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 4:1). A small amount of aragonite was identified in precipitates.

(c): Ca-dolomite ($d_{104} = 2.9388$ Å, 41.8 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 5:1).

(d): Ca-dolomite (d_{104} = 2.9305 Å, 46.7 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 8:1).



Fig. 2. XRD patterns of synthetic HMC from control experiments with synthetic calcite seeds

(0.2 g/L). Peaks correspond to: A: aragonite; C: synthetic calcite; H: HMC.

(a): Aragonite and HMC ($d_{104} = 3.0128$ Å, 8.4 mol% MgCO₃) synthesized in control solutions

(Mg:Ca = 3:1).

(b): Aragonite and HMC ($d_{104} = 3.0078$ Å, 9.5 mol% MgCO₃) synthesized in control solutions

(Mg:Ca = 4:1).

(c): Aragonite and HMC ($d_{104} = 3.0027$ Å, 11.5 mol% MgCO₃) synthesized in control solutions (Mg:Ca = 5:1).

(d): Aragonite and HMC ($d_{104} = 2.9831$ Å, 18.5 mol% MgCO₃) synthesized in control solutions





Fig. 3. XRD patterns of synthetic Ca–Mg carbonates induced by inactive biomass of *M*. *barkeri* (65±7 mg/L). Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond to: A: aragonite; C: calcite seeds; H: HMC.

(a): HMC ($d_{104} = 3.0215$ Å, 4.7 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 3:1). A small amount of aragonite was identified in precipitates.

(b): HMC ($d_{104} = 3.0004$ Å, 12.5 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 4:1). A small amount of aragonite was identified in precipitates.

(c): HMC ($d_{104} = 2.9590$ Å, 28.6 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 5:1).

(d): HMC ($d_{104} = 2.9568$ Å, 30.0 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 8:1).



Fig. 4. XRD patterns of synthetic Ca-Mg carbonates induced by inactive biomass of M.

barkeri (161±17 mg/L). Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond

to: A: aragonite; C: calcite seeds; D: Ca-dolomite; H: HMC; M: monohydrocalcite.

(a): HMC ($d_{104} = 2.9611$ Å, 26.7 mol% of MgCO₃) synthesized in inactive biomass-bearing

solutions (Mg:Ca = 3:1). A small amount of monohydrocalcite was observed in precipitates.

(b): Ca-dolomite ($d_{104} = 2.9373$ Å, 42.7 mol% of MgCO₃) synthesized in inactive biomass-bearing

solutions (Mg:Ca = 4:1). Monohydrocalcite was identified in precipitates.

(c): Ca-dolomite ($d_{104} = 2.9256$ Å, 49.3 mol% of MgCO₃) synthesized in inactive biomass-bearing solutions (Mg:Ca = 5:1). Monohydrocalcite was identified in precipitates.

(d): No crystalline Ca–Mg carbonate precipitation was observed in inactive biomass-bearing solutions (Mg:Ca = 8:1). Monohydrocalcite was identified in precipitates.



Fig. 5. The MgCO₃ contents in synthetic Ca–Mg carbonates as a function of inactive biomass concentration and initial Mg:Ca ratio in experimental solutions.



Fig. 6. XRD patterns of synthetic Ca-Mg carbonates induced by EPS (27±5 mg/L) of M.

barkeri. Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond to: A: aragonite;

C: calcite seeds; D: Ca-dolomite; H: HMC; M: Monohydrocalcite.

(a): HMC (d_{104} = 3.0186 Å, 5.8 mol% of MgCO₃) synthesized in EPS-bearing solutions (Mg:Ca =

3:1). A small amount of aragonite was identified in precipitates.

(b): HMC (d_{104} = 2.9537 Å, 32.1 mol% of MgCO₃) synthesized in EPS-bearing solutions (Mg:Ca

= 4:1).

(c): Ca-dolomite ($d_{104} = 2.9284$ Å, 48.0 mol% of MgCO₃) synthesized in EPS-bearing solutions

(Mg:Ca = 5:1). A small amount of monohydrocalcite was identified in precipitates.

(d): Ca-dolomite ($d_{104} = 2.9281$ Å, 48.2 mol% of MgCO₃) synthesized in EPS-bearing solutions (Mg:Ca = 8:1). A small amount of monohydrocalcite was identified in precipitates.



Fig. 7. Comparison of the catalytic strength of inactive biomass (113±12 mg/L) and EPS

(25±7 mg/L). Approximately 25 mg EPS can be extracted from 113 mg biomass following our extraction procedure.



Fig. 8. SEM images of synthetic dolomite. "C" and "D" stand for calcite seed and precipitated dolomite, respectively.

(a): SEM image of dolomite nano-crystals synthesized in EPS-bearing solutions growing on the surface of a euhedral calcite seed. Arrows indicate precipitated dolomite.

(b): A close up of the image in (a) showing that dolomite occurred as extremely small nanocrystals.

(c): SEM image showing a calcite seed enclosed by dolomite nano-crystals.



Fig. 9. TEM examinations of synthetic dolomite.

(a): TEM image of dolomite nano-crystals synthesized in EPS-bearing solutions growing on the surface of a calcite seed. Arrows indicate precipitated dolomite.

(b): TEM image of dolomite synthesized in EPS-bearing solutions. Dolomite occurred as nanocrystals with a size of ~10-20 nm. Inset is an X-ray EDS spectrum of the synthetic dolomite that contained ~48 mol% of MgCO₃.

(c): SAED pattern of the dolomite in (b). The diffraction arcs suggested that there were low-angle grain boundaries among dolomite nano-crystals. No super-lattice reflections like (003) and ($\overline{1}05$) were observed on the SAED pattern. Therefore, the synthetic dolomite was disordered.

(d): High-resolution TEM image from synthetic dolomite. No supper-lattice fringes like (003) and ($\overline{1}05$) were observed. Inset is a [010] zone axis FFT pattern of the image. No super-lattice reflections were shown on the FFT indicating that the synthetic dolomite was fully disordered.



Fig. 10. XRD patterns of synthetic Ca–Mg carbonates induced by DCP after EPS extraction.

Synthetic calcite (0.2 g/L) was used as seed crystals. Peaks correspond to: A: aragonite; C: calcite

seeds; D: Ca-dolomite; H: HMC; M: monohydrocalcite.

(a): Mg-calcite ($d_{104} = 3.0301$ Å, 2.2 mol% of MgCO₃) synthesized in DCP-bearing solutions

 $(87\pm9 \text{ mg/L}, \text{Mg:Ca} = 3:1)$. Aragonite was identified in precipitates.

(b): Mg-calcite ($d_{104} = 30035$ Å, 11.3 mol% of MgCO₃) synthesized in DCP-bearing solutions

 $(87\pm9 \text{ mg/L}, \text{Mg:Ca} = 4:1)$. Aragonite was identified in precipitates.

(c): Mg-calcite ($d_{104} = 2.9980$ Å, 13.2 mol% of MgCO₃) synthesized in DCP-bearing solutions

 $(87\pm9 \text{ mg/L}, \text{Mg:Ca} = 5:1)$. Aragonite was identified in precipitates.

(d): Two phases of Ca-Mg carbonates were precipitated in DCP-bearing solutions (87±9 mg/L,

Mg:Ca = 8:1). One is HMC (d_{104} = 2.9657 Å, 25.4 mol% of MgCO₃); the other Ca-dolomite (d_{104} = 2.9303 Å, 46.9 mol% of MgCO₃). Monohydrocalcite was identified in precipitates.

