P/Ca in carbonates as a proxy for alkalinity and phosphate levels

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Abstract

Understanding mechanisms, rates, and drivers of carbonate formation provides insight into the chemical evolution of Earth's oceans and atmosphere. We paired geological observations with elemental and isotope geochemistry to test potential proxies for calcium-to-alkalinity ratios (Ca:ALK). Across diverse carbonate facies from Pleistocene closed-basin lakes in Owens Valley, CA, we observed less $\delta 44/40$ Ca variation than theoretically predicted (>0.75 these systems. Carbonate clumped isotope disequilibria implied rapid carbonate growth—kinetic isotope effects, combined with the diverse carbonate minerals present, complicated the interpretation of [?] $\delta 44/40$ Ca as a paleo-alkalinity proxy. In contrast, we observed that high phosphate concentrations are recorded by shoreline and lake bottom carbonates formed in eleven Pleistocene lakes at orders of magnitude greater concentrations than marine carbonates. Because the maximum phosphate content of water depends on Ca:ALK, we propose that carbonate P/Ca can inform phosphate levels and thereby Ca:ALK of aqueous environments in the carbonate record.

P/Ca in Carbonates as a Proxy for Alkalinity and Phosphate Levels

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Key Points:

- Calcium-to-alkalinity ratios (Ca:ALK) in water determine rates and mechanisms of carbonate salt formation
- Shoreline tufa carbonate facies is a useful indicator of low Ca:ALK aqueous chemistry in the rock record
- Calcium isotope proxy for Ca:ALK in these materials is complicated by rate and mineralogical impacts on carbonate-water fractionation
- Carbonate P/Ca provides another means to assess Ca:ALK via the phosphate content of ancient aqueous environments

1 Abstract

2 Understanding mechanisms, rates, and drivers of past carbonate formation provides insight into 3 the chemical evolution of Earth's oceans and atmosphere. We paired geological observations with 4 elemental and isotope geochemistry to test potential proxies for calcium-to-alkalinity ratios 5 (Ca:ALK). Across diverse carbonate facies from Pleistocene closed-basin lakes in Owens Valley, CA, we observed less $\delta^{44/40}$ Ca variation than theoretically predicted (>0.75‰) for the very low 6 7 Ca:ALK in these systems. Carbonate clumped isotope disequilibria implied rapid carbonate growth-kinetic isotope effects, combined with the diverse carbonate minerals present, 8 complicated the interpretation of $\delta^{44/40}$ Ca as a paleo-alkalinity proxy. In contrast, we observed that 9 10 the high phosphate concentrations are recorded by shoreline and lake bottom carbonates formed 11 in eleven Pleistocene lakes at orders of magnitude greater concentrations than in marine 12 carbonates. Because the maximum phosphate content of water depends on Ca:ALK, we propose 13 that carbonate P/Ca can inform phosphate levels and thereby Ca:ALK of aqueous environments in 14 the carbonate record.

15

16 Plain Language Summary

17 Carbonate minerals record information about the local environments in which they form, for 18 example along the margins of a lake or on the seabed, as well as the aqueous and atmospheric 19 chemistry at the time of mineralization (e.g. pCO₂, pH). This information is recorded in both the 20 chemical signatures and textures carbonate rocks acquire during precipitation. Formation of calcium carbonate requires both a source of calcium (Ca^{2+}) and carbonate (CO_3^{2-} , or, carbonate 21 22 alkalinity [ALK]). The ratio of calcium to alkalinity (Ca:ALK) in lake or ocean water influences 23 carbonate saturation state, the rate of carbonate formation, the textures that carbonate will develop, 24 and the chemical signatures recorded by the carbonate mineral. In this study, we tested several 25 approaches to identify very low Ca:ALK chemistry in modern and ancient lakes. We found that 26 water with extremely low levels of calcium and high levels of alkalinity forms carbonates with a 27 characteristic "tufa" texture in stratigraphic positions tied to riverine and groundwater sources of 28 Ca^{2+} . Moreover, the phosphate concentrations in these rocks were orders of magnitude higher than 29 carbonates that precipitate under the high Ca:ALK conditions of modern oceans. Together the 30 results illustrated that identification of carbonate tufa textures in the rock record and phosphate 31 measurements of carbonate rocks can be used to study the ancient environments and mechanisms

32 by which carbonate rocks formed.

33 **1 Introduction**

Geochemical signatures recorded by carbonate minerals are used to reconstruct ancient environmental and atmospheric conditions. However, the rates and mechanisms of carbonate formation, which are set by the geochemistry (and often biology) of the depositional environment, impact those materials. It is often challenging to determine the processes and environments responsible for carbonate precipitation in the geological record—particularly in strata devoid of fossils, as occurs in Precambrian sedimentary basins.

40 Broadly, carbonate formation is determined by a balance of cations (e.g. Ca^{2+} or Mg^{2+}) and 41 the carbonate anion ($CO_3^{2^-}$), which scales with carbonate alkalinity (ALK = $HCO_3^{-} + 2CO_3^{2^-}$) in 42 systems where pCO₂ is largely fixed and carbonate is the primary buffer, e.g. seawater and most 43 lakes. Carbonate formation typically occurs between two end-member regimes: high alkalinity in excess of Ca²⁺ (ALK>>Ca, as in alkaline lakes), and vice versa (modern marine). Processes that source or consume cations (herein generalized as Ca²⁺) and alkalinity set the carbonate saturation state of a solution ($\Omega = \frac{[Ca^{2+}][ALK]}{K_{sp}*}$, where K_{sp}^* is the solubility product of CaCO₃ at a given temperature, pressure, and composition), and determine whether carbonate is likely to precipitate ($\Omega > 1$) or dissolve ($\Omega < 1$). The continuum of calcium-to-alkalinity ratios (Ca:ALK) that produce carbonate phases in natural environments varies by orders of magnitude. Where a given environment sits on that continuum impacts how carbonates get made.

51 Microbial metabolisms can contribute to carbonate formation and dissolution through the 52 production and consumption of alkalinity and dissolved inorganic carbon (DIC; Vasconcelos & 53 McKenzie, 1997; Folk & Chafetz, 2000). Photosynthesis and aerobic respiration increase and 54 decrease Ω , respectively, by consuming and producing DIC (Δ DIC) with little to no change to 55 ALK (Δ ALK; Bergmann et al., 2013; Higgins et al., 2009). The effect of these processes on Ω , 56 however, depends on the Ca:ALK of the environment. In high Ca:ALK environments (modern 57 marine Ca:ALK=4.1), Ω depends most on the production and consumption of DIC by microbial 58 carbon cycling because Ca^{2+} is not limiting. In contrast, Ca^{2+} -starved, hyperalkaline environments, like Mono Lake, California (Ca:ALK= 10^{-4}), are less sensitive to localized Δ DIC because Δ ALK 59 60 is relatively insignificant compared to total ALK. Carbonate production from very low Ca:ALK water is controlled by processes that impact Ca²⁺ supply and cycling. For example, nearly all 61 62 carbonate formed in Mono Lake and other alkaline, closed-basin Pleistocene lakes can be 63 described as shoreline-proximal facies (e.g. tufa towers and shoreline crusts) wherein carbonate production is tied to external Ca^{2+} input from rivers and emergent groundwaters. In these settings, 64 distal lake bottom sediments tend to be comparatively carbonate-poor. The shoreline-associated 65 66 carbonate facies characteristic of low Ca:ALK systems can be useful geological indicators of this 67 carbonate production regime, as long as they are preserved with sufficient stratigraphic and 68 sedimentologic detail. However, diagnostic shoreline facies are rarely exposed in sedimentary basins. Due to the incompleteness of the record, we aimed to develop complementary geochemical 69 70 proxies for Ca:ALK conditions that could be paired with field observations to assess paleo-71 alkalinity in ancient carbonate-producing systems.

72 73

1.1 Geochemical proxies for Ca:ALK in ancient environments

74 Based on a study of Ca isotope distillation in hyperalkaline Mono Lake (Nielsen & 75 DePaolo 2013), Blättler and Higgins (2014) presented a framework in which Ca isotopes in evaporite salts could be used to reconstruct relative alkalinity. They found a notably large Ca 76 isotope range ($\Delta \delta^{44/40}$ Ca>1.5‰) preserved within CaSO₄ salts in one evaporite system. From these 77 78 measurements and experiments, they derived a relationship between $\Delta\delta^{44/40}$ Ca of evaporite 79 minerals and cation-to-anion ratio of the aqueous solution to reconstruct $Ca^{2+}:SO_4^{2-}$ ratios in 80 ancient evaporite sequences. By analogy to the Ca-SO₄ system and gypsum precipitation, Blättler 81 et al. (2017) applied the same Ca isotope principles to evaluate how seawater carbonate chemistry 82 might have changed over Earth history. Theoretically, when ALK greatly exceeds calcium 83 concentration (Ca:ALK<0.75), like in Mono Lake, distillation driven by net evaporation should yield a $\Delta \delta^{44/40}$ Ca > 0.75‰ within measurements from one evaporative carbonate sequence. The 84 85 carbonate record, however, has additional mechanics that complicate $\Delta\delta^{44/40}$ Ca as a paleoalkalinity proxy. First, evaporitic environments often precipitate multiple carbonate minerals (e.g. 86 87 aragonite, vaterite, calcite, dolomite), and sometimes admixtures within co-occurring textures, 88 each with unique Ca isotope fractionation factors (e.g. Gussone et al., 2005). Second, mechanisms 89 and rates of carbonate precipitation are controlled by chemistry, hydrology, and biology in aqueous

90 systems. The rate of carbonate precipitation can change based on the relative contributions of these

91 factors, thereby driving variable expression of kinetic isotope effects (KIEs) in calcium and other 92 isotope systems in the resultant carbonate record.

93 There is a widely appreciated relationship between dissolved phosphate concentrations and 94 DIC in modern alkaline lakes (e.g. Toner & Catling, 2020). In typical marine environments, 95 phosphate abundance ($\leq 1 \mu$ M in surface waters) is set by biological consumption, organic matter 96 decomposition, sedimentary burial and release of P scavenged by ferric iron oxides, and 97 formation/dissolution of fluorapatite (Ruttenberg & Berner, 1993). With evolving atmospheric and 98 oceanic redox chemistry and evolution of microfauna, marine P concentrations are thought to have 99 fluctuated throughout Earth history, potentially to higher concentrations due to limited P 100 scavenging under lower pO₂ conditions (e.g. Precambrian) or when expanses of deep ocean were 101 anoxic (e.g. Cretaceous). However, phosphate concentrations can reach 100 mM in closed-basin 102 lakes in quasistatic hydrological equilibrium (inflow approximately equal to evaporation).

103 Phosphate is typically a limiting nutrient due to biological uptake. However, the phosphate-104 concentrating mechanisms in alkaline lakewater greatly outpace biological consumption even in 105 the extremely productive Mono Lake $(10^{9}-10^{10} \text{ cells} \cdot \text{liter}^{-1}$ [Humayoun et al., 2003] versus surface 106 seawater, $\sim 10^8$ cells liter⁻¹ [Whitman et al., 1998]). This is in part due to feedbacks on productivity 107 by fixed nitrogen species which are often limiting in these systems (Herbst, 1998; Jellison & 108 Melack, 2001). Further, in low Ca:ALK systems, Ca²⁺ is titrated by carbonate mineralization 109 before low-solubility phosphate salts (e.g. apatite, vivianite) precipitate, which allows phosphate 110 to accumulate in solution.

In this study, we hypothesized that phosphate should incorporate more-or-less proportional to its aqueous concentration in the carbonate phases precipitated from Mono, Searles, and other Pleistocene closed-basin lakes. We further proposed that carbonate P/Ca could be used to determine phosphate levels and relative alkalinity in the carbonate rock record.

115 2 Geologic setting, Samples and Methods

116 2.1 Field geology, facies descriptions, and mineralogy

Pleistocene Owens Valley, California, hosted a series of glacial lakes interconnected by the Owens River and fed by streams off of the Sierra Nevada range (Fig. 1). Today, these lakes exist either as restricted basins (e.g. Mono Lake) or as predominantly dry lake beds with seasonal wetting and drying (e.g. Searles Lake and Death Valley). The Searles dry lake bed and the relict shoreline facies of both Searles Lake and Pleistocene Lake Russell (proto-Mono Lake) are useful testing grounds for paleo-alkalinity proxies because they comprise geologically young carbonates 123 that have not yet been impacted by potential geochemical complications for which carbonate rocks 124 are notoriously sensitive (i.e. burial-related heating or recrystallization).



Figure 1. Interconnected alkaline lakes of Owens Valley, California, adapted from Smith (1979). The pale and dark teal shading marks the paleo- and modern shorelines, respectively. Line dashing denotes the water chemistry of the Sierran drainages that feed Owens River. Sampling sites are marked with stars.

Three field expeditions to Mono Lake and Searles Valley were conducted during 2019 (Figs. S1, S2; Table S1), during which diverse carbonate facies were collected (Fig. S3) and a stratigraphic section of the Pleistocene Wilson Creek Formation was measured (Fig. S4). During summer 2018, a one-meter sediment gravity core was collected in collaboration with the International Geobiology Course from Mono Lake at Station 6 (USGS), in the deepest part of the lake (42.5 m). This core contains sediment deposited over the past ~150 years, and was subsampled every 5 cm.

143 Carbonate mineralogy was determined by x-ray diffraction (XRD) at the California 144 Institute of Technology (Text S1). An aragonite-calcite calibration curve (Fig. S5) was used to 145 quantify samples of mixed mineralogy. Peak areas were integrated at 20 of 25.6° to 26.7° for 146 aragonite and 28.7° to 30.0° for calcite.

147 2.1 Calcium isotopes

148 Approximately 5 mg of bulk carbonate were processed for calcium isotope analysis 149 following the methods of Blättler et al. (2018). Carbonate minerals were dissolved in 0.1N buffered acetic acid, and centrifuged to separate insoluble material. The resulting supernatant was diluted 150 to approximately 30 ppm Ca²⁺, and purified by ion chromatography using an automated Thermo-151 152 Dionex ICS-5000+ with a fraction collector. The purified calcium solutions were diluted in 2% 153 nitric acid to a concentration of 2 ppm calcium for mass spectrometry (Text S2). Isotopic ratios of 154 calcium were analyzed at Princeton University with a Thermo Neptune Plus multi-collector 155 inductively coupled plasma mass spectrometer (ICP-MS) with an ESI Apex-IR sample 156 introduction system. Corrections for mass interferences and standardization procedures are described in Text S2. $\delta^{44/42}$ Ca values were calculated relative to modern seawater by sample-157 standard bracketing. External precision on $\delta^{44/40}$ Ca values is $\pm 0.14\%$ (2 σ), derived from the long-158

159 term reproducibility of the carbonate standard SRM-915b treated as an unknown sample (n=199

- 160 from 2014 to 2018).
- 161 2.2 Clumped (Δ_{47}) isotopes

162 The carbonate clumped isotope thermometer is based on the energetics of "clumping" of the rare, heavy isotopes of carbon and oxygen (¹³C and ¹⁸O) at low temperatures (Eiler, 2007). 163 164 This geothermometer relies on the measurement of the mass-47 anomaly (Δ_{47}) defined as the 165 enrichment of the ¹³C¹⁸O¹⁶O isotopologue of CO₂ relative to a stochastic distribution of the heavy all CO₂ molecules: $\Delta_{47} = \left(\frac{{}^{47}R}{{}^{47}R^*} - 1\right) \times 1000$ where $^{47}\mathbf{R}$ = 166 isotopes among $[^{13}C^{16}O^{18}O^{+12}C^{17}O^{18}O^{+13}C^{17}O_2]/[^{12}C^{16}O_2]$ and * denotes a stochastic intramolecular distribution 167 of isotopes. Carbonate clumped isotope measurements were made at the California Institute of 168 169 Technology over three analytical sessions from March 2019 to January 2020 (Table A3) following 170 the analytical and data reduction procedures of Ingalls et al. (2020) (Text S3).

171 We used clumped isotopes in this study to investigate potential disequilibrium in 172 carbonates forming rapidly in alkaline systems. Departures from carbonate clumped isotope 173 equilibrium are most sensitive to parameters that impact the DIC composition (Watkins & Hunt, 2015). For example, experiments and natural samples have shown Δ_{47} depletions in carbonates 174 175 formed from waters affected by HCO₃⁻ dehydration and dehydroxylation and Δ_{47} enrichments in 176 carbonates affected by CO₂ hydration and hydroxylation (e.g., Saenger et al., 2012; Hill et al., 177 2014; Daeron et al., 2019), with a greater effect in high pH. Disequilibrium can also be driven by 178 rapid carbonate precipitation rates because of the slow interconversion between CO₂ and HCO₃-179 in the precipitating solution (Guo, 2020; Watkins & Hunt, 2015). Sample isotopic heterogeneity 180 due to carbonate precipitating at a range of conditions that promote variable Δ_{47} disequilibrium can 181 cause larger variance in replicate measurements and significant deviation in $T(\Delta_{47})$ from 182 precipitation temperatures (Guo, 2020). Importantly, a carbonate system out of isotopic 183 equilibrium with respect to one set of equilibria may also yield disequilibrium behavior in other 184 isotope systems, e.g. Ca isotopes, and inform other geochemical data impacted by kinetics (e.g. 185 Thiagarajan et al., 2020).

186 2.3 Phosphorus-to-calcium ratios

187 Phosphate concentrations were measured by the malachite green method (Ohno & Zibilske, 188 1991; Van Veldhoven & Mannaerts, 1987) colorimetric assay with a Biotek Cytation5 plate reader. 189 Approximately 25 mg of carbonate powder was dissolved in 0.33 M acetic acid and ultrasonicated 190 for >4 hours. A weak acid was chosen to avoid leaching of organic phosphorus and other 191 phosphorus-bearing minerals like apatite. Three unique dilutions of each sample solution with 192 three replicates of each dilution were pipetted into a well plate, for a total of at least nine 193 absorbance readings per sample. An Abcam phosphate standard (ab65622) was used to generate 194 calibration curves of 0 to 25 μ M P for each plate. 30 μ l of a proprietary formulation of malachite 195 green and ammonium molybdate was added to each well and incubated in the dark for 30 minutes 196 to facilitate the complexation of the heteropoly molybdophosphate that gives an absorption band at a wavelength of 650 nm. The dissolved CaCO₃ mass, volume of solution, and nmol P measured
by absorbance were used to calculate P/Ca molar ratios (Text S4).

199 **3 Results and Discussion**

200 3.1 Field observations to identify closed-basin chemistry without geochemical tools

201 In very low Ca:ALK systems, the loci of carbonate formation are almost entirely restricted 202 to zones where Ca^{2+} -charged freshwater inputs mix with Ca^{2+} -starved and DIC-rich lake water. 203 Tufa 'towers', ranging from meter- to tens of meters-scale (Fig. S3k), grew directly out of alluvial 204 fan and deltaic deposits in the Wilson Creek Formation and at sites of spring discharge at both 205 Mono and Searles (Fig. S3i).

Shoreline tufas were found draping igneous bedrock and volcanics at both sites (Fig. S3h)—these were generally more laterally extensive than the towers and, in places, exhibited reeflike morphologies. Tufa textures were subdivided into nodular, columnar or dendritic, and porous growth habits with fine laminations that could be identified on both the micro- and macroscopic scale. Tufas at both locations were variably cemented, and often encased in a laminated crust. These textures and their proposed formation mechanisms have been described elsewhere (e.g.,

212 Dunn, 1953; Guo & Chafetz, 2012).

Multiple tufa fabrics identified in this study can be morphologically linked to relatively rapid mineralization under low Ca:ALK conditions. We identified casts of brine fly pupae casings and tens-of-meters-scale reefs of carbonate-coated algal filaments (Fig. S3b). The *in vivo* preservation of fauna and cellular morphology by carbonate prior to decay indicated that precipitation occurred on biological rather than geological timescales.

218 Minimal carbonate precipitation occurs in the water column at Mono Lake. Most water 219 column carbonate formation is thought to occur rapidly when the chemocline overturns and 220 slightly Ca²⁺-enriched, lower salinity, oxygenated surface water mixes with anoxic, alkaline bottom water to create transient $\Omega > 1$ conditions (Nielsen & DePaolo, 2013). The sediment core 221 222 from Mono Lake was carbonate-poor and organic-rich (~5-10 wt.% total organic carbon; Fig. 223 S3L). We hypothesized that the finely laminated muddy sediment should yield an organic-rich 224 shale under compaction and diagenesis—a lithotype not typically targeted for study of ancient 225 carbonates or for carbonate-based proxy materials, and thus, geochemical signals recorded in these 226 sediments may be overlooked in geological studies. Finely laminated lake bottom sediments from 227 both the Wilson Creek Formation and Searles were also carbonate-poor (Fig. S3g), suggesting 228 that—as occurs today in Mono Lake—relatively little water column carbonate precipitation 229 occurred in the Pleistocene lakes.

230 Slightly acidic closed-basin lakes, such as Andean Laguna Negra with Ca-Na-Cl water, 231 can also form carbonate at shoreline freshwater-lakewater mixing zones. Laguna Negra's Ω is 232 limited by low pH rather than calcium. Carbonates described at Laguna Negra are morphologically 233 and genetically associated with the microbial mats that form in mixing zones due to increased pH 234 and reduced salinity (Gomez et al., 2014). Laguna Negra microbialites and oncoids form wrinkled 235 and clotted textures in micritic to sparry laminae, which are easily differentiated from porous tufa 236 fabrics. Thus, identification of tufa-like textures in stratigraphic positions that indicate riverine and 237 groundwater inputs (e.g. on fans, within deltas, or near shorelines) can be used to help identify low 238 Ca:ALK systems in the record. However, chemical proxies are necessary to quantitatively estimate 239 Ca:ALK in geological systems with insufficient preservation or outcrop exposure.

240 241

3.2 Ca isotope range as a proxy for Ca:ALK

242 The magnitude of calcium isotope fractionation between a carbonate mineral and water is 243 determined by mineralogy, precipitation rate, and the Ca:ALK of the solution. When Ca:ALK<0.75, dissolved Ca²⁺ is near-quantitatively removed by carbonate formation during 244 evaporitic concentration of solutes, yielding >0.75% of Ca distillation and, thus, $\Delta \delta^{44/40}$ Ca_{water} 245 (Fig. 2a; Nielsen & DePaolo, 2013). To test if the predicted $\Delta \delta^{44/40}$ Ca could be recovered in 246 247 evaporitic carbonate salts that formed in the Pleistocene Owens Valley lake systems, we measured 248 the carbonate facies found in modern Mono Lake and Pleistocene Lakes Russell (Wilson Creek 249 Formation) and Searles, which presumably precipitated from similarly low Ca:ALK waters. 250 However, our $\delta^{44/40}$ Ca_{carb} measurements yielded total ranges of 0.68‰ and 0.52‰ from Pleistocene Lakes Russell and Searles, respectively—values below the 0.75‰ threshold expected 251 252 for low Ca:ALK systems (Fig. 2b). The diverse carbonates sampled in this study did not capture 253 the anticipated isotopic distillation behavior.

254 Carbonate precipitated within the water column of the most open part of the lake where Ca^{2+} is lowest, and evaporative concentration of Ca^{2+} is necessary for mineralization, should yield 255 the greatest isotopic distillation. As such, we predicted we would measure >0.75% $\Lambda\delta^{44/40}$ Ca 256 driven by evaporative precipitation from the Mono Lake sediment core. Subsamples were collected 257 258 every 5 cm to simulate relatively fine sampling practices for very fine-scaled chemostratigraphy with the goal of detecting chemical perturbations. However, we found a range of $\delta^{44/40}$ Ca of 0.47‰ 259 260 within the core. A finer-scale sampling and/or analytical technique may be required to resolve 261 evaporative distillation events.

262 We can better predict and identify Rayleigh behavior in a system that precipitates the full 263 evaporite mineral sequence (i.e., carbonate, gypsum, halite). However, Mono Lake is undersaturated with respect to gypsum, and thus, only accumulates carbonate salts. In the absence 264 265 of other evaporite minerals, it is difficult to identify evaporative versus equilibrium precipitation. Because we were unable to recover the modeled $\Delta \delta^{44/40}$ Ca within a diverse range of carbonate 266 facies in modern sediments prior to diagenetic modification, it may be exceedingly difficult to 267 recover Rayleigh distillation of Ca isotopes in ancient lacustrine sediments. While the perfect 268 269 stratigraphic section preserving multiple evaporitic carbonate sequences might exist, the difficulty 270 of recovering such sequences in modern and recent settings where we have absolute constraints on 271 Ca:ALK suggested that Ca isotope range may be better interpreted when employed concurrently 272 with an additional tool for reconstructing Ca:ALK.

273



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Figure 2. Stable isotope effects of low Ca:ALK water. (a) Expected Rayleigh distillation of Ca 275 276 isotopes during evaporative precipitation of aragonite (green) and calcite (purple). Vertical bars plot $\delta^{44/40}$ Ca ranges from Searles calcite and aragonite, and the Mono-Russell system. Ca ranges 277 were projected to the mineral-specific Rayleigh curves to determine the fraction of Ca removed 278 279 during evaporative mineral precipitation (arrows), or degree of distillation under model assumptions. The starting $\delta^{44/40}$ Ca_{water} of -0.6‰sw is 0.1‰ higher than the most distilled Mono 280 Lake water reported in Nielsen & DePaolo (2013) and the upper limit of the $\delta^{44/40}$ Ca value of water 281 that could precipitate our lowest calcite $\delta^{44/40}$ Ca values. (b) Ca isotope data from a modern Mono 282 283 Lake core and Pleistocene Lakes Russell and Searles. The gray arrow marks the theoretically predicted range evaporative precipitation from low Ca:ALK water should exceed. Points in panels 284 B and C are colored by percent aragonite. Green and purple bars mark $\Delta\delta^{44/40}$ Ca binned by 285 286 aragonite (>60%) or calcite, respectively. (c) Large intra- (error bars) and intersample carbonate 287 clumped isotope variability, as well as $T(\Delta_{47})$ exceeding the modern annual range of Mono Lake 288 surface water temperatures (light blue box) and predicted cooler Pleistocene temperatures in 289 Owens Valley (teal), suggests Δ_{47} disequilibrium.

290

Variable mineralogy and precipitation kinetics tend to expand the total $\Delta \delta^{44/40}$ Ca beyond 291 the range driven by Rayleigh distillation. First, aragonite and calcite have different $^{44/40}$ Ca α_{carb} -292 293 water fractionation factors (Gussone et al., 2005). Each carbonate phase that precipitates from the same $\delta^{44/40}$ Ca_{water} yields different $\delta^{44/40}$ Ca_{carb} values(Fig. 2a). This is important when considering 294 295 an ancient carbonate system that may have formed as a mixed-mineral assemblage but has been diagenetically replaced by calcite and/or dolomite; a large $\Delta \delta^{44/40}$ Ca value resulting from different 296 297 mineralogical fractionations could be misinterpreted as evaporative distillation. In Mono and 298 Searles Lakes, we identified both aragonite and calcite with pseudomorphic textures of primary 299 cold-water carbonates, ikaite and monohydrated calcite (Fig. S3e). We considered $\Delta\delta^{44/40}$ Ca of 300 each carbonate phase separately to avoid confounding mineralogical and distillation effects. The Mono-Russell system had a $\Delta \delta^{44/40}$ Ca_{cal} of 0.56‰ and $\Delta \delta^{44/40}$ Ca_{arag} of 0.47‰. Searles Lake yielded 301 $\Delta \delta^{44/40}$ Ca_{cal} of 0.32‰ and $\Delta \delta^{44/40}$ Ca_{arag} of 0.53‰. 302

Additionally, we suspected that some amount of $\Delta \delta^{44/40}$ Ca could be explained by varying 303 304 KIEs-driven by differences in carbonate precipitation rates between fabrics, and between 305 individual tufa samples (Lemarchand et al., 2004). Tufas from the Wilson Creek Formation 306 (Russell), and Mono and Searles Lakes yielded $T(\Delta_{47})$ values ranging from -7.1±4.3°C to 307 39.5±4.5°C (Fig. 2; Table A2). This amount of $T(\Delta_{47})$ variability is geologically implausible, both 308 within individual samples (i.e. large error bars capture variability in measurements from the same 309 tufa) and between tufas formed within each basin under similar climate conditions. Furthermore, 310 Owens Valley tufas have never experienced high-temperature burial or hydrothermal alteration, 311 which drives $T(\Delta_{47})$ to higher apparent temperatures. Some of the tufas yield warmer apparent 312 temperatures than modern summertime (JJA) surface water temperatures (Schneider et al., 2009; 313 Fig. 2). Additionally, we anticipated that carbonates formed during the Pleistocene glacial periods 314 should have yielded colder formation temperatures than modern. We attributed the unusually warm $T(\Delta_{47})$ values and sample heterogeneity to variably expressed KIEs driven by fast tufa carbonate 315 316 formation. Although magnitude of KIEs are independent, both Δ_{47} and Ca isotopes are susceptible to precipitation rate-driven KIEs (Thiagarajan et al., 2020), and thus we hypothesized that some 317 amount of the tufa $\delta^{44/40}$ Ca variability captured kinetics in ways that masked the expected patterns 318 319 of Ravleigh distillation.

320

321 3.3 High P/Ca in carbonates formed from low Ca:ALK water

322 Carbonates that formed within modern Mono Lake, Pleistocene Lakes Russell and Searles, 323 and other Pleistocene lakes vielded P/Ca ratios up to three orders of magnitude greater than marine 324 carbonate (Fig. 3). All carbonate facies from Searles Lake displayed P/Ca values from 0.432±0.019 325 to 3.177±0.177 mmol/mol. Carbonates from the one-meter Mono Lake core ranged from 326 0.674±0.033 to 1.301±0.112 mmol/mol. The highest P/Ca values were from the Pleistocene Wilson Creek Formation, which ranged from 1.192±0.082 to 13.995±0.794 mmol/mol. Carbonate 327 328 pebble coatings from a lag pavement in a deltaic fan at the base of the Wilson Creek Formation 329 vielded 3.209 mmol P/mol Ca. From sedimentological observations, we interpreted that these 330 coated clasts formed in the wave spray/swash zone during early lake-level rise when the lake was 331 still undergoing cycles of wetting and drying typical of very shallow, saline lakes. Given these 332 values, Lake Russell may have had a low Ca:ALK early when lake levels were still low and 333 dissolved solute levels were highly sensitive to evaporation.

334 In marine materials, the majority of existing P/Ca data were derived from coral skeletons 335 (Anagnostou et al., 2011; Chen et al., 2019), with the highest P/Ca concentrations in deep-water species D. dianthus. Chen et al. (2019) showed that P was incorporated into coral aragonite in 336 337 proportion to its concentration in surrounding seawater. Biogenic coral aragonite populated the 338 lowest mode of P/Ca, ranging from 0.003 to 0.118 mmol/mol—lower values than biogenic calcite 339 or abiotic marine aragonite and calcite (Fig. S6). Aragonitic, calcitic, and dolomitic marine 340 cements and ooids and biogenic calcite comprised the second lowest P/Ca mode from 0.049 to 341 0.184 mmol/mol (Fig. 3a, S6). There was no statistical difference between biogenic calcite and 342 abiotic marine carbonate P/Ca, which was interpreted as phosphate partitioning without strong 343 dependencies on biogenicity and carbonate mineralogy.

344 Marine carbonates and those from Pleistocene to modern alkaline lakes formed distinct 345 P/Ca populations with minimal overlap (Fig. 3a). The minor overlap seen between marine and 346 nonmarine carbonate data could be due to the complexities of impurity incorporation into 347 carbonate minerals (Gutjahr et al., 1996; Nielsen et al., 2013) as well as natural variability in both 348 dissolved phosphate concentrations and Ca:ALK in both marine and terrestrial systems. 349 Nonetheless, the striking bimodality of the P/Ca data between high and low Ca:ALK systems 350 presented a means by which to differentiate between end-member chemical environments (Fig. 351 3a).



352

Figure 3. Phosphorus-to-calcium ratios of marine and closed-basin lake carbonates. (a) Distribution of marine (purple) and nonmarine (yellow) carbonate P/Ca measured in this study and some marine measurements using the same method in the literature (Anagnostou et al., 2011; Chen et al., 2019). (b) Nearly all carbonate P/Ca from restricted basins are substantially greater than marine values.

358 We assessed if phosphate enrichments in Owens Valley lacustrine carbonates were more 359 related to chemical weathering of Sierran bedrock specifically, rather than Ca:ALK, by analyzing 360 shoreline tufa from 10 sites with idiosyncratic basin geology (Table A1): the Eocene Green River 361 Formation and Pleistocene to modern lake basins from Tibet and western North America (Fig. 3). 362 All tufa P/Ca ratios were highly elevated above median marine values. The mean P/Ca values from 363 Balung Co, San Agustin, Lake Surprise, and Lake Chewaucan were comparable with the extreme 364 P/Ca values from Owens Valley, demonstrating that the extremely high phosphate levels found in 365 Mono Lake water and partitioned into tufa carbonate are non-unique and shoreline tufa facies can 366 be a meaningful indicator of elevated phosphate levels and possibly low Ca:ALK chemistry in the 367 carbonate record.

Elevated P/Ca values of 0.211±0.027 to 0.533±0.094 mmol/mol observed in seven tufas, stromatolites, and cements from the Eocene Green River Formation demonstrated that carbonateassociated phosphate enrichments could be preserved through burial diagenesis. It remains to be tested how robust this P/Ca signal is in deeper time, but we have seen that the high phosphate levels that are correlated with Ca:ALK in these systems can be captured in the carbonate minerals they produce.

4 Conclusions

The ability to reconstruct alkalinity at specific times and basins in Earth's past allows us to better understand the processes responsible for the generation of the carbonate record, and evaluate secular changes in seawater chemistry over Earth history. Geological observations of textures, fabrics, facies, and stratal geometry of carbonate-rich units provide useful constraints on ancient 379 Ca:ALK based on characteristic modes of carbonate formation in low or high Ca:ALK systems. 380 In the Pleistocene lake system of Owens Valley, we found that carbonate formation is tied closely to processes that add Ca^{2+} to the water, as compared to high Ca:ALK systems (e.g. marine basins), 381 382 wherein carbonate formation is more tied to biogeochemical processes that impact the distribution 383 of DIC and alkalinity. We also examined potential geochemical proxies for reconstructing 384 Ca:ALK. Ca-isotope range has previously been used to reconstruct Ca:ALK of ancient environments based on Rayleigh distillation behavior within the Ca²⁺-starved water column of 385 386 Mono Lake. However, we were unable to recover the theoretically predicted > 0.75% range from carbonates formed in Ca²⁺-starved waters in Mono, Russell, and Searles Lakes. We hypothesized 387 388 that mineralogical differences (i.e. aragonite vs. calcite) and rate dependence of $\alpha_{carb-water}$ complicated interpretations of $\Delta\delta^{44/40}$ Ca in the diverse carbonate phases that form in these 389 390 hyperalkaline lake systems. An unreasonable range of $T(\Delta_{47})$ values within individual samples and 391 between samples from the same basin, as well as much warmer temperatures than expected from 392 carbonates that precipitated during glacial periods, supported the view that isotopic disequilibrium 393 and precipitation kinetics are important in these systems.

394 Based on our understanding of the mechanics that lead to high phosphate concentrations in 395 alkaline systems, we studied carbonate-associated P/Ca as a potential proxy for the phosphate 396 content and Ca:ALK in ancient aqueous environments. Carbonates that formed in closed-basin 397 lakes under Ca²⁺-starved conditions vielded orders of magnitude greater P/Ca ratios than modern 398 and ancient marine carbonate. Phosphate was incorporated into carbonates relatively proportional 399 to the Ca:ALK of the system. Questions remain about how robust P/Ca is to post-depositional 400 processes that impact the geochemistry of ancient carbonates. However, we found that P/Ca was 401 robust through burial diagenesis and partial silicification of carbonates in the Eocene Green River 402 Formation, and posited that-along with geological constraints-P/Ca has the potential to be 403 useful in identifying phosphate levels and Ca:ALK in ancient environments.

404 Acknowledgments, Samples, and Data

405 Supporting information detailing methods and all data may be found in the Supplemental 406 Information, and were archived in the Open Science Framework (doi: 10.17605/OSF.iO/N9Y3H). 407 Samples collected as a part of this study were registered with IGSNs, listed in Table S1. The authors would like to thank Christine Chen, Adam Hudson, Dan Ibarra, Max Lloyd and Theodore 408 409 Present for contributing sample material from other paleo-lakes and marine carbonates, and the 410 International Geobiology Course (supported by the Agouron Institute, Simons Foundation, NASA, 411 and Caltech) for access to Mono Lake sediment cores. Funding for the work was provided by the 412 Simons Foundation Collaboration on the Origins of Life (WWF) and Barr Foundation Postdoctoral 413 Fellowship to MI.

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