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Early diagenetic constraints on Permian seawater chemistry from the Capitan Reef

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Abstract

The Capitan Reef Complex in West Texas is famous for its high prevalence of early marine cements, unusual for a Phanerozoic platform, leading some to suggest that Precambrian styles of carbonate sedimentation enjoyed a Permian encore. Here, we use patterns of stable Ca, Mg, C and S isotopes to better understand the environmental driver(s) of the enigmatic cementation. We find that calcite that is the most enriched in ⁴⁴Ca has δ^{34} S values that approach the inferred composition of Permian seawater sulfate. Microbial sulfate reduction in pore fluids must have been spatially and temporally coincident with recrystallization of primary carbonate phases, such that substantial ³⁴S-enriched sulfate was incorporated into diagenetic calcite under relatively closed-system conditions. Moreover, the magnitude of ³⁴S-enrichment of carbonates relative to seawater was strongly influenced by local diagenetic conditions, with fluid-buffered early marine cements, shelf, reef, and upper slope preserving more seawater-like S isotope ratios than the more sediment-buffered lower slope. Some samples are far more 34 S-enriched relative to seawater than those from modern sites in similar depositional environments, possibly responding to specific combinations of sedimentary parameters (e.g., grain size, porosity, organic matter rain rate). Additionally, the sulfate concentration in the Delaware Basin might have been slightly lower than modern levels, leading to more extensive isotopic evolution of sulfate in pore waters during carbonate recrystallization. Based on the data and a numerical model of carbonate recrystallization, we suggest that one driver of the extensive seafloor cement precipitation in the Capitan Reef Complex was a Permian water column $[Ca^{2+}]$: $[SO_4^{2-}]$ ratio somewhere between 1 and modern seawater. © 2022 Elsevier Ltd. All rights reserved.

Keywords: Carbonate diagenesis; Ca isotopes; Mg isotopes; S isotopes; Sulfate reduction

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1. INTRODUCTION

Our knowledge of the history of Earth's surface oxidation state owes much to measurements of the sulfur isotope composition (δ^{34} S) of sulfur-bearing phases in marine sedimentary rocks (Garrels and Lerman, 1981; Berner, 1989; Canfield, 2001; Canfield and Farquhar, 2009). Rock-derived δ^{34} S_{sulfate} values approximate seawater $\delta^{34}S_{sulfate}$ (Claypool et al., 1980; Paytan et al., 1998; Kampschulte and Strauss, 2004; Rennie et al., 2018), while rock-derived $\delta^{34}S_{sulfide}$ values record the signal of microbial sulfate reduction, which discriminates against the heavy isotope, ³⁴S (Leavitt et al., 2013; Fike et al., 2015). The magnitude of isotopic offsets between coeval $\delta^{34}S_{sulfate}$ and $\delta^{34}S_{sulfide}$ values (Δ_{pvr}) have long been thought to reflect the relative magnitude of global burial fluxes of oxidized (i.e., sulfates) vs. reduced sulfur (i.e., sulfides) (Canfield and Teske, 1996; Fike et al., 2006). Together, timecalibrated records of $\delta^{34}S_{sulfate}$ and Δ_{pyr} offer the potential for semi-quantitative assessment of $[SO_4^{2-}]_{seawater}$ using simple steady state models that relate the rate of change of rock-derived $\delta^{34}S_{sulfate}$ to the size of the global sulfate reservoir (Kah et al., 2004; Planavsky et al., 2012; Algeo et al., 2015). However, whereas seawater $\delta^{34}S_{sulfate}$ is wellconstrained through the Mesozoic and Cenozoic Eras by a combination of relatively continuous marine barite (Paytan et al., 1998; Paytan, 2004), taxon-specific carbonate-associated sulfate (CAS) (Burdett et al., 1989; Rennie et al., 2018), and episodic massive sulfate evaporites (Claypool et al., 1980), the Paleozoic and Precambrian seawater $\delta^{34}S_{sulfate}$ record is comparatively patchy and uncertain (Fike et al., 2015). The patchiness is due to the subduction of most pre-Jurassic oceanic crust (and deepsea sediments hosting barite and pelagic fossils). The uncertainty in the seawater $\delta^{34}S_{sulfate}$ record is due to 'noise' in the most temporally continuous and ubiquitous seawater $\delta^{34}S_{sulfate}$ record, bulk-rock CAS (Present et al., 2020).

Recent work has shown that the 'noise' in the bulk-rock $\delta^{34}S_{CAS}$ record, which might superficially be interpreted as rapid oscillations in seawater $\delta^{34}S_{sulfate}$, is likely largely due to bulk-rock CAS being an inconsistent recorder of seawater $\delta^{34}S_{sulfate}$. Firstly, stratigraphic bulk-rock $\delta^{34}S_{CAS}$ variability can, in some cases, be attributed to depositional environment or diagenetic conditions (Richardson et al., 2019a; Murray et al., 2021). Secondly, coeval carbonates from different depositional settings record markedly different bulk-rock $\delta^{34}S_{CAS}$ values (Present et al., 2019), despite the presumed homogeneity of seawater $\delta^{34}S_{sulfate}$. Thirdly, different minerals (e.g., dolomite vs. calcite) and components in carbonate rocks (e.g., bioclasts vs. different types of cement) can preserve wildly different $\delta^{34}S_{CAS}$ values (Present et al., 2015; Present et al., 2019) and CAS concentrations (Present et al., 2015; Richardson et al., 2019b; Rose et al., 2019), responding to their local conditions of formation and diagenesis. Finally, the 'noise' in bulk-rock $\delta^{34}S_{CAS}$ data increases with sample age, implying that postdepositional processes (i.e., early and late diagenesis) can affect bulk-rock $\delta^{34}S_{CAS}$ (Present et al., 2020). More promisingly, other studies have found that primary $\delta^{34}S_{CAS}$ values can be preserved through burial diagenesis and hydrothermal dolomitization (Fichtner et al., 2017; Schurr et al., 2021). In summary, the degree to which bulk-rock $\delta^{34}S_{CAS}$ records seawater $\delta^{34}S_{sulfate}$ therefore is highly dependent on carbonates' environment of deposition, mineralogy, component make-up, and post-depositional history.

To increase the utility of the existing bulk-rock $\delta^{34}S_{CAS}$ record, it would be beneficial to have a unifying geochemical framework for understanding the diagenetic controls on

bulk-rock $\delta^{34}S_{CAS}$. Ideally, different diagenetic endmembers (pristine and altered) would purposely be sampled to map out the entire phase space. The advantage of such an approach is that it can be applied systematically at the bulk-rock level without detailed facies/sequence stratigraphic context or the presence of specific carbonate components (e.g., bioclasts), both of which are absent and/or underconstrained through much of the carbonate record. Here, using diagenetically altered carbonates, we show that combined bulk-rock $\delta^{34}S_{CAS}$ values with magnesium ($\delta^{26}Mg$) and calcium ($\delta^{44/40}Ca$) isotope ratios, in addition to minor/major element ratios (e.g., Sr/Ca, Mn/Ca, Fe/Ca). provide a powerful geochemical toolkit for understanding marine carbonate diagenesis. These systems together distinguish between restricted basinal (i.e., evaporitic) and openocean depositional settings (Blättler et al., 2018; Higgins et al., 2018), which affect the degree to which $\delta^{34}S_{CAS}$ resembles seawater $\delta^{34}S_{sulfate}$ (Fike et al., 2015). They are also sensitive to the nuances of carbonate recrystallization and closed vs. open system behaviour during early marine diagenesis (Blättler et al., 2015; Ahm et al., 2018; Higgins et al., 2018; Ahm et al., 2019; Jones et al., 2020; Staudigel et al., 2021; Crockford et al., 2021), and are therefore a function of local depositional and diagenetic conditions another major control on $\delta^{34}S_{CAS}$ (Richardson et al., 2019a; Murray et al., 2021) - and the chemical composition of the overlying water column.

To test whether δ^{26} Mg and/or $\delta^{44/40}$ Ca can be useful for quantitatively predicting how well $\delta^{34}S_{CAS}$ resembles seawater $\delta^{34}S_{\text{sulfate}}$, we apply these analyses to the micro-drilled, late Guadalupian-age Capitan Reef Complex samples of (Present et al., 2019), for which $\delta^{34}S_{CAS}$, overlying seawater $\delta^{34}S_{sulfate}$, depositional facies, petrographic context, other geochemical indices ($\delta^{13}C_{carb}$, $\delta^{18}O$), and mineralogy are already well-constrained. We characterize the relationships among samples with different depositional and diagenetic histories and assess the mechanisms and conditions that generate these relationships. Here, we also adapt an existing diagenetic model (Ahm et al., 2018), to explore how established diagenetic indices can be used to disentangle diagenetic effects on $\delta^{34}S_{CAS}$, constrain Permian seawater chemistry, and potentially explain anomalous features of the Capitan Reef Complex such as the abundant and ubiquitous early marine cements that are reminiscent of Precambrian carbonates (Grotzinger and Knoll, 1995). Moving forward, correlations between any mineral- or texture-specific suite of $\delta^{34}S_{CAS}$ values and $\delta^{26}Mg$ and/or $\delta^{44/40}Ca$ form a basis for assessing the degree to which $\delta^{34}S_{CAS}$ measurements in other, less well studied locations capture contemporaneous seawater $\delta^{34}S_{sulfate}$. This information could be used to screen the bulk-rock $\delta^{34}S_{CAS}$ record through Earth history for samples whose $\delta^{34}S_{CAS}$ values have been altered by diagenesis and do not reflect seawater $\delta^{34}S_{sulfate}$.

2. MATERIALS & METHODS

2.1. Isotopic notation

For the purposes of this study, we use delta notation to describe the deviation in per mil (%) of the isotopic

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composition of samples from known reference standards (Coplen, 2011). For sulfur isotopes, $\delta^{34}S_{\text{sample}} = [(^{34}S/$ ${}^{32}S)_{sample}/({}^{34}S/{}^{32}S)_{V-CDT}] - 1$, where V-CDT is the Vienna Canyon Diablo Troilite reference standard (Ding et al., 1999). For carbon isotopes, $\delta^{13}C_{\text{sample}} = [({}^{13}C/{}^{12}C)_{\text{sample}}]$ $(^{13}C/^{12}C)_{V-PDB}] - 1$, where V-PDB is the Vienna-Pee Dee Belemnite reference standard. For oxygen isotopes,
$$\begin{split} \delta^{18}O_{sample} &= [(^{18}O/^{16}O)_{sample}/(^{18}O/^{16}O)_{V-PDB}] - 1. \text{ For magnesium isotopes, } \delta^{26}Mg_{sample} &= [(^{26}Mg/^{24}Mg)_{sample}/(^{26}Mg/^{24}Mg/^{24}Mg)_{sample}/(^{26}Mg/^{24}Mg/^{24}Mg)_{sample}/(^{26}Mg/^{24}Mg/^{24}Mg)_{sample}/(^{26}Mg/^{24}Mg/^{24}Mg/^{24}Mg/^{24}Mg)_{sample}/(^{26}Mg/^{24}Mg$$
 $({}^{26}Mg/{}^{24}Mg)_{DSM3}] - 1$, where DSM3 is the Dead Sea Magnesium reference standard (Galy et al., 2003). For calcium δ⁴⁴/⁴⁰Ca_{sample} $\left[\left(\frac{^{44}\text{Ca}}{^{40}\text{Ca}}\right)_{\text{sample}}\right]$ isotopes, = $({}^{44}Ca/{}^{40}Ca)_{SW}] - 1$, where SW is IAPSO modern seawater (Hippler et al., 2003). For reference, IAPSO is +1.86% relative to NIST915a (Jochum et al., 2005), the other most used Ca reference standard.

2.2. Samples

The sample set for this study features carbonates from the late Guadalupian-age Capitan Reef Complex in the Permian Basin of West Texas and New Mexico, one of the best preserved and best studied ancient mixed siliciclastic and carbonate platform and slope systems. Depositional facies from shelf to basin are correlated by detailed biostratigraphy and sequence stratigraphic models (Silver and Todd, 1969; Tyrrell Jr, 1969; Borer and Harris, 1991; Osleger, 1998; Kerans and Tinker, 1999; Sarg et al., 1999; Wilde et al., 1999; Lambert, 2002; Rush and Kerans, 2010), crucially allowing carbonates deposited from the same water column but in different depositional environments to be compared. The carbonates record a range of early-late diagenetic processes in their petrographic and geochemical characteristics (Mazzullo and Cys, 1977; Schmidt, 1977; Yurewicz, 1977; Given and Lohmann, 1986; Garber et al., 1989; Mruk, 1989; Scholle et al., 1992; Mutti and Simo, 1994; Mazzullo, 1999; Melim and Scholle, 2002; Budd et al., 2013; Frost et al., 2013; Bishop et al., 2014; Wu et al., 2020). Notably, syn- and eogenetic marine diagenetic processes dominated the texture of Capitan Reef Complex limestones and dolomites. Primary fibrous, bladed, isopachous, and micritic cements comprise most of the rock volume of reef and fore-reef (Cys et al., 1977; Mazzullo and Cys, 1977; Grotzinger and Knoll, 1995), and voluminous precipitates form allochems and void-filling cements in peritidal shelf-top strata (Kendall, 1969; Esteban and Pray, 1977; Assereto and Kendall, 1977; Esteban and Pray, 1983; Chafetz et al., 2008). Stratal geometries of dolomite show that platform interior-derived brines drove dolomite replacement of precursor aragonitic and calcitic sediments in concert with syndepositional accommodation cycles (Adams and Rhodes, 1960; Kendall, 1969; Mutti and Simo, 1994; Melim and Scholle, 2002; Koša and Hunt, 2006; Garcia-Fresca et al., 2012). Neomorphic dolomite textures include fabric-retentive mimetic dolomite and fabric-selective non-mimetic dolomite (Mutti and Simo, 1993; Mutti and Simo, 1994; Melim and Scholle, 2002; Frost et al., 2013), and primary aragonite cements were replaced by neomorphic calcite (Mazzullo, 1980; Chafetz et al., 2008). Later generations of percolating fluids caused

coarser dolomite crystallization along permeable sandstones (Melim and Scholle, 1989), Neptunian fracture systems, and eogenetically fractured tepee and sheet crack structures (Kendall, 1969; Melim and Scholle, 1989; Koša and Hunt, 2006; Budd et al., 2013; Frost et al., 2013). Rock fabric was largely unchanged during burial diagenesis, although porosity was modified by anhydrite and calcite cement precipitation and dissolution (Schmidt, 1977; Garber et al., 1989; Mruk, 1989; Scholle et al., 1992; Budd et al., 2013; Loyd et al., 2013; Frost et al., 2013).

Contemporaneous seawater $\delta^{34}S_{sulfate}$ value is constrained to $\sim 8-10\%$ VCDT by the composition of local and more remote mid-late Permian-age evaporite deposits (Thode et al., 1961; Holser and Kaplan, 1966; Claypool et al., 1980; Sarg, 1981; Hill, 1990; Leslie et al., 1997; Kirkland et al., 2000; Present et al., 2020). Present et al. (2019) collected carbonate hand-samples along a shelfbasin transect through the Yates, Tansill, Capitan and Bell Canyon formations (Fig. 1A) in McKittrick Canyon, Guadalupe Mountains, National Park, Texas, and sub-sampled specific textures/phases (e.g., different generations of cement, or minerals; Fig. 1B-C) using a handheld rotary micro-drill. For this study, we micro-drilled $259 \sim 1-3$ mg sub-samples from the same spots already sub-sampled by (Present et al., 2019), based on the assumption that the newly sampled material would be chemically and mineralogically equivalent.

2.3. Standards

To correct our isotopic data to international reference standards, we processed various isotopic reference materials along with our samples. For Ca isotopes, these were: seawater (collected at Woods Hole Oceanographic Institution), EN-1 from the USGS (modern *Tridacna* shell, Enewetak lagoon, Marshall Islands), NIST SRM 915b calcium carbonate, the 'PUA' in-house aragonite standard from Princeton University, and an in-house aragonite standard (San Salvador Sand; 'SSS'). For Mg isotopes, these were: seawater, the 'RUB' in-house calcite standard from Ruhr University Bochum (Riechelmann et al., 2018), a high purity Mg ICP standard obtained from Princeton University, and an in-house dolomite standard from the University of Chicago collection ('IHD').

2.4. Wet chemistry and ion chromatography (Dionex)

For each carbonate sample or standard, 1 mg was added to 1 mL of a 0.1 M acetic acid, 0.05 M ammonium hydroxide solution and treated ultrasonically for 4–8 hours to ensure complete dissolution of carbonate materials. For carbonate samples, aliquots of solution were diluted using the same acetic acid/ammonium hydroxide solution to generate solutions with $[Ca^{2+}]$ of 100–200 ppm or $[Mg^{2+}]$ of 10–25 ppm. Sample aliquots of 25 µL were then autosampled and processed with a Thermo/Dionex ion chromatography system, using 30 mM methanesulfonic acid as the eluent (25 mM for seawater). To minimize crosscontamination between sample vials and fraction collection vials, we 'chased' the sample injection with a 25 µL injection



Fig. 1. (A) Stratigraphy of Guadalupian-age strata of the Capitan Reef Complex in west Texas, after Present et al. (2019), and references therein. Sampling locations of Present et al. are marked in orange. Samples were taken along a single composite sequence boundary so that samples would be roughly age equivalent. Marked above the uppermost composite sequence boundary are the approximate extents of the shelf, reef, and slope facies tracts, along with the symbols used to represent them in subsequent figures. (B) Different generations of carbonate cement (e.g., botryoids – 'bot', isopachous cements – 'iso', spar) in the Capitan Reef. The dashed white line marks the boundary between the botryoidal cements and later generations of infilling cement, including spar (ringed by the dashed black line). (C) Mimetically dolomitized pisoid rudstone, from the shelf facies tract – scale bar is in mm.

of ultrapure water, designed to clean the autosampler needle and tubing prior to in-line fraction collection (Razionale et al., 2021). Ca and Mg were then isolated from samples by collecting the eluent in timed windows. Ca and Mg peak areas were used to calculate molar Mg/(Ca + Mg) ratios.

Ca and Mg solutions were transferred to Teflon vials and dried down on a hotplate at ~150 °C; samples were then treated with 200 μ L 20 N nitric acid for ~30 minutes to oxidize and volatilize any organic compounds and dried down again. After using the Ca or Mg peak areas as assessed by the in-line conductivity detector during ion chromatography to calculate the mass of Ca or Mg in each vial, the appropriate volume of 2% nitric acid was added to each vial to standardize the [Ca] or [Mg] of each sample to ~1 ppm and ~150 ppb, respectively.

2.5. Multi-collector mass spectrometry methods

Ca and Mg isotope ratios were measured using a Thermo Neptune XT multi-collector inductively-coupled plasma mass spectrometer (MC-ICP-MS). For Ca isotope $(\delta^{44/42}Ca)$ analyses, samples were introduced using an ESI Apex- Ω desolvating nebulizer. Sample-standard bracketing was used to correct for instrumental mass fractionation. Medium- or high-resolution modes were used to avoid interferences such as ArHH⁺ and to allow stable measurement of the low-mass shoulder over > 12-hour-long runs.

For all sample/standard measurements, we used a Faraday cup at 43.5 amu to monitor ${}^{87}Sr^{++}$ and corrected for Sr by assuming that the contributions of ${}^{84}Sr^{++}$, ${}^{86}Sr^{++}$ and ${}^{88}Sr^{++}$ to ${}^{42}Ca$, ${}^{43}Ca$, and ${}^{44}Ca$ are related to ${}^{87}Sr^{++}$ by their natural abundances. The effect of [Ca] on ${}^{64/42}Ca$ was established using a 5-point calibration between 0.9 and 1.1 ppm and corrected for using the ratio of the intensity of the mass 44 ion beam for each sample to that of the prior analysis of the bracketing standard, whose [Ca] was known and constant throughout the run. ${}^{64/42}Ca$ values were calculated from ${}^{64/42}Ca$ using established kinetic fractionation laws (Young et al., 2002) and assuming no radiogenic ${}^{40}Ca$ excess. Long-term external reproducibility was $\pm 0.12\%$ (2σ , n = 44) for NIST SRM 915b, which had an average ${}^{64/44}Ca$ of -1.12% (Razionale et al., 2021), in line with published values.

For Mg isotope analyses, samples were introduced using a dual-path glass cyclonic spray chamber and samplestandard bracketing was used to correct for instrumental mass fractionation. Due to the absence of significant isobaric interferences at masses 24, 25 and 26, low resolution mode was used. The effect of [Mg] on δ^{26} Mg was established using a 5-point calibration between 135 and 165 ppb, and corrected for using the ratio of the intensity of mass 24 for each sample to that of the prior analysis of the bracketing standard, whose [Mg] was known and constant throughout the run. Long-term external reproducibility was $\pm 0.10\%$ (2σ , n = 34) for IHD, $\pm 0.11\%$ For both Ca and Mg, we used triple-isotope plots to ensure the mass-dependence of all measured raw data (see Table S1 for raw isotope data).

2.6. Quadrupole ICP-MS methods

A suite of major and trace element abundances was measured using a Thermo Scientific iCAP RQ ICP-MS. Kinetic energy discrimination (KED) mode, which employs He gas and limits polyatomic interferences, was used to measure Fe, whereas standard (STD) mode was used to measure Ca. Sr and Mn were analyzed in both STD and KED mode. Samples and standards were adjusted to 10 ppm Ca in 2% nitric acid prior to analysis to mitigate matrix related effects. Additionally, samples were spiked with equal amounts of an internal standard solution containing Sc, In, and Y to correct for instrumental drift during analysis. A five-point calibration curve was generated for each element at the start of each run and used to calculate elemental concentrations. One calibration standard and the calibration blank were analyzed intermittently throughout each run for quality control, in addition to one duplicate sample analysis every 15 samples. Data are reported as molar ratios to Ca + Mg.

2.7. Numerical model

Sulfate was added to a previously-published numerical model simulating early marine diagenesis of shallow water carbonates along a one-dimensional advective flow path (Ahm et al., 2018). The model consists of a prescribed series of boxes numbered from 1 (closest to the water column) to 'n' (farthest along the flow path). Fluid flow is assumed to occur by advection rather than diffusion, based on evidence that fluid flow in the Capitan Formation was density driven (Melim and Scholle, 2002). Advection and recrystallization occur at a constant rate until 100% of the primary mineral (e.g., aragonite) has been converted to diagenetic mineral (e.g., calcite). The eventual chemical compositions of diagenetic minerals vary between box 1 and 'n' because the fluid becomes progressively more chemically evolved by all chemical reactions further along the flow path. Thus, the chemistry of diagenetic minerals in box 1 can be described as more fluid-buffered (i.e., determined by the chemistry of the infiltrating fluid, in this case seawater) and the chemistry of diagenetic minerals in box 'n' can be described as more sediment-buffered (i.e., determined by the chemistry of the recrystallizing primary mineral plus any other reactions happening in pore waters). Sulfate is supplied by advection and primary mineral dissolution and then is lost from the fluid components of the system by incorporation into diagenetic mineral (as CAS) and at a constant rate via microbial sulfate reduction (MSR) – i.e., organic matter is assumed to be present in non-limiting amounts. We assumed that all sulfide was immediately lost from the system as pyrite. Parameters for the model were derived from a

combination of published experimental data, including distribution coefficients and fractionation factors between aqueous sulfate and aragonite- and calcite-bound CAS (Barkan et al., 2020). Initial conditions (seawater chemistry) and rates of recrystallization and MSR were varied to generate the best visual fit between model and data. A list of all key assumptions made is given in Table 1.

We conducted a sensitivity analysis to explore the model response (particularly the differences between the $\delta^{44/40}$ Ca and $\delta^{34}S_{CAS}$ values of fluid- and sediment-buffered diagenetic calcite) to a range of inputs, including initial mineral-ogy (aragonite vs. high-Mg calcite), $[SO_4^{2-}]_{seawater}$, $[Ca^{2+}]_{seawater}$, MSR rate, and the Ca isotopic fractionation factor of recrystallization (see Discussion below and supplementary material for more details).

3. RESULTS

A total of 249 sub-samples were analyzed. These samples had Mg/(Ca + Mg) ratios (mol/mol) ranging from 0.003 (low-Mg calcite) to 0.5 (pure dolomite). Present et al. (2019) used a combination of field relationships and petrography to map all hand samples and subsamples onto the detailed facies framework for the Capitan Reef (Tinker, 1998). Here we use a simplified version of that facies framework plus Mg/(Ca + Mg) (this study) as context for comparisons between δ^{13} C, δ^{18} O, and δ^{34} S_{CAS} (Present et al., 2019), and $\delta^{44/40}$ Ca and δ^{26} Mg (this study). Generally, all samples can be assigned to either early marine cements (EMC, i.e., originally-aragonitic botryoids or isopachous cements; Fig. 1B), later diagenetic sparry cements (i.e., equant or poikilitic calcite), or to sedimentary textures from the shelf, reef, or slope facies tracts (Fig. 1A). Shelf samples came from supratidal barrier island/tepee-pisolite, shoreface, shoal, and outer shelf shallow subtidal depositional environments of the Yates and Tansill formations; inner shelf evaporitic lagoonal facies were not sampled. Reef samples came from a shelf margin (but not raised barrier) reef in a deep subtidal-possibly mesophotic-reef of the Capitan Formation. Slope samples of the Capitan and Bell Canyon formations were forereef carbonate debrites and turbidites shed from the reef, which interfingered with basinal clastics. Dolomitization in the shelf, reef, and forereef/ slope was dominantly fabric-retentive with subordinate fabric-destructive recrystallization, including mimetic replacement of aragonite marine cements and fossils (Melim and Scholle, 2002; Frost et al., 2013).

3.1. Magnesium isotopes

Measured δ^{26} Mg values range from -3.41 to -1.41%(Table S1). Generally, samples that are classified as low-Mg calcite [Mg/(Ca + Mg) < 0.04] have more negative and variable δ^{26} Mg values [avg. = $-2.75 \pm 0.34\%$ (1 σ , n = 73)], whereas samples with Mg/(Ca + Mg) > 0.04 (variable mixtures of dolomite and calcite) have more positive and less variable δ^{26} Mg values [avg. = $-1.88 \pm 0.21\%$ (1 σ , n = 162)]. When plotted against Mg/(Ca + Mg), δ^{26} Mg values monotonically increase, plateauing at high Mg concentrations (Fig. 2).

Table 1 List of key model input parameters and literature justification.

Parameter	Value(s)	Reference
u (advection rate)	0.1 m y^{-1}	(Henderson et al., 1999)
MSR rate constant	$3e-3 y^{-1}$	(Canfield, 1991) and refs. therein; this study.
Recrystallization rate constant	$5e-6 y^{-1}$	(Higgins et al., 2018; Staudigel et al., 2021)
[Mg ²⁺] _{seawater}	45 mM	(Farkaš et al., 2007)
[Ca ²⁺] _{seawater}	13 mM	(Farkaš et al., 2007)
[DIC] _{seawater}	3 mM	(Farkaš et al., 2007)
[SO ₄ ²⁻] _{seawater}	13 to 20 mM	$[SO_4^{2-}]$ in excess relative to $[Ca^{2+}]$; (Horita et al., 2002)
$\delta^{26}Mg_{seawater}$	-0.3 to -0.1‰	This study.
$\delta^{44/40}$ Ca _{seawater}	-0.4 to -0.2‰	This study.
$\delta^{13}C_{\text{fluid}}$	-2%	This study.
$\delta^{34}S_{seawater}$	+8 to 10%	(Present et al., 2019) and refs. therein.
$\epsilon^{26}Mg_{aq-cc}$	-2.58%	(Li et al., 2015)
$\epsilon^{26}Mg_{aq-arag}$	-1.15%	(Wang et al., 2013)
$\epsilon^{26}Mg_{aq-dol}$	−2 to −1.81‰	(Li et al., 2012; Higgins et al., 2018)
$\epsilon^{44/40}$ Ca _{aq-arag} (primary)	-1.6%	(Higgins et al., 2018)
$\epsilon^{44/40}$ Ca _{ag-calc} (primary)	-1.0%	(Gussone et al., 2005)
$\epsilon^{44/40}$ Ca _{aq-calc} (diagenetic/	0 to −0.5‰	(Fantle and DePaolo, 2007; Bradbury and Turchyn, 2018; Mills et al., 2021)
recrystallization)		
$\epsilon^{34}S_{SO4-CAS(cc)}$	+1.7%	(Barkan et al., 2020)
$\epsilon^{34}S_{SO4-CAS(arag)}$	+1.0%	(Barkan et al., 2020)
$\epsilon^{34}S_{SO4-CAS(dol)}$	0%0	Similarity between dolomite and evaporite δ^{34} S (Present et al., 2019)
Kd _{CAS(cc)} (partition coeff.)	16e-6	(Barkan et al., 2020)
Kd _{CAS(arag)}	8e-6	(Barkan et al., 2020)
ESO4-H2S	70‰	(Eldridge et al., 2016)

3.2. Calcium isotopes

Measured $\delta^{44/40}Ca$ values range from -1.73 to -0.46% (Table S1; Fig. 3). Samples consisting predominantly of calcite [Mg/(Ca + Mg) < 0.25] have on average slightly more positive $\delta^{44/40}Ca$ values [$-0.88 \pm 0.19\%$ (1 std. dev., n = 133)] than those consisting predominantly of dolomite [$-1.02 \pm 0.19\%$ (1 std. dev., n = 112)]. When $\delta^{44/40}Ca$ values are plotted against $\delta^{26}Mg$ values (Fig. 3), the dolomiterich samples define a limb with relatively invariant $\delta^{26}Mg$ and more variable $\delta^{44/40}Ca$, whereas low-Mg calcites show substantial variability in both parameters and plot entirely distinctly from dolomites in the Ca-Mg phase space. Samples with 0.04 < Mg/(Ca + Mg) < 0.25 generally fall between the calcites with more positive $\delta^{26}Mg$ and $\delta^{44/40}Ca$ values and the center of the dolomite cluster.

3.3. Major/minor element ratios

Measured Sr/(Ca + Mg), Mn/(Ca + Mg) and Fe/(Ca + Mg) ratios range from 2.3 × 10⁻⁶ to 6.6×10^{-4} , 4.1×10^{-7} to 1.3×10^{-3} , and 0 to 5.9×10^{-3} mol mol⁻¹, respectively (n = 248; Table S1; Fig. 4). Samples that are predominantly calcite [Mg/(Ca + Mg) < 0.25] have on average higher and more variable Sr/(Ca + Mg) ratios [$1.7 \times 10^{-4} \pm 1.3 \times 10^{-4}$ mol mol⁻¹ (1σ , n = 135)] than samples that are predominantly dolomite [$6.0 \times 10^{-5} \pm 4.1 \times 10^{-5}$ mol mol⁻¹ (1σ , n = 113)] (Fig. 4A). Calcite-dominated samples have on average similar Mn/(Ca + Mg) ratios [$1.5 \times 10^{-4} \pm 2.0 \times 10^{-4}$ mol mol⁻¹ (1σ , n = 135)] to dolomite-dominated samples [$1.3 \times 10^{-4} \pm 1.7 \times 10^{-4}$ mol mol⁻¹ (1σ , n = 113)] (Fig. 4B). Calcite-dominated samples have on average lower Fe/(Ca + Mg) ratios $[1.6 \times 10^{-4} \pm 3.1 \times 10^{-4} \text{ mol} \text{mol}^{-1}$ (1 σ , n = 135)] than dolomite-dominated samples $[3.1 \times 10^{-4} \pm 5.8 \times 10^{-4} \text{ mol} \text{ mol}^{-1}$ (1 σ , n = 113)] (Fig. 4B). Within the calcite-dominated samples, Sr/(Ca + Mg), Mn/(Ca + Mg) and Fe/(Ca + Mg) are highest for a subset of samples from the slope facies tract (specifically, the toe-of-slope) (Fig. 4). The toe-of-slope samples also have more negative $\delta^{44/40}$ Ca values than most of the other calcite-dominated samples (Fig. 4A).

3.4. Comparison of $\delta^{44/40}Ca$ and $\delta^{26}Mg$ with facies, $\delta^{13}C,$ $\delta^{18}O,$ and $\delta^{34}S_{CAS}$

Sample δ^{26} Mg values become more negative and more variable along a shelf-basin transect (Fig. 5). This trend is primarily driven by the increasing relative frequency of samples dominated by low-Mg calcite in basinward facies tracts (Fig. 5A). An additional driver of the basinward variability is the fact that dolomite-dominated samples in the slope facies tract are ²⁶Mg-enriched on average relative to dolomite-dominated samples from shelf and reef facies tracts (Fig. 5A). When δ^{26} Mg values are plotted versus $\delta^{13}C,\ \delta^{18}O,\ and\ \delta^{34}S_{CAS}$ values (Fig. 5B-D), several commonalities emerge. Dolomite-dominated samples form a tight and distinct cluster in all three plots. Calcitedominated samples (with the obvious exception of the spar facies tract) generally define a limb of variable δ^{26} Mg and relatively invariant δ^{13} C, δ^{18} O, and δ^{34} S_{CAS} in all three plots, with the magnitude of the δ^{26} Mg deviation from the dolomitic cluster to first order negatively correlating with Mg/(Ca + Mg). Slope and spar facies tracts generally plot farthest from the dolomitic cluster in the δ^{13} C, δ^{18} O, and $\delta^{34}S_{CAS}$ axes.



Fig. 2. δ^{26} Mg values for carbonate samples from Capitan Reef, plotted against Mg/(Ca + Mg). Symbols correspond to sample facies/textures and are color coded by Mg/(Ca + Mg). Dashed lines and arrows indicate the x-axis positions of upper limit for Mg in low-Mg calcite, ideal stoichiometric dolomite, and a 50:50 mix of those two mineral end members, respectively.



Fig. 3. $\delta^{44/40}$ Ca values for carbonate samples from Capitan Reef, plotted against δ^{26} Mg. Symbols correspond to sample facies/textures and are color coded by Mg/(Ca + Mg).



Fig. 4. Minor/major element ratios for carbonate samples from Capitan Reef. (A) Sr/Ca + Mg plotted against $\delta^{44/40}$ Ca values, (B) Fe/ (Ca + Mg) plotted against Mn/(Ca + Mg). Symbols correspond to sample facies/textures and are color coded by Mg/(Ca + Mg).

Sample $\delta^{44/40}$ Ca values show a large degree of variability for both dolomite- and calcite-dominated samples, but calcite-dominated samples are on average enriched in ⁴⁴Ca relative to dolomite-dominated samples – this is true for all facies tracts where both minerals are present (Fig. 6A). When $\delta^{44/40}$ Ca values are plotted against δ^{13} C, δ^{18} O, and δ^{34} S_{CAS} values (Fig. 6B-D), the data fall into three clusters defined by mineralogy and facies: dolomitedominated samples, sparry calcite, and non-sparry calcitedominated samples. When samples with chemistry clearly influenced by meteoric or burial fluids during later diagenesis [i.e., $\delta^{18}O < -2\%$ and negative $\delta^{13}C$; (Present et al., 2019)] are excluded, broad trends are visible between $\delta^{44/40}$ Ca and the other variables within the non-sparry calcite-dominated cluster, some of which are clearly defined by the transect from shelf-reef-slope facies tracts. Notably, $\delta^{44/40}$ Ca and $\delta^{34}S_{CAS}$ have a significant negative relationship (R = -0.31, p-value = 0.0023; Fig. S1A), with the most ³⁴S-enriched and ⁴⁴Ca-depleted samples belonging to the slope facies tract, and the reverse being true for the shelf facies tract. When EMCs are excluded, the correlation between $\delta^{44/40}$ Ca and $\delta^{34}S_{CAS}$ becomes clearer $(R = -0.49, p-value = 1.3 \times 10^{-5}; Fig. S1B).$

4. INTERPRETATION AND DISCUSSION

The Mg and Ca isotope trends we observe, when coupled with existing geochemical data and facies context (Present et al., 2019), shed new light on the diagenetic histories of the various sedimentary components of the Capitan Reef complex. After interpreting the Mg and Ca isotope data, we will discuss the broader implications for the chemistry of the Delaware Basin and environmental drivers of the precipitation of the early marine cements.

4.1. Magnesium and calcium isotopic constraints on the diagenetic history of the Capitan Reef

We interpret the relationship between samples' $\delta^{26}Mg$ values and Mg/(Ca + Mg) (Fig. 2) as indicative of a mixing trend between two isotopically distinct, albeit noisy, endmembers: low-Mg calcite and dolomite. While only a minority of samples entirely escaped the pervasive dolomitization of the Capitan Reef Complex, some samples (mostly from the reef and slope facies) are recrystallized calcite – these samples display the most δ^{26} Mg variance. In contrast, the δ^{26} Mg variance in the dolomites is small, which we interpret to mean that dolomite formation was predominantly seawater-buffered with respect to Mg (i.e., only a minor component of the dolomite Mg was sourced from precursor carbonate minerals). Even when $[Mg^{2+}]$ is high, Rayleigh distillation during dolomitization causes the δ^{26} Mg value of the dolomitizing fluid to become progressively more positive (Bialik et al., 2018), so the most negative dolomite δ^{26} Mg value is the most informative regarding the δ^{26} Mg value of the starting fluid. The most negative δ^{26} Mg value of dolomite was $\sim -2.1\%$ and the isotopic fractionation (ϵ) between aqueous Mg²⁺ and dolomite is -2 to -1.8% (Li et al., 2015; Higgins et al., 2018). Our data therefore suggest that the δ^{26} Mg value of the Delaware Basin seawater was between -0.3 and -0.1%. This is similar to a prior estimate of -0.1% by Li et al. (2015), which was based on their experimentally derived dolomite fractionation factor and compiled dolomite δ^{26} Mg values. If calcite formed directly from seawater with a δ^{26} Mg value of -0.3 to -0.1% at equilibrium, expected δ^{26} Mg values would be between -2.9 and -2.7% (Li et al., 2012; Li et al., 2015), which is broadly consistent with the average δ^{26} Mg value of calcite measured in this study,



Fig. 5. Comparison of δ^{26} Mg (this study) with existing data (Present et al., 2019). (A) δ^{26} Mg histograms by facies group; (B-D) δ^{26} Mg versus δ^{13} C, δ^{18} O, and δ^{34} S_{CAS}. Symbols correspond to facies tracts and are color coded by Mg/(Ca + Mg).

 $-2.75 \pm 0.34\%$, perhaps indicating that the majority of recrystallized calcites precipitated from a fluid that was minimally evolved from seawater, with respect to δ^{26} Mg. However, the most 26 Mg-depleted calcites in the slope and reef likely inherited some of their Mg from primary calcite bioclasts (i.e., recrystallized in more sediment-buffered conditions), which in general have larger δ^{26} Mg fractionations from seawater (Mavromatis et al., 2013; Higgins

and Schrag, 2015). Additionally, there is sedimentological evidence that some dolomitization of primary carbonates started before other primary carbonates recrystallized to low-Mg calcite in the reef and slope (Mutti and Simo, 1994; Melim and Scholle, 2002; Present et al., 2019), which could have made the samples that escaped dolomitization more likely to retain something closer to their precursor δ^{26} Mg upon their later recrystallization.



Fig. 6. Comparison of $\delta^{44/40}$ Ca (this study) with existing data (Present et al., 2019). (A) $\delta^{44/40}$ Ca histograms by facies group; (B-D) $\delta^{44/40}$ Ca versus δ^{13} C, δ^{18} O, and δ^{34} S_{CAS}. Symbols correspond to facies tracts and are color coded by Mg/(Ca + Mg).

Some δ^{26} Mg variability correlates with facies. For example, the dolomite-dominated samples of the slope facies tract are ~0.2‰ enriched in ²⁶Mg relative to the average dolomite-dominated sample from the shelf facies tract (Fig. 5A). This is likely to reflect more sediment-buffered conditions in the slope facies, which along with upstream

precipitation of ²⁶Mg-depleted phases may have led to ²⁶Mg-enrichment of the dolomitizing fluid by Rayleigh distillation and slightly elevated δ^{26} Mg of dolomites along the fluid flow path.

The $\delta^{44/40}$ Ca values paint a more complex picture of diagenesis in the Capitan Reef. Firstly, both calcite- and

dolomite-dominated samples display a broad range of $\delta^{44/40}$ Ca values (Fig. 3), indicating that the mineralogical endmembers reflect a variety of different conditions or initial compositions. Additionally, dolomite-dominated samples can be variably depleted in ⁴⁴Ca relative to calcitedominated samples in the same facies group - particularly on the shelf, where dolomitization was most pervasive (Fig. 6). This observation indicates a larger and more variable isotopic fractionation between seawater Ca²⁺ and dolomites than seawater Ca^{2+} and calcites. One possibility is that the primary sediment was a local source of Ca during dolomitization along the fluid flow path, through the substitution of Mg²⁺ for Ca²⁺, in addition to the influx of seawater Ca^{2+} . However, the shelf dolomites and calcites were likely precipitated in an open-system environment where Ca transport was advection-dominated [i.e., by density-driven refluxing brines (Melim and Scholle, 2002)], and relatively fluid-buffered with respect to $\delta^{44/40}$ Ca. As a result, it is possible that the variable calcite-dolomite $\delta^{44/40}$ Ca offset reflects varying extents of dolomitization and/or variation in the fractionation factor during dolomitization, driven by variable precipitation rates or $Ca^{2+}:CO_3^{2-}$ ratios in the pore-fluids (Tang et al., 2008; Mills et al., 2021). Variability in the Ca isotope fractionation factor during dolomitization is supported by the relatively invariant $\delta^{34}\bar{S}_{CAS}$ and $\delta^{26}Mg$ values of the dolomites (Present et al., 2019), indicating that dolomites were fluid-buffered by seawater-derived brines.

Variability in $\delta^{44/40}$ Ca values within the non-sparry calcite-dominated samples appears to correspond to variation in Sr/(Ca + Mg), δ^{13} C, and δ^{34} S_{CAS}, which maps onto facies tracts (Fig. 7). Interpreting these observations with a

diagenetic model, modified from Ahm et al. (2018), allows for the determination of and detailed comparison between fluid- versus rock-buffered diagenetic calcite endmembers - in addition to their percent recrystallization [based on Sr/(Ca + Mg)] – in a multi-proxy phase space. Here, we show that this approach provides an internally consistent, mechanistic, and semi-quantitative understanding of $\delta^{34}S_{CAS}$ behavior during early marine diagenesis along a shelf-basin transect. Additionally, the richness of the data allows us to disentangle the primary seawater signal (e.g., the concentration and isotopic composition of seawater SO_4^{2-}), allowing for a greater understanding of the ancient sulfur cycle. We used the results of a sensitivity analysis to determine suitable values for various input parameters (initial mineralogy, [SO₄²⁻]_{seawater}, [Ca²⁺]_{seawater}, MSR rate, the number of boxes ('n'), and the Ca isotopic fractionation factor for recrystallization; Fig. S2-7).

Fig. 7 shows crossplots of $\delta^{44/40}$ Ca and $\delta^{34}S_{CAS}$ (Fig. 7A), and $\delta^{13}C$ and Sr/(Ca + Mg) (Fig. 7B), overlain by model-data fits of the compositions of fully recrystallized diagenetic calcite precipitated in a range of fluid- to sediment-buffered conditions. In both plots, different facies plot on a continuum between modeled fluid- and sedimentbuffered diagenetic endmembers, close to the fully recrystallized limb, with very small contributions from the remaining primary precursor mineral. Specifically, samples from the slope facies tract plot closest to the more sedimentbuffered endmember (box n), shelf samples plot closest to the fluid-buffered endmember (box 1), and reef samples plot in an intermediate position. The data indicate that Permian seawater had a $\delta^{44/40}$ Ca value of greater than -0.4%



Fig. 7. Diagenetic model fits of, (A) $\delta^{44/40}$ Ca and δ^{34} S_{CAS}, and (B) δ^{13} C and Sr/(Ca + Mg), for non-sparry, calcite-dominated samples. The blue shaded region labeled 'SW' indicates the inferred composition of Delaware Basin seawater/brine; the white star indicates the expected composition of primary aragonite given empirical fractionation factors. Note that the displayed primary aragonite δ^{13} C composition is a maximum estimate – primary aragonite in shelf settings was likely several per mil enriched relative to reef and slope settings (Rivers et al., 2019; Present et al., 2019; Smith et al., 2020). The two black curves indicate the evolution of sediment- and fluid-buffered endmembers during progressive aragonite neomorphism; Gray curves are recrystallization deciles (10–90%) and the blue line indicates full recrystallization, i.e., diagenetic calcite. All these samples are ~100% recrystallized, so the spread in the data is not caused by varying degrees of recrystallization.

(IAPSO), likely closer to -0.2% (Fig. 7A). This is consistent with estimates of Permian seawater $\delta^{44/40}$ Ca from evaporites (Blättler and Higgins, 2014).

EMCs plot slightly away from the main shelf-reef-slope trend in Fig. 7A (notably ⁴⁴Ca-depleted relative to seawater, but with the most seawater-like $\delta^{34}S_{CAS}$), albeit closest to the shelf samples. This indicates that although these cements were likely fluid-buffered with respect to δ^{34} S (as expected for crystals growing directly in contact with seawater), a parameter other than the degree of fluidsediment buffering generated the $\delta^{44/40}$ Ca variability. EMCs either. (1) were additionally subject to larger or more variable isotopic fractionations between primary mineral and fluid (Tang et al., 2008; Mills et al., 2021), (2) did not fully recrystallize (Chafetz et al., 2008), or (3) had a negative isotopic fractionation from the diagenetic fluid during recrystallization (Bradbury and Turchyn, 2018; Mills et al., 2021). Neither explanation 1 nor 2 is strongly supported by the Sr/ (Ca + Mg) ratios of EMCs, which are very low, albeit slightly higher than those of reef and shelf samples (Fig. 7B). Therefore, explanation 3 [involving an inferred negative fractionation between pore water fluid and diagenetic mineral, -0.2 to -0.5% (Bradbury and Turchyn, 2018; Mills et al., 2021); Fig. 7A, S7] is the most plausible for the $\delta^{44/40}$ Ca values of the EMCs.

4.2. Constraints on Permian seawater [Ca²⁺]:[SO₄²⁻]

The offset between the sulfur isotope ratio of CAS in recrystallized carbonates and that of seawater ($\Delta^{34}S_{CAS}$ seawater) is predominantly a function of the relative rates/timescales of carbonate recrystallization versus MSR below the sediment-water interface (Rennie and Turchyn, 2014). In modern sediments, the rates of both recrystallization and organoclastic MSR have been observed to decay exponentially with increasing burial depth (Middleburg, 1989; Fantle and DePaolo, 2007). The rates of sulfate reduction [mostly a function of the initial concentration of labile organic matter; (Boudreau and Westrich, 1984)] and transport of new sulfate into pore waters control the timescale of sulfate drawdown in the pore fluid. The timescale of carbonate recrystallization is typically on the order of millions of years for typical deep-sea sediments (Fantle et al., 2010), and approaching 1 Myr for more rapidly accumulating platform sediments (Higgins et al., 2018; Staudigel et al., 2021). Evidence from the geologic record suggests much shorter timeframes (sub-100 Kyr) timeframes for early diagenetic dolomitization and lithification (Manche and Kaczmarek, 2019; Ning et al., 2020), and stratal geometries of dolomite in the Capitan Reef Complex indicate that dolomitization occurred on the timescale of high frequency sequences (sub-Myr) (Mutti and Simo, 1993; Mutti and Simo, 1994; Garcia-Fresca et al., 2012; Wu et al., 2020). An ideal ratio exists between the rates of MSR and recrystallization that will result in maximally elevated $\delta^{34}S_{CAS}$ values, with a maximal $\Delta^{34}S_{CAS-seawater}$ of between 1 and 3% for three modern sites in slope settings (Rennie and Turchyn, 2014). Ratios that are either too high or too low would result in very little incorporation of ³⁴Senriched sulfate into diagenetic calcite because pore water sulfate would either be consumed prior to substantive recrystallization (recrystallization/MSR ratio too low) or would not diverge from the isotopic composition of seawater sulfate prior to complete recrystallization (recrystallization/MSR ratio too high). In addition to recrystallization and MSR, the magnitude of $\Delta^{34}S_{CAS-seawater}$ in geologic carbonates also depends on other factors that influence the drawdown of sulfate in pore waters, including $[SO_4^2]_{seawater}$ and the depth of the redox-cline, as a function of bottom water $[O_2]$.

Given that the Capitan Reef Complex samples span a shelf-basin transect, we expected them to capture a large range (from fully fluid- to more sediment-buffered) of δ^{34} - S_{CAS} and $\delta^{44/40}$ Ca values. If seawater $\delta^{34}S_{sulfate}$ was + 9% (Present et al., 2019), $\Delta^{34}S_{CAS}$ -seawater ranges from 1 to 10% for the calcite-dominated samples (excluding sparry cements) (Fig. 7A). The smallest $\Delta^{34}S_{CAS-seawater}$ values occur in EMCs on the shelf and the largest occur in the slope facies tract (particularly the toe-of-slope). This sample set therefore provides a new opportunity to set constraints on the major ion seawater chemistry of the Delaware Basin. We use a numerical model to explore the key parameters that determine the geochemical spread of data in the δ^{34} - $S_{CAS}/\delta^{44/40}$ Ca phase space. To explain the observed ³⁴Senrichment (Fig. 7A), MSR and recrystallization must overlap in time and space (Rennie and Turchyn, 2014). The model predicts that the relative rates of MSR and recrystallization control the spread of $\delta^{34}S_{CAS}$ values (but not $\delta^{44/40}$ Ca values; Fig. S5) as a higher rate of MSR results in greater ³⁴S-enrichment of residual sulfate along the fluid flow path. Alternatively, lowering seawater (and pore water) $[SO_4^{2-}]$ could produce a similar pattern because of the effect of reservoir size on Rayleigh distillation (Fig. S2). Despite being highly productive environments, carbonate platforms typically preserve low organic matter concentrations (Geyman and Maloof, 2019), mostly due to high rates of aerobic respiration near the sediment-water interface (Bowles et al., 2014). Similarly, these rocks have very little preserved organic matter [e.g., the Capitan Fm. contains on average < 0.5 wt.% organic matter; (Mazzullo and Cys, 1983)]. Thus, it is hard to imagine the Capitan Reef having supported high net rates of sulfate reduction throughout the sediment column, and it seems likely that low seawater [SO₄²⁻] contributed to the spread of the δ^{34} -S_{CAS} data. Holding all other model input parameters (rates, concentrations and isotopic compositions) constant and at reasonable values based on literature estimates (see Table 1), we varied $[SO_4^{2-}]$ between 13 and 52 mM such that the $[Ca^{2+}]$: $[SO_4^{2-}]$ ratio varied between 1.0 [upper constraint from evaporite fluid inclusions (Horita et al., 2002)] and 0.25. We find that a $[Ca^{2+}]$: $[SO_4^{2-}]$ ratio approaching 1 (likely close to 0.8) gives the best model-data fit (Fig. 8) as the modeled fully recrystallized diagenetic calcite spans the range of both the $\delta^{44/40}$ Ca and $\delta^{34}S_{CAS}$ data with an appropriate slope. Our [Ca²⁺]:[SO₄²⁻] ratio estimate is at the upper end of estimates from fluid inclusions (Horita



Fig. 8. Illustration of how lowering $[Ca^{2+}]$; $[SO_4^{2-}]$ _{seawater} (by raising $[SO_4^{2-}]$ _{seawater}) affects predictions for the $\delta^{34}S_{CAS}$ and $\delta^{44/40}Ca$ values by decreasing the slope of recrystallized/diagenetic calcite. For these conditions, a $[Ca^{2+}]$; $[SO_4^{2-}]$ _{seawater} between 0.5 and 1 (i.e., 13 mM < $[SO_4^{2-}]$ _{seawater} < 26 mM) is most consistent with the data – this is because a high $[SO_4^{2-}]$ _{seawater} results in slower drawdown and isotopic distillation of pore water $[SO_4^{2-}]$ relative to the rate of recrystallization, minimizing the degree of ³⁴S-enrichment of diagenetic CAS relative to seawater-SO_4^{2-}. If a smaller MSR rate constant than that assumed here [3e-3 y⁻¹ (Canfield, 1991)] were used, the corresponding slopes of the model outputs would be shallower, and vice versa (see Fig. S5). If a more positive Ca fractionation factor of recrystallization than that assumed here [-0.2‰ (Mills et al., 2021)] were used, the model outputs would be shifted to more positive $\delta^{44/40}Ca$ values, with slightly shallower slopes, and vice versa (see Fig. S7).

et al., 2002), and roughly two to three times that observed in the modern ocean – however it is not so high as to be inconsistent with the presence of sulfate evaporites and sulfate-rich fluid inclusions in the Delaware Basin (Leslie et al., 1997; Horita et al., 2002).

4.3. Implications for the origins of the early marine cements of the Capitan Reef Complex

Our interpreted $[Ca^{2+}]$: $[SO_4^{2-}]$ value for Permian seawater could help explain the anomalous presence of abundant early marine (botryoidal and isopachous) cements in the Capitan Reef Complex – a style of cementation famously compared to Precambrian examples by Grotzinger and Knoll (1995). Those authors suggested that upwelling anoxic deep waters could have provided a source of alkalinity for seafloor carbonate precipitation. Due to the biological pump, upwelling waters would be expected to have more ¹³C-depleted dissolved inorganic carbon (DIC) than surface waters – however, we see no evidence of contribution of a ¹³C-depleted DIC pool to any subset of the carbonates deposited during this time interval (excluding those impacted by burial diagenesis). Instead, we infer that primary carbonates must have had a δ^{13} C value of ~+5 – 7% [and possibly several per mil higher than this in evaporitic, supratidal shelf environments due to the greater degree of restriction (Rivers et al., 2019)], consistent with interpretations from previous studies (Given and Lohmann, 1985; Mutti and Simo, 1994; Smith et al., 2020) (Fig. 7B). The inferred enriched δ^{13} C value of platform aragonite is consistent with carbonate precipitation from a DIC pool that was ¹³C-enriched due to either photosynthesis (Geyman and Maloof, 2019; Present et al., 2019) and/or CO₂ degassing caused by a combination of evaporation and carbonate precipitation in a restricted setting (Stiller et al., 1985; Beeler et al., 2020). To explain the differences in $\delta^{13}C$ between sediment- and fluid- buffered endmembers (Fig. 7B), DIC in pore waters must have been substantially ¹³C-depleted relative to the water column [as expected from organic matter remineralization; (Meister et al., 2019)]. Based on the $\delta^{44/40}$ Ca, $\delta^{34}S_{CAS}$ and $\delta^{13}C$ data for the

Capitan Reef samples, rather than invoking an alkalinity influx for the extensive cement precipitation, we posit that an elevated $[Ca^{2+}]$: $[SO_4^{2-}]$ similar to that of the late Proterozoic contributed to the favorability of carbonate cement precipitation. As sulfate inhibits the precipitation of calcite and aragonite (Reddy and Nancollas, 1976; Sjoberg, 1978; Busenberg and Niel Plummer, 1985; Walter, 1986; Burton and Walter, 1987; Mucci et al., 1989), a lower [SO₄²⁻]_{seawater} in the Delaware Basin would have increased the kinetic favorability of early marine carbonate cement precipitation. In addition, a combination of the restrictedness of the depositional setting (Garber et al., 1989) and low sea level (Ridgwell, 2005; Smith et al., 2021) may have further increased the basin wide saturation state of aragonite. Higher temperatures in the shallow waters overlying the shelf and reef facies would have particularly favored the precipitation of aragonite over calcite (Burton and Walter, 1987). However, on the shelf, higher rates of aerobic respiration might have lowered the aragonite saturation state below the sediment-water interface (Bergmann et al., 2013), opposing the preservation of aragonite and favoring early diagenetic dolomitization (Rivers et al., 2021).

5. CONCLUSIONS

The $\delta^{34}S_{CAS}$ values of the Permian Capitan Reef carbonates are highly sensitive to the degree of fluid- vs. sediment-buffering of the environment of carbonate precipitation, which can be assessed using other carbonate-based geochemical proxies such as $\delta^{44/40}Ca,\ \delta^{26}Mg$ and Sr/(Ca + Mg). Because sulfur isotopes are sensitive to microbial sulfur cycling processes such as MSR, whereas other proxies are mostly subject to recrystallization of primary carbonate minerals, we can constrain the spatial and/or temporal overlap in the processes of MSR and carbonate recrystallization from the correlations between the proxies. Most notably, for non-sparry calcites, $\delta^{44/40}$ Ca and $\delta^{34}S_{CAS}$ are negatively correlated, a relationship that maps onto facies context. In contrast, dolomites consistently record seawater $\delta^{34}S_{sulfate}$, despite having variable $\delta^{44/40}$ Ca values (likely due to variable precipitation rates or $Ca^{2+}:CO_3^{2-}$ ratios). The relative lack of $\delta^{26}Mg$ variability

in the dolomites emphasizes that dolomitization occurred during early, fluid-buffered diagenesis, locking in the signal of seawater/brine $\delta^{34}S_{sulfate}$. Our results demonstrate that $\delta^{44/40}$ Ca and δ^{26} Mg can be powerful tools for identifying fluid-buffered rocks, which - so long as the fluid is syndepositional seawater – may record seawater $\delta^{34}S_{sulfate}$, even in the absence of facies context. Furthermore, we show that the richness of the data in the Ca-S isotopic phase space enables semi-quantitative assessment of water column $[Ca^{2+}]$: $[SO_4^{2-}]$ ratios. For the Capitan Reef, an estimated $[Ca^{2+}]$: $[SO_4^{2-}]$ ratio of ~1 is the most parsimonious explanation for the presence of ubiquitous pre-Cambrian-like early marine cements. Finally, looking to the broader rock record prior to the availability of pelagic drill core sediments, we can use a paired $\delta^{44/40}$ Ca - δ^{26} Mg approach to identify early diagenetic dolomites and the most fluidbuffered calcites, both of which are likelier than sedimentbuffered calcites to record seawater $\delta^{34}S_{sulfate}$ in their δ^{34} -S_{CAS} values.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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RESEARCH DATA

All data presented in this study can be found in the attached file named Research Data. The model code is available from the authors upon request.

APPENDIX A. SUPPLEMENTARY MATERIAL

Supplementary material to this article can be found online at https://doi.org/10.1016/j.gca.2022.04.027.

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