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Calcium isotopes in deep time: Potential and limitations

Nikolaus Gussone^{a,*}, Anne-Sofie C. Ahm^b, Kimberly V. Lau^c, Harold J. Bradbury^d

^a Institut für Mineralogie, Universität Münster, Corrensstraße 24, D-48149 Münster, Germany

^b Department of Geosciences, Princeton University, Guyot Hall, Princeton, NJ, USA

^c Department of Geology and Geophysics, University of Wyoming, Laramie, WY, USA

^d Department of Earth Sciences, University of Cambridge, Cambridge, UK

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ABSTRACT

Calcium is an essential element in the biogeochemical cycles that regulate the long-term climate state of Earth. The removal of CO₂ from the ocean-atmosphere system is controlled by the burial of carbonate sediments (CaCO₃), ultimately linking the global calcium and carbon cycles. This fundamental link has driven the development of the stable calcium isotope proxy with application to both ancient skeletal and non-skeletal bulk carbonate sediments. Calcium isotope ratios ($\delta^{44/40}$ Ca) have been used to track long-term changes in seawater chemistry (e.g., aragonite vs. calcite seas) and to elucidate short-term climatic perturbations associated with mass extinction events. However, developments in the calcium isotope proxy have shown that $\delta^{44/40}$ Ca values in carbonate minerals also are sensitive to changes in precipitation rates, mineralogy and diagenesis, thereby complicating the application of the proxy to the reconstruction of global cycles. First, inorganic carbonate precipitation experiments have demonstrated that carbonate $\delta^{44/40}$ Ca values are sensitive to precipitation rates with higher rates generally leading to larger fractionation. Second, $\delta^{44/40}$ Ca values are sensitive to carbonate mineralogy with inorganic aragonite and calcite being on average \sim 1.5‰ and \sim 0.9‰ depleted relative to contemporaneous seawater, respectively. The effects of both changes in carbonate mineralogy and precipitation rates affect primary and secondary minerals, but are particularly pronounced during carbonate diagenesis where relatively slow rates of recrystallization and neomorphism can lead to significant changes in bulk sediment $\delta^{44/40}\text{Ca}$ values. Third, changes in faunal composition expressed in skeletal fossil archives can lead to large changes in carbonate $\delta^{44/40}$ Ca values that are decoupled from changes in global cycles. Nevertheless, when these factors are appropriately considered the application of calcium isotopes in ancient carbonate sediments becomes a powerful tool for understanding biogeochemical processes that operate over many scales; from diagenetic changes within the sediment pore-space, to regional changes across ancient carbonate platforms, and to global changes in seawater chemistry through time. Importantly, the processes that contribute to variability in carbonate $\delta^{44/40}$ Ca values are likely to impact other carbonate-bound proxies, highlighting the potential for calcium isotopes as a tool to better understand the variability of other isotope systems.

1. Introduction

Variations in the Ca isotope composition of rocks, minerals, fossils, soils and fluids have attracted the interest of researchers since the origins of mass spectrometry. As a major constituent in magmatic, metamorphic and sedimentary rocks, Ca is involved in the global cycling of rocks; it plays an important role during melting and crystallization, metamorphism and metasomatism, authigenesis and recrystallization, as well as rock alteration and weathering. In addition, Ca plays an important role in the metabolic pathways of organisms and as a stoichiometric component of numerous biominerals. In particular, the mass production of $CaCO_3$ in the ocean provides an important link

between the global Ca and C cycles which regulates the long-term climate state on Earth.

Calcium isotope ratios have been used to identify sources and quantify budgets in geochemical cycles on various spatial and temporal scales, ranging from molecular to solar system processes and from seconds to billions of years. Calcium has six stable or extremely longlived isotopes (⁴⁰Ca to ⁴⁸Ca), spanning a relative mass-difference of 20% (e.g., Haynes et al., 2017) and mass-dependent fractionation between these isotopes can be induced by chemical, physical and biological processes. During the last two decades, the increased precision and accuracy of analytical techniques has resulted in greater resolution of natural isotopic variability and contributed to an increased

* Corresponding author.

E-mail address: nikolaus.gussone@uni-muenster.de (N. Gussone).

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understanding of mass-dependent Ca isotope systematics. In particular, high-precision records can reveal small scale Ca isotope fluctuations in deep time-records, which may not be resolved in less precise data. These seemingly discrepant results can further lead to different concepts of mechanisms, governing Ca isotope fractionation on global and or local scale. Nevertheless, a comparison of different analytical approaches is not within the scope of this contribution, as a separate article within this special issue is dedicated to the development of analytical methods (Chakrabarti et al., this issue) and consequently not further discussed here.

In the literature, mass dependent differences of Ca isotope ratios are mostly reported as ⁴⁴Ca/⁴⁰Ca or ⁴⁴Ca/⁴²Ca ratios, expressed as either $\delta^{44/40}$ Ca [%] = ((⁴⁴Ca/⁴⁰Ca)_{sample}/(⁴⁴Ca/⁴⁰Ca)_{standard} - 1) · 1000 or $\delta^{44/42}Ca \ [\%] = (({}^{44}Ca/{}^{42}Ca)_{sample}/({}^{44}Ca/{}^{42}Ca)_{standard} - 1) \cdot 1000, re$ spectively. Because of the different relative mass differences between ⁴⁴Ca and ⁴⁰Ca or ⁴⁴Ca and ⁴²Ca, $\delta^{44/40}$ Ca and $\delta^{44/42}$ Ca differ from each other. The relation between both δ -values differs, if kinetic or equilibrium isotope fractionation is assumed. For kinetic isotope fractionation the conversion follows $\delta^{44/40}$ Ca = $\delta^{44/42}$ Ca · $(\ln(m^{44}$ Ca/m^{40}Ca))/ (ln(m⁴⁴Ca/m⁴²Ca)) and can be approximated by $\delta^{44/40}Ca~[\%]$ = $\delta^{44/42}\text{Ca}$ \cdot 2.05, and for equilibrium isotope fractionation $\delta^{44/40}$ Ca = $\delta^{44/42}$ Ca · $(1/m^{44}$ Ca - $1/m^{40}$ Ca)/ $(1/m^{44}$ Ca - $1/m^{42}$ Ca), which can be approximated by $\delta^{44/40}$ Ca [‰] = $\delta^{44/42}$ Ca · 2.10, with m^xCa being the exact atomic masses of the respective Ca isotopes (e.g. Heuser et al. 2016). In addition, it should be noted that there is currently no consensus on the normalizing standards for Ca isotope δ values. However, based on recommendations of the IUPAC (Coplen et al., 2002), most studies report Ca isotope data either relative to SRM 915a or to modern seawater, with seawater being +1.88% ($\delta^{44/40}$ Ca) and 0.92‰ ($\delta^{44/42}$ Ca) relative to SRM 915a (e.g., Heuser et al., 2016). Here, we report data as $\delta^{44/40}$ Ca relative to SRM 915a.

Early Ca isotope research focused on processes taking place early in Earth history and the solar system. Favoured topics included high temperature processes, such as mass-dependent isotope fractionation during condensation and evaporation, nucleosynthetic anomalies and the branched ⁴⁰K decay to ⁴⁰Ar and ⁴⁰Ca. For example, early research investigated the Ca isotope variability of the oldest preserved components formed in the solar system (Ca-aluminium-rich inclusions), radiogenic ⁴⁰Ca signatures in ancient magmatic rocks and age dating of old terrestrial and lunar rocks (e.g., Bermingham et al., 2018; Kreissig and Elliott, 2005; Marshall and DePaolo, 1989; Russell et al., 1978; Shih et al., 1993; Antonelli et al., this issue; Valdes et al., this issue).

The consequences of mass-dependent Ca isotope fractionation for low temperature processes have focused on reconstruction of paleoenvironments and geochemical cycling of Ca through time. In the low temperature environment, the coupling between Ca and carbonate has received special attention, ultimately connecting Ca with the global C cycle and climate. Consequently, a large proportion of studies dealing with Ca isotopes are directly or indirectly aimed at constraining oceanic Ca—C cycling. This link has mainly been explored through paleo-seawater reconstructions of Ca isotope ratios and Ca budget modelling that is constrained by estimates of sources and sinks from the modern (see Section 2.2). In addition to the link with C cycling, Ca isotopes can contribute to a better understanding of sulphate and phosphate cycling, for example, in evaporitic basins and upon formation of phosphorite deposits (Blättler and Higgins, 2014; Soudry et al., 2006).

The focus of this article is to discuss the potential and challenges of using Ca isotopes to study Ca cycling throughout Earth history. We highlight how increasing understanding of processes that fractionate Ca isotopes has led to a re-examination of existing approaches and the development of new concepts. The intention of this article is not to produce a fixed recipe to generate Ca isotope records in deep time. Rather, we encourage the reader to reconsider and validate basic assumptions, potentially influencing parameters and model prerequisites for the respective regional setting, sedimentary facies and investigated time period. The continuing evaluation of fundamental assumptions and model parameters is required for future developments in Ca isotope research. The newly gained or refined knowledge on Ca isotope fractionation processes or characteristics of Earth's reservoirs point to both potentials for novel applications and to complications for established approaches, which need to be taken into account for meaningful reconstructions.

In this paper, we review methods of tracking Ca isotope evolution through Earth history by modelling the oceanic Ca isotope budget, based on different archives (e.g., biominerals, bulk carbonate, phosphate, gypsum and barite). We discuss the methods, assumptions, and issues associated with each technique, and then provide an overview of the published Ca isotope records. Finally, we discuss the potential of Ca isotope ratios, through these methods, for providing new insights into seawater Ca isotope variations, carbonate diagenesis and mineralogy.

2. Global Ca cycling throughout Earth history

2.1. The oceanic Ca isotope budget

When interpreting the Ca isotope record in deep time, the original goal was to reconstruct the ocean budget of Ca, in order to quantify global changes in the input and output fluxes. One of the major tools for interpreting changes in the global Ca cycle through deep time is the use of a box model. Calcium concentration and isotope box models have been used to reconstruct the modern oceanic Ca budget in a range of studies (e.g., De La Rocha and DePaolo, 2000; Fantle and Tipper, 2014; Heuser et al., 2005; Schmitt et al., 2003; Tipper et al., 2010; Zhu and Macdougall, 1998). A box model simulates the response of the system to changes in the fluxes and isotopic composition of inputs and outputs. The basic principle for a simple box model is that the total quantity of the element of interest (in this case Ca: M_{Ca}) within the box is dependent on the sum of the fluxes of the inputs (F_{inputs}) and outputs ($F_{outputs}$).

$$\frac{dM_{Ca}}{dt} = \sum_{inputs} F_{inputs} - \sum_{outputs} F_{outputs}$$
(1)

Modern seawater Ca concentrations are ~10 mmol/kg and indicate an estimated residence time of 1.1×10^6 years (Broecker and Peng, 1982), significantly longer than the ocean mixing time. Consequently, Ca concentrations and isotope ratios are likely homogenous throughout the ocean (Zhu and Macdougall, 1998), because carbonate precipitation in shallow water and dissolution at depth is not sufficiently large compared to the Ca inventory to create a significant isotope gradient in the ocean.

The addition of isotopes to Eq. (1) adds an extra degree of complexity and allows for a tighter constraint to be placed upon the system (Eq. (2)).

$$M_{Ca}\frac{d\delta_{SW}}{dt} = \sum F_{inputs} \left(\delta_{inputs} - \delta_{SW}\right) - \sum F_{output} \Delta_{sed}$$
(2)

where δ_{SW} refers to the isotopic composition of seawater, δ_{inputs} is the isotopic compositions of the input flux from the ocean, and the isotopic offset for the outputs is expressed as Δ_{sed} , representing the average fractionation factor on the formation of the calcium-bearing minerals relative to seawater (see Section 2.2).

Box modelling is addressed in greater detail by Druhan et al. (2020) in this issue. Using Eqs. (1) and (2), it is possible to evaluate the impact on the Ca seawater reservoir and isotope composition from changes in the F_{inputs} through the weathering of carbonate and silicate rocks, and from changes in the $F_{outputs}$ through transient perturbations to carbonate burial. Similarly, changes in both δ_{inputs} and in the fractionation factor of the outputs (Δ_{sed}) can also be evaluated. However, to accurately predict the response to seawater $\delta^{44/40}$ Ca values from transient perturbations in the oceanic Ca budget, carbonate precipitation (and hence, $F_{outputs}$) must be tied to alkalinity and coupled to similar mass balance equations for the oceanic C budget through carbonate weathering and the precipitation of carbonate from the ocean (e.g., Komar

and Zeebe, 2016).

To investigate changes in the ratio of the input to output flux as a function of Ca isotope seawater changes, Eq. (2) can be rearranged to Eq. (3) (Fantle and DePaolo, 2005).

$$\frac{\sum F_{outputs}}{\sum F_{inputs}} = \frac{1}{\Delta_{sed}} \left[\frac{M_{Ca}}{\sum F_{inputs}} \frac{d\delta_{SW}}{dt} + (\delta_{inputs} - \delta_{SW}) \right]$$
(3)

Eq. (3) highlights the factors that impact the calculated ratio of the inputs and outputs ($\Sigma F_{outputs}/\Sigma F_{inputs}$) of the system. These include, besides the isotopic characterizations of Ca sources and sinks and the quality of δ_{SW} reconstructions, the assumed residence time ($M_{Ca}/\Sigma F_{inputs}$) (e.g., Gussone and Friedrich, 2018).

The following sections discuss the sources and sinks for the oceanic Ca budget and the uncertainties associated with them. Due to the variety of necessary assumptions, it is the uncertainties related to the fluxes and isotopic compositions that limits the potential of simple box model approaches.

2.2. Signatures and fluxes of oceanic Ca sources

The reconstruction of the oceanic Ca budget through time is challenging and requires a detailed understanding of the Ca content and isotope composition of the involved sources and isotope fractionation mechanisms occurring along all major Ca transport paths. While our understanding of the modern Ca cycle has improved, there are still uncertainties and unknowns in both sources and sinks (Griffith et al., this issue) and it is uncertain to what degree modern estimates can be extended into the past. Below, we highlight the most important sources and discuss how their uncertainties can affect Ca budget reconstruction.

At present, the Ca input to the ocean is dominated by three main Ca sources - hydrothermal input, riverine runoff, and submarine ground-water discharge (Fig. 1). While the relative size of these Ca input fluxes is still uncertain, the isotopic composition of riverine and hydrothermal inputs is relatively well defined with average $\delta^{44/40}$ Ca values of ~0.9‰ and 0.64‰, respectively (e.g., Amini et al., 2008; Fantle and Tipper, 2014; Tipper et al., 2016). Rivers that drain basaltic terrains can be dominated by weathering of hydrothermal calcite with higher $\delta^{44/40}$ Ca values (e.g., Jacobson et al., 2015; Moore et al., 2013). The isotopic composition of groundwater and spring waters is less constrained and shows significant variability ranging from 0.2 to 2.1‰ (e.g., Holmden et al., 2012; Shao et al., 2018; Tipper et al., 2016).

Given the relatively small isotopic difference between input fluxes (e.g., 0.2–0.3‰ difference between hydrothermal and riverine Ca input), estimating the contribution of the different Ca sources during past times presents a major challenge. A strategy to approximate the ratio of continental to hydrothermal input of Ca includes the coupling of $\delta^{44/40}$ Ca records with inferred seawater 87 Sr/ 86 Sr (e.g., McArthur et al., 2001) and Mg/Ca (e.g., Hardie, 1996) ratios through time (Wang et al., 2019). However, currently there are no proxies to distinguish



Fig. 1. Simplified model for the oceanic Ca budget. Main Ca sources of the ocean are displayed on the left side, while main sinks are plotted on the right. Some reservoirs may act as sources and sinks (indicated by arrows), e.g., redissolved carbonate sediments.

between past groundwater and riverine inputs. Furthermore, the flux of Ca from groundwater discharge may have varied significantly through time and space and between glacial and interglacial periods, complicating its proper implementation into models (e.g., Milliman, 1993).

A further source of isotopically light Ca that has been proposed is a flux of diagenetic fluids into the ocean (e.g., Fantle and Higgins, 2014; Sun et al., 2016). The basic principle of the hypothesis is that during recrystallization, the primary low $\delta^{44/40}$ Ca values of the biogenic carbonates are increased leading to a mobilization of isotopically light Ca which can potentially be released into the open ocean. This flux may have been more important prior to the evolution of pelagic skeletal carbonates in the Mid-Mesozoic (Ridgwell and Zeebe, 2005), when carbonate deposition was concentrated in shallow-water platforms. The contribution of this source to the global Ca cycle is currently unconstrained.

In addition to the relative proportion of the fluxes, the $\delta^{44/40}$ Ca value of the continental input may have changed through time. For example, the isotopic composition of the exposed rocks, the evolution of crust, and the fractionation of Ca isotopes on the continents may have been temporally variable. The last mechanism is demonstrated by an offset (~0.2‰) between the average $\delta^{44/40}$ Ca of rivers and weathered rocks (e.g., Tipper et al., 2006; Fantle and Tipper, 2014; Tipper et al., 2016). The higher riverine $\delta^{44/40}$ Ca values suggests an actively growing ⁴⁰Ca-enriched continental Ca reservoir, composed of vegetation, secondary minerals and/or an exchangeable Ca pool (e.g., Tipper et al., 2016). Although little is known about the development of the continental reservoir over time and the degree of which a steady-state has been reached (if it evolved steadily, or if phases of growth were interrupted by phases of decay), it is likely that before its establishment, continental runoff was less fractionated relative to the exposed continental crust. In addition, in the early eons of Earth history, other transport processes, such as dust transport (Fantle et al., 2012; Ewing et al., 2008), may have played more important roles than today. However, a recent compilation of $\delta^{44/40}$ Ca values from Precambrian carbonate rocks indicates that the long-term Ca input flux had a nearconstant isotopic composition equal to the value of bulk silicate earth (Blättler and Higgins, 2017). While independent constraints for some parameters exist (e.g., hydrothermal input), other parameters are still poorly constrained. Strategies to cope with these complications could include the development of independent proxies to track changes in groundwater input, diagenetic flux and the evolution of the continental Ca reservoir. Before such proxies are established, Ca isotope budget reconstructions require a critical assessment of known and unknown input parameters and their associated uncertainties, leading to a full discussion of how the unconstrained parameters may affect the isotope record.

2.3. Isotope fractionation during deposition of oceanic Ca sinks and formation of archives

Marine carbonate sediments are the dominant sink for Ca from the ocean and are on average 1‰ lighter than seawater (Fig. 2). It was originally debated whether this fractionation is constant or dependent on temperature, and consequently, if variations in the $\delta^{44/40}$ Ca record reflect changes in the Ca budget or paleo-temperatures (e.g., Nägler et al., 2000; Skulan et al., 1997). Although it is now established that Ca isotopes have little sensitivity to temperature (generally < 0.03%/°C, e.g., compilation in Gussone and Heuser, 2016), this initial debate initiated a series of detailed calibration studies. These studies identified parameters that influence Ca isotope fractionation in inorganic mineral phases and in biogenic minerals, including species-specific Ca isotope fractionation (e.g., Böhm et al., 2006; Gussone et al., 2003; Lemarchand et al., 2004; Marriott et al., 2004; Sime et al., 2005). One of the major findings from these studies was that CaCO₃ mineralogy has a large effect on the fractionation (Gussone et al., 2005) and must be taken into account when interpreting and modelling Ca isotope records of the past



Fig. 2. Range of $\delta^{44/40} Ca$ values of different carbonate materials acting as main Ca sinks of the ocean.

Carbonate $\delta^{44/40}$ Ca values coloured by Sr/Ca ratios from a range of synthetic and modern to Neogene sediments (grey is used for studies where Sr concentrations are not reported). For comparison with other materials, the synthetic carbonates are reported as their fractionation re-calculated relative to modern seawater: (1) fractionation of inorganic aragonite from laboratory experiments (AlKhatib and Eisenhauer, 2017b; Dietzel et al., 2004; Gussone et al., 2003), (2) modern skeletal calcite composed of bivalves, brachiopods, and molluscs (Farkaš et al., 2007b; Immenhauser et al., 2005; Steuber and Buhl, 2006). (3) platform aragonite surface sediments (bulk and ooids, Higgins et al., 2018), (4) pelagic sediments (~0-10 Myrs) composed of foraminiferas (Fantle, 2015; Fantle and DePaolo, 2007; Gussone and Heuser, 2016; Kısakürek et al., 2011), coccoliths (Gussone et al., 2007; Langer et al., 2007), and dinoflagellates (Gussone et al., 2010), (5) modern biogenic skeletal aragonite corals (Chen et al., 2016; Gothmann et al., 2016; Inoue et al., 2015), (6) authigenic aragonite clathrites (Teichert et al., 2005), (7) high-magnesium calcite from Site 1131 (Higgins et al., 2018) and high-magnesium foraminifers (Gussone et al., 2016), (8) inorganic calcite from laboratory experiments with varying precipitation rates (AlKhatib and Eisenhauer, 2017a; Tang et al., 2008b; Lemarchand et al., 2004), (9) authigenic carbonates from the Miocene Monterey Formation and the northern South China Sea (Blättler et al., 2015; Wang et al., 2012, 2014), (10) synthetic and natural ikarite (Gussone et al., 2011), (11) platform dolomites from the Great and Little Bahamas Bank (Ahm et al., 2018; Higgins et al., 2018), (12) modern hardground carbonate cements from the Enewetak Atoll (Erhardt et al., 2020b; Erhardt et al., 2020a). For comparison, dashed vertical lines indicate the $\delta^{44/40}$ Ca values of Bulk Silicate Earth (BSE) and modern seawater (SW) and black triangles indicate the mean value and standard deviation for each group.

(Fig. 2). Moreover, these studies showed that Ca isotopes values in inorganic carbonates are especially sensitive to mineral precipitation rates with important implications for the future application of this proxy (Lemarchand et al., 2004; Nielsen et al., 2012; Tang et al., 2008b).

Due to the complications of both mineralogy and precipitation rates, the biomineral record plays a critical role in reconstructing changes in the Ca budget of the recent past and for defining Δ_{sed} (the average Ca isotope fractionation between seawater and oceanic Ca sink). Additionally, Ca isotope systematics of inorganic and biogenically

induced carbonate minerals are important to study ancient oceans, evaporitic basins, and diagenesis. Here, we summarize Ca isotope fractionation in both inorganic and biogenic carbonates.

2.3.1. Inorganic mineral phases

The physico-chemical processes that control Ca isotope fractionation during inorganic mineral precipitation have been investigated with precipitation experiments and modelling studies (e.g., AlKhatib and Eisenhauer, 2017a; Lemarchand et al., 2004; Tang et al., 2008b). The apparent Ca isotope fractionation between crystal and fluid ($\Delta^{44/40}$ Ca) observed in natural and synthetic mineral phases is the result of thermodynamic, kinetic and/or disequilibrium isotope effects. Equilibrium isotope fractionation is controlled by the difference in Ca bond strength between the solid phase and the fluid. In general, the heavy isotopes (i.e. ⁴⁴Ca) are enriched in the stronger bonds. This relationship is expressed by an inverse relationship between $\Delta^{44/40}$ Ca and crystal Ca coordination, where lower coordination numbers are associated with shorter and stronger bonds, and has been observed in carbonates, sulphates and phosphates (e.g., Colla et al., 2013; Griffith et al., 2008b; Gussone et al., 2005, 2011; Harouaka, 2011; Hensley, 2006) as well as silicate minerals (e.g., Ryu et al., 2011; Huang et al., 2010). However, there are deviations from this general trend that may be related to the fact that the Ca-O bond strength in the minerals depend not only on the Ca coordination but also on other factors, i.e. Ca isotope fractionation between mineral and dissolved Ca depends not only on the Ca bonds in the solid, but also on the speciation of Ca in the fluids (Colla et al., 2013; Moynier and Fujii, 2017).

The variability in $\Delta^{44/40}$ Ca of a single phase, such as the range of \sim 2‰ for calcite formed at different conditions (Fig. 2), reflects kinetic isotope fractionation during mineral precipitation (e.g., Lemarchand et al., 2004; Marriott et al., 2004; Reynard et al., 2011; Tang et al., 2012). The degree of kinetic Ca isotope fractionation is controlled by precipitation rate and indirectly related to other factors such as the temperature-dependent speciation of carbonate ions (Lemarchand et al., 2004). In general, higher calcite saturation leads to higher precipitation rates and increased isotope fractionation (Nielsen et al., 2012). However, the saturation state can alter the relationship between precipitation rate and $\Delta^{44/40}$ Ca. A positive relationship between precipitation rate and $\Delta^{44/40}$ Ca has been reported for experiments using highly supersaturated solutions that likely formed via a soluble metastable amorphous calcium carbonate (ACC) precursor phases (e.g., Lemarchand et al., 2004; Nielsen et al., 2012; Teng et al., 2017). In contrast, calcite formed at lower saturation and presumably without an ACC precursor has an inverse relationship, showing an increase in isotope fractionation with increasing precipitation rate (AlKhatib and Eisenhauer, 2017a; Tang et al., 2008b). Furthermore, the degree of Ca isotope fractionation also seems to depend on the stoichiometry between Ca and the respective anion as the availability of Ca determines whether kinetic or thermodynamic isotope fractionation effects dominate (Harouaka et al., 2014; Nielsen et al., 2012).

Calcite synthesis experiments demonstrate a larger variability in kinetic Ca isotope fractionation than the variability found in early diagenetic cements and authigenic minerals, suggesting that conditions on the seafloor are closer to equilibrium. For example, $\delta^{44/40}$ Ca values of modern marine authigenic aragonite and ikaite are on average about ~0.7% and 1.3, respectively (Gussone et al., 2011; Teichert et al., 2005) and the $\delta^{44/40}$ Ca values of calcite cements and dolomites are close to the seawater value compared to the range of synthetic calcite (Fig. 2, Blättler et al., 2015; Higgins et al., 2018; Steuber and Buhl, 2006; Wang et al., 2012, 2014). Nevertheless, partly recrystallized sediments can span the range between biogenic carbonates and seawater/pore-fluid (e.g., Farkaš et al., 2016).

2.3.2. Ca isotope fractionation in biominerals

Biominerals are important archives recording the $\delta^{44/40}$ Ca fluctuations of seawater throughout the Phanerozoic. Understanding Ca isotope fractionation during biomineral formation is of great relevance, because biogenic carbonates are the main oceanic Ca sink from the Phanerozoic ocean and dominantly define Δ_{sed} . As Δ_{sed} is one of the key parameters for modelling the Ca budget (see Sction 2.1), an understanding of the Ca isotope fractionation of biominerals is required to calculate paleo- $\delta^{44/40}$ Ca_{seawater} values through time.

Applied Δ_{sed} for a given time interval are based on estimates for the taxon- or mineral-specific Ca isotope fractionation, weighted by their global abundance in the sediments. Potential bias by preferential preservation of certain taxa or sedimentary facies (e.g., deep sea vs. shelf) and shifts in Δ_{sed} due to large changes in environmental conditions may need to be implemented into Ca budget modelling, but have so far not been equally considered.

Calcium isotope fractionation as a function of different environmental parameters have been studied for various taxa. This includes the main contributors to the present-day global Ca export production, namely corals (e.g., Böhm et al., 2006; Chen et al., 2016; Gothmann et al., 2016; Inoue et al., 2015), coccolithophores (e.g., Langer et al., 2007; Mejia et al., 2018) and foraminifers (e.g., Griffith et al., 2008b; Gussone and Heuser, 2016; Kasemann et al., 2008; Sime et al., 2005). In addition, taxa were studied that were considered as archives for the reconstruction of $\delta^{44/40} Ca_{seawater}$ records, such as brachiopods (e.g., Farkaš et al., 2007a; von Allmen et al., 2010), mussels (e.g., Hippler et al., 2013; Ullmann et al., 2013), ostracods (Gussone and Greifelt, 2019), sponges (Gussone et al., 2005) and dinoflagellates (Gussone et al., 2010). These calibrations revealed offsets in Ca isotope fractionation between taxa, demonstrating the advantages of species-specific records as a basis for oceanic Ca budget modelling (e.g., Sime et al., 2007). However, while differences in $\delta^{44/40}$ Ca between different species, or between species and cements, present a complication they also provide the chance to monitor diagenetic alteration and provide insight into biomineralisation processes (Steuber and Buhl, 2006).

Calcium isotope fractionation is also variable within a taxon and can be dependent on several parameters related to biomineralisation and environmental factors. For instance, the typical ~0.5‰ Ca isotope difference between calcite and aragonite found in inorganic CaCO₃ is not featured in all CaCO₃ biominerals (Fig. 2). It is apparent in different species of sclerosponges and coralline algae (Blättler et al., 2014; Gussone et al., 2005), but it is not observed in corals and bivalves (e.g., Inoue et al., 2018; Taubner et al., 2012; Hiebenthal, 2009). The differences in fractionation are suggested to result from different cellular Ca transport pathways (Gussone et al., 2006) and can also explain the insignificant dependency of Ca isotope fractionation on environmental factors that control the growth rate (e.g., Inoue et al., 2015; Langer et al., 2007). While Ca isotope fractionation during inorganic mineral formation is highly dependent on precipitation rates, Ca:CO₃ stoichiometry, supersaturation, carbonate chemistry and salinity, these parameters have overall a relatively small impact on Ca isotope fractionation in corals, coccolithophores and foraminifers (e.g., Inoue et al., 2015; Kısakürek et al., 2011; Langer et al., 2007; Roberts et al., 2018). However, Ca isotope fractionation appears to be substantially reduced at low calcite saturation states in cultured coccolithophores (Gussone et al., 2007; Meija et al., 2018), a process that may also be responsible for anomalously small Ca isotope fractionation found in planktic and benthic foraminifers and ostracods collected from water masses with low temperatures and calcite saturation (Gussone et al., 2009; Gussone and Filipsson, 2010; Gussone et al., 2016; Gussone and Greifelt, 2019).

The effect of temperature on Ca isotope fractionation in biominerals has been heavily debated and intensively studied, but the observed temperature dependence of $\sim 0.2\%$ /°C in a few taxa remains enigmatic, because large and small temperature sensitivities have been reported for the same taxa (e.g., Immenhauser et al., 2005; Nägler et al., 2000). Contrasting results have been reported for two planktic foraminifer species *Globigerinoides sacculifer* (culture, core top and downcore records) and *Neogloboquadrina pachyderma* (sinistral) (core top and plankton net samples) (e.g., Gussone et al., 2004, 2009; Heuser et al., 2005; Hippler et al., 2006, 2009; Nägler et al., 2000; Sime et al., 2005) and fossil rudists from the Cretaceous (Immenhauser et al., 2005; Steuber and Buhl, 2006). Nevertheless, the majority of taxa demonstrate temperature dependencies below 0.03%/°C (e.g., Gussone and Heuser, 2016 and refs therein). These are unsuited for paleothermometry but may need to be considered if large temperature fluctuations have occurred that can lead to a measurable shift in the $\delta^{44/40}$ Ca of the biomineral archive (Gussone and Friedrich, 2018). In the case of large global temperature changes, this would also lead to a shift in Δ_{sed} , which may need to be considered in the model parameters.

Despite the progress that has been made to understand and quantify the influence of different processes on Ca isotope fractionation in different minerals and taxa, there still remain open questions and observations that are discrepant with the current understanding of Ca isotope fractionation that are pending further investigation.

2.4. Mass independent isotope effects

Mass-independent isotope variability can affect the determination of stable isotope fractionation, by either contributing directly to one of the isotope masses of interest (e.g., ⁴⁰Ca), or to a commonly spiked isotope mass (e.g., ⁴⁸Ca). In contrast to solar system material that is not fully homogenized, nucleosynthetic anomalies are not a significant source of uncertainty for terrestrial Ca isotope records. However, radiogenic ⁴⁰Ca, expressed as $\varepsilon_{Ca} = (({}^{40}Ca/{}^{44}Ca)_{sample}/({}^{40}Ca/{}^{44}Ca)_{mantle} - 1) \cdot 10^4$ or $\varepsilon_{Ca} = (({}^{40}Ca/{}^{42}Ca)_{sample}/({}^{40}Ca/{}^{42}Ca)_{mantle} - 1) \cdot 10^4$ can build up in old terrestrial K-rich reservoirs. Radiogenic ${}^{40}Ca$ is produced from the branched decay of 40 K to 40 Ar (electron capture) and 40 Ca (β - decay), a branching ratio of 10.67% ⁴⁰Ar and 89.33% ⁴⁰Ca (Nägler and Villa, 2000) and a decay constant $\lambda(^{40}K_{\beta}) = 4.962 \cdot 10^{-10} \cdot \text{yr}^{-1}$ (Steiger and Jäger, 1977). When old continental crust is weathered, it may provide a local radiogenically-enriched 40 Ca source. For instance, reported ε_{Ca} values reach 19,000 for muscovite mineral separates, 40 for bulk rocks in Archean pegmatites from the Jack Hills region (Fletcher et al., 1997), and 3 to 26 in Archean pelagic sediments (Nelson and McCulloch, 1989).

In contrast to old terrestrial reservoirs, in situ ⁴⁰Ca production in young K-poor and Ca-rich reservoirs, such as carbonate sediments, is not sufficient to alter the mass-dependent $\delta^{44/40}$ Ca signal. However, the contribution of radiogenic ⁴⁰Ca-enriched Ca sources may need to be considered in special catchment areas. For example, Archean to Early Proterozoic gypsum deposits in Australia have an inherited ε_{Ca} signature between 2 and 6 that would translate into a bias of up to -0.6%in δ -notation (Nelson and McCulloch, 1989). Radiogenic ⁴⁰Ca enrichment was also reported for natural soils, granitic rocks and weathering experiments (e.g., Ryu et al., 2011; Farkaš et al., 2011; Marshall and DePaolo, 1989). K-rich evaporite minerals are another potential source of radiogenic ⁴⁰Ca and contain ε_{Ca} values between 3 and 700 (e.g., Baadsgaard, 1987; Heumann et al., 1979).

Most studied evaporites show a lower degree of ⁴⁰Ca ingrowth than predicted from the K/Ca ratio and age, implying that dating of evaporites using K–Ca system is hindered by Ca release during salt metamorphism and recrystallization events (e.g., Baadsgaard, 1987). Similarly, K-rich authigenic minerals such as sanidine and glauconite are generally less enriched in ⁴⁰Ca than predicted due to Ca mobilization during diagenetic reactions (Cecil and Ducea, 2011; DePaolo et al., 1983; Gopalan and Kumar, 2008). Although ε_{Ca} does not provide a stratigraphic age, it can reveal information about the hydrologic and thermal history of a sedimentary basin. More importantly, the observed deficits in ⁴⁰Ca of the K-rich minerals demonstrate that ⁴⁰Ca-enriched fluids are indeed mobilized during such diagenetic events and can locally act as radiogenically-enriched Ca sources.

Potential variations in ε_{Ca} can complicate inferences of mass-dependent Ca isotope variability, but can also provide additional information that can be used to better constrain the global Ca cycle. For instance, it was suggested, based on the absence of significant ε_{Ca}

variations in marine carbonates throughout the Phanerozoic that the oceanic Ca cycling is not dominated by the weathering of K-enriched continental crust and that mantle derived crustal rocks play a more important role than previously thought (Caro et al., 2010). In contrast, an offset of 1.3 ε units between mantle and seawater was recently described, suggesting a contribution of 10–20% hydrothermal Ca and 80–90% from the upper crust (Antonelli et al., 2018).

2.5. Behaviour of Ca isotopes during carbonate diagenesis

One of the main limitations in using marine carbonate sediments to reconstruct the evolution of global biogeochemical cycles through time is the susceptibility of carbonate minerals to diagenesis. After deposition, primary carbonate sediments recrystallize and react with porefluids which may alter their primary chemical composition. For example, carbonate platform sediments are susceptible to early diagenesis due to significant fluid flow from the advection (fluid-buffered) of both seawater and freshwater below the seafloor. The effects of early diagenesis in platform sediments are therefore important to constrain, in particular, prior to the evolution of pelagic skeletal carbonates in the Mid-Mesozoic where the nature and location of carbonate deposition was concentrated in shallow water platforms (Ridgwell and Zeebe, 2005). Early seafloor diagenesis also takes place in pelagic sediments deposited in deeper environments, but fluid-flow rates in these settings tend to be low (diffusion dominated) and diagenesis is therefore often sediment-buffered.

To avoid complications caused by carbonate diagenesis, different techniques exist to test for mineralogical and diagenetic changes in carbonate sediments (e.g., petrographic microscopy, scanning electron microscopy, cathodoluminescence, X-ray diffraction and micro Raman spectroscopy) and geochemical techniques such as O or Sr isotopes (e.g., Veizer et al., 1999). While these techniques are particularly useful for assessing alteration of fossil specimens (e.g., Gothmann et al., 2015), it is more challenging to evaluate diagenetic alteration of bulk carbonate sediments. Furthermore, these techniques do not provide a quantitative evaluation of the extend of alteration (fluid- vs. sedimentbuffered) of individual proxies. Calcium isotope measurements in bulk carbonate sediments provide a powerful tool to evaluate the degree of diagenetic alteration in both shallow carbonate platforms (Fantle and Higgins, 2014; Higgins et al., 2018; Ahm et al., 2018) and deeper environments (Fantle and DePaolo, 2006; Fantle and DePaolo, 2007; Fantle et al., 2010) by quantifying the degree of diagenetic alteration and the amount of fluid that has interacted with the sediment during recrystallization.

As a major constituent in carbonate sediments, Ca isotopes are inherently resistant to diagenetic alteration. Carbonate $\delta^{44/40}$ Ca values are only reset under conditions where there is sufficient supply of Ca²⁺ by fluid advection or diffusion to overwhelm Ca in the sediment (Higgins et al., 2018; Fantle and Higgins, 2014; Fantle and DePaolo, 2007; Fantle et al., 2010). Due to the relatively high concentrations of Ca in natural fluids (modern seawater = 10.28 mmol/kg), sedimentary $\delta^{44/40}$ Ca values are sensitive to alteration during early marine diagenesis (Higgins et al., 2018). For example, alteration of $\delta^{44/40}$ Ca values has been observed in carbonate sediments in the Great Bahama Bank where subsurface circulation of seawater contributes to early neomorphism of primary aragonite to more stable carbonate phases such as low magnesium calcite or dolomite (Higgins et al., 2018). In these settings, primary aragonite with low $\delta^{44/40}$ Ca values is deposited on the bank top and is subsequently recrystallized into secondary calcite and dolomite with higher $\delta^{44/40}$ Ca values (Ahm et al., 2018; Higgins et al., 2018).

The behaviour of $\delta^{44/40}$ Ca values during early marine diagenesis is a consequence of the rate dependence of Ca isotope fractionation in carbonate minerals (Blättler et al., 2015; Fantle and Higgins, 2014; Higgins et al., 2018; Tang et al., 2008b). The slow precipitation rates associated with recrystallization within the sediment pore-space do not

appreciably fractionate Ca isotopes ($\alpha \sim 1-0.9995$, Bradbury and Turchyn, 2018; DePaolo, 2011; Fantle and DePaolo, 2007; Jacobson and Holmden, 2008; Fantle, 2015). In contrast, the rates associated with precipitation of primary carbonate minerals in the surface ocean are orders of magnitude higher and can lead to significant Ca isotope fractionation (on average about -1.5% for aragonite and -0.9% for calcite, Gussone et al., 2005; Marriott et al., 2004, see Fig. 2 for the observed range of $\delta^{44/40}$ Ca values). As a result, primary carbonate minerals, or carbonate sediments that have preserved their geochemical fingerprint during sediment-buffered diagenesis, tend to record generally lower $\delta^{44/40}$ Ca values than carbonate sediments that have been significantly altered during early marine fluid-buffered diagenesis (e.g., dolomites and diagenetic cements, Fig. 2). Although early marine diagenesis is ubiquitous in carbonate sediments deposited in the marine realm, other types of diagenesis may contribute to secondary alteration (e.g., late-stage diagenesis and meteoric diagenesis). The behaviour of $\delta^{44/40}$ Ca values during different types and stages of diagenesis will depend on the specific composition of the diagenetic fluid and the primary sediment.

In studies of deep sea sediments, the lack of appreciable fractionation of Ca isotopes during near-equilibrium carbonate mineral dissolution and precipitation has led to the extensive use of pore-fluid Ca isotope measurements in studies of carbonate dissolution, precipitation, and recrystallization. In sediment-buffered systems with little fluid advection or diffusion, pore-fluid Ca $^{2+}$ will approach the $\delta^{44/40}$ Ca value of the carbonate sediment over a length scale that is proportional to the recrystallization rate (e.g., Bradbury and Turchyn, 2018; DePaolo, 2011; Fantle, 2015; Fantle and DePaolo, 2007; Fantle and DePaolo, 2006; Fantle, 2010; Turchyn and DePaolo, 2011). Slow recrystallization rates are recorded by pore-fluid $\delta^{44/40}$ Ca profiles that are out of equilibrium from the surrounding carbonate sediment (e.g., Turchyn and DePaolo, 2011), whereas fast recrystallization rates are recorded by pore-fluid $\delta^{44/40}$ Ca profiles that approach the isotopic composition of the sediment over short length-scales (shallow depths, e.g., Fantle, 2015; Fantle and DePaolo, 2007). In reactive-transport models, porefluid Ca isotope measurements are often combined with measurements of Sr concentrations and Sr isotopes to quantify carbonate recrystallization rates (Richter and DePaolo, 1987, 1988). In addition, by pairing pore-fluid $\delta^{44/40}\mbox{Ca}$ value with other geochemical proxies (e.g., Sr concentrations, δD , $\delta^{18}O$, and chloride concentrations), it is possible to identify different interstitial water masses such as glacial seawater stored deep within modern carbonate platforms (Blättler et al., 2019).

Calculations of recrystallization rates based on pore-fluid $\delta^{44/40}\mbox{Ca}$ value can be complicated in systems that are affected by other processes than net carbonate recrystallization. For example, recrystallization rates may be underestimated in settings with high sedimentation rates where rapid burial effectively transports recently deposited pore-fluids to a greater relative depth in the sediment column, giving the appearance of larger length scales and slower recrystallization rates (Huber et al., 2017). The precipitation of authigenic carbonate minerals in organic-rich settings have also been observed to produce large changes in pore-fluid $\delta^{44/40}$ Ca values (Blättler et al., 2015; Bradbury and Turchyn, 2018; Teichert et al., 2005, 2009). In these settings, it has been suggested that pore-fluid $\delta^{44/40}$ Ca values should increase in response to a small Ca isotope fractionation during precipitation of authigenic carbonates (Teichert et al., 2005, 2009). In contrast, it has also been suggested that during organic matter remineralisation and NH4⁺ production, Ca is released due to ion exchange from clay minerals, contributing to a decrease in pore-fluid $\delta^{44/40}$ Ca values (Ockert et al., 2013; Teichert et al., 2009).

As Ca is a major cation in carbonate sediments, diagenetic processes that affect $\delta^{44/40}$ Ca values will also affect other carbonate bound proxies. Early marine diagenesis in carbonate platform sediments from the Bahamas has been observed to produces distinct trends between $\delta^{44/40}$ Ca values and Sr/Ca ratios in limestone and $\delta^{44/40}$ Ca and δ^{26} Mg values in dolomites (Ahm et al., 2018; Higgins et al., 2018). Sediments that have recrystallized under fluid-buffered conditions show high $\delta^{44/40}$ Ca values, low δ^{26} Mg values, and low Sr/Ca ratios (e.g., dolomites from the Bahamas, Fig. 2). In contrast, sediments that have recrystallized under sediment-buffered conditions are labelled by low $\delta^{44/40}$ Ca values, high δ^{26} Mg values and high Sr/Ca ratios (Ahm et al., 2018; Higgins et al., 2018). By quantifying these geochemical signatures using diagenetic models, $\delta^{44/40}$ Ca values can be used to identify fluid- from sediment-buffered endmembers and thereby constrain the degree of alteration of other carbonate-bound proxies (Ahm et al., 2018, 2019). In contrast to reactive-transport models that aim to constrain length-scales and recrystallization rates, the strengths of endmember models that compare multiple carbonate proxies in cross-plot space is that the degree of covariation is independent of advection and recrystallization rates (Ahm et al., 2018).

Covariation between $\delta^{44/40}$ Ca values and Sr concentrations has also been observed in fossil specimens and marine cements from Cretaceous limestone (Steuber and Buhl, 2006). The lowest $\delta^{44/40}$ Ca values and highest Sr concentrations were found in well preserved skeletal components, and the highest $\delta^{44/40}$ Ca values and the lowest Sr concentrations were found in marine cements. Similar to the samples from the Bahamas, the variation of Ca isotope values and Sr concentrations observed in the Cretaceous limestones are interpreted to reflect the replacement of biological calcite by diagenetic marine calcite that records the composition of seawater at the time of alteration (Steuber and Buhl, 2006). Combined, these results suggest that early marine cements and dolomites that contain high $\delta^{44/40}$ Ca values (fluid-buffered) may offer a yet to be explored archive of ancient seawater chemistry.

3. Variation of oceanic Ca isotopes through time

Changes in seawater Ca isotopes through time are of interest because flux imbalances may reflect changes in paleo-environmental conditions. Imbalances in the Ca isotope budget have been argued to reflect changes in the input fluxes—in other words, the weathering flux of dissolved Ca in rivers (De La Rocha and DePaolo, 2000; Fantle and DePaolo, 2005; Kasemann et al., 2005)—and relative changes in the output flux through CaCO₃ burial (see Section 2.1). Further, ocean acidification would dramatically decrease the CaCO₃ burial flux, potentially leading to the dissolution of carbonate sediments, followed by enhanced continental weathering, higher seawater alkalinity, and increased carbonate precipitation (Payne et al., 2010). Because the Ca cycle is directly linked to the C cycle through CaCO₃ precipitation and dissolution, seawater Ca isotope records have also been promoted as a proxy for constraining C cycle perturbations (Payne et al., 2010).

There are several issues that can complicate the interpretation of seawater Ca isotopes as a record of Ca cycle imbalance. In order to significantly impact the seawater Ca isotope value, the perturbation must be of sufficient magnitude and duration, given the long residence time of Ca in the modern ocean (Section 2.1). Conversely, perturbations to the Ca isotope cycle that are significantly longer than the residence time will not result in equivalently long changes in the seawater Ca isotope composition, since the output flux on long time scales must be equivalent to the input flux (Blättler and Higgins, 2017). As a result, long-term shifts in Ca isotope values likely reflect changes in the fractionation of Ca isotopes of the output fluxes, not flux imbalances. Moreover, modelling of Ca and C cycles suggests that interpreting Ca isotope excursions that result from a transient C cycle perturbation requires coupling of Ca and C through the carbonate system to correctly interpret these records (Komar and Zeebe, 2016).

Tracking the evolution of seawater isotopes through time is also complicated by the variable isotope fractionation that occurs during the precipitation of Ca-bearing minerals (summarized in Section 2.3). Challenges with variable fractionation have been identified for bulk carbonate records across the geologic record (Ahm et al., 2019; Farkaš et al., 2016; Husson et al., 2015; Jost et al., 2014; Lau et al., 2017), with aragonite vs. high-magnesium calcite, and diagenesis, identified as important drivers of the bulk carbonate Ca isotope signature (summarized in Section 2.3). Moreover, the bulk carbonate sediment fractionation can be spatially variable depending on the mixture of different carbonate components inherent to a depositional environment (e.g., Silva-Tamayo et al., 2018; Wang et al., 2019). One approach to reduce the uncertainty associated with Ca isotope fractionation into bulk CaCO₃ rocks is to utilize other archives. For example, the skeletal remains of brachiopods, belemnites, and foraminifera have a known mineralogy and are often more resistant to post-depositional alteration (Veizer et al., 1999), and have been a focus for many records (Blättler et al., 2012: Farkaš et al., 2007a, 2007b: Heuser et al., 2005: Sime et al., 2007). Other common Ca-bearing authigenic marine minerals, such as barite (Griffith et al., 2008a), evaporites (Blättler et al., 2017; Blättler et al., 2018) and phosphates (Schmitt et al., 2003; Soudry et al., 2006; Arning et al., 2009), have also been used to reconstruct seawater Ca isotope signatures.

In this section, we will describe records of Ca isotopes through Earth history and their implications for interpretations for these data. In particular, we will highlight the complexities of Ca isotope incorporation into $CaCO_3$, and the implications for interpreting Ca and other biogeochemical cycles.

3.1. Transitional periods in Earth history and short time events

3.1.1. Carbon cycle perturbations and mass extinction events

Due to the link between the Ca and C cycles, Ca isotope records have been applied to major carbon cycle perturbations with the intent to test the degree of environmental impact. Examples of past climate changes have been a particular focus because periods of transiently higher atmospheric pCO_2 are expected to exhibit a cascade of environmental effects, such as increased continental weathering, ocean acidification, and oceanic anoxia—all related to carbon cycle feedbacks that counteract changes in pCO_2 (Archer, 2005; Berner, 2004). As weathering, acidification, and anoxia would individually impact the Ca cycle through direct changes in the riverine Ca flux, ocean pH, and alkalinity, Ca isotopes are thought to be an ideal proxy for the Ca and carbonate systems Fig. 6. Many of these periods also correspond to major extinctions, with the trigger for extinctions often attributed to the environmental changes that are listed above.

For example, the end-Permian mass extinction (ca. 251.9 Ma), which saw the loss of over 80% of marine genera, is generally linked to the emplacement of the Siberian Traps, a large igneous province that released an estimated $30,000 \times 10^{15}$ g of carbon as a result of volcanic degassing and volatilization of the coal- and limestone-rich strata (reviewed in Burgess et al., 2014; Payne and Clapham, 2012). The boundary is coincident with a large negative carbon isotope excursion, likely related to the release of mantle-derived and other ¹³C-depleted sources (e.g., Cui et al., 2015). Payne et al. (2010) observed a negative $\delta^{44/40}$ Ca excursion of ~0.3‰ in shallow marine platform carbonates in the Dajiang section, south China that they attributed to a combination of increased continental weathering and acidification, which reduced the carbonate precipitation and burial flux. A Ca isotope excursion of similar magnitude, coinciding with the δ^{13} C shift, was also recorded in conodont fossils composed of apatite from Meishan in south China (Hinojosa et al., 2012), though this negative excursion is absent in Meishan carbonates (Wang et al., 2019). Furthermore, a boundary negative shift in $\delta^{44/40}$ Ca has also been observed in carbonate successions from Italy, Turkey, and Oman, although the size of the shift and the absolute values differ among sites (Silva-Tamayo et al., 2018). Different mixtures of calcite and aragonite at each site are hypothesized to cause this variability among the $\delta^{44/40}$ Ca datasets, but it was noted that the overall reproducibility of a negative $\delta^{44/40}$ Ca shift in multiple sections and replicated in conodont apatite supported a secular change in Ca cycling in these environments, rather than a diagenetic fingerprint (Silva-Tamayo et al., 2018).

Several other studies have added insight into Permian-Triassic Ca

isotope records. First, Komar and Zeebe (2016) noted that coupling Ca and C cycles in numerical box models is a critical step in elucidating the drivers of Ca cycle behaviour. Because changes in Ca cycling via weathering or CaCO3 precipitation are directly linked with ocean carbonate chemistry, the predicted negative excursion that results from acidification is relatively small (< 0.1%). These authors instead propose that volcanism, a reduction in biological carbon export, and Ca isotope fractionation that varies depending on the seawater $[CO_3^{2^-}]$ are the cause for the negative shift, although the relationship between fractionation and $[CO_3^{2^-}]$ is controversial (AlKhatib and Eisenhauer, 2017a; Lemarchand et al., 2004; Tang et al., 2008b). Alternatively, the negative $\delta^{44/40}$ Ca shift at the extinction boundary at Daijang parallels a negative shift in $\delta^{88/86}$ Sr whereas 87 Sr/ 86 Sr is invariant, leading to the hypothesis that a seawater shift in $\delta^{44/40}$ Ca was caused by a major regression that exposed and weathered shallow shelfal carbonates (Wang et al., 2019).

A study of Lower-to-Middle Triassic $\delta^{44/40}$ Ca from platform carbonates in south China and Turkey identified an additional challenge. Lau et al. (2017) observed that $\delta^{44/40}$ Ca data from the two sections do not exhibit similar absolute values nor patterns through this time interval, which is hypothesized to reflect different CaCO₃ mineralogies as well as variable recrystallization (fluid- and sediment-buffered). These syndepositional and early diagenetic changes may be accounted for via additional geochemical observations, such as comparison with Sr concentrations (Fig. 3). Inconsistent $\delta^{44/40}$ Ca datasets in multiple carbonate sections across the Middle-to-Late Permian boundary illustrate the ability of local effects, whether CaCO₃ mineralogy, diagenesis, or other conditions, to impact the geological $\delta^{44/40}$ Ca value (Jost et al., 2014).

Similar to the Permian-Triassic boundary, the interplay between changes in seawater chemistry, $CaCO_3$ mineralogy, and diagenesis was observed for the end-Triassic extinction and attributed to volcanism and carbon release associated with the Central Atlantic Magmatic Province (CAMP). A negative $\delta^{44/40}Ca$ excursion of ~0.8‰ was observed in shallow marine carbonate rocks from Italy (Jost et al., 2017). Because this excursion is much larger than can be explained only by an imbalance in the Ca and C cycles, these authors attribute this shift to a combination of acidification (seawater variability) and changes in CaCO₃ mineralogy (Ca isotope fractionation).

Transient Cenozoic volcanic events have also been investigated with $\delta^{44/40}Ca$ data. In the Cenozoic, negative $\delta^{13}C$ excursions—also known as hyperthermals-are much shorter than in preceding time intervals (on the order of 10^4 to 10^5 yr). A study of $\delta^{44/40}$ Ca values in marine barite and bulk carbonate across the Eocene-Oligocene Transition (EOT, ca. 34 Ma) observed a negative shift of \sim 0.4‰ in the bulk carbonate, but no change in the barite record (Griffith et al., 2011). This multimineral approach suggests that there was no change in seawater $\delta^{44/40}$ Ca, and that bulk carbonate $\delta^{44/40}$ Ca values are reflecting other effects from variable isotopic fractionation of different biogenic and diagenetic phases. During the most significant Cenozoic hyperthermal, the Paleocene-Eocene Thermal Maximum (PETM, ca. 55 Ma), $\delta^{44/40}$ Ca in marine barite and bulk carbonate were also compared (Griffith et al., 2015). Similar to the EOT, the carbonate and barite records do not show agreement—with the bulk carbonate $\delta^{44/40}$ Ca data interpreted as a diagenetic fingerprint of acidification within the sedimentary column. This study proposed a new mechanism-sediment dissolution-for generating apparent stratigraphic shifts in bulk carbonate $\delta^{44/40}$ Ca. As an alternative hypothesis, Fantle and Ridgwell (in press) suggested that the observed Ca isotope signal of the carbonate may result from an increase in $\delta^{44/40}$ Ca_{bulk carbonate} caused by the formation of authigenic carbonates due to a saturation overshoot in response to ocean acidification events.

Together, these studies highlight the evolution of $\delta^{44/40}Ca$ data in the Phanerozoic, and the increasing recognition that while seawater Ca was likely to have been impacted during these events, the resulting impact on $\delta^{44/40}Ca$ was small and may have been dwarfed by local

processes, including CaCO₃ mineralogy, secondary diagenesis, sediment dissolution, and a host of other factors. Some of these processes may have been global (e.g., PETM), resulting in potentially reproducible signals that are not a reflection of seawater Ca and C cycle change. Additionally, the reproducibility of the end-Permian $\delta^{44/40}$ Ca negative shift—despite some variability—suggests that there likely is a global signal that can be extracted from these types of carbon cycle perturbations. Importantly, apparent global shifts in $\delta^{44/40}$ Ca can still be attributed to local process on carbonate platforms (i.e. mineralogy and early diagenesis) that record wide-spread changes in the cycling of Ca and C in shallow restricted environments related to changes in sea level, climate and oxygenation.

3.1.2. Oceanic anoxic events

Perturbations in the global Ca-C-cycle also took place during Mesozoic Oceanic Anoxic Events (OAE). Similar to mass extinction events, Ca isotope observations differ considerably between individual OAEs. For the Pliensbachian-Toarcian transition and the Toarcian anoxic event (183 Ma), negative $\delta^{44/40}$ Ca excursions of up to 0.5‰ were reported (Brazier et al., 2015). Similarly, negative excursions were shown by Blättler et al. (2011) for Cretaceous OAE 1a (~123 Ma) and 2 (~93.9 Ma). Both studies argue that the Ca isotope fractionation remained more or less constant during the OAE and bulk carbonate $\delta^{44/40}$ Ca patterns reflect changes in seawater isotope composition caused by an increased weathering flux. In contrast, Du Vivier et al. (2015) reported positive $\delta^{44/40}$ Ca excursions for OAE2 in the order of 0.1 to 0.2‰, obtained from high-precision TIMS analyses ($2\sigma \sim 0.04$) from the Eastbourne chalk samples, which was not captured by the previously measured, less precise data based on multi-collector ICP-MS from the same section (Blättler et al., 2011). Du Vivier et al. (2015) attribute their positive Ca excursion to a decrease in the magnitude of the Ca isotope fractionation factor of the archive-material related to an increase in Ca:CO₃ ratios of seawater during transient ocean acidification.

It should be noted that other studies have suggested that a reduction in Ca isotope fractionation during phases of ocean acidification may not exclusively be related to an inferred increase in the Ca:CO₃ ratio. For instance, recently it has been suggested that the formation of authigenic carbonates can be a mechanism that significantly impact bulk sediment $\delta^{44/40}\mbox{Ca}$ (Fantle and Ridgwell, in rev./this issue). In addition, as biominerals show taxon-specific fractionation characteristics often different from non-skeletal carbonate minerals, sediment composition can also play an important role (faunal composition, ratio of skeletal to nonskeletal; Fig. 2). In particular, culture experiments show that reduced seawater Ca and CO₃²⁻ concentrations are associated with reduced Ca isotope fractionation in coccolithophores (Gussone et al., 2007; Mejia et al., 2018) and increasing Ca isotope fractionation is observed with increasing seawater Ca:CO₃ ratios in foraminifera (Roberts et al., 2018). Although culture experiments are difficult to extend to natural settings, it is likely that sediments containing different contributions of skeletal and non-skeletal carbonates may show considerably different Ca isotope profiles, highlighting the importance of detailed sample characterization.

3.1.3. Glaciations in the Neoproterozoic and Paleozoic

As there are no established biostratigraphic frameworks and a general paucity of absolute ages, in the Neoproterozoic (\sim 1000–541 Ma) carbon isotope stratigraphy has been widely used to create age models (e.g., Cox et al., 2016; Halverson et al., 2005; Hoffman and Schrag, 2002). The Neoproterozoic carbonate carbon isotope record is characterized by generally high background values of \sim 5–10‰ interrupted by dramatic negative excursions with values down to -15%. Intriguingly, most of these negative excursions have been linked to the onset and aftermath of globally expansive glaciations. However, it is still widely debated whether these enigmatic negative excursions record changes in the global carbon cycle (e.g.,

Bjerrum and Canfield, 2011; Rothman et al., 2003; Tziperman et al., 2011), synchronous shifts in local conditions on ancient platforms (Ahm et al., 2019; Swart, 2008), or diagenetic events (Derry, 2010; Knauth and Kennedy, 2009). Combining carbon isotope records with measurements of $\delta^{44/40}$ Ca values offers a tool to assess the origin of the Neoproterozoic carbon isotope excursions.

The first published Neoproterozoic Ca isotope record is from carbonate rocks bracketing the last Snowball Earth event - the Marinoan glaciation (Kasemann et al., 2005). By combining $\delta^{44/40}\text{Ca}$ values with both boron and magnesium isotope measurements from the Ombaatiie and Majeberg formations in Namibia, Kasemann et al. (2005, 2014) aimed to estimate the perturbation in atmospheric pCO_2 levels, seawater pH, and alkalinity inputs associated with a Snowball Earth event. In the glacial aftermath, these authors found a ~0.7‰ negative Ca isotope excursion in the basal Ediacaran cap carbonates. A large postglacial negative Ca isotope excursion has also been found in Brazil, NW Canada, and China (Silva-Tamayo et al., 2010; Sawaki et al., 2014). Similarly, the observations from the younger Marinoan glaciation have been reproduced for the older Snowball Event-the Sturtian glaciation-that also show a negative Ca isotope excursion during the glacial aftermath (Silva-Tamayo et al., 2010). These pioneering studies attributed the large changes in $\delta^{44/40}$ Ca values to transient changes in the post-glacial marine Ca cycle, with the negative excursion caused by Ca²⁺ weathering inputs exceeding Ca²⁺ removal through carbonate precipitation.

New observations that document the behaviour of bulk sediment $\delta^{44/40}$ Ca values in modern carbonate platforms (Higgins et al., 2018) have inspired a reevaluation of the Neoproterozoic Ca isotope records (Ahm et al., 2019; Husson et al., 2015; Wei et al., 2019). These studies have demonstrated that the effects of carbonate mineralogy (aragonite, calcite, and dolomite) and diagenesis (fluid- and sediment-buffered) can produce large stratigraphic changes in $\delta^{44/40}$ Ca values that are independent of changes in the global Ca and carbon cycles. In addition, modelling studies have demonstrated that the combined changes in weathering rates and ocean acidification are incapable of producing change in seawater $\delta^{44/40}$ Ca values of more than ~0.3‰ (Komar and Zeebe, 2016) thereby raising questions about the original interpretations of Neoproterozoic Ca isotope variability.

More recently, new research has attributed the large changes in $\delta^{44/40}\mbox{Ca}$ values in the Neoproterozoic to changes in carbonate mineralogy and diagenesis. For example, $\delta^{44/40}$ Ca values in the post-glacial basal Ediacaran cap carbonates are spatially variable recording a range from $\sim -0.2\%$ to 1.4% both regionally and globally (Ahm et al., 2019). This geochemical variability can be explained by early diagenetic dolomitization of aragonite along a spectrum of fluid to sedimentbuffered diagenetic conditions (Ahm et al., 2019). Driven by the postglacial sea-level rise, aragonite sediments from the outer platform environments were dolomitized under fluid-buffered conditions (in reaction with seawater), whereas aragonite sediments on the inner platform were dolomitized under more sediment-buffered conditions and in reaction with glacial meltwater (Ahm et al., 2019) or mixtures between seawater and meltwater (Wei et al., 2019). Furthermore, a numerical diagenetic model combining $\delta^{44/40}\mbox{Ca}$ values, Sr/Ca ratios, magnesium isotopes and carbon isotopes (Ahm et al., 2018, 2019) showed that it is possible to extract the chemical composition of the dolomitizing fluid (glacial seawater) and the primary mineral (platform aragonite). This study highlights the application of Ca isotopes for constraining geochemical signals of a wide range of elements in ancient bulk carbonate sediments.

Glacial intervals associated with changes in both Ca and carbon isotopes have also been documented in the Early Paleozoic. In the Monitor Range in Central Nevada, the end-Ordovician Hirnantian glaciation is marked by a positive carbon isotope excursion of up to $\sim 7\%$ and a large negative Ca isotope excursion of $\sim 0.5\%$ (Holmden et al., 2012). This apparent synchronicity of Ca and C isotope excursions is inconsistent with what is known about global Ca and C geochemical cycles in the modern ocean because the residence time of Ca is longer than that of C (Holmden et al., 2012). Instead, these authors attributed the changes in Ca isotopes across the Hirnantian glaciation to restriction between ocean and epeiric seas with increased isotopically light Ca-inputs by submarine ground water discharge. Kimmig and Holmden (2017) combined $\delta^{44/40}$ Ca values with magnesium isotope measurements across the Hirnantian glacial interval and showed that the negative Ca isotope values and positive magnesium isotope values can be attributed to changes in aragonite abundance. Although the Ordovician ocean generally is classified as a 'calcite sea', the precipitation of primary aragonite in carbonate platform environments has also been identified by low $\delta^{44/40}$ Ca values and high Sr/Ca ratios in sections from Anticosti island, correlating with the Hirnantian glacio-eustatic sealevel fall (Jones et al., 2020).

Similar to the end-Ordovician interval, the Late Silurian is associated with major climatic changes, abrupt cooling, and global sea-level fluctuations. However, while Late Silurian marine carbonates record a large positive carbon isotope excursion (~8.5‰), there is no parallel excursion in Ca isotopes. Bulk sediment $\delta^{44/40}$ Ca values are constantly low (~0.3‰) during the initial stages of the carbon isotope excursion and subsequently increase to ~1‰ in parallel with the decrease in carbon isotope values back towards 0‰ (Farkaš et al., 2016). These authors identified a linear relationship between Sr concentrations and $\delta^{44/40}$ Ca values, suggesting that changes in precipitation rates, carbonate mineralogy (aragonite to calcite), and/or diagenesis (fluid- to sediment-buffered) may have been related to the changes in Ca isotope values and the decoupling from the carbon isotope record (Fig. 3).

The observed relationship between glacial intervals and stratigraphic changes in Ca isotope ratios in the Neoproterozoic and Early Paleozoic points to a link between climate, sea-level changes, carbonate mineralogy, and diagenesis. Changes in sea level associated with glaciation are capable of producing globally synchronous changes in local platform environments by increasing restriction, changing local surface water chemistry, and changing rates of subsurface fluid flow and carbonate diagenesis (e.g., size of freshwater lens, groundwater discharge, and buoyancy driven seawater recirculation). Importantly, the pre-Mesozoic bulk carbonate record is composed of sediments derived from carbonate platforms that are not always reliable archives of open ocean conditions (Higgins et al., 2018; Swart, 2008; Swart and Eberli, 2005). With these new insights, Ca isotopes provide an important tool to disentangling the local processes that operate on and within carbonate platforms from global changes in seawater chemistry.

3.2. Long-term trends

Long-term records of $\delta^{44/40}$ Ca have been generated from a number of sedimentary archives. However, a comparison between different Neogene records demonstrates large differences depending on the analysed sample material (Fig. 4). Overall, records based on bulk carbonate sediments (De La Rocha and DePaolo, 2000; Fantle and DePaolo, 2005) display larger variability compared to barite (Griffith et al., 2008a), phosphate (Schmitt et al., 2003) and foraminiferal records (Heuser et al., 2005; Sime et al., 2007). The differences among individual archives are likely a result of both diagenesis and speciesspecific variability in the Ca isotope fractionation factor (Fig. 4).

A Late Mesozoic dataset of $\delta^{44/40}$ Ca in skeletal carbonates, belemnites, and brachiopods was interpreted to reflect changes in global seawater $\delta^{44/40}$ Ca values through time driven by changing input fluxes of Ca (Farkaš et al., 2007b). Subsequently, a Phanerozoic compilation of $\delta^{44/40}$ Ca in skeletal carbonates (brachiopods, belemnites, and planktonic foraminifera) captured a positive shift of ~0.7‰ from the Ordovician until today, with short-term oscillations superimposed over this first-order trend (Farkaš et al., 2007a). However, box modelling results do not support variations in Ca mass balance to explain the overall trend. Instead, changing Ca isotope fractionation of carbonate sediments, related to oscillating calcite-aragonite seas (e.g., Stanley,



Fig. 3. Scatterplots of $\delta^{44/40}$ Ca vs. Sr/Ca ratios for published bulk carbonate records.

The Sr/Ca ratios (mmol/mol) are shown on a linear scale in the left panel and on a logarithmic scale in the right panel. Mg/Ca ratios are shown on a colour ramp, where blue symbols have higher Mg/Ca and red symbols have lower Mg/Ca. Precambrian data from Husson et al. (2015), Blättler et al. (2017), Pruss et al. (2018), Ahm et al. (2019), Wei et al. (2019). Aragonite data from Jost et al. (2014, 2017), Lau et al. (2017), Silva-Tamayo et al. (2018), Wang et al. (2019). Calcite data from Holmden (2009), Griffith et al. (2015), Farkaš et al. (2016), Kimmig and Holmden (2017), Jones et al. (2020). The terms calcite sea and aragonite sea refer to periods in Earth history in which the chemical composition of the ocean water promoted precipitation of calcite and aragonite, respectively (e.g., Hardie, 1996). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

2006), was proposed as a primary driver of seawater $\delta^{44/40}$ Ca through time. This first-order positive shift between the Early Silurian and Late Devonian/Early Carboniferous has also been confirmed by condont $\delta^{44/40}$ Ca data (Le Houedec et al., 2017). Blättler et al. (2012) also advocated the hypothesis that the mode of carbonate precipitation may be responsible for first-order trends in $\delta^{44/40}$ Ca, and further proposed that variability in seawater $\delta^{44/40}$ Ca due to CaCO₃ mineralogy became dampened after the advent of pelagic calcification in the Mesozoic. The carbonate $\delta^{44/40}$ Ca record was recently extended to the Archean (ca. 3 Ga; Blättler and Higgins, 2017), with values that predominantly record values similar to Bulk Silicate Earth (BSE). This observation reflects the fact that skeletal CaCO₃ does not appear in the fossil record until the Ediacaran, and that partitioning of $\delta^{44/40}$ Ca in pelagic vs. shallow carbonate sinks was not apparent in the Precambrian.

In addition to linking $\delta^{44/40}$ Ca values with the evolution of the carbonate system through time, there is a close relationship between $\delta^{44/40}$ Ca and seawater sulphate concentrations. In a study of Phanerozoic $\delta^{44/40}$ Ca, Farkaš et al. (2007a) proposed that short-term variability might have been related to the ratios of Ca to bicarbonate and sulphate in rivers, as well as dolomite precipitation. Subsequent studies of $\delta^{44/40}$ Ca in anhydrite (CaSO₄) and gypsum (CaSO₄·2H₂O) evaporites have been used to reconstruct the relative concentration of

Ca and sulphate in the past-a key clue into the evolution of the seawater sulphate pool, and thus, ocean oxygenation. Briefly, this method is based on the Ca isotope fractionation between evaporites and seawater, leading to Rayleigh-type distillation of the fluid under conditions where available Ca^{2+} < sulphate (Blättler and Higgins, 2014). In contrast, in conditions where sulphate limits Ca sulphate precipitation, there will be little consumption of the Ca in the fluid and $\delta^{44/40}\text{Ca}$ values will remain lower. This method has been used to identify periods of sulphate-rich and Ca-rich oceans in the Phanerozoic (Blättler and Higgins, 2014), in the Neoproterozoic (Blättler et al., 2020), and has also extended to a 2 Ga sedimentary evaporite sequence to constrain the concentration of marine sulphate to > 10 mmol/kg as additional evidence of significant ocean oxygenation following the Great Oxidation Event (Blättler et al., 2018). As another creative approach, the $\delta^{44/40}$ Ca in 2.7 to 1.9 Ga evaporitic sedimentary carbonates exhibited very limited variability (Blättler et al., 2017). Based on experimental calibrations, the Ca-to-alkalinity ratio was reconstructed with evidence for oceans that were less alkaline than previously proposed, providing new constraints on ocean pH and atmospheric pCO₂ in the Archean and Paleoproterozoic that can be used to test hypotheses for the so called Faint Young Sun Paradox.



Fig. 4. Comparison of different archives recording $\delta^{44/40}$ Ca_{seawater} variation through time.

Most records suggest an increase in $\delta^{44/40}$ Ca of seawater from the Miocene towards the Holocene. but timing and magnitude of the variation differ. Because of the larger isotope fractionation, the barite records (Griffith et al., 2008a, 2011) are related to the secondary $\delta^{44/40}$ Ca axis, which is offset by 1‰. The $\delta^{44/40}$ Ca records of carbonate material, foraminifers from Heuser et al. (2005) and Sime et al. (2007) and bulk carbonate (Griffith et al., 2011: dotted), Fantle and DePaolo (2007: long dashed, 2005: short-dashed), including the data of De La Rocha and DePaolo (2000) is related to the primary y-axis. Marine phosphorites show an offset between peloidal phosphorites (Schmitt et al., 2003) and crusts (Arning et al., 2009) of about 0.5%. Overall, records based on a limited number of taxa seem to show less scatter compared to bulk samples.

4. Outlook and future directions

As discussed in the previous sections, the application and interpretation of Ca isotopes for deep-time records have evolved tremendously since early papers were published in the late 1990s and early 2000s. Here, we summarize our view on the revised approaches for using Ca isotopes. First, we discuss the importance of identifying and selecting biogenic (skeletal) materials for reconstructing seawater $\delta^{44/40}$ Ca through time. Second, we propose that new discoveries about bulk carbonate $\delta^{44/40}$ Ca may be useful for elucidating the variability in depositional conditions, including carbonate mineralogy and fluid interactions, in particular in combination with other proxies such as Sr concentrations and Mg isotopes. As Ca isotope systematics continue to be refined, it can be anticipated that these new approaches will lead to reinterpretation and assessments of deep-time records.

4.1. Selection of archives

The selection of appropriate archives is of particular importance for the $\delta^{44/40}Ca_{sw}$ reconstruction ($\delta^{44/40}Ca_{sw}$) and a critical determination of the best available archive may be required for different time intervals and sedimentary facies. For relatively young records, suitable archives can be identified using calibrations based on culture experiments and samples collected from the environment, while for older eras, modern analogues are not readily available.

Archives selected from the depositional record need to balance the requirements in terms of fractionation characteristics, availability (abundance, continuity throughout the record and ability to isolate), and preservation. Possible limitations of archives with suitable fractionation patterns include aragonite corals, which are susceptible to recrystallisation, and coccolithophores which are difficult to isolate, because of their small size. The further back in time, the more pronounced the uncertainties become and simultaneously, the potential options for archives become more limited.

The use of biomineral archives is complicated when measuring extinct species where the fractionation characteristics need to be approximated using cross calibration with modern taxa, as no direct calibrations are possible. If the dominant fractionation mechanisms remained invariable throughout Earth history, this approach may result in consistent $\delta^{44/40}Ca_{sw}$ records. However, identifying suitable archives

becomes more complicated considering potential shifts in fractionation characteristics over very long time periods, such as due to changing ocean chemistry and/or adaptation of biomineralisation strategies. Although the effect of environmental parameters on Ca isotope fractionation can be tested in laboratory experiments, short-term experiments may not capture the effect of major environmental changes over long time periods, as taxa may behave differently, generating stress-induced proxy signals when exposed to environmental stress in lab experiments. Moreover, evolutionary adaption may occur on longer time scales during slow natural changes, leading to a different response to environmental changes. In addition, the composition of ocean water is not fully constrained in terms of temperature and chemical composition (e.g., dissolved inorganic carbon (DIC), stoichiometry (Ca:SO₄, Ca:CO₃), Ω , Ca²⁺, etc), which adds further uncertainties for estimating $\Delta_{\rm sed}$ and $\Delta^{44/40}$ Ca of the studied archives.

Depending on the availability of archives, different strategies have been applied to obtain $\delta^{44/40}$ Ca_{sw} records, such as monospecific records or records with a limited number of different related taxa. Compared to bulk carbonate $\delta^{44/40}$ Ca records, artefacts caused by faunal/floral shifts and species-specific fractionation can be largely reduced (e.g., Heuser et al., 2005; Sime et al., 2007). To further minimize potential artefacts in the paleo- $\delta^{44/40}$ Ca_{sw} record caused by differential fractionation patterns of taxa, the application of parallel records of different taxa/ archives was applied (e.g., Brazier et al., 2015; Gussone and Friedrich, 2018). Another strategy is to avoid carbonate archives and instead use passive tracers, such as barite (e.g., Griffith et al., 2008a) and phosphates (e.g., Arning et al., 2009; Schmitt et al., 2003; Soudry et al., 2006). In this context, the combination of passive and bulk tracers (Fantle, 2010; Hinojosa et al., 2012; Fantle and Tipper, 2014) is of special interest, as this approach may be used to identify changes in the fractionation factor (Δ_{sed}). As the dominant Ca sink, a potential complication of using biogenic carbonates could be if a long-term shift in $\delta^{44/40} Ca_{sw}$ was caused by a shift in $\alpha_{bio\text{-}carb}$ (the Ca isotope fractionation of biogenic carbonates relative to seawater), which would then not be apparent in the recorded carbonate $\delta^{44/40}$ Ca. In contrast, a change in the fractionation factor of a passive tracer would be reflected by a change of $\delta^{44/40}$ Ca recorded in the archive, even though $\delta^{44/40}$ Ca_{sw} may have remained more or less constant. However, not all CaCO₃-forming taxa fractionate in the same way as the bulk CaCO₃ output, due to the taxon-specific Ca isotope fractionation. Consequently, depending on



Fig. 5. Approaches using the combined $\delta^{44/40}$ Ca-Sr/Ca systematics to constrain the paleo seawater composition. A: The offset between inorganic calcite formed from modern seawater and ancient diagenetically overprinted calcite indicates the difference between modern and paleo $\delta^{44/40}$ Ca_{seawater} (Tang et al., 2008b; Farkaš et al., 2016). B: Determination of paleo $\delta^{44/40}$ Ca_{seawater} and Sr/Ca using the intercept of the fractionation arrays defined by ostracods and calcite cements or foraminifers (Gussone and Greifelt, 2009).



Fig. 6. Coupled Ca and C box model results.

The predicted range in carbonate δ^{13} C, carbonate $\delta^{44/40}$ Ca, seawater $\delta^{44/40}$ Ca (grey lines), and seawater omega (Ω) are shown for four different forcings, illustrated in the left hand panels: (A) an increase in volcanic degassing (solid line) and subsequent riverine Ca delivery (dashed line), (B) a shift to globally more prevalent aragonite precipitation, (C) increase in alkalinity, and (D) increase in the hydrothermal Ca flux. Note that the seawater and carbonate $\delta^{44/40}$ Ca covary, except if there is a global shift in CaCO₃ mineralogy (example B). Model adapted from Jost et al. (2017) and Silva-Tamayo et al. (2018).

their abundance and contribution to the CaCO₃ export production, some CaCO₃-forming taxa may act as either passive or bulk tracers. A recently suggested concept to determine past $\delta^{44/40}$ Ca_{sw} is the intercept method, which utilizes the different slopes of taxa in the $\delta^{44/40}$ Ca-Sr/Ca space. Assuming that the positions of the fractionation arrays stay constant and individuals only shift their position on the respective slopes due to environmental or physiological control, the Sr/Ca and $\delta^{44/40}$ Ca of past seawater can be estimated from the intercept of two fractionation arrays of species/cements featuring different $\delta^{44/40}$ Ca-Sr/Ca partitioning characteristics (Fig. 5B, Gussone and Greifelt, 2019). This approach is complementary to an earlier approach using the offset between diagenetically overprinted calcite and the modern fractionation array of inorganic calcite and accordingly the Yaxis intercept of the regression trough the ancient calcites (Fig. 5A, Tang et al., 2008b; Farkaš et al., 2016).

For much of Earth's history, as the availability of easily calibrated archives becomes scarcer, it becomes increasingly important to assess and discuss the limitations and uncertainties related to the use of the selected biogenic archive. Even deeper in time, prior to the advent of skeletal biomineralisation, or during major environmental perturbations that resulted in discontinuous skeletal records, using only biominerals becomes impossible or challenging. As a result, bulk carbonate sediments may become more important options.

4.2. Bulk carbonate sediments

As highlighted previously, combining bulk carbonate Ca isotopes with other carbonate-bound proxies, such as carbon isotopes, can provide unique constraints on the carbon cycle. For example, if seawater $\delta^{44/40}$ Ca can be captured from the bulk carbonate record, estimates of the CO₂ fluxes that lead to coupled perturbations to C and Ca cycles can be inferred (Fig. 6; e.g., Komar and Zeebe, 2016; Payne et al., 2010). However, it has been shown that large changes in bulk Ca isotope records cannot be driven purely by changes in global fluxes (< 0.15%) or ocean acidification (carbonate ion effect < 0.15%), which combined can only account for a maximum perturbation of ~0.2-0.3‰ (Komar and Zeebe, 2016). Calcium isotope perturbations that are larger than feasible are indicative of local depositional conditions that impacted Ca isotope fractionation or subsequent diagenetic recrystallization and resetting. These local and diagenetic effects can also modulate the associated carbon isotope records but by combining C and Ca isotope models it may be possible to isolate the potential effects of acidification and mineralogy (Fig. 6; Jost et al., 2017). Changes in local depositional conditions can possibly occur on a global scale if they are related to a widely expressed perturbation such as acidification or sea-level related diagenesis (Ahm et al., 2019; Griffith et al., 2015). Moreover, local factors can act to amplify or depress concurrent seawater $\delta^{44/40}$ Ca perturbations.

For example, the systematic covariation between $\delta^{44/40}$ Ca values and Sr/Ca ratios in diagenetic limestones, and $\delta^{44/40}$ Ca and δ^{26} Mg values in dolomites, can provide additional insights into the origins and preservation of δ^{13} C values in carbonates rocks. As the ratio of Ca and carbon are broadly similarly abundant in seawater and carbonates, their behaviour during fluid- and sediment-buffered early marine diagenesis is expected to be similar (Ahm et al., 2018). Interpreted through the lens of diagenesis, stratigraphic excursions in δ^{13} C values that correlate with stratigraphic excursions in $\delta^{44/40}$ Ca, Sr/Ca ratios, and δ^{26} Mg values may reflect temporal changes in the style of early dolomitization/diagenesis (fluid- and sediment-buffered) and not necessarily changes in the δ^{13} C values of dissolved inorganic carbon (DIC) in global seawater.

The Sr/Ca ratio in carbonates has long been a subject of investigation because its partitioning into carbonate minerals is controlled by several factors, such as carbonate mineralogy (e.g., Kinsman, 1969), paleo sea surface temperatures (e.g., Smith et al., 1979; Rosenthal et al., 1997; Gagan et al., 1998), productivity (e.g., Weinbauer and Velimirov, 1995; Stoll and Schrag, 2000; Stoll and Schrag, 2001), sea-level change (Stoll and Schrag, 1998), and fluid geochemistry (Langer et al., 2006). Here, we focus on the breadth of research that indicates that Sr/Ca variability is a signature of carbonate recrystallization (e.g., Kinsman, 1969; Brand and Veizer, 1980; Richter and Liang, 1993; Stoll and Schrag, 1998; Fantle and DePaolo, 2006; Tang et al., 2008a).

To illustrate the potential of coupling $\delta^{44/40}$ Ca with Sr/Ca, arrays of $\delta^{44/40} \text{Ca}$ values and Sr/Ca ratios in the bulk carbonate rock record (Fig. 3) are not readily explained by changes in seawater $\delta^{44/40}$ Ca and Sr/Ca values (e.g., Lau et al., 2017). Instead, the overall inverse relationship is more likely related to combined effects of fluid- and sediment-buffered diagenesis and variations in carbonate mineralogy (Husson et al., 2015; Lau et al., 2017; Ahm et al., 2018). This relationship is most evident for $\delta^{44/40}$ Ca and Sr/Ca data from the Precambrian, and for time intervals with aragonite-dominated CaCO₃ precipitation inferred by changing seawater Mg/Ca ("aragonite seas," Hardie, 1996). This trend agrees with the observation that aragonite tends to have lower $\delta^{44/40}$ Ca values and higher Sr/Ca ratios (Gussone et al., 2005; Kinsman, 1969), and that recrystallization would result in higher $\delta^{44/40}$ Ca values and lower Sr/Ca ratios with greater fluid-rock interaction. Because the aragonite-sea data are dominated by intervals with known volcanic CO₂ perturbations, it is possible that a sampling bias results in common trends. However, the similarity to the Precambrian data may indicate common drivers.

In contrast, the relationship between bulk carbonate $\delta^{44/40}$ Ca values and Sr/Ca ratios for time intervals with calcite-dominated CaCO₃ precipitation show greater variability (Fig. 3). Indeed, a significant positive correlation is observed for a carbonate-poor PETM deep-sea core (Griffith et al., 2015), which may indicate that a unique diagenetic regime characterizes predominantly siliciclastic $\delta^{44/40}$ Ca records. Data from the Late Silurian exhibit a linear inverse relationship between $\delta^{44/40}$ Ca and Sr/Ca, potentially reflecting variable precipitation rates and not diagenesis (Farkaš et al., 2016). Because aragonite precipitation is not as dominant in times of calcite seas, it is possible that other factors, besides the aragonite diagenetic pathway, are being observed. Nonetheless, the range in $\delta^{44/40}$ Ca values and Sr/Ca ratios for calcite seas is generally comparable to data for aragonite seas. Besides the overlapping $\delta^{44/40}\text{Ca}$ ranges of the different Ca-carbonates, it can also reflect the potential for aragonite to still precipitate in calcite seas, likely because individual taxa do not change the mineralogy of their biogenic carbonate production despite changing seawater Mg/Ca (Kimmig and Holmden, 2017).

The range in $\delta^{44/40}\mbox{Ca}$ values that result from variations in carbonate mineralogy and rate-dependent Ca isotope fractionation during diagenesis are significantly larger than plausible changes in seawater $\delta^{44/40}$ Ca values associated with transient perturbations to the global Ca cycle (Fig. 6; Blättler and Higgins, 2017; Husson et al., 2015; Komar and Zeebe, 2016). Therefore, large stratigraphic changes in $\delta^{44/40}$ Ca values, that covary with other carbonate bound proxies may not only reflect changes in global weathering rates or other Ca cycle imbalances (e.g., Kasemann et al., 2005, 2014; Silva-Tamayo et al., 2010). Instead, it is likely that large stratigraphic changes in $\delta^{44/40}\text{Ca}$ values are associated with changes in carbonate mineralogy and diagenesis (fluidand sediment-buffered). For example, large stratigraphic changes in both $\delta^{44/40}\mbox{Ca}$ values and Sr concentrations that are consistent with sediment-buffered preservation of former aragonite has been observed across the Permian-Triassic boundary (Lau et al., 2017), the Late Silurian (Farkaš et al., 2016), the end-Ordovician glaciation (Holmden et al., 1998; Kimmig and Holmden, 2017; Jones et al., 2020), the Ediacaran Shuram excursion (Husson et al., 2015), and the Marinoan cap carbonate sequence (Ahm et al., 2019) (see Section 3.2 for more details). Although this new application of Ca isotopes is distinct from initial interpretations, bulk carbonate $\delta^{44/40}Ca$ is emerging as a new tool for evaluating diagenetic processes and CaCO_3 mineralogy for a depositional environment—conditions that have been difficult to characterize using other methods.

5. Conclusion

Due to the increasing number of unknown variables-particularly as one investigates further back in geologic time-obtaining meaningful environmental or geological information through Earth's history presents unique challenges that can be probed using Ca isotopes. Here we have highlighted several different processes that affect the Ca isotope composition of seawater, and in addition, the $\delta^{44/40}$ Ca of different archives. For bulk sediment data, each record is unique in terms of age, coeval ocean chemistry, paleogeography, sediment composition, depositional and diagenetic history, availability of archives and preservation. Consequently, the approaches for using Ca isotopes for both skeletal and non-skeletal carbonate archives have evolved in the last several decades, as the growing body of research has led to the reevaluation of fundamental assumptions about the factors driving Ca isotope variability that guide how this proxy is applied and interpreted. Taking these aspects into consideration, Ca isotope variability has a great potential to shed new light into the evolution of the Earth system in deep time, as well as the diagenetic pathway of the carbonate rocks that record the geochemical clues that can be used to address these questions.

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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