Distinguishing the combined vegetation and soil component of $\delta^{13}C$ variation in speleothem records from subsequent degassing and prior calcite precipitation effects

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Received: 28 September 2022 – Discussion started: 10 October 2022
Revised: 9 September 2023 – Accepted: 25 September 2023 – Published: 4 December 2023

Abstract. The carbon isotopic signature inherited from soil and epikarst processes may be modified by degassing and prior calcite precipitation (PCP) before its imprint on speleothem calcite. Despite laboratory demonstration of PCP effects on carbon isotopes and increasingly sophisticated models of the governing processes, to date, there has been limited effort to deconvolve the dual PCP and soil–epikarst components in measured speleothem isotopic time series. In this contribution, we explore the feasibility, advantages, and disadvantages of using trace element ratios and $\delta^{44}Ca$ to remove the overprinting effect of PCP on measured $\delta^{13}C$ to infer the temporal variations in the initial $\delta^{13}C$ of drip water prior to degassing and PCP. In nine examined stalagmites, the most widely utilized PCP indicators Mg/Ca and $\delta^{44}Ca$ co-vary as expected. However, Sr/Ca does not show consistent relationships with $\delta^{44}Ca$ so PCP is not the dominant control on Sr/Ca. From $\delta^{44}Ca$ and Mg/Ca, our calculation of PCP as $f_{Ca}$, the fraction of initial Ca remaining in solution at the time the stalagmite layer is deposited, yields multiple viable solutions depending on the assumed $\delta^{44}Ca$ fractionation factor and inferred variation in $D_{Mg}$. Uncertainty in the effective fractionation of $\delta^{13}C$ during degassing and precipitation contributes to uncertainty in the absolute value of estimated initial $\delta^{13}C$. Nonetheless, the trends in initial $\delta^{13}C$ are less sensitive to these uncertainties. In coeval stalagmites from the same cave spanning the 94 to 82 ka interval, trends in calculated initial $\delta^{13}C$ are more similar than those in measured $\delta^{13}C$ and reveal a common positive-anomaly initial $\delta^{13}C$ during a stadial cooling event. During deglaciations, calculated initial $\delta^{13}C$ implies a trend of greater respiration rates and higher soil $CO_2$, although the higher interglacial drip water saturation favors more extensive PCP. Initial $\delta^{13}C$ can be estimated for active and fossil speleothems from a range of settings, wherever there is confidence that Mg/Ca and/or $\delta^{44}Ca$ provides a quantitative indication of past changes in PCP. Further study of Mg partitioning in speleothems will improve the robustness of Mg/Ca as a PCP proxy.

1 Introduction

Changes in vegetation productivity and soil processes significantly influence the $\delta^{13}C$ of drip water. The $\delta^{13}C$ of $CO_2$ in the soil and karst reflects the relative contributions of isotopically light respired $CO_2$ and isotopically heavier atmospheric $CO_2$. Conditions which favor higher vegetation productivity, including deeper-rooted trees (Meyer et al., 2014), and faster rates of heterotrophic and autotrophic respiration in soils will lead to higher soil $pCO_2$ and a lower $\delta^{13}C$ of $CO_2$. In contrast, in less productive and slower respiring systems, the
atmospheric CO$_2$ and its higher $\delta^{13}$C will be more significant C sources in the soil. This soil and vegetation signature imparted to the drip water is also imprinted on speleothem $\delta^{13}$C. In the temperature range characterizing the middle and high latitudes, respiration rates and vegetation density are highly sensitive to temperature, and speleothem $\delta^{13}$C has been exploited to serve as a temperature proxy in midlatitude speleothems (Genty et al., 2003, 2006). Sufficient soil moisture is also required for high soil CO$_2$ (Romero-Mujallli et al., 2019). In addition to its direct effect on the $\delta^{13}$C of soil CO$_2$, higher soil CO$_2$ concentrations also lead to more open system dissolution of karst host rock carbonate, which further contributes to lower $\delta^{13}$C of the drip water dissolved inorganic carbon (DIC).

Superimposed on soil and karst process are in-cave processes which subsequently modify the $\delta^{13}$C of DIC and thereby speleothem $\delta^{13}$C, since coupled degassing and precipitation of CaCO$_3$ progressively enriches the $\delta^{13}$C of the remaining dissolved inorganic carbon. This process has been demonstrated in lab experiments (Polag et al., 2010; Hansen et al., 2019) and in “farmed calcite” in cave environments (Mickler et al., 2004, 2006) and can be modeled as a Rayleigh process. Extensive quantitative modeling of this processes has been conducted in purpose built models (Mühlinghaus et al., 2007, 2009; Scholz et al., 2009; Sade et al., 2022) and embedded in broader models (Owen et al., 2018). Our interest here is to extend the previous analysis by relating the evolution of the $\delta^{13}$C of bicarbonate to the evolution of the Ca in solution. In this way, we can evaluate the viability of using trace element and Ca isotopic data as independent indices of the progression of coupled CO$_2$ degassing and prior calcite precipitation (PCP).

In this contribution, we explore the potential and limitations of simple, easy to use approaches to distinguish temporal variations in $\delta^{13}$C attributable to soil and epikarst vs. in-cave processes. We acknowledge that effects to recover the absolute initial $\delta^{13}$C will be challenging in most applications. Therefore we focus on the simpler goal of separating the temporal trends in PCP from those due to soil and epikarst components within a given stalagmite. We present new analyses of PCP proxies $\delta^{44}$Ca and Mg/Ca on nine stalagmites from a single cave system which experienced the same climate regime and represent a range of climate states from glacial through interglacial. Using these stalagmites as examples, we evaluate approaches for estimating the quantitative extent of PCP and degassing in fossil stalagmites in which initial drip water chemistry and partition coefficients are not independently constrained. For three time intervals including Termination II (TII), Marine Isotope Stage (MIS) 5b–c, and the last deglaciation, we present $\delta^{13}$C time series and compare trends of coeval $\delta^{13}$C records with and without deconvolution of in-cave as well as soil and epikarst processes.

2 Background on evolution of $\delta^{13}$C with progressive CO$_2$ degassing and prior calcite precipitation

The $\delta^{13}$C value of dissolved inorganic carbon in drip water, dominantly bicarbonate, will evolve with the coupled precipitation of calcite and evolution of CO$_2$ (g) from the drip water. Because the C removed by CO$_2$ degassing is isotopically much lighter than the bicarbonate pool, the $\delta^{13}$C value of the dissolved bicarbonate increases. The net total fractionation, $\varepsilon_{\text{tot}}$ is the weighted mean (by the number of C atoms) of the individual fractionations between bicarbonate and CO$_2$ (g) and between calcite and bicarbonate, respectively (Scholz et al., 2009):

$$\varepsilon_{\text{tot}} = 1/2 \frac{\varepsilon_{\text{CO}_2(g)/\text{HCO}_3^-}}{\varepsilon_{\text{CaCO}_3(s)/\text{HCO}_3^-}} = 1/2 \frac{\varepsilon_{\text{CaCO}_3(s)/\text{HCO}_3^-} + 1/2 \varepsilon_{\text{CaCO}_3(s)/\text{HCO}_3^-}}{\varepsilon_{\text{CaCO}_3(s)/\text{HCO}_3^-}}.$$  (1)

The first of these fractionations is strongly temperature-dependent. The second bicarbonate–calcite fractionation factor showed negligible temperature dependence in seeded experiments (Romanek et al., 1992) but modest temperature dependence in earlier studies and which were used in compilations (Emrich et al., 1970; Rubinson and Clayton, 1969; Mook and Rozanski, 2001). Thus, the net fractionation evolves from $-3.5\%$ at 25°C to $-5.2\%$ at 5°C using temperature-sensitive calcite fractionation factors, but from $-3.5\%$ at 25°C to $-4.6\%$ at 5°C when the calcite fractionation factor is invariant. The evolution of the $\delta^{13}$C of the bicarbonate in drip water has been modeled by a Rayleigh process, in which the dependence of the $\delta^{13}$C of bicarbonate is a function of the remaining HCO$_3^-$ in solution (Mühlinghaus et al., 2007, 2009). Newer models have added the effect of isotopic exchange between cave air and the drip water (Hansen et al., 2017). Finally, advection–diffusion–reaction models simulate the dynamic speleothem formation from a flowing thin film, including the evolution from degassing, precipitation, and exchange with atmosphere (Sade et al., 2022). This latter model confirms that the evolution of $\delta^{13}$C of speleothem calcite is very closely correlated to the extent of PCP once $> 20\%$ of initial DIC has precipitated.

Previous modeling has focused on the evolution of $\delta^{13}$C bicarbonate as a function of the remaining DIC. In contrast, here we focus on the evolution of $\delta^{13}$C bicarbonate as a function of the Ca remaining in the solution. The Ca remaining in solution allows us to estimate the extent of calcite precipitation, and from this we estimate the degree of degassing (and carbonate precipitation) the solution has undergone prior to deposition of the speleothem. We target Ca as the indicator because trace element ratios and Ca isotopic systems may preserve proxy information about the Ca remaining in solution. CaveCalc (Owen et al., 2018) simulates this process and examples are given in Fig. 1a. We therefore relate the Rayleigh-driven progressive enrichment in $\delta^{13}$C of bicarbonate with the progressive depletion of Ca in the drip water, $f_{\text{Ca}}$.
Figure 1. (a) Simulation of the equilibrium evolution of δ^{13}C in calcite with increasing PCP for three initial δ^{13}C values of DIC corresponding to different pCO2 of soil, with the soil CO2 isotopic composition following the typical Keeling line (e.g., Pataki et al., 2003). Calculations were completed in CaveCalc (Owen et al., 2018). The fraction of initial Ca remaining at the time of speleothem deposition (f_{Ca}) is the index of PCP. (b) Example simulation of the evolution of δ^{13}C in calcite from a single initial δ^{13}C of DIC following Eq. (3); an equilibrium degassing and precipitation fractionation slope A (−5‰) is contrasted with two possible kinetically enhanced fractionation slopes (−8‰ and −11‰).

is the fraction of initial Ca remaining in solution. This relationship is

\[ \delta^{13}C_{\text{init}} = \delta^{13}C_{\text{meas}} - A \cdot \ln(f_{\text{Ca}}), \]

in which \( \delta^{13}C_{\text{init}} \) is the composition for a case of negligible degassing and CaCO3 precipitation (\( f_{\text{Ca}} = 1 \)), and \( \delta^{13}C_{\text{meas}} \) corresponds to the \( \delta^{13}C \) at a given value of \( f_{\text{Ca}} \) (Table 1 provides a summary of abbreviations used in this paper). The term \( A \) describes the rate of evolution of \( \delta^{13}C \) with progressive degassing and CaCO3 precipitation. We subsequently refer to term \( A \) as the degassing slope. A range of values have been proposed for \( A \), from equilibrium precipitation and degassing in CaveCalc to greater kinetically enhanced fractionation during degassing suggested by some laboratory (Hansen et al., 2019) and field (Mickler et al., 2019) studies (Figs. 1b and S1 in the Supplement). The support for each of these values and implications are discussed at length in Sect. 5.3. To test the influence of different degassing slopes on the magnitude of the PCP effect, we complete a sensitivity analysis with a range of values for the parameter \( A \).

2.2 Indicators of PCP

2.2.1 Mg/Ca and Sr/Ca

As discussed previously (Stoll et al., 2012), the signal of PCP imprinted on a stalagmite includes calcite precipitation truly prior to landing of the water drop on the stalagmite surface, as well as the “extent of precipitation” occurring from the drop on the stalagmite surface before it is displaced from the active growth axis by flow. As the combined signal is manifest in the stalagmite, hereafter we discuss both true PCP and the extent of precipitation processes under the term “PCP”.

Mg/Ca is the most widely applied indicator of degassing and PCP. Due to the low partition coefficient of Mg in calcite (e.g., as low as 0.012; Day and Henderson, 2013), the precipitation of calcite leads to an increase in the solution Mg/Ca. The Rayleigh equation (Eq. 2) relates the fraction of initial Ca remaining in solution (\( f_{\text{Ca, MgCa}} \)) to the initial drip water Mg/Ca (e.g., prior to degassing) and measured solution Mg/Ca when the partition coefficient \( D \) is known.

\[ f_{\text{Ca, MgCa}} = \frac{\text{Mg/Ca}_{\text{speleothem}}}{\text{Mg/Ca}_{\text{initial}} \cdot D_{\text{Mg}}} \]

Figure 2a illustrates an example evolution. Mg/Ca is most robust as a quantitative indicator of PCP in a given stalagmite when the initial drip water Mg/Ca (prior to degassing) remains constant and when there is minimal variation in the partition coefficient of Mg. In cave analogue laboratory experiments, \( D_{\text{Mg}} \) increases by about 17 % with a 10 °C temperature increase (Day and Henderson, 2013). Additionally, in farmed calcite a 60 %–70 % increase in the \( D_{\text{Mg}} \) was observed as calcite Mg/Ca increases from 10 to 30 mmol mol⁻¹ (Wassenburg et al., 2020). If \( D_{\text{Mg}} \) increases with solution Mg/Ca, it would amplify the Mg/Ca due to increasing PCP. This dependence of \( D_{\text{Mg}} \) on Mg/Ca has also been shown in some non-cave analogue laboratory experiments (Alkhathib et al., 2022). For inference of PCP, the measured Mg in the stalagmite should be fully in the calcite and not in detrital minerals, a criterion which can be effectively checked using other detrital-sensitive indicators like Al/Ca.

A formulation similar to Eq. (1) can be made for other divalent cations with partition coefficients significantly less than 1, such as Sr/Ca and Ba/Ca. The coherence of changes in Mg/Ca, Sr/Ca, and to a lesser extent Ba/Ca...
Table 1. List of variables and abbreviations.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>A</td>
<td>degassing slope from Eq. (2)</td>
</tr>
<tr>
<td>$\Delta_{\text{calcite-dissolved}}^{44}\text{Ca}$</td>
<td>Ca isotope fractionation factor ($^{44}\text{Ca}/^{40}\text{Ca}$)</td>
</tr>
<tr>
<td>$\delta^{13}\text{C meas}$</td>
<td>measured stalagmite $\delta^{13}\text{C}$ for a given sample</td>
</tr>
<tr>
<td>$\delta^{13}\text{C init}$</td>
<td>$\delta^{13}\text{C}$ expected in stalagmite prior to degassing and PCP as calculated by Eq. (2)</td>
</tr>
<tr>
<td>$\delta^{13}\text{C}_{\text{DIC init}}$</td>
<td>$\delta^{13}\text{C}$ expected in drip water prior to degassing and PCP</td>
</tr>
<tr>
<td>$f_{\text{Ca}}$</td>
<td>modeled fraction of the initial Ca remaining at the time of speleothem carbonate deposition</td>
</tr>
<tr>
<td>$f_{\text{Ca},\delta\text{Ca}}$</td>
<td>proportion of Ca remaining in solution determined using calcium isotope measurements on the stalagmite</td>
</tr>
<tr>
<td>$f_{\text{Ca},\text{MgCa}}$</td>
<td>proportion of Ca remaining in solution determined using Mg/Ca measurements on the stalagmite</td>
</tr>
<tr>
<td>$f_{\text{Ca}}(\text{fit})$</td>
<td>proportion of Ca remaining in solution, calculated from Mg/Ca measurements on the stalagmite with adjustments for compatibility with $f_{\text{Ca},\delta\text{Ca}}$</td>
</tr>
<tr>
<td>$f_{\text{Ca}}^{\text{o}}$</td>
<td>$f_{\text{Ca},\text{MgCa}}$ with AF = 1 and B = 1</td>
</tr>
<tr>
<td>B</td>
<td>scaling factor to estimate bedrock Mg/Ca from minimum stalagmite measured Mg/Ca, as in Eq. (6)</td>
</tr>
<tr>
<td>AF</td>
<td>attenuation factor to transform $f_{\text{Ca},\text{MgCa}}$ into $f_{\text{Ca}}(\text{fit})$ as in Eq. (7)</td>
</tr>
<tr>
<td>Mg/Ca initial</td>
<td>inferred bedrock Mg/Ca and initial drip water Mg/Ca</td>
</tr>
<tr>
<td>Mg/Ca min</td>
<td>minimum Mg/Ca measured in a given stalagmite</td>
</tr>
<tr>
<td>Mg/Ca meas</td>
<td>measured stalagmite Mg/Ca for a given sample</td>
</tr>
<tr>
<td>scenario A1, scenario A2</td>
<td>scenario for which DCa is constant, whilst factors B and AF are adjusted to fit $f_{\text{Ca},\text{MgCa}}$ to $f_{\text{Ca},\delta\text{Ca}}$</td>
</tr>
<tr>
<td>scenario A3</td>
<td>scenario for which DCa is variable, whilst factors B and AF are adjusted to fit $f_{\text{Ca},\text{MgCa}}$ to $f_{\text{Ca},\delta\text{Ca}}$</td>
</tr>
<tr>
<td>scenario “full”</td>
<td>scenario for which AF = 1 and B = 1, not fit to $f_{\text{Ca},\delta\text{Ca}}$</td>
</tr>
<tr>
<td>$D_{\text{Mg}}$</td>
<td>partitioning coefficient of Mg in calcite</td>
</tr>
<tr>
<td>$D_{\text{Mg}}^{\text{o}}$</td>
<td>partitioning coefficient of Mg in calcite implied by $f_{\text{Ca},\text{MgCa}}$ with AF = 1 and B = 1</td>
</tr>
<tr>
<td>$D_{\text{Mg}}(\text{fit})$</td>
<td>partitioning coefficient of Mg in calcite implied by $f_{\text{Ca}}(\text{fit})$ as in Eq. (8)</td>
</tr>
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</table>

is often interpreted to signify that PCP is the controlling process (Sinclair, 2011; Wassenburg et al., 2020). Deviations from expected PCP control may reflect variation in the partition coefficients of Sr (or Ba). Increased DSr with a higher growth rate or saturation state is widely seen in non-cave analogue laboratory experiments (Tang et al., 2008a; Tesoriero and Pankow, 1996; Lorens, 1981). An up to 5-fold increase in DSr is simulated for a 2 order of magnitude increase in growth rate (Nielsen et al., 2013) based on ion-by-ion growth models and non-cave analogue experiments. According to this model, the magnitude of the growth rate dependence depends on the forward and reverse reaction rates, which are sensitive to temperature and solution chemistry. Additionally, in-cave measurements suggest that the DSr increases with increasing calcite Mg/Ca ratio. A tripling of calcite Mg/Ca from 1.4 to 4.2 mmol mol$^{-1}$ leads to an 18 % increase in DSr, and a tripling of Mg/Ca from 10 to 30 mmol mol$^{-1}$ leads to an 80 % increase in DSr (Wassenburg et al., 2020). A similar increase in DSr with higher Mg/Ca, observed in laboratory growth in a seawater-like ma-
the effect of the fractionation factor on the calculated $f$. Also shown with a dashed line is the “index” approximation produced by Eq. (5).

![Figure 2](https://doi.org/10.5194/cp-19-2423-2023)

**Figure 2.** (a) Example variation in Mg/Ca with $f_{Ca}$ from Rayleigh simulations of progressive calcite precipitation at 6 °C (lowermost curve) and 14 °C (uppermost curve) using the $D_{Mg}$ of Day and Henderson (2013); an example is shown for Mg/Ca drip water 400 mmol mol$^{-1}$. Also shown with a dashed line is the “index” approximation produced by Eq. (5). (b) Example estimation of $f_{Ca}$ from $\delta^{44}Ca$, illustrating the effect of the fractionation factor on the calculated $f_{Ca}$ for two examples of constant fractionation and one case in which the fractionation factor systematically evolves from $\Delta_{calcite}$-dissolved of $-1.37$ to $-1.08$ with increasing PCP and concomitant slowing of the growth rate. The example shown is for a bedrock $\delta^{44}Ca$ of 0.58.

The use of Mg/Ca (or Sr/Ca) as a quantitative indicator of PCP also requires that the initial drip water Mg/Ca attained by karst rock dissolution prior to PCP remain constant for a given drip water site and stalagmite over the time period of interest. Yet, in karst rocks composed of heterogeneous carbonate mineralogy, there is potential for variation in initial drip water trace element to Ca ratios. In karst rocks composed of both dolomite and calcite, the slower dissolution kinetics of dolomite may lead to greater relative dolomite contribution with longer water residence times, leading to higher drip water Mg/Ca and higher drip water Mg/Sr with longer water residence times (Tremaine and Froelich, 2013; Fairchild and Treble, 2009). In young reef limestone, primary aragonite and high-Mg calcite dissolve more rapidly than diagenetic low-Mg calcite, potentially contributing to temporal evolution of initial, undegassed drip water ratios over time due to progressive meteoric diagenesis of primary phases (Sinclair et al., 2012). Where bedrock is heterogeneous, it can be useful to define the initial (undegassed, without PCP) drip water chemistry as resulting from the mixture of the multiple bedrock components, e.g., the bedrock mixing fraction (Tremaine and Froelich, 2013). In a synthesis of published drip water time series, in 30 out of 32 drip locations (from 11 different cave systems), bedrock mixing fractions remained constant to within 3 %, and the temporal variations in Mg/Ca were driven by variable extent of PCP (Tremaine and Froelich, 2013). This compilation suggests that in a large majority of investigated settings, for an individual drip, seasonal and interannual variations in water residence time do not significantly affect the initial Mg/Ca attained through karst dissolution. In the cave system from which our example stalagmite records derive, drip water monitoring similarly suggests stable bedrock mixing fractions at each drip location (Kost et al., 2023b).

In monitoring studies, a constant drip water Mg/Sr ratio over periods of strongly varying drip rate and water residence time is an indicator of constancy in the proportion of dolomite–calcite dissolution. When partition coefficients of Mg and Sr in speleothems are constant over time, constancy of Mg/Sr ratio of speleothems may be a useful indicator that PCP, rather than differential dissolution, is the main process responsible for Mg/Ca variations (Tremaine and Froelich, 2013). However, variable Mg/Sr in stalagmites may also be caused by strong variations in the partition coefficients of Sr, an interpretation supported by the dependence of stalagmite Mg/Sr and inferred DSr on growth rate in many stalagmites (Sliwinski et al., 2023). In such cases, variation in stalagmite Mg/Sr ratios does not necessarily indicate variation in the drip water Mg/Sr ratios or variation in the dolomite and calcite relative contribution. Thus, we suggest that monitoring of drip water systems may serve as the most meaningful check of the potential stability of the bedrock mixing fraction and initial drip water trace element to Ca ratios.

### 2.2.2 Calcium isotopes

$\delta^{44}Ca$ has also been employed as an indicator of PCP, since there is a fractionation of Ca isotopes during the dehydration of the ion for it to be incorporated in the crystal lattice. The lighter isotope desolvates more rapidly and is thereby preferentially incorporated in the calcite lattice (Hofmann et al., 2012). Consequently, with progressive removal of Ca into
calcite, the remaining dissolved Ca evolves to an isotopically heavier composition (Owen et al., 2016) (Fig. 2b). The quantitative calculation of $f_{\text{Ca}}$ from calcium isotopes ($f_{\text{Ca}}$ vs $\delta^{44}\text{Ca}$) is most robust when the Ca isotopic composition of the host rock is known, initial $\delta^{44}\text{Ca}$ from dissolution is constant (i.e., always congruent), and the fractionation factor for Ca isotopes during dehydration and calcite precipitation is constant and known. The fractionation factor may be expressed as $\alpha$ or as $\Delta_{\text{calcite-dissolved}}$.

Previous applications of $\delta^{44}\text{Ca}$ in speleothem studies have focused on the Holocene and have monitored cave drip water and farmed calcite to calculate the dissolved calcite fractionation factor in that particular cave setting and drip site (Owen et al., 2016; de Wet et al., 2021). Studies then apply this constant fractionation factor to calculate $f_{\text{Ca}}$ values in stalagmite. For fossil, non-active stalagmites, the fractionation factor cannot be measured in farmed calcite and must be assumed or based on information from farmed calcite in the same or similar cave settings. A summary of field and experimental fractionation factors is given in Table S1 in the Supplement. In one monitoring study, $\Delta_{\text{calcite-dissolved}}$ of farmed calcite varied among different drips in a modern cave (de Wet et al., 2021). Models of desolvation and attachment during crystal growth suggest that at higher net precipitation rates, the effective fractionation is greater (DePaolo, 2011). This has been observed in non-cave analogue laboratory experiments (Tang et al., 2008b) and simulated in models which suggest that a 2 order of magnitude increase in net precipitation rates leads to a decrease in $\Delta_{\text{calcite-dissolved}}$ from $-0.75\%_e$ to $-1.45\%_e$ (Mills et al., 2021). According to this model, the magnitude of the growth rate dependence – specifically the absolute range of growth rate at which $\Delta_{\text{calcite-dissolved}}$ varies strongly – depends on the forward and reverse reaction rates, which are sensitive to temperature and solution chemistry (DePaolo, 2011). For a given growth rate, lower temperatures strengthen the water solvation structure, leading to greater fractionation. A laboratory study suggests that a 10$^\circ$ cooling leads to a $\Delta_{\text{calcite-dissolved}}$ decrease of $0.17\%_e$ (Tang et al., 2008b). Yet over a seasonal 7.5$^\circ$C cave air temperature cycle in Heshang Cave, no seasonal difference in $\Delta^{44-40}\text{Ca}$ was resolved within analytical error (Owen et al., 2016).

2.2.3 Previous comparisons of PCP indicators

Both $\delta^{44}\text{Ca}$ and Mg/Ca have been simultaneously employed to study Holocene stalagmites (HS4 in Heshang Cave) (Owen et al., 2016; Li et al., 2018) and last interglacial stalagmites from northern India (Magiera et al., 2019). These studies report good qualitative agreement in estimated $f_{\text{Ca}}$, $\delta^{44}\text{Ca}$ and $f_{\text{Ca,Mg}}$. Relative to glacial–interglacial transitions, the Holocene and last interglacial experienced more stable temperatures and potentially also more stable drip water initial oversaturation, with both factors potentially contributing to constant calcite water $\Delta^{44}\text{Ca}$ fractionation factors. Heshang Cave contains dolomite host rock, and varying water residence times could potentially alter bedrock mixing fraction, altering the initial Mg/Ca of drip water prior to degassing. However, if this process were significant, it would lead to deviation in the $f_{\text{Ca}}$ calculated from Mg/Ca compared to that calculated from $\delta^{44}\text{Ca}$. The consistency of $f_{\text{Ca}}$ estimates from Mg/Ca and $\delta^{44}\text{Ca}$ suggests that variations in water–rock contact times did not result in appreciable variation in the bedrock mixing fraction and initial Mg/Ca from dissolution.

2.3 Limitations of the Hendy test for degassing correction

The sampling of calcite precipitated from the same initial drip water but with a progressive degree of CO$_2$ degassing and calcite removal has often been attempted through extraction of samples in a single growth layer at successive distances along the growth axis (Hendy, 1971). In theory, the evolution of Mg/Ca and $\delta^{13}\text{C}$ could be examined along a growth layer. However, in practice it is difficult and often impossible to follow a single growth layer laterally along the stalagmite, especially since growth layer thickness decreases with increasing distance from the main growth axis (Dorale and Liu, 2009). Furthermore, simulations suggest that for a drip interval from the 1960s, a flow distance in excess of 3 cm from the main axis is required to generate even a 1$\%_e$ enrichment of $\delta^{13}\text{C}$ through PCP and CO$_2$ degassing (Mühlinghaus et al., 2009). Finally, because the drip waters experiencing a lot of degassing are less saturated as they flow off axis than drips with little degassing, the off-axis growth may be biased towards times of low degassing and low PCP. For these reasons, it is often impractical to test relationships between $\delta^{13}\text{C}$ and Mg/Ca using the Hendy test.

3 Methods

3.1 Geochemical models of speleothems

We employ CaveCalc for simulations of the equilibrium fractionation of carbon isotopes. The CaveCalc package employs the PHREEQC geochemical model to simulate the initial dissolution of karst limestone in equilibrium with a given volume of soil gas of specified $p$CO$_2$ and $\delta^{13}\text{C}$ and the subsequent equilibration of this solution with a cave atmosphere of specified $p$CO$_2$, leading to precipitation of CaCO$_3$. Relevant for this study, the CaveCalc package calculates the initial $\delta^{13}\text{C}$ of bicarbonate and initial Ca concentration of the drip water resulting from karst dissolution and the step-wise evolution of both parameters as well as the $\delta^{13}\text{C}$ of the precipitated calcite as the solution equilibrates with the cave atmosphere. These CaveCalc simulations allow us to relate the Rayleigh-driven progressive enrichment in $\delta^{13}\text{C}$ of bicarbonate with the progressive depletion of Ca in the drip water in equilibrium conditions. CaveCalc employs the
temperature-dependent calcite–HCO$_3^-$ fractionation as calculated by Mook and Rozanski (2001).

Independent from CaveCalc, a modified version of the I-STAL model (Stoll et al., 2012) is used to simulate PCP variation resulting from changes in initial drip water saturation state and changes in drip interval. The effect of variations in initial drip water Ca, temperature, and drip interval are explored at conditions of cave pCO$_2$ fixed at ventilated, near interglacial atmospheric concentrations (300 ppm). In I-STAL calculations, the temperature sensitivity of $D_{Mg}$ follows Day and Henderson (2013).

3.2 Analysis of fossil stalagmites

We compile existing data and report new data on nine stalagmites from NW Spain with growth periods during the Holocene, during late MIS 5, and during the penultimate deglaciation (Tables S2 and S3). Reported stalagmites are from La Vallina Cave, whose setting and lithology are described in a detailed previous monitoring study (Kost et al., 2023b). The cave is hosted in Carboniferous limestones of the Barcaliente Formation. Because the section dips at 80°, different sectors of the cave sample stratigraphically different portions of the limestone, which have significant heterogeneity in Mg/Ca and some heterogeneity in Sr/Ca and other trace elements. Thus, different drip locations can feature different initial trace element ratios due to congruent dissolution of limestones of differing composition. Despite site-to-site heterogeneity, individual drips monitored over an 18-month period show very limited temporal variation in Mg/Sr ratios despite order of magnitude differences in drip rate, suggesting that in this cave, individual drips sample a relatively stable bedrock dissolution source and maintain a constant bedrock mixing fraction. For most stalagmites, age models are previously published, including those for GAE, GAL, and GLO (Stoll et al., 2013, 2015), as well as GAR and GUL (Stoll et al., 2022). For BEL, GLD, and ROW, age model constraints are given in Table S2. The slowly growing fossil stalagmites consist of dense calcite and show no evidence of columnar porous fabrics. Stalagmite calcite for GAR and GAL consist of predominantly columnar compact to columnar open fabrics (Sliwinski and Stoll, 2021). Similar fabrics are confirmed in GUL and GAE (Sliwinski et al., 2023). Samples ROW, BEL, and NYM exhibit growth rates (12–40 µm yr$^{-1}$) similar to GAR, GAL, GUL, and GAE and feature similar dense, nonporous macroscopic textures. The much more slowly growing GLO and GLD likewise feature dense, compact calcite comparable to sample GLA with a similar growth rate and are described as columnar and columnar open (Kost et al., 2023a, b).

For geochemical analyses, we sampled approximately 1 mg of powder. Trace element ratios (Mg/Ca, Sr/Ca) were determined by dissolution of powders in 2 % HNO$_3$ and analysis on an Agilent 8800 inductively coupled mass spectrometer (ICP-MS) at ETH Zurich in collision mode. On splits of the same powders, we measured $\delta^{13}$C and $\delta^{18}$O using a Thermo Fisher Scientific Gas Bench II with methods previously described and precision of 0.08 ‰ for both isotopes (Breitenbach and Bernasconi, 2011).

In eight stalagmites, we selected several intervals of contrasting Mg/Ca to additionally measure $\delta^{44}$Ca. From the same aliquot used for Mg/Ca analysis, ~500 µg of powder was dissolved in distilled 2M HNO$_3$. An automated Ca-Sr separation method (PrepFAST MC, Elemental Scientific, Omaha, NE, USA) was used to separate Ca from Sr, Mg, and other matrix elements to avoid isobaric interferences with the multi-collector inductively coupled mass spectrometer (MC-ICP-MS). SRM 915b solutions were purified in parallel with the samples to provide a combined column chemistry and analytical accuracy assessment. Ca isotope ratios were determined using a Nu Instruments MC-ICP-MS (the University of Oxford) with a desolvating nebulizer as described previously (Reynard et al., 2011). Solutions were run at 10±1 ppm concentration, and the samples were measured with standard sample bracketing. A minimum of five analyses were conducted on each aliquot. $\delta^{44}/^{40}$Ca is reported normalized to NIST SRM 915a and was calculated from measured $\delta^{44/42}$Ca as $\delta^{44/40}$Ca = $\delta^{44/42}$Ca $\cdot$ ((43.956–39.963)/(43.956–41.959)) (Hippler et al., 2003). To determine accuracy and external precision, secondary standards NIST SRM 915b and HP-Snew (in-house standard) were used. Uncertainty in Ca isotope data is quoted as the t-distribution-derived 95 % confidence interval on the mean of repeat measurements calculated using either the standard deviation or all repeat measurements on each sample.

Additionally, we incorporate published $\delta^{13}$C, $f_{Ca}\_\delta$Ca, and Mg/Ca data from Heshang stalagmite HS4 (Owen et al., 2016; Noronha et al., 2014).

3.3 Calculation of $f_{Ca}$ from $\delta^{44}$Ca, Mg/Ca, and Sr/Ca

Mg/Ca measurement is much faster than $\delta^{44}$Ca, and therefore it is common to have Mg/Ca measured for every $\delta^{13}$C$_{init}$ measured in the stalagmite. We therefore carry out examples employing Mg/Ca as the preferred measurement for deriving a continuous record of $\delta^{13}$C$_{init}$. At the same time, using $\delta^{44}$Ca to derive the PCP-corrected $\delta^{13}$C$_{init}$ may be more robust because the $\delta^{44}$Ca of the initial solution can be derived from the measured bedrock, and the isotopic composition of the bedrock is expected to be more homogeneous than Mg/Ca, leading to less uncertainty in the initial solution, and because Ca is a major element.

We calculate $f_{Ca}\_\delta$Ca from $\delta^{44}$Ca measurements using a constant bedrock $\delta^{44}$Ca equal to average bedrock (0.58 ‰) reported previously (Lechleitner et al., 2021). Use of the highest or lowest measured bedrock $\delta^{44}$Ca leads to a ±0.05 range in absolute $f_{Ca}\_\delta$Ca values. Because bedrock $\delta^{44}$Ca is not expected to change at a given location over time, we do not expect this factor to contribute to temporal variations in $\delta^{44}$Ca. The $\Delta_{calcite-dissolved}$ for these fossil stalag-
mites is not independently constrained. Therefore, we complete a sensitivity analysis using the range of \( \Delta_{\text{calcite-dissolved}} \) observed in laboratory calcite growth and ion-by-ion growth models. Table S1 lists the four values of \( \Delta_{\text{calcite-dissolved}} \) which we evaluate.

Inference of the \( f_{\text{Ca}_{-}\text{MgCa}} \) from Mg/Ca using the Rayleigh formula requires knowledge of the partition coefficient and the initial drip water Mg/Ca, which are both difficult to infer for fossil stalagmites.

Because the \( D_{\text{Mg}} \) is very low and the exponent in the Rayleigh formula (Eq. 3) is therefore very close to \(-1\), the following provides a close approximation for \( f_{\text{Ca}} \):

\[
f_{\text{Ca}_{-}\text{MgCa}} = \frac{\text{MgCa}_{\text{initial}}}{\text{MgCa}_{\text{sample}}}. \quad (4)
\]

For a \( D_{\text{Mg}} \) of 0.025, this approximation deviates from the Rayleigh equation by 0.01 at \( f_{\text{Ca}} = 0.36 \) and deviates by lesser degrees for higher \( f_{\text{Ca}_{-}\text{MgCa}} \) (e.g., dashed gray line in Fig. 2a).

Because of heterogeneity in cave bedrock, we calculate \( f_{\text{Ca}_{-}\text{MgCa}} \) assuming there may be differences in the initial drip water Mg/Ca for each stalagmite location. Time series data for each stalagmite indicate a range of Mg/Ca values. In a first approach, we assume that the minimum Mg/Ca of a stalagmite corresponds to a solution of negligible degassing and PCP, that the \( D_{\text{Mg}} \) is constant, and therefore that the numerator can be approximated by the minimum Mg/Ca for the stalagmite:

\[
f_{\text{Ca}_{-}\text{MgCa}} = \frac{\text{MgCa}_{\text{min}}}{\text{MgCa}_{\text{sample}}}. \quad (5)
\]

By default in this approximation, the maximum \( f_{\text{Ca}_{-}\text{MgCa}} \) calculated for any stalagmite is 1. Therefore, we consider the consequences for calculated \( f_{\text{Ca}_{-}\text{MgCa}} \) if the measured minimum Mg/Ca in the stalagmite corresponds to precipitation from a solution which has already experienced PCP. We assess this impact with a scaling factor \( B \) (between 0.5 and 1) to approximate the initial bedrock dissolution Mg/Ca from the minimum Mg/Ca measured in the stalagmite and apply a constant \( B \) for the stalagmite:

\[
\text{MgCa}_{\text{initial}} = \text{MgCa}_{\text{min}} \cdot B. \quad (6)
\]

\( f_{\text{Ca}_{-}\text{SrCa}} \) may also be calculated from Sr/Ca measurements in an approach analogous to Mg/Ca. Because of the higher D Sr, Eq. (5) is a less accurate approximation for Sr; e.g., for a D Sr of 0.1, there would be a 0.04 deviation of the \( f_{\text{Ca}_{-}\text{SrCa}} \) by Eq. (5) \( (f_{\text{Ca}} = 0.30) \) compared to the Rayleigh formula at \( f_{\text{Ca}_{-}\text{SrCa}} = 0.265 \).

Finally, we also evaluate the possibility of temporal variation in \( D_{\text{Mg}} \) in a given stalagmite, testing the scale of variation in \( D_{\text{Mg}} \) which would be consistent with \( f_{\text{Ca}} \) \( \delta^{13} \text{C} \) estimations from \( \delta^{44} \text{Ca} \). In this \( f_{\text{Ca}} \) (fit), the curvature or slope of the relationship in Fig. 2a is modified by an attenuation factor AF.

\[
f_{\text{Ca}}(\text{fit}) = B - \left( \frac{B - \text{MgCa}_{\text{initial}}}{\text{AF}} \right) \quad (7)
\]

The relative change in \( D_{\text{Mg}} \) needed for the fit is given as

\[
\frac{D_{\text{Mg}}(\text{fit})}{D_{\text{Mg}}(\text{fit})} = \frac{f_{\text{Ca}}(\text{fit}) \text{MgCa}_{\text{init}}}{f_{\text{Ca}}(\text{fit}) \text{MgCa}_{\text{init}}(\text{fit})} \quad (8)
\]

where \( f_{\text{Ca}}^{\text{fit}} \) indicates the \( f_{\text{Ca}_{-}\text{MgCa}} \) for the full scenario, with \( \text{AF} = 1 \) and \( B = 1 \). This AF gives a linear increase in \( D_{\text{Mg}} \) with Mg/Ca; however, other forms of dependence could also fit the data, since we have \( \delta^{44} \text{Ca} \) for only two points for most of the analyzed stalagmites.

### 3.4 Estimation of \( \delta^{13} \text{C}_{\text{init}} \) from \( \delta^{13} \text{C}_{\text{meas}} \) and estimated \( f_{\text{Ca}} \)

We use Mg/Ca to derive continuous PCP-corrected \( \delta^{13} \text{C}_{\text{init}} \), but we employ the few existing \( \delta^{44} \text{Ca} \) measurements for each stalagmite to validate and adjust the precise relationship between Mg/Ca and PCP yielding \( f_{\text{Ca}}(\text{fit}) \) in order to achieve a more robust estimate of \( \delta^{13} \text{C}_{\text{init}} \). In detail, we use the Mg/Ca to calculate \( f_{\text{Ca}} \) in four different ways: \( f_{\text{Ca}_{-}\text{MgCa}} \) from the minimum Mg/Ca and constant \( D_{\text{Mg}} \) (termed “full” with both \( B \) and \( \text{AF} = 1 \)) and additionally from three other \( f_{\text{Ca}}(\text{fit}) \) values using Eq. (7) (termed A1, A2, and A3) in which the parameters \( B \) and \( \text{AF} \) are adjusted to provide \( f_{\text{Ca}_{-}\text{MgCa}} \) compatible with \( f_{\text{Ca}_{-}\delta^{13} \text{C}} \). For each stalagmite, the values of \( B \) and \( \text{AF} \) employed for A1, A2, and A3 are given in Table 2.

For each of these four possible \( f_{\text{Ca}} \) scenarios, we additionally calculate \( \delta^{13} \text{C}_{\text{init}} \) using three possible values of the degassing slope (A) in Eq. (3): \(-5\%\), \(-8\%\), and \(-11\%\) (e.g., in Fig. 2b). This exercise illustrates the consequences of a range of possible equilibrium and disequilibrium fractionation behaviors.

### 4 Results

#### 4.1 PCP indicators in stalagmites

Within the subsamples measured for both Mg/Ca and \( \delta^{44} \text{Ca} \), the \( \delta^{44} \text{Ca} \) of measured samples ranges from \(-0.49 \) to \(+0.42 \). In all of these subsets, the Mg/Ca and \( \delta^{44} \text{Ca} \) positively correlate (Fig. 3). In all but two stalagmites, Sr/Ca and \( \delta^{44} \text{Ca} \) positively correlate. In GUL and GAL, Sr/Ca and \( \delta^{44} \text{Ca} \) negatively correlate.

In the full geochemical sampling of the nine examined speleothems from La Vallina Cave, Mg/Ca variation in a given stalagmite ranges between 1.7-fold and 3.5-fold, with the exception of GAE (12.2-fold range). Six of the nine stalagmites have Mg/Ca variations of 2-fold to 3-fold (Table S3). The 1.7- to 3.5-fold range could be fully explained
Table 2. Parameters for elaboration of $f_{Ca}$ based on $\delta^{44}Ca$ and Mg/Ca.

<table>
<thead>
<tr>
<th>Scenarios</th>
<th>Full B1</th>
<th>A1</th>
<th>A2</th>
<th>A3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$B$</td>
<td>$B$</td>
<td>$\Delta^{44}Ca$</td>
<td>$B$</td>
</tr>
<tr>
<td>GAL</td>
<td>1</td>
<td>0.75</td>
<td>-0.66</td>
<td>0.95</td>
</tr>
<tr>
<td>GUL</td>
<td>1</td>
<td>0.75</td>
<td>-1.08</td>
<td>0.95</td>
</tr>
<tr>
<td>GLO</td>
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<td>0.8</td>
<td>-0.66</td>
<td>0.75</td>
</tr>
<tr>
<td>GAR</td>
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<td>0.85</td>
<td>-1.08</td>
<td>1</td>
</tr>
<tr>
<td>ROW</td>
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<td>0.9</td>
<td>-0.66</td>
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</tr>
<tr>
<td>NYM</td>
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<td>-0.86</td>
<td>1</td>
</tr>
<tr>
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<td>-1.08</td>
<td>0.95</td>
</tr>
<tr>
<td>GAEL*</td>
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<td>0.75</td>
<td>-1.08</td>
<td>1</td>
</tr>
<tr>
<td>GLD</td>
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<td></td>
<td>0.95</td>
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</tr>
<tr>
<td>HES</td>
<td></td>
<td></td>
<td>1</td>
<td>1.2</td>
</tr>
</tbody>
</table>

* Fit with Mg/Ca min substituted by the Mg/Ca 5th percentile value of 6 mmol.

Figure 3. Mg/Ca (orange squares on left axes, in mmol mol$^{-1}$) and Sr/Ca (blue circles on right axes, mmol mol$^{-1}$) vs. $\delta^{44}Ca$ (‰) for paired samples from each stalagmite.

by PCP even with no change in $D_{Mg}$, since a 3.5-fold range requires the initial drip water Ca concentration to be greater than 3.5 times the saturation Ca concentration for the cave conditions. Changes in the Mg/Ca of the initial drip water prior to degassing from varying bedrock mixing fraction or an enhanced Mg/Ca ratio due to increased fluid inclusion density are not required to attain the range of Mg/Ca in seven of the eight of the stalagmites. With the exception of Holocene GAL, minimum Mg/Ca generally occurs during the glacial and stadials (Table S3). In GAE, the basal 0.5 cm of the stalagmite features an unusually low Mg/Ca ratio, which is 12-fold lower than the maximum ratio. If driven solely by PCP, this range would require drip water Ca in excess of 300 ppm during warm periods to drive such a large range in PCP. In this stalagmite, the Mg/Ca of this basal portion may reflect a different initial state of the fracture network and mineral surface age over time. GAE was sampled growing on a large block fallen from the ceiling, and first growth after block emplacement may reflect flow through newly opened fracture networks.

The total range in Sr/Ca across each speleothem is between 1.7-fold and 4-fold in all speleothems except GAE (6) (Table S3). The strongest positive correlation between Sr/Ca and Mg/Ca is found in GLO, GLD, and BEL. Strong negative correlation occurs in GUL, and modest negative correlation occurs in GAL and GAR.
Figure 4. (a) The degree of PCP which is possible based on the initial drip water saturation state simulated by CaveCalc for the initial Ca (ppm), soil $pCO_2$ for 150 L gas volume, and reaction temperature. The lowest $f_{Ca}$ is defined as that which would correspond to an instantaneous calcite deposition rate equivalent to 4 $\mu$m yr$^{-1}$, as simulated in I-STAL with Dreybrodt (Romanov et al., 2008) kinetics. (b) The $f_{Ca}$ simulated for a fixed drip interval of 300 s but variable initial saturation (indexed by Ca concentration). In one case shown with circles, the variable initial Ca corresponds to a progressive decline in temperature from 18 to 5 $^\circ$C, simulating soil $pCO_2$ limited by temperature. In a second case shown with green crosses, variable initial Ca corresponds to constant temperature, simulating soil $pCO_2$ limited by moisture at constant temperature. Simulations assume Dreybrodt kinetics (Romanov et al., 2008) executed in I-STAL with a PCP enhancement factor of 3. (c, d) Illustration of the nonlinearity of PCP relative to drip interval, simulated with Dreybrodt kinetics at a temperature of 12 $^\circ$C, initial Ca 90 ppm, $d = 0.01$, and a PCP enhancement factor of 3. The drip interval range of maximum PCP sensitivity will vary with modeled temperature and PCP enhancement parameters.

4.2 Calculated $f_{Ca}$ from PCP indicators

4.2.1 Expected ranges in $f_{Ca}$

Because highly oversaturated drip waters have a greater potential for PCP than minimally oversaturated drip waters, $f_{Ca}$ can vary over a wider range in settings with high oversaturation (e.g., warm climates with higher soil CO$_2$ and initial drip water Ca; Fig. 4a). In contrast, minimum degassing and high $f_{Ca}$ will be favored by a very low oversaturation state of the drip, even for slow drip rates (Fig. 4a and b). Minimal degassing is also favored by colder temperatures. In the middle and high latitudes, low oversaturation of drip waters and low PCP are more likely during cold glacial or stadial time periods when CO$_2$ in soils is depressed due to low temperatures (e.g., Fig. 4b). We consider this expected association of colder climates and higher $f_{Ca}$ in the quantitative interpretation of $f_{Ca}$ indices. Alternatively, when soil $pCO_2$ and drip water initial saturation are regulated by the moisture limitation of soils, then $f_{Ca}$ varies over a narrower range at a constant drip interval (Fig. 3b). We simulate $f_{Ca}$ as low as 0.15 in the case of an initial Ca concentration of 137 ppm.

In addition to the oversaturation state, PCP is also dependent on the degassing time. PCP can occur in air-filled voids above the cave as well as on walls and ceilings of the cave prior to the landing of drip water on the studied stalagmite. Here, we discuss the integrated PCP, regardless of where along the flow path PCP has occurred. The susceptibility of a given speleothem to PCP may depend on the geometry of the flow path. Temporal variations in PCP in a given stalagmite are expected to depend on the flow (e.g., drip rate) as well as on the oversaturation. Because the relationship between drip interval and PCP is known to be highly nonlinear (Fig. 4c and d), different coeval stalagmites often have asynchronous variations in PCP indicators or contrasting magnitudes of PCP variation.

4.2.2 $\delta^{44}Ca$

The absolute $f_{Ca,\delta Ca}$ calculated from a given $\delta^{44}Ca$ depends on the choice of the calcite dissolved fractionation factor (Table S1). Given bedrock estimated at 0.58 ‰, the measured $\delta^{44}Ca$ in some samples would imply $f_{Ca,\delta Ca}$ higher than 1 for the $\Delta^{44}Ca$ fractionation factor corresponding to
Figure 5. (a, b) $f_{Ca_{\delta}Ca}$ (crosses) as well as $f_{Ca_{Mg}Ca}$ and $f_{Ca_{fit}}$ (both with lines) vs. measured Mg/Ca. Crosses show $f_{Ca_{\delta}}$ calculated from $\delta^{44}Ca$ according to different $\Delta^{44}Ca$ fractionation factors, vertically ordered according to fractionation factor as given in the legend. Gray curves show calculation of $f_{Ca_{Mg}Ca}$ from Mg/Ca assuming Mg/Ca$_{min}$ reflects undegassed drip water (“full”, upper gray line) or that undegassed drip water is lower than the minimum Mg/Ca by a factor of 0.65 to 0.95 (“full adj.”, lower gray line; value in Table 2). Pink, blue, and green lines illustrate scenarios A1, A2, and A3, which are potential relationships between $f_{Ca_{fit}}$ vs. Mg/Ca consistent with $f_{Ca_{\delta}}$ estimates according to Eq. (7) and fit parameters in Table 2. (c, d) The variation in $D_{Mg}$ implied by scenarios in (a) and (b), calculated as in Eq. (8) and assuming constant congruent bedrock dissolution to yield a constant initial undegassed Mg/Ca ratio of drip water. A 10°C temperature increase would cause $D_{Mg}/D_{o}$ to reach 1.22 according to laboratory experiments (Day and Henderson, 2013).

the slowest laboratory growth rates ($-0.66$). For a given stalagmite, choosing a $\Delta^{44}Ca$ corresponding to slower laboratory growth rates ($-0.66$, $-0.86$) yields a wider range of calculated $f_{Ca_{\delta}Ca}$ than choosing $\Delta^{44}Ca$ corresponding to faster laboratory growth rates ($-1.08$, $-1.37$). The largest range in $f_{Ca_{\delta}Ca}$ is found in GAE, and for a given $\Delta^{44}Ca$, the lowest range in $f_{Ca_{\delta}Ca}$ values are in GAL and GLO (Fig. 5, Appendix A). Because the fractionation factor depends on growth rate and the solution characteristics, it may not be constant throughout the growth period of a stalagmite if there are significant changes in the growth conditions. For a given stalagmite, a slower growth rate during periods of low initial drip water oversaturation might be characterized by less fractionation, e.g., a $\Delta^{44}Ca$ which is closer to 0, compared to periods of stalagmite growth from solutions with greater oversaturation.

4.2.3 Mg/Ca and Sr/Ca

Using Eq. (4) and the minimum Mg/Ca for each stalagmite as an estimation of a nondegassed $f_{Ca_{Mg}Ca} = 1$, the high-Mg/Ca stalagmites BEL and ROW exhibit a range in $f_{Ca_{Mg}Ca}$ from 0.95 to 0.6 (Fig. 5; Appendix A) and GLO exhibits a similar range (Appendix A). As mentioned previously, in GAE, the total Mg/Ca range exceeds that expected from PCP if the minimum Mg/Ca in the basal growth phase is used. We complete a sensitivity analysis calculating $f_{Ca_{Mg}Ca}$ of GAE with the 5th percentile value rather than minimum, which is equivalent to the minimum Mg/Ca observed in the upper 82 cm of the stalagmite, including the period of interest presented here. GAR exhibits the highest range in $f_{Ca_{Mg}Ca}$ estimated from measured Mg/Ca. A smaller range of $f_{Ca_{Mg}Ca}$ and the lowest $f_{Ca_{Mg}Ca}$ are calculated from Eq. (6) when the $B$ factor is < 1.
4.2.4 Comparison of f<sub>Ca</sub> estimations

We explored which combinations of Δ<sup>44</sup>Ca and assumptions of B and AF lead to coherent estimations of f<sub>Ca</sub>,δCa and f<sub>Ca</sub>,MgCa of yield f<sub>Ca</sub>(fit). We find that in many stalagmites, with AF = 1 the Mg/Ca leads to a wider range in f<sub>Ca</sub>,MgCa than f<sub>Ca</sub>,δCa (e.g., Figs. 5 and S1). Covariation in D<sub>Mg</sub> with Mg/Ca is one process which could cause AF to be greater than 1.

Estimates of f<sub>Ca</sub>,δCa and f<sub>Ca</sub>,MgCa for stalagmites BEL, ROW, and NYM are consistent with no systematic variation in D<sub>Mg</sub> (i.e., an AF ≈ 1) if the calcite dissolved fractionation factor Δ<sup>44</sup>Ca varies by 0.2‰ between the sample of lowest and highest PCP (scenario A3; Fig. 5, Table 2). Alternatively, if Δ<sup>44</sup>Ca is constant (scenarios A1, A2), then f<sub>Ca</sub>,δCa and f<sub>Ca</sub>,MgCa estimates are consistent only when AF is 1.25 to 3, which implies a systematic variation in D<sub>Mg</sub> with Mg/Ca (Fig. 5). For other stalagmites (Fig. S1), if the calcite dissolved fractionation factor Δ<sup>44</sup>Ca varies by 0.2‰ between the sample of lowest and highest PCP (Fit A3), some variation in D<sub>Mg</sub> is still required (AF 1.5–2.2); for the assumption of constant Δ<sup>44</sup>Ca in each stalagmite, a larger range of AF is required (1.5 to 3, with a single higher value of 4 required for GAR). For GAE, only a Mg/Ca min of 6 mmol mol<sup>−1</sup>, which is the 5th percentile value, enabled calculation of a PCP factor consistent with Δ<sup>44</sup>Ca.

The concomitant changes in D<sub>Mg</sub> for these scenarios range from 1 (e.g., constant D<sub>Mg</sub>) to over a 2.5-fold increase in D<sub>Mg</sub> over the range of Mg/Ca in the stalagmite (Fig. 5). GAL and GAR have the most significant increases in D<sub>Mg</sub>, whereas BEL, NYM, and ROW feature the lowest. Most scenarios require an increase in D<sub>Mg</sub> larger than that expected from reasonable (5 to 10°) temperature dependence of D<sub>Mg</sub> alone; for example, according to an experimental cave analogue calculation, a 10° warming would lead to a D<sub>Mg</sub>/D<sub>Mg</sub> of 1.22 (Day and Henderson, 2013). Holocene stalagmite GAL features a large simulated range in D<sub>Mg</sub> despite only limited regional temperature change.

We acknowledge that we have limited f<sub>Ca</sub>,δCa for each stalagmite, and therefore our particular scenarios are intended to illustrate potential compatible solutions but do not cover all the possible ranges. The variation in D<sub>Mg</sub> may be exaggerated because datasets with higher numbers of paired δ<sup>44</sup>Ca and Mg/Ca such as GAL show coherency but also some scatter around this relationship, which may be interpreted as a steeper or shallower relationship in our limited dataset.

4.3 Estimation of δ<sup>13</sup>C<sub>init</sub> in measured speleothems

Following the use of scenarios A1, A2, and A3 to generate the f<sub>Ca</sub> used in Eq. (3), we generated time series records of δ<sup>13</sup>C<sub>init</sub> for eight stalagmites spanning three time periods of interest – the late glacial to the Holocene, the penultimate glacial to the interglacial transition, and a stadial event occurring between 87 and 85 ka BP. For clarity in time series figures, we illustrate the three scenarios only for the −8‰ degassing slope. For one scenario (A1) we illustrate the range in δ<sup>13</sup>C<sub>init</sub> for degassing slopes from −5‰ to −11‰. This range is variable over time in a given stalagmite because the choice of degassing slope (−5‰, −8‰, −11‰) impacts the δ<sup>13</sup>C<sub>init</sub> significantly for low f<sub>Ca</sub> and to a lesser degree for higher f<sub>Ca</sub>.

We begin by comparing the range of δ<sup>13</sup>C<sub>init</sub> calculated for growth during interglacials and present the time series trends for selected individual stalagmites.
The calculated $\delta^{13}C$ exhibits a decrease over the deglaciation. The measured $\delta^{13}C$ features a very stable average value punctuated by two positive anomalies, while Mg/Ca exhibits a decrease over the deglaciation. The $\delta^{13}C$ features a significant negative shift between 136 and 130 ka. With this significant change in trends among the early glacial and the Holocene, as the calculated $\delta^{13}C$ of respired end-members in temperate broadleaf (TB), temperate conifer (TC), and boreal (B) ecosystems (Pataki et al., 2003) illustrated as vertical ranges. For each stalagmite, the whisker plots show distribution of $\delta^{13}C$ for three $f_{Ca}$ scenarios consistent between Mg/Ca and $\delta^{44}Ca$ (A1, A2, A3; blue shading) as well as the $f_{Ca}$ derived from the full unadjusted Mg/Ca record (gray shading) for three different degassing slopes. Shown are the median as well as the upper and lower quartile, and whiskers show 99 % and 1 % for the four different $f_{Ca}$ scenarios and three different slopes of degassing fractionation. The dashed green line gives the upper limit of calcite expected to form from drip waters in the temperate conifer ecosystem; interglacial $\delta^{13}C$ is not expected to be more negative than this value.

4.3.3 Deglaciation

For the Holocene stalagmite GUL, there is a significant temporal trend of Mg/Ca increase from the Bølling–Allerød (B/A) into the Holocene (Figs. 8 and S3). Measured $\delta^{13}C$ was comparable in the B/A and the Holocene, but for all scenarios and degassing slopes, the $\delta^{13}C$ exhibits greater contrast between the early glacial and the Holocene, as the calculated $\delta^{13}C$ of the Bølling–Allerød (B/A) is more positive than that of the Holocene.

A similar steepening of the temporal trend in $\delta^{13}C$ occurs for stalagmite GLD spanning the penultimate deglaciation. GLD measured $\delta^{13}C$ features a very stable average value punctuated by two positive anomalies, while Mg/Ca exhibits a decrease over the deglaciation. The $\delta^{13}C$ features a significant negative shift between 136 and 130 ka. With this significant change in trends among $\delta^{13}C$, correlation coefficients drop well below 0.5 for two scenarios at the −8 slope of degassing and correlations approach 0 at the −11 % slope (Figs. 9 and S4).

5 Discussion

5.1 Interpretation of $\delta^{13}C_{\text{init}}$ and coherence of coeval speleothem signals and regional climate

The calculated $\delta^{13}C_{\text{init}}$ is formulated to reflect the isotopic signal of DIC prior to significant evolution via degassing and PCP. Warmer interglacial climates characterized by higher soil $pCO_{2}$ promote more negative $\delta^{13}C_{\text{DIC}}$ and increase the degree of drip water saturation, which promotes greater PCP for a given drip interval (Fig. 4). For some locations with drip intervals very sensitive to PCP, these processes may exert comparable magnitude but opposing influence on measured stalagmite $\delta^{13}C$ and lead to nearly constant measured $\delta^{13}C$ despite significant changes in temperature (Martrat et al., 2007) and vegetation (Goni et al., 2008; Tzedakis et al., 2018). We suggest that such a process may have contributed to the similar average measured $\delta^{13}C$ in GUL in late glacial and Holocene time intervals (Fig. 6) and also to the similar glacial and interglacial measured $\delta^{13}C$ in GLD (Fig. 7). Both of these stalagmites are characterized by large increases in Mg/Ca in the interglacial, indicating significant enhancements of PCP.

For stalagmite GUL growing during the last deglaciation, all of the examined calculations of $\delta^{13}C_{\text{init}}$ yield a temporal pattern more consistent with independent evidence of cooler regional temperatures during the late glacial B/A compared to the early Holocene (Fig. 7) (Martrat et al., 2007; Ausín et al., 2019; Cacho et al., 1999). The aggregate $\delta^{13}C_{\text{init}}$ of both stalagmites spanning TI (GAL and GUL) exhibits a stronger correlation with regional sea surface temperature (SST) than the measured $\delta^{13}C$ (Fig. 10).

Because the extent of degassing and PCP can be strongly conditioned by the drip rate and drip chemistry supplying an individual stalagmite, the extent and temporal variations of PCP may differ significantly among coeval stalagmites, leading to contrasting measured $\delta^{13}C$ among coeval stalag-
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Figure 7. Stadial event in the time interval 94 to 82 ka in ROW, GLO, and GAE. (a) The δ¹³Cinit for three /Ca scenarios (A1 and A2 solid lines, A3 dashed line) for each stalagmite; the line shows the result for a degassing fractionation slope of −8‰, and shading for the A3 scenario shows the range using other degassing slopes of −5‰ (more positive) and −11‰ (more negative). Diamond symbols on the red curve indicate the position of U/Th dates and the uncertainty in the age. (b) The SST record from the Iberian Margin (Martrat et al., 2007). (c) The measured δ¹³C for the three stalagmites; small symbols indicate measured samples. (d) Mg/Ca for the three stalagmites; small symbols indicate measured samples. A box-and-whisker plot is given in Fig. S2. In (a) through (c), yellow bars highlight the inferred position of the stadial cooling event in each record, given uncertainty in the age model.

Figure 8. Glacial to Holocene transition in GUL. (a) The measured δ¹³C for GUL. (b) The δ¹³Cinit for three /Ca scenarios; the line shows the result for a degassing fractionation slope of −8‰, and shading for the A3 scenario shows the range using other degassing slopes of −5‰ (more positive) and −11‰ (more negative). (c) The alkenone-based SST for the southern Iberian Margin (Cacho et al., 1999). (d) The Mg/Ca for GUL. At the base the position of U/Th age points and their uncertainty are indicated with diamond symbols. A box-and-whisker plot is given in Fig. S3. Green shading highlights the B/A period.

Stalagmite GAE features a long-term trend towards negative measured δ¹³C not observed in GLO or ROW, and GAE also features a significant decrease in Mg/Ca (Fig. 7). The calculated δ¹³Cinit of GLO does not alter the trend seen in measured δ¹³C. However, the δ¹³Cinit of ROW features the positive anomaly typical of stadial cooling. Significantly, despite the strong long-term trend in measured δ¹³C, the calculated δ¹³Cinit of GAE also resolves a positive anomaly consistent with the timing of the GS22 cooling. These more coherent temporal trends are resolved in the three records regardless of the /Ca scenario employed.

These results for multiple coeval stalagmites suggest that calculation of δ¹³Cinit has the potential to improve repro-
ducibility among coeval records and resolve signals of important regional changes in temperature and vegetation effects in the soil and epikarst environment.

5.2 Coherency of quantitative PCP estimations and impact on deconvolving PCP effects in $\delta^{13}C$

5.2.1 Quantitative PCP indicators in this sample set

The calculation of $\delta^{13}C_{\text{init}}$ relies on quantitative estimation of PCP. PCP should exert a similar effect on ratios of divalent cations with low partition coefficients like Mg/Ca and Sr/Ca. However, many published Mg/Ca and Sr/Ca records do not exhibit strong positive correlation (Sinclair, 2011). In our dataset, the greater correlation of Mg/Ca and Sr/Ca in some of the high-Mg/Ca stalagmites may reflect the proportionally greater effect of calcite Mg/Ca on the $D_{\text{Sr}}$ in higher-Mg/Ca stalagmites (e.g., Wassenburg et al., 2020). A 3-fold reduction in Ca due to PCP is attainable for reasonable degrees of soil $pCO_2$ and drip water saturation in our interglacial climates, and therefore a 3-fold range in Mg/Ca or Sr/Ca due to PCP is possible. Over a 3-fold range in Mg/Ca in low-Mg/Ca stalagmites (e.g., 2 to 6 mmol mol$^{-1}$ Mg/Ca), existing compilations suggest that the $D_{\text{Sr}}$ increases by 20%. Yet, for a 3-fold increase in Mg/Ca in the high-Mg/Ca stalagmites (e.g., 10 to 30 mmol mol$^{-1}$ Mg/Ca) the $D_{\text{Sr}}$ would increase by 80%. Because Mg/Ca and $\delta^{44}\text{Ca}$ are always positively correlated, but Sr/Ca and $\delta^{44}\text{Ca}$ are not positively correlated in all samples, we conclude that growth-rate-driven variations in $D_{\text{Sr}}$ can strongly overprint the PCP-driven trends in Sr/Ca in some stalagmites. For a given $f_{\text{Ca}}$, faster growth rates would promote a lower $\delta^{44}\text{Ca}$ and a higher $D_{\text{Sr}}$, fueling inverse correlation. Because Ca and Sr are dominantly bedrock-sourced in this setting, this reversal from the expected PCP relationship cannot be attributed to variable contribution of other element sources but must reflect the operation of processes at the solution–calcite interface. Similarly, deviation of Sr/Ca and Mg/Ca from the PCP slope, or variation in the Sr/Mg ratio, does not conclusively require variation in non-bedrock sources in either element but could reflect partitioning effects which for Sr are decoupled from PCP, as suggested by strong growth rate dependence of Sr partitioning in stalagmites from this cave (Sliwinski et al., 2023).

In our dataset, temperature variation in $D_{\text{Mg}}$ could explain some, but not all, of the apparent amplification in the Mg/Ca-based $f_{\text{Ca}}$ compared to $\delta^{44}\text{Ca}$-based $f_{\text{Ca}}$. Significant amplification is observed within the Holocene when temperature variation is limited. Additionally, in many cases
the required range in \(D_{\text{Mg}}\) is larger than the 22% increase predicted from reasonable-amplitude (e.g., < 10¹⁰) temperature increases according to laboratory calibrations (Day and Henderson, 2013). One candidate explanation for higher-magnitude changes in \(D_{\text{Mg}}\) may be the up to 2-fold increase in the \(D_{\text{Mg}}\) with increasing calcite Mg/Ca (Wassenburg et al., 2020). This effective partitioning increase may reflect true lattice partitioning and/or increased Mg in fluid inclusions. The existing field data do not precisely constrain the slope of this increase in \(D_{\text{Mg}}\) or define if the slope is indeed linear. For future absolute estimation of \(f_{\text{Ca}}\) from Mg/Ca, improved constraints on the variation of \(D_{\text{Mg}}\) are required from laboratory and field studies. In field monitoring studies, true variation in \(D_{\text{Mg}}\) can be most accurately determined when observations adequately control for both the initial solution Mg/Ca and the integrated water Mg/Ca during calcite precipitation, which may be higher than the initial Mg/Ca if a significant amount of the Ca is precipitated on the stalagmite from each solution aliquot. Future field monitoring and farmed calcite studies should also evaluate whether calcite fabrics could provide independent evidence of variation in \(D_{\text{Mg}}\), since some stalagmites show a relationship between fabrics and Sr partitioning (Frisia et al., 2022).

In addition to variation in \(D_{\text{Mg}}\), temporal changes in the Mg/Ca of undegassed drip water could also contribute to discrepancies between \(f_{\text{Ca}}\) from \(\delta^{44}\text{Ca}\) and Mg/Ca (e.g., the requirement for AF > 1). We observe only a greater, never lower, magnitude of Mg/Ca change than that expected from changes in \(f_{\text{Ca}}\) alone. In the La Vallina Cave system, drip water monitoring shows very constant drip water Mg/Sr ratios seasonally despite order of magnitude changes in drip water flow rates (Fig. 12 in Kost et al., 2023b). This suggests that despite variation in water residence times, there is not widespread or significant variation in bedrock mixing fraction.

5.2.2 Suggested approach for fossil stalagmites

Mg/Ca is a widely measured PCP-sensitive indicator. However, it can be unreliable in host rocks which feature significant components of widely varying solubility and Mg/Ca if the bedrock mixing fraction varies over time. This can produce variation in the initial drip water Mg/Ca attained by bedrock dissolution before evolution of drip water by PCP. Large temporal variations in drip water Mg/Sr ratios at a single drip during monitoring may be one sign of varying bedrock mixing fraction. Mg/Ca can also be unreliable as a PCP indicator if a large fraction of the Mg in the stalagmite is present in detrital minerals rather than in the calcite. Stalagmite Mg/Ca which covaries with Al/Ca or Si/Ca may indicate an important detrital component of Mg. In these two situations, it is not recommended to use Mg/Ca for quantitative PCP estimation.

In settings where bedrock mixing fraction is stable and detrital minerals are not a significant source of Mg in the stalagmite, the quantitative estimation of \(f_{\text{Ca}}\) nonetheless entails uncertainty. For stalagmites in which only Mg/Ca is measured and \(\delta^{44}\text{Ca}\) is not determined, the constancy or range of variation in \(D_{\text{Mg}}\) will represent one source of uncertainty in the calculation of \(\delta^{13}\text{C}_{\text{init}}\). The calculation of \(f_{\text{Ca}}\) from Mg/Ca can be adjusted by an attenuation factor as in Eq. (7). An attenuation factor \(> 1\) is often required if measured Mg/Ca leads to calculation of a larger \(f_{\text{Ca}}\) range than expected. For example, \(f_{\text{Ca}} < 0.2\) would require initial drip water Ca concentrations > 100 ppm. If a bedrock adjustment factor in Eq. (7) is used, it may need to be coupled with an attenuation factor (other than 1) so that absolute \(f_{\text{Ca}}\) does not reach values lower than expected for the cave and climatic context. This uncertainty will be reduced as future research better constrains the variation of \(D_{\text{Mg}}\) with Mg/Ca.

5.3 What is the magnitude of the PCP+degassing effect on \(\delta^{13}\text{C}\)?

5.3.1 Do interglacial stalagmites constrain the degassing \(\delta^{13}\text{C}\) slope?

For interglacial stalagmites, we can estimate the lowest expected \(\delta^{13}\text{C}_{\text{init}}\) and use this limit to evaluate the range of reasonable degassing slopes. The \(\delta^{13}\text{C}_{\text{DIC}}\) of drip water has been measured under forested sections of the cave, with sample collection techniques to limit degassing (Kost et al., 2023b). From this, we estimate the \(\delta^{13}\text{C}_{\text{init}}\) for calcite forming under similar drip water \(\delta^{13}\text{C}_{\text{DIC}}\). Additionally, we hypothesize that pre-anthropogenic land use at this location would have led to soil \(p\text{CO}_2\) in the 8000 to 10 000 ppm range during the mid-Holocene and previous interglacials (Lechleitner et al., 2021). We therefore calculate the composition of calcite forming from DIC in equilibrium with open-system dissolution (e.g., 150 L gas volume and resulting 3% dead carbon percentage) of limestone by 10 000 ppm soil \(\text{CO}_2\) of isotopic composition corresponding to modern respired end-members in three ecosystems (Pataki et al., 2003) (Fig. 6).

We find that interglacial stalagmites cannot rule out any of the investigated degassing slopes. The stalagmites from our monitored cave which include long Holocene growth phases are GAL and GUL, which require low to moderate attenuation factors (1.3 to 3) for calculation of \(f_{\text{Ca}}\) from Mg/Ca. When the \(f_{\text{Ca}}\) calculated from Mg/Ca includes the estimated AF for coherency with \(\delta^{44}\text{Ca}\) as in Table 2, then the −8% slope for all GAL and GUL samples falls within the range of calcite composition expected from modern drip water with limited degassing (Fig. 6). With the −11% slope, the median \(\delta^{13}\text{C}_{\text{init}}\) approaches −17%, which, although higher than modern drip water predictions, is still compatible with DIC derived from equilibration with respired \(\text{CO}_2\) from temperate broadleaf and conifer ecosystems (Pataki et al., 2003). With the −5% slope, the median \(\delta^{13}\text{C}_{\text{init}}\) is slightly below that implied for calcite on the basis of modern monitoring. Results for stalagmites spanning the last interglacial
also indicate that the −8‰ to −11‰ slopes yield δ¹³C<sub>init</sub> coherent with expected interglacial calcite. BEL, which requires little attenuation, best matches expected range with −11‰. But, our comparison does reveal that unreasonable corrections which are more negative than expected for the respired CO₂ component result from the combination of the −11‰ degassing slope and $f_{Ca}$ calculated from measured Mg/Ca without the use of AF (scenario “full”) for some low-Mg/Ca stalagmites.

A similar analysis for the Holocene Heshang Cave, for which both Mg/Ca and $f_{Ca}$ data are available, yields δ¹³C<sub>init</sub> with the −5‰ slope of −12‰ and with the −11‰ slope δ¹³C<sub>init</sub> of −15‰, both within the range of calcite composition expected from high-CO₂ soils with respired CO₂ composition from temperature broadleaf or tropical ecosystems (Pataki et al., 2003) (Fig. 6), consistent with the C3 evergreen broadleaf vegetation overlying Heshang Cave (Li et al., 2014).

Thus, the available interglacial datasets so far do not preclude the kinetically enhanced slope of −11‰ for fractionation during degassing and PCP. In fact, for the last interglacial, for many stalagmites, the higher slopes yield δ¹³C<sub>init</sub> more consistent with expected interglacial soil pCO₂. Although the degree of open vs. closed dissolution cannot be inferred for last interglacial samples, if it is in the range of Holocene and late glacial samples from this and nearby caves (Lechleitner et al., 2021), the δ¹³C<sub>init</sub> is not expected to differ by more than 1‰ from that calculated for the relatively open 10,000 ppm soil CO₂ CaveCalc model.

### 5.3.2 Previous laboratory and field calibrations

The CaveCalc simulation assumes that the loss of carbon from drip water is attained via CO₂ degassing and calcite precipitation in a 1 : 1 ratio, with close coupling of the processes, and that fractionation is occurring at equilibrium according to laboratory-estimated fractionation factors. Empirical laboratory and field results are sensitive to true degassing and PCP as well as the isotopic consequences of exchange between drip water and cave atmosphere. We are aware of one published field study in which drip water was sampled before and after a ~3 m transit across a subvertical flowstone surface in a cave with average temperature of 22.5°C (Mickler et al., 2019). In this transect, Ca and DIC were directly measured and δ¹³C DIC was determined from total evolution as CO₂ in flushed vials. Ca concentrations were on average 20% lower after the ISST flowstone transit, and the median estimated degassing slope was −9.9‰ in the eight monthly paired samples. The slopes estimated on individual sampling dates (ranging from −7.7 to −13.7) did not correlate with drip water oversaturation or the estimated ratio of Ca to DIC decrease on those dates. The slope in this study is more than double that of CaveCalc at comparable temperatures. One interpretation is that the rapid initial phases of degassing occur at kinetically enhanced fractionation factors (Mickler et al., 2019). A similar experiment conducted on one day in La Vellina Cave was also consistent with a slope of −11‰ based on sampling at a cave temperature of 12°C and modest saturation (55 ppm Ca in drip water vs. 21 ppm for equilibrium at cave temperature and pCO₂ ppm) (Kost et al., 2023b). During this sampling, cave air was ventilated to nearly atmospheric CO₂ concentrations and isotopic composition. These laboratory and field results suggest that, given current available data, the CaveCalc slopes may need to be considered the minimum slopes required to correct for degassing and PCP effects on carbon isotopes.

The laboratory experiment best simulating the coupled evolution of δ¹³C DIC, calcite δ¹³C, and $f_{Ca}$ allowed highly supersaturated solutions to degas and precipitate CaCO₃ as they flowed along a marble plate, with electrical conductivity measurements estimating the degree of Ca depletion along the flow path in a parallel experimental setup with flow over a glass plate (Hansen et al., 2019). The δ¹³C of DIC was measured by precipitation in SrCO₃ and analysis as solid phase. The slope A in the 10 experiments, based on the evolution of δ¹³C of DIC, ranged from −4.63‰ to −13.2‰ from −5.9‰ to −16‰ based on evolution of δ¹³C of calcite, although precise estimation of the latter slope is complicated by the measurement of precipitated calcite δ¹³C and conductivity in separate experimental setups which may have experienced slightly different locations of calcite precipitation (Sade et al., 2022). Many of these estimated experimental fractionations are considerably higher than those predicted by CaveCalc.

A dynamical model (Sade et al., 2022) was developed to simulate the laboratory experiments (Hansen et al., 2019). This model derived a single relationship between PCP and DIC evolution using $f_{DIC}$ in contrast to our calculated index based on Ca consumption, $f_{Ca}$. In this model formulation, the degassing slope relative to ln($f_{DIC}$) averages −8.7‰ from $f = 1.0$ to $f = 0.3$, but the slope is not constant. Rather, at $f_{DIC}$ between 0.8 and 1, there is an inverse slope as net DIC evolution is modeled to lead to more negative, rather than more positive, δ¹³C of speleothem calcite. Over the subsequent phase of PCP ($f_{DIC}$ 0.8 to 0.3), the average modeled degassing slope relative to $f_{DIC}$ would be −10.1‰; however, it flattens to −4‰ in the range of $f_{DIC}$ 0.33 to 0.4. If this model is representative of cave flowstone formation, it may suggest that some of the calculated differences in degassing slopes in field experiments may result from measurements of solutions sampled in different stages of drip water Ca and DIC evolution. Alternatively, it is possible that some differences in degassing slopes reflect different degrees of coupling between degassing and CaCO₃ precipitation. The influence of factors inhibiting CaCO₃ precipitation, potentially including elevated Mg concentrations or certain dissolved organic compounds, on the degassing slope remains to be investigated.

If there were significant variations in the degassing slope experienced by a given stalagmite over time, it could compli-
cate efforts to estimate trends in $\delta^{13}C_{\text{init}}$ using Eq. (2). Yet, our speleothem time series are broadly consistent with the narrow range of variation of degassing slope within a given stalagmite, since correction of $-5\%$ in one part of the time series and $-11\%$ in another would alter the trends and reduce coherence among the records in Fig. 7. This coherence with a single slope may reflect the limited range of variation in $f_{\text{Ca}}$ inferred for our stalagmites. For example, for all stalagmites except GAE, the A2 scenarios and the majority of data from A3 scenarios fall between $f_{\text{Ca}}$ of 0.4 and 0.8. If coincident with $f_{\text{DIC}}$ of 0.4 to 0.8, these samples would all be within the range of similar slope ($-10\%$ to $-11.5\%$) predicted by the dynamical model simulation. Sampled stalagmite growth may often be biased to periods when $f_{\text{Ca}} > 0.4$, which lead to higher precipitation rates and more calcite deposition. The consistency of our interglacial data with steeper degassing slopes than CaveCalc may reflect the reality that much calcite precipitation will happen when cave $pCO_2$ is much lower than drip water $pCO_2$, a situation in which drip water exchange with isotopically heavier cave air could be a significant contributor to a steep degassing slope.

A future calculation of the dynamical model prediction (Sade et al., 2022) of $\delta^{13}C$ evolution relative to $f_{\text{Ca}}$, rather than $f_{\text{DIC}}$, as a PCP indicator would provide a helpful reference for future calculation of $\delta^{13}C_{\text{init}}$. This could elucidate whether a nonlinear relationship should be used to calculate $\delta^{13}C_{\text{init}}$ from $\ln(f_{\text{Ca}})$ and $\delta^{13}C_{\text{meas}}$. It would clarify if calcite precipitated at high $f_{\text{Ca}}$, analogous to that precipitated at high $f_{\text{DIC}}$, should be excluded from the correction because of compensating effects of kinetic fractionation factors and evolution of $f_{\text{Ca}}$ in the range of high $f_{\text{Ca}}$. Scenarios employing a $B \leq 0.8$ by default have a maximum $f_{\text{Ca}}$ largely outside the range of inverse slope.

5.3.3 Suggested approach

Trends in $\delta^{13}C_{\text{init}}$ are less sensitive to the choice of the degassing slope but the absolute value of $\delta^{13}C_{\text{init}}$ is strongly dependent on the choice of degassing slope. At the moment, the absolute $\delta^{13}C_{\text{init}}$ can be reconstructed with low confidence. However, the reconstructed $\delta^{13}C_{\text{init}}$ can be used to rule out combinations of $f_{\text{Ca}}$ scenarios and degassing slopes which lead to “overcorrection” to $\delta^{13}C_{\text{init}}$ values which are more negative than DIC equilibrated with a reasonable respired soil $pCO_2$ composition for the ecosystem type. Until further constraints to assess the slope of $\delta^{13}C$ and Ca co-evolution during degassing and PCP which characterizes typical cave environments are available, a sensitivity analysis employing a range of degassing slopes will provide the greatest transparency for assessing $\delta^{13}C_{\text{init}}$.

6 Conclusions

We provide a first analysis of the potential estimation of speleothem $\delta^{13}C_{\text{init}}$, the $\delta^{13}C$ which characterized drip water DIC prior to significant degassing and PCP. $\delta^{13}C_{\text{init}}$ is proposed as a useful interpretable variable derived from speleothem isotope measurements because trends in $\delta^{13}C_{\text{init}}$ are expected to be more reproducible than measured $\delta^{13}C$ among coeval stalagmites from a given region and because $\delta^{13}C_{\text{init}}$ is more sensitive to the vegetation, soil, and epikarst processes which in many regions may be sensitive to temperature in addition to moisture. Calculation of $\delta^{13}C_{\text{init}}$ for a given sample or growth increment requires a quantitative indicator of the extent of PCP experienced in that growth increment and knowledge of the general rate of change in $\delta^{13}C$ with progressive degassing and PCP.

In fossil stalagmites, the extent of PCP, as the Rayleigh $f_{\text{Ca}}$, can be approximated for a particular growth increment using the measured Mg/Ca and the minimum Mg/Ca measured in the entire stalagmite. $f_{\text{Ca}}$ can also be estimated from $\delta^{44}Ca$ given reasonable choices of the calcite dissolved fractionation factor. In several of the stalagmites examined here, the $f_{\text{Ca}}$ calculated from Mg/Ca shows an amplified range compared to that calculated from $\delta^{44}Ca$. An increase in $D_{\text{Mg}}$ with increasing Mg/Ca, as proposed in previous studies (Wassenburg et al., 2020), is one viable explanation for this systematic trend. This observation warrants further investigation to improve confidence in future estimates of $f_{\text{Ca}}$ from Mg/Ca, which is the most widely available PCP indicator for stalagmites.

At the moment, there is uncertainty in the degassing slope or rate of change in $\delta^{13}C$ with progressive degassing and PCP. Values may range from $-5\%$ to $-11\%$. The stalagmites investigated here could be consistent with any value in this range. Because these different possible degassing slopes affect the absolute value of the calculate $\delta^{13}C_{\text{init}}$, currently there is uncertainty in the exact value. However, the trends in $\delta^{13}C_{\text{init}}$ are less sensitive to the choice of degassing slope.

Despite these uncertainties, $\delta^{13}C_{\text{init}}$ provides more consistent time series among coeval stalagmites and with regional climate records. In one example $\delta^{13}C_{\text{init}}$ reconciles three divergent measured stalagmite $\delta^{13}C$ records in the 94 to 82 ka time interval, yielding three $\delta^{13}C_{\text{init}}$ time series which feature a pronounced positive anomaly corresponding to the regional cooling of Greenland Stadial 22. In western Europe, over the warming trend of deglaciations, the trend towards more negative $\delta^{13}C_{\text{DIC}}$ due to higher soil respiration and soil CO$_2$ may be fully or partially compensated for in stalagmite $\delta^{13}C$ due to the increased PCP from the more oversaturated drip waters. The calculation of $\delta^{13}C_{\text{init}}$ reveals the trend of increasing respiration rates and soil $pCO_2$. Over the last deglaciation, the $\delta^{13}C_{\text{init}}$ matches the millennial structure in regional SST more closely than the measured $\delta^{13}C$.

As better constraints emerge on the degassing slope and on Mg partitioning, $\delta^{13}C_{\text{init}}$ estimates will become more precise and should improve the utility of the high volume of stalagmite $\delta^{13}C$ measurements made simultaneously with $\delta^{18}O$ in all labs.
Appendix A

Table A1. Measured \(\delta^{44}\)Ca, Mg/Ca, and Sr/Ca, as well as the calculated \(f_{Ca}\) for differing \(\Delta^{44}\)Ca and \(B\) (bedrock factors).

<table>
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<th>(\delta^{44/40})Ca (%e)</th>
<th>Mg/Ca (mmol mol(^{-1}))</th>
<th>Sr/Ca (mmol mol(^{-1}))</th>
<th>(f_{Ca}) from (\Delta^{44})Ca (%e)</th>
<th>(f_{Ca}) from Mg/Ca (B)</th>
<th>(f_{Ca}) from Sr/Ca (B)</th>
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<td>3.12</td>
<td>0.034</td>
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<tr>
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<td>0.032</td>
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<td>0.49 0.39 0.29</td>
<td>0.23 0.18 0.14</td>
</tr>
<tr>
<td>GAE_60.4 nd</td>
<td>0.13</td>
<td>17.59</td>
<td>0.115</td>
<td>0.73 0.62 0.56 0.51</td>
<td>0.12 0.10 0.07</td>
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<td>GAR_b2_035.0</td>
<td>−0.25</td>
<td>7.42</td>
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<td>1.29 0.96 0.79 0.67</td>
<td>0.33 0.26 0.20</td>
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<tr>
<td>GAR_b6_131.5</td>
<td>−0.49</td>
<td>2.42</td>
<td>0.073</td>
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<td>1.00 0.80 0.60</td>
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<tr>
<td>GLO_b1_17.4</td>
<td>0.23</td>
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<td>0.78 0.62 0.47</td>
<td>0.55 0.44 0.33</td>
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<tr>
<td>GLO_b2_08.2</td>
<td>0.42</td>
<td>17.24</td>
<td>0.068</td>
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<td>0.30 0.24 0.18</td>
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<tr>
<td>GUL-2.3</td>
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<td>2.19</td>
<td>0.062</td>
<td>1.11 0.86 0.72 0.63</td>
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<td>0.41 0.33 0.25</td>
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<td>0.06</td>
<td>3.69</td>
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<tr>
<td>NYM_331</td>
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<td>NYM_457.5</td>
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<td>0.69 0.56 0.42</td>
</tr>
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</table>

Data availability. Data and sample Excel sheets for calculations are archived in the ETH Research Collection at https://doi.org/10.3929/ethz-b-000630754 (Stoll et al., 2023).

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/cp-19-2423-2023-supplement.

Author contributions. HMS conceived the study, conducted calculations, prepared figures, and wrote the text in discussion with CD and FL. CD completed Ca isotope measurements. FL and LE conducted CaveCalc simulations. OK and JS assisted with trace element analysis and sampling for Ca isotopes and dating. DS contributed to interpretation. HC and CPM completed new chronology.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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Acknowledgements. Heather M. Stoll acknowledges ETH core funding and grant ETH-13 18-1. Franziska Lechleitner was supported by SNSF grant P400P2_180789. Chris Day and Ca isotope measurements were supported by the John Fell Oxford University Press Research Fund under grant no. 0007911. Franziska Lechleitner was supported by SNSF grant P400P2_180789. Carlos Pérez-Mejías and Hai Cheng acknowledge the NSFC (41888101, 42050410317) and Postdoctoral Science Foundation of China (2020M683452). We thank the ETH Climate Geology lab manager Madalina Jaggi and student assistants Tim Loefel and Romain Alosius for assistance with stable isotope and trace element sampling. We thank the ETH fall 2021 paleoclimate course students and course assistant Pien Anjewierden for preparing samples from GLD, as well as fall 2019 paleoclimate course students and Romain Alosius for assistance with stable isotope and trace element sampling. We thank the ETH fall 2021 paleoclimate course students and course assistant Pien Anjewierden for preparing samples from GLD, as well as fall 2019 paleoclimate course students for initiating the study of GUL.

Financial support. This research has been supported by the Eidgenössische Technische Hochschule Zürich (grant no. ETH-13 18-1), the Schweizerischer Nationalfonds zur Förderung der Wissenschaftlichen Forschung (grant no. P400P2_180789), and the National Natural Science Foundation of China (grant nos. 41888101, 42050410317, and 2020M683452).

Review statement. This paper was edited by Dominik Fleitmann and reviewed by two anonymous referees.

https://doi.org/10.5194/cp-19-2423-2023
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