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Research Article

Using Carbon Isotope Equilibrium to Screen Pedogenic Carbonate Oxygen Isotopes: Implications for Paleoaltimetry and Paleotectonic Studies

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Stable isotope compositions of pedogenic carbonates (δ¹³C carb, δ¹⁸O carb) are widely used in paleoenvironmental and paleoaltimetry studies. At the same time, both in vertical stratigraphic sections and in horizontal transects of single paleosols, significant variability in δ¹⁸O carb values is observed well in excess of what could reasonably be attributed to elevation changes. Herein, a new screening tool is proposed to establish which pedogenic carbonate δ¹⁸O carb compositions reflect formation in isotopic equilibrium with environmental conditions through the use of the co-occurring δ¹³C org composition of carbonate-occluded or infiltrate organic matter, where Δ¹³C = δ¹³C carb – δ¹³C org. Based upon 51 modern soils from monsoonal, continental, and Mediterranean moisture regimes, Δ¹³C = +15.6 ± 1.1‰ (1σ), which closely matches theoretical predictions for carbonates formed at carbon isotope equilibrium through Fickian diffusion. Examples from both disequilibrium and equilibrium cases in the geologic record are examined, and it is shown that previous δ¹⁸O carb records used to infer Cenozoic uplift in southwestern Montana do not provide any constraint on paleo-elevation because >90% of the pedogenic carbonate isotopic compositions are out of equilibrium. Guidelines for future paleoaltimetry studies include collection of both vertical stratigraphic sections and lateral transects, of at least three nodules per horizon, petrographic screening of nodules for diagenesis, collection of at least one independent proxy for paleoclimate or paleovegetation, and screening δ¹⁸O carb values using Δ¹³C measured for each paleosol.

1. Introduction

In many environments, soils form carbonate (CaCO₃) nodules in response to seasonal drying of HCO₃⁻-rich interstitial water, which leads to carbonate supersaturation (e.g., [1, 2]). While formation of pedogenic carbonates is most common in arid to subhumid environments (<750 mm yr⁻¹; [3]), under strongly seasonal climatic regimes, pedogenic carbonate may form under much wetter conditions in a mean annual sense. As a result, pedogenic carbonate is nearly ubiquitous in the Phanerozoic geologic record at least at the regional scale (e.g., Figure 19 of [4]). Because the C and O isotopic composition of pedogenic carbonate reflects the isotopic composition of fluid that it precipitated from the time of their formation (e.g., [5, 6]), pedogenic carbonates have been widely used for reconstructing past environmental conditions.

Carbon isotopic compositions (δ¹³C carb) have been used to reconstruct the composition of paleovegetation (e.g., [7]; Fox and Koch, 2003) and have been used in concert with C isotopic compositions of co-occurring organic matter (δ¹³C org) to reconstruct paleo-CO₂ levels ([8–10] and many others) and paleoproductivity (e.g., [11, 12]). Values for δ¹³C carb can reflect up to three sources of CO₂ (e.g., [4]): (1) open-system oxidation of soil organic matter, (2) atmospheric CO₂, and (3) CO₂ derived from preexisting carbonate. In general, pedogenic carbonate derived from only the first component of CO₂ is restricted to swampy or wetland environments (e.g., [6, 13]) and pedogenic carbonate formed from all three components only occurs in places where the soil parent material was derived from lacustrine or marine limestones. Thus, most pedogenic carbonates represent a two-component mixture of CO₂ derived from oxidation of soil organic matter and atmospheric CO₂.
Oxygen isotopic compositions ($\delta^{18}O_{\text{carb}}$) of pedogenic carbonates reflect both the temperature of precipitation and $\delta^{18}O$ composition of the fluid that the carbonates crystallized from (e.g., [14]), and as a result, have been used to reconstruct both of those variables by assuming the other, through an independent constraint on one variable or other, and more recently, through the use of “clumped isotope” measurements, which produce temperatures that are independent of the starting fluid isotopic composition. The underlying assumption in this is that the carbonates formed in equilibrium with their environmental conditions (e.g., reviewed by Quade et al. [15]). While a few studies have focused on $\delta^{18}O_{\text{carb}}$ for purely climatological purposes (e.g., [16]), $\delta^{18}O_{\text{carb}}$ has most often been used to consider tectonic questions. When an air mass rises over a topographic impediment, the meteoric $\delta^{18}O$ value of the remaining water vapor becomes progressively more negative due to Rayleigh distillation (e.g., [17]). As a result, $\delta^{18}O_{\text{carb}}$ values of pedogenic carbonates that formed at higher elevations are isotopically more negative relative to those formed at lower elevations. Because the relationship between elevation and meteoric water $\delta^{18}O$ can be estimated either globally [18] or on a region-to-region basis (e.g., [19–21]), changes in $\delta^{18}O_{\text{carb}}$ through a stratigraphic succession can be used to back-calculate elevation change through time.

1.1. Challenges of Using $\delta^{18}O_{\text{carb}}$ for Paleoaltimetry Studies. A variety of potential challenges of using $\delta^{18}O_{\text{carb}}$ data as a tool for deciphering paleoaltimetry have been raised including how pure the carbonate mineral phase is (e.g., 100% calcite versus other mixtures), variations in the amount of meteoric precipitation at the time of carbonate formation, variations in the $\delta^{18}O$ composition of the meteoric water in space, time, and seasonally, changes in climatic temperatures through time, among others (for an in-depth review, see [15]). As a result of these complications, in many terrestrial carbonate data series, there is a substantial spread in the measured $\delta^{18}O_{\text{carb}}$ values. For example, Figure 1 shows a long-term $\delta^{18}O_{\text{carb}}$ record compiled from carbonates in Montana and Idaho that has been used to interpret a significant uplift event between 50 and 30 Ma ago (data from [22]), in which, for some five million year time bins, the spread in $\delta^{18}O_{\text{carb}}$ exceeds 13‰. Two different potential interpretive frameworks are shown using those data within the context of the global $\delta^{18}O$ lapse rate (~2.8‰ km$^{-1}$; [18]): one that assumes that the maximum range between high and low $\delta^{18}O_{\text{carb}}$ values represents maximum possible uplift of ~4.5 km, and a second one that uses the mean $\delta^{18}O_{\text{carb}}$ values before and after a depositional hiatus from 32–24 Ma ago to infer just 0.6 km of uplift. To be clear, neither of these scenarios was favored by the authors, but they illustrate the scale of the potential uncertainty in using all of the measured $\delta^{18}O_{\text{carb}}$ data without regard to any consideration of “good” versus “bad” data points.

A second issue that was raised recently by Hyland and Sheldon [23] concerns the stratigraphic fidelity of single vertical stratigraphic sections. Those authors measured $\delta^{18}O_{\text{carb}}$ values from a single paleosol over a multi-km lateral transect and found a range of ~8‰ (Figure 2). The star in Figure 2 gives the value of the previous paleosol in the vertical succession, and the circles give the potential paleoelevation change that one could interpret in spite of ~5 m difference in stratigraphic height between the two pedogenic carbonate-bearing paleosols. The range in interpretations is from little or no change to nearly two km of instantaneous uplift. Which, if any, of those results seem credible?

While the recent development of “clumped isotope” geochemistry of pedogenic carbonates promises to improve paleoaltimetry studies (e.g., reviewed by [15]) and to alleviate some of the issues raised above, it is a comparatively higher cost (both analytically and in terms of time) technique and requires significantly larger samples sizes. In addition, because massive archives of conventional $\delta^{18}O_{\text{carb}}$ already exist, having tools to screen them directly would substantially improve future paleoaltimetry studies as well as making it possible to refine existing interpretations. Here, I will propose a new interpretive framework for understanding $\delta^{18}O_{\text{carb}}$ values from pedogenic carbonates to improve terrestrial paleoclimate reconstructions and paleoaltimetric reconstructions.

2. Methods

To refine the use of $\delta^{18}O_{\text{carb}}$ values from paleosols in paleoaltimetry studies, I will look at carbon isotope equilibrium as a screening tool. Modern soil samples were collected from nine separate pits from two different soil series in Arizona, USA. The Guvo soil series ($n = 5$) represents a calcic Aridisol formed at a mean annual temperature (MAT) of 21.7°C and a mean annual precipitation of 232 mm yr$^{-1}$. The Delthorny soil series ($n = 4$) represents a calcic Aridisol formed at a MAT of 21.6°C and an MAP of 304 mm yr$^{-1}$. Most of the precipitation for both soil series is delivered in the spring,
as both sites are located within the monsoon belt for the southwestern US. Both soil series preserve pedogenic carbonate as nodules and as coatings on rock fragments. Bulk soil was collected at 2–3 sample depths within the upper 30 cm of each profile as well as intact roots. For all but one of the soil pits where there was little carbonate present, three carbonate nodules were collected from each soil pit. Samples for organic isotope analysis were decarbonated in weak HCl (2%), dried, homogenized, and measured into tin capsules for analysis. Organic carbon isopes (δ\textsuperscript{13}C\textsubscript{org}) were measured at the University of Michigan’s Stable Isotope Lab on a Delta V+ IRMS coupled to a Costech ECS 4010 elemental analyzer; results were calibrated against IAEA sucrose and caffeine standards and are presented relative to the PDB scale. External reproducibility was better than 0.1‰ for both standards and replicates. Carbonate stable isotope compositions (δ\textsuperscript{13}C\textsubscript{carb} and δ\textsuperscript{18}O\textsubscript{carb}) were obtained on micro-drilled samples of micritic carbonate (n = 3 per nodule) that were analyzed on a Thermo 253 IRMS coupled to a Kiel IV autosampler; results were calibrated against NBS-19 and internal standards, and results are presented relative to the PDB scale for C and the SMOW scale for oxygen. Extern regional reproducibility was better than 0.04‰ for both standards and replicates.

Precipitation δ\textsuperscript{18}O estimates for the Guvo and Delthorny soil pit sites were obtained using the Online Isotopes in Precipitation Calculator (OIPC; [24]), using latitude, longitude, and elevation for each site. The OIPC uses data from IAEA precipitation monitoring sites and interpolates for sites located between observational record sites. Previously published data for modern soils come from Cerling and Quade [25] and Tabor et al. [6], and the geologic data for discussion come from [22, 26, 27] and [23, 28]. All of the new data are compiled in the Supplemental Materials (available here).

### 2.1. Interpretational Framework

Cerling [2, 9] presented a foundational understanding of the stable isotope values of pedogenic carbonates, and an updated discussion of subsequent work can be found in a recent review paper by Sheldon and Tabor [4]. Briefly though, the δ\textsuperscript{13}C\textsubscript{carb} value of pedogenic carbonates should reflect a mixture of up to three components of CO\textsubscript{2} as described above. By general practice, three-component (i.e., with inherited carbonate) pedogenic carbonate is avoided, because it is extremely difficult to deconvolve the contribution of CO\textsubscript{2} from inherited carbonate from the authigenic sources. In other words, paleosols with carbonate in their parent material are avoided. Isotopic compositions of two-component pedogenic calcites (i.e., those derived from soil-respired CO\textsubscript{2} and atmospheric CO\textsubscript{2}) will reflect the depth position within the soil [2], and studies of in situ CO\textsubscript{2} mixing in soils reflect one-dimensional Fickian diffusion (various; [1, 2]). Below a characteristic depth, both modelled and observed to be >30 cm, soil isotopic CO\textsubscript{2} and δ\textsuperscript{13}C\textsubscript{carb} values should be in isotopic equilibrium. Figure 3 illustrates this reaction pathway, along with the processes responsible for the observed fractionations. For pedogenic carbonates formed prior to ~30 Ma ago when C\textsubscript{4} photosynthesis first evolved [29], they all should reflect the distribution on the left-hand column, which depicts plants using the C\textsubscript{3} photosynthetic pathway. More recent carbonates are more complicated and could reflect either a pure C\textsubscript{3} ecosystem, a mixed C\textsubscript{3}-C\textsubscript{4} ecosystem, or for the last ~8 Ma, potentially a nearly pure C\textsubscript{4} ecosystem.

For paleosols that have both carbonates and organic matter, it should be possible to demonstrate whether they have formed in isotopic equilibrium. Based upon the theoretical framework in Figure 3, that relationship should be as follows:

\[
\Delta^{13}C = \delta^{13}C_{\text{carb}} - \delta^{13}C_{\text{org}} = +15\%o.
\]

Soils that have formed in C isotope equilibrium should also be in O isotope equilibrium; thus, Δ\textsuperscript{18}O can be used as a screening tool to determine which δ\textsuperscript{18}O\textsubscript{carb} values can be reliably interpreted.

### 3. Results

Figure 4 shows δ\textsuperscript{13}C\textsubscript{org} as a function of depth for the five Guvo soil series pits. Each pit is most negative at the surface and reaches a characteristic equilibrium value below a depth of 20 cm. This is consistent with Fickian diffusion as described in Cerling’s [2] theoretical framework as well as
observed in the studies of in situ measurements of soil gas $\delta^{13}C$ and coexisting $\delta^{13}C_{\text{org}}$ [1, 2, 25]. Measured $\delta^{13}C_{\text{carb}}$ values for the Guvo soil series ranged from $-4.0$ to $-6.2\%_{\text{o}}$, and standard deviations of replicate analyses of multiple nodules from the same pit ranged from $\pm 0.19\%_{\text{o}}$ to $\pm 0.43\%_{\text{o}}$, with a mean of $\pm 0.33\%_{\text{o}}$. $\Delta^{13}C$ ranged from 16.5 to 17.8. Measured $\delta^{13}C_{\text{carb}}$ values for the Delthorny soil series ranged from $-2.8$ to $-4.8\%_{\text{o}}$, and standard deviations of replicate analyses of multiple nodules from the same pit ranged from $\pm 0.19\%_{\text{o}}$ to $\pm 0.58\%_{\text{o}}$, with a mean of $\pm 0.42\%_{\text{o}}$. $\Delta^{13}C$ ranged from 16.2 to 17.1.

Figure 5 plots $\delta^{13}C_{\text{carb}}$ versus $\delta^{13}C_{\text{org}}$ for each of the nine soil pits as well as the data from two-component calcite soils that were previously published by Tabor et al. [6] from California sites with a Mediterranean climate (winter-wet) and from Cerling and Quade [25] from a variety of western US and Midwestern US sites that reflect an array of precipitation regimes, including both monsoonal and continental (summer-wet) sites. Regardless of the climatic moisture regime, there is a strong correlation ($r = 0.96$) between $\delta^{13}C_{\text{carb}}$ and $\delta^{13}C_{\text{org}}$ that is highly significant ($p < 10^{-26}$). The empirically derived $\Delta^{13}C$ value is $+15.3\%_{\text{o}}$ (i.e., intercept in Figure 5), which closely matches the theoretical $\Delta^{13}C$ value given in equation (1). The mean $\Delta^{13}C$ value for all 51 soils is $+15.6\%_{\text{o}}$ (1 $\sigma$), which also closely matches equation (1), and will be used as the threshold for equilibrium herein.

4. Discussion

4.1. Interpreting Nonequilibrium $\Delta^{13}C$ Values. Cotton et al. [26] used $\delta^{13}C_{\text{org}}$ and phytoliths (fossil plant biosilica) to reconstruct the proportion of $C_4$ plants present in southwestern Montana during the late-middle Miocene (~10.2–8.9 Ma ago), with both types of proxies indicating up to 20% $C_4$. However, pedogenic carbonates from the same stratigraphic section indicated up to 50% $C_4$ plants and had $\delta^{18}O_{\text{carb}}$ values...
Differences in surface meteoric precipitation. With Δ13C values from pedogenic carbonates collected below those depths should be in isotopic equilibrium with their coexisting soil organic matter’s δ13Corg value.

Figure 4: Individual soil pit δ13Corg depth profiles of five Guvo Soil Series soils. Differences in surface δ13Corg and δ13Corg at depth reflect differences in the overlying vegetation, but each of the five profiles becomes asymptotic below ~20 cm in the profile, so δ13Ccarb values from pedogenic carbonates collected below those depths should be in isotopic equilibrium with their coexisting soil organic matter’s δ13Corg value.

Figure 5: Plot of the relationship between δ13Ccarb and δ13Corg for samples collected >30 cm below the soil surface. The results from different regions and different climatic regimes all fall along the same regression line and the calculated Δ13C values closely matches the theoretical value given in equation (1). While δ13Corg actually drives δ13Ccarb, by convention Δ13C is δ13Ccarb − δ13Corg, so that is how the data have been regressed here. The solid black line represents the linear regression, the dashed lines the 95% confidence interval, and the grey lines the 95% prediction confidence interval.

Figure 6: Plot of δ13Ccarb versus Δ13C. The grey box spans Δ13C values +13.4–17.84‰, which reflects the measured mean Δ13C value of +15.62‰ ± 2σ (2.22‰). The solid black line reflects the slope of all carbonates formed from one-, two-, and three-components of CO2 from Tabor et al. [6]. The data from Cotton et al. [26] are given by black stars and represent either carbonates formed under extremely evaporatively enriched conditions in terms of meteoric water or from three-component CO2. In this case, because the entire stratigraphic section is calcite-cemented (grey dashed line; δ13Ccarb = −3.2 ± 0.2‰) and the calcite cement is isotopic indistinguishable from the pedogenic carbonates, one would conclude that the pedogenic carbonates reflect this inheritance or recrystallization.

4.2. Screening δ18Ocarb Values Using Δ13C. Returning to Figure 2, the open question was which, if any, of the pedogenic carbonate δ18Ocarb values could be used from the Wasatch Formation paleosols [23, 28, 30] to determine paleoelevation. Measured Δ13C values along the single paleosol transect ranged from 12.4–19.1‰ and were accompanied by δ18Ocarb values that ranged from 18.5–26.3‰ (SMOW). Using the empirically determined Δ13C value of +15.6 ± 1.1 (1σ), it is clear that only one of the four paleosol profiles from the lateral transect was formed in isotopic equilibrium (Figure 2). In this case, paleosol also corresponds to the most geologically reasonable scenario, namely, of little to no uplift between the closely separated (stratigraphically speaking) paleosols. Hyland and Sheldon [23, 28] noted hydromorphic features locally along the transect that were confirmed by rock magnetic properties as likely reflecting localized ponding or a water table that was near to the surface of soils at least ranging from 13.4 to 14.5‰ (SMOW), which are indistinguishable from the values in Figure 1 for that time period. Using the empirically defined Δ13C value of +15.6 ± 1.1 (1σ), it is possible to determine which of these two vegetation scenarios is most likely to be correct, but also whether the δ18Ocarb values provide a meaningful constraint on meteoric precipitation. With Δ13C values of 18.2–22.4, all of the pedogenic carbonate values are too positive relative to their coexisting organic matter to be considered in equilibrium (Figure 6) for the framework that is established here. Therefore, the pedogenic carbonate-based paleovegetation results cannot be considered viable, and the δ18Ocarb values are unlikely to reflect meteoric precipitation. Indeed, as those authors noted, the δ18Ocarb values were indistinguishable between the pedogenic carbonate nodules and interstitial cements from the same section (Figure 6). However, in the absence of that information, the same conclusion could be drawn using Δ13C values to screen for equilibrium (Figure 6).
seasonally during their formation. Transect positions with evidence for hydromorphy correspond to both of the $\Delta^{13}C$ values that fall below the ±2σ envelope around the empirically derived equilibrium value. This is consistent with Tabor et al.’s [6] observations of the effects of water-logging on modern soils and with $\delta^{13}C_{\text{org}}$ values that reflect one-component calcite. Overall, 18 of 24 sets of paired organic matter-pedogenic carbonate measurements in the main vertical section of Hyland and Sheldon [28] were formed in isotopic equilibrium (Figure 7(a)). By screening out the nonequilibrium values, the maximum range in $\delta^{18}O_{\text{carb}}$ values drops to ~6.5‰ from ~11‰. Using the global $\delta^{18}O$ lapse rate of Poage and Chamberlain [18], this would reflect a difference of ~1.6 km between the estimates of paleoelevation. (b) This panel shows the data from Gutierrez and Sheldon [27]. Using the $\Delta^{13}C$ screening criteria outlined in the text, nine of ten carbonates formed in equilibrium. The dashed line in both panels represents the empirically observed mean $\Delta^{13}C$ value of +15.6‰; the solid lines represent ±2σ (2.2‰).

Based upon the close match of theoretical equilibrium framework and the empirical observations, and the additional observations available at both of these sites that confirm the conditions under which nonequilibrium fractionation is observed, I conclude that measuring $\Delta^{13}C$ values provides a powerful screening tool for understanding $\delta^{18}O_{\text{carb}}$ Values. Recognizing samples that are out of equilibrium will make it possible to refine the interpretation of samples that are in equilibrium and should substantially reduce the potential uncertainty in paleoaltimetry studies.

There are other applications where paleosols with non-equilibrium $\Delta^{13}C$ values can provide valuable insights into soil productivity (e.g., [4]), soil drainage as outlined above (e.g., [6]), or to reconstruct MAP in arid systems [31]. Accompanying $\delta^{18}O$ values from paleosols with nonequilibrium $\Delta^{13}C$ values are still useful for understanding diageneric alteration (see below) and mixing of multiple pedogenic and postburial water sources.

4.3. Validating the Meaning of $\delta^{18}O_{\text{carb}}$ Values. To validate that $\delta^{18}O_{\text{carb}}$ values reflect source water $\delta^{18}O$ compositions, an ideal dataset would represent pedogenic carbonates that formed very close to their meteoric water source and that the meteoric water source would be relatively homogeneous throughout the year. Gutierrez and Sheldon [27] reported on a series of paleosols from the Upper Jurassic Vega Formation of Spain. The Vega Formation represents a dinosaur footprint-bearing floodplain environment; it is underlain by the ammonite-bearing shale-limestone couplets of the Rodiles Formation and is overlain by the mixed estuarine deposition of the Tereñes Formation [32]. Thus, the Vega Formation is unambiguously near sea level and near its meteoric water source. Among the paleosols with co-occurring organic matter, nine of ten are in isotopic equilibrium when screened using $\Delta^{13}C$ (Figure 7(b)) and have a mean $\delta^{18}O_{\text{carb}}$ value of 26.25 ± 0.65‰ (1σ). Using the reconstructed MAT of 13 ± 4°C [27] and measured $\delta^{18}O_{\text{carb}}$ value, the meteoric water $\delta^{18}O$ composition was −4.8 ± 0.9‰ (Figure 8; height of grey box includes ±2σ). Pedogenic carbonates typically form seasonally, with the duration of moisture deficit necessary and timing of first annual moisture deficit necessary to form pedogenic carbonate both dependent on the climatic regime (e.g., [5, 33–35]), so they could potentially form over anywhere from less than a month to the whole year depending upon the local conditions. Based upon OIPC estimates for the site today, the mean annual $\delta^{18}O_{\text{water}}$ is −6.3‰ and seasonally adjusted (i.e., March–September, representing the most likely season of carbonate formation; [5]) mean $\delta^{18}O_{\text{water}}$ is −4.1 ± 2.3‰ (Figure 8). The mean annual value is within the 2σ uncertainty of the reconstructed value and the seasonally adjusted predicted range of precipitation $\delta^{18}O$ encompasses the reconstructed value (Figure 8), suggesting that regardless of which metric is more representative of Jurassic meteoric precipitation at the site, the pedogenic
carbonate $\delta^{18}O_{\text{carb}}$ values clearly reflect their source water. Furthermore, the $\delta^{18}O_{\text{carb}}$ data exhibit low variability (i.e., low $\sigma$) throughout the stratigraphic section indicating little variation in the source moisture composition, which is consistent with Gutierrez and Sheldon [27] paleoclimatic and paleoenvironmental reconstruction based upon independent paleoclimatic proxies for MAT and MAP from major element oxide transfer functions that indicated relatively constant paleoclimatic conditions.

4.4. Implications for Montana Paleoelevation Record. As outlined above, one of the goals herein is to provide better constraints on long-term $\delta^{18}O_{\text{carb}}$ records that have been used to reconstruct paleoaltimetry and paleoelevation. Figure 1 shows a large dataset of pedogenic carbonate $\delta^{18}O_{\text{carb}}$ values from southwestern Montana [22] that can be reassessed in light of the new $\Delta^{13}C$ criterion for isotopic equilibrium. For most of the sites in that database, only $\delta^{13}C_{\text{carb}}$ values were published, but independent constraints from paleovegetation studies can be used to constrain $\delta^{13}C_{\text{org}}$. While differences in the value of $\delta^{13}C_{\text{atm}}$ would change the absolute values expected for $C_3$ plants a la Figure 3, reconstructions of Cenozoic $\delta^{13}C_{\text{atm}}$ are typically centered around $-6\%$, and show relatively little variability [36], and Retallack [37] reconstructed long-term climatic stability in the region, so the expected $C_4$ plant $\delta^{13}C_{\text{org}}$ value can be considered relatively constant. As discussed in

Interpretational Framework section, prior to the evolution of $C_4$ plants, only the empirical $\Delta^{13}C$ criterion is needed to screen for isotopic equilibrium. However, following the evolution of $C_4$ plants, $\delta^{13}C_{\text{org}}$ and $\delta^{13}C_{\text{carb}}$ would both be shifted toward more positive values if any $C_4$ plants were present. Based upon the studies of phytoliths [38] and paleosol C isotopes [26, 39, 40], $C_4$ plants comprised up to 20% of ecosystems in southwestern Montana for the whole of the Neogene. To be conservative, then, all pre-24 Ma pedogenic carbonates are assumed to have formed in a pure-$C_3$ ecosystem, and all post-24 Ma pedogenic carbonates are assumed to have formed in an ecosystem with 20% $C_4$ plants. Even using this relaxed criterion, just 29 of 308 paleosols described by Chamberlain et al. [22] formed in isotopic equilibrium (Figure 9(a)). If one allows for $\sim50\%$ more $C_4$ plants than were observed in the previous studies or for plant water stress equivalent to semiarid conditions and relaxes the high end uncertainty in $\Delta^{13}C$ by another 1.5‰ (alternatively, this could be thought of as incorporating $\geq +3\sigma$), then still just 73 of the paleosols were formed in isotopic equilibrium. Looking at just those culled data (Figure 9(b)), there is no discernible pattern in $\delta^{18}O_{\text{carb}}$ for any significant elevation change over the past $\sim40$ Ma. Instead, the data would be consistent with little or no post-Laramide uplift of the region. At the same time, there are only two pre-24 Ma data points that represent equilibrium conditions, so a more conservative conclusion would be that while there could have been a significant uplift event between 50 and 30 Ma ago, the $\delta^{18}O_{\text{carb}}$ record can neither support nor exclude that possibility, so other types of data will be necessary to confirm that hypothesis.

Could Cenozoic climate change be masking an elevation change? Warmer climatic conditions would make $\epsilon_{\text{CO}_2}$-calcite smaller than the $+10.5\%$ given in Figure 3 [41], which would lower the expected $\Delta^{13}C$ below the empirically observed value. Rather than increasing the range of potential isotopic equilibrium values, this would actually reduce the mean and range further, indicating that even fewer of the pedogenic carbonates from Montana formed in isotopic equilibrium. Similarly, even if $\Delta^{13}C_{\text{atm}}$ was closer to one of the extreme ends of its geologically plausible range of $-5$ to $-6.5\%$, for any additional soils that could represent equilibrium conditions, an equal number would be lost.

Two plausible explanations exist to explain why so few of the pedogenic carbonates from Montana appear to have formed in isotopic equilibrium: (1) significant evaporative enrichment and water stress or (2) diagenetic alteration. While quantitative paleoclimatic reconstructions do not exist for the entirety of the record, for at least the past 40 Ma, southwestern Montana has fluctuated within a relative narrow range of subhumid to semiarid conditions [37], and prior to the Miocene, warm-wet indicator plants such as palms and gingkers are frequently recorded by phytolith assemblages [38, 42], with drier-adapted vegetation such as grasses emerging to dominance from the Miocene onward. Thus, the part of the record in which isotopic disequilibrium due to water stress is most plausible as an explanation is actually the part of the record where the pedogenic carbonates instead reflect isotopic equilibrium most frequently, so this
indicated by text for discussion), the white circles would also represent equilibrium conditions. However, in both cases, little or no change in elevation is likely observed shifts in and crustal diagenetic water sources. In that sense, the large higher \( \Delta^{13}C \) noted that in some cases, pedogenic carbonates had source water, which has subsequently been identified in other modern studies as well (reviewed by Quade et al. [15]). Breecker et al. [1] used observations of instrumented modern soils in New Mexico to argue for a spring to warm-season bias in pedogenic carbonate formation, and Passey et al. [44] used clumped isotopes to confirm a warm season bias for soils from East Africa. Quade et al. [15] proposed that this observation of warm-season bias reflected summertime ground heating above air temperatures. However, Tabor et al. [6] found that under some conditions, pedogenic carbonate stable oxygen isotope values did not reflect a warm season bias and that the season of moisture availability was more important than the time of the highest temperatures. Gallagher et al. [5] recently extended this approach using a combination of clumped isotope data and hydrological cycle modeling to show that pedogenic carbonate isotopic compositions could reflect anything from sub-MAT to warm seasonal bias, depending on the seasonality of the moisture regime and amount of vegetative covering, which in turn impacts ground heating. In none of these studies though, was the extreme spread observed in \( \delta^{18}O_{\text{carb}} \) values from the Montana record (Figure 1), so as discussed above, extensive diagenetic alteration is the only plausible explanation. This may be true of many pedogenic carbonate records that have been used for paleoaltimetry purposes and where extremely wide spreads in \( \delta^{18}O_{\text{carb}} \) values are observed. Many of those studies could be usefully reappraised using \( \Delta^{13}C \) to screen for carbonates that formed in isotopic equilibrium.

Based upon the results in this study and other recent studies, it is appropriate to propose revised guidelines for future paleoaltimetry studies that include collection of both vertical stratigraphic sections and lateral transects [23], description and classification of paleosol type [4, 45], collection of at least three nodules per horizon and petrographic screening of nodules for micrite rather than spar [26], collection of at least one independent proxy for paleoclimate or paleovegetation [46], and screening \( \delta^{18}O_{\text{carb}} \) values using \( \Delta^{13}C \) measured for each paleosol (this study). While this approach will necessitate significantly more work than some previous studies have undertaken, it will minimize uncertainty in paleoaltimetry reconstructions.

**Figure 9:** Montana paleoelevation revisited. (a) Plot of \( \delta^{18}O_{\text{carb}} \) versus \( \Delta^{13}C \) for the past 75 Ma in southwestern Montana (data from [22]). Based upon the theoretical and empirical constraints from Figures 3 and 5, the data set has been literally decimated, with only 29 of 308 (9.4%) pedogenic carbonates were formed in isotopic equilibrium and most of the data are extremely out of equilibrium (note the scale change versus Figure 8). (b) Plot of the remaining pedogenic carbonates that formed in isotopic equilibrium (black circles). Almost all of the structure in the dataset present in Figure 1 is gone. If one allows for evaporative due to water stress on the plants producing the organic matter of +1.5‰ (see text for discussion), the white circles would also represent equilibrium conditions. However, in both cases, little or no change in elevation is indicated by \( \delta^{18}O_{\text{carb}} \) values, and there are essentially no pre-Miocene data points that can be used to assess the previously identified elevation change between 50 and 30 Ma ago identified by Chamberlain et al. [22].
and make it possible to explain both carbonates formed in equilibrium and those that are out of equilibrium mechanistically.

5. Conclusions

Many studies that attempt to reconstruct paleoaltitudes either to understand paleotectonic or paleo-geodynamic changes in the geologic record rely on pedogenic carbonates. However, many of those carbonate records have δ18O_{carb} variability that exceeds reasonable elevation changes or which would conflict with independent lines of evidence. Herein, I have proposed that carbon isotope equilibrium as defined by Δ¹³C can be a useful screening tool for δ¹⁸O_{carb} values. Case studies from the Miocene and Eocene where there are independent constraints on the paleoenvironment were used to establish evidence of both equilibrium formation and disequilibrium, and Jurassic paleosols were used to establish that pedogenic carbonates in the geologic record should reflect their meteoric source water just as they do in modern soils. Using the new screening tool, a long-term record of δ¹⁸O_{carb} values that has been used to reconstruct paleoolevation and paleotectonic changes in the northern Rocky Mountains of the US was reassessed and >90% of the published isotopic analyses do not pass the isotopic equilibrium test. For the samples that do, no significant change in elevation is indicated for the past 24 Ma. With virtually all of the pre-32 Ma showing significant alteration, it may be that the signal of uplift is best defined by looking for isotopic disequilibrium. However, this precludes the use of δ¹⁸O_{carb} values for quantitative paleoolevation reconstruction. Other previously published records likely need a similar reassessment. Finally, protocols are suggested for future paleoaltimetry studies to overcome many of the challenges that are outlined herein. Future work will endeavor to understand with more clarity the broad departures in δ¹⁸O_{carb} variability in the paleosol record as compared to modern systems and to evaluate its potential utility as a chemical proxy of burial environment.

Data Availability

New data discussed in the paper are available as part of the Supplementary Materials.

Conflicts of Interest

The author declares that there is no conflict of interest regarding the publication of this paper.

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Supplementary Materials

Supplementary materials consist of two supplemental data tables. (Supplementary Materials)

References


