Single-species dinoflagellate cyst carbon isotope fractionation in coretop sediments: environmental controls, CO₂-dependency and proxy potential

5 Joost Frieling^{1‡}, Linda van Roij¹, Iris Kleij¹, Gert-Jan Reichart^{1,2} & Appy Sluijs¹

1. Department of Earth Sciences, Faculty of Geosciences, Utrecht University, Princetonlaan 8, 3584CB Utrecht, The Netherlands

2. NIOZ Royal Netherlands Institute for Sea Research and Utrecht University, Texel, The Netherlands

‡ now at: Department of Earth Sciences, University of Oxford, South Parks Road, Oxford, OX1 3AN, Oxford, United Kingdom

10 Correspondence to: J. Frieling (joost.frieling@earth.ox.ac.uk)

Abstract. Sedimentary bulk organic matter and various molecular organic components exhibit strong CO₂-dependent carbon isotope fractionation relative to dissolved inorganic carbon sources. This fractionation (ε_p) has been employed as proxy for paleo-*p*CO₂. Yet, culture experiments indicate that CO₂-dependent ε_p is highly specific at genus and even species level, potentially hampering the use of bulk organic matter and non-species-specific organic compounds. In recent years, significant

- 15 progress has been made towards a CO₂ proxy using controlled growth experiments with dinoflagellate species, also showing highly species-specific ε_p values. These values were, however, based on motile specimens and it remains unknown whether these relations also hold for the organic-walled resting cysts (dinocysts) produced by these dinoflagellate species in their natural environment. We here analyze dinocysts isolated from core-tops from the Atlantic Ocean and Mediterranean Sea, representing several species (*Spiniferites elongatus*, *S. (cf.) ramosus*, *S. mirabilis*, *Operculodinium centrocarpum* sensu Wall
- 20 & Dale (1966) (hereafter referred to as *O. centrocarpum*) and *Impagidinium aculeatum*) using Laser ablation nano Combustion – Gas Chromatography – Isotope Ratio Mass Spectrometry (LA/nC/GC-IRMS). We find that the dinocysts produced in the natural environment are all appreciably more ¹³C-depleted compared to the cultured motile dinoflagellate cells, implying higher overall ε_p values and, moreover, exhibit large isotope variability. Where several species could be analysed from a single location, we often record significant differences in isotopic variance and offsets in mean δ^{13} C values between
- 25 species, highlighting the importance of single-species carbon isotope analyses. The most geographically expanded dataset, based on *O. centrocarpum*, shows that $ε_p$ correlates significantly with various environmental parameters. Importantly, *O. centrocarpum* shows a CO₂-dependent $ε_p$ above ~240 µatm *p*CO₂. Similar to other marine autotrophs, relative insensitivity at low *p*CO₂ is in line with active carbon concentrating mechanisms at low *p*CO₂, although we here cannot fully exclude that we partly underestimated $ε_p$ sensitivity at low *p*CO₂ values due to the relatively sparse sampling in that range. Finally, we use the
- 30 relation between ε_p and pCO₂ in O. centrocarpum to propose a first pCO₂ proxy based on a single dinocyst species.

1 Introduction

Stable carbon isotope fractionation in marine autotrophs is governed for a large part by the carbon fixing enzyme RubisCO (e.g. Farquhar et al., 1989; Roeske and O'Leary, 1984), which implies most marine organic matter and therefore sedimentary marine organic matter is strongly ¹³C-depleted with respect to the dissolved inorganic carbon (DIC) source ($CO_2(aq)$, HCO_3^-

- or CO_3^{2-}), with the stable carbon isotope composition ($\delta^{13}C$) of organic matter ranging from -10 to -30‰ (Freeman and Hayes, 1992). While many groups of marine autotrophs show clear CO_2 -dependent carbon isotope fractionation (ϵ_p), the exact relation strongly varies between marine phytoplankton groups, genera and cell morphologies (Popp et al., 1998; Boller et al., 2011, 2015; Brandenburg et al., 2022). Still, because of the assumed CO_2 -dependency of RubisCO fractionation, bulk marine organic matter and more specific organic compounds of marine autotrophs (e.g. lipids biomarkers) have been proposed and applied as
- 40 pCO_2 proxies over the past decades (Freeman and Hayes, 1992; Naafs et al., 2016). The application of these pCO_2 proxies (e.g. Bijl et al., 2010; Pagani et al., 2011; Schoon et al., 2011; Witkowski et al., 2018) has provided constraints on past atmospheric pCO_2 and earth system sensitivity beyond the ice core record (e.g. Pagani et al., 2010; PALAEOSENS, 2012). However, many of the organic compounds used for CO₂ reconstructions such as alkenones (e.g. Pagani, 2013), phytane (e.g.
- Witkowski et al., 2018), porphyrins (e.g. Freeman and Hayes, 1992) or bulk organic matter (e.g. Hayes et al., 1999) are not unique to a single species, genus and sometimes not even a group of organisms. This implies that reconstructions based on these compounds integrates interspecific differences in CO₂-dependency, which complicates the interpretation of such proxy records. Secondly, even if specific compounds derive from a single species or genus, they intrinsically derive from a multitude of individual organisms, differing in shape and size, affecting isotopic fractionation and hence limiting the accuracy of such CO₂ reconstructions.
- 50 Part of the uncertainties and biases in carbon isotope fractionation can be circumvented if the carbon isotopic fractionation of individual fossils can be analyzed. In recent years, significant progress has been made towards a CO₂ proxy based on the stable carbon isotope fractionation in organic walled dinoflagellate cysts (Burkhardt et al., 1999; Hoins et al., 2015, 2016a, 2016b; Wilkes et al., 2017). A fraction (~15%) of modern dinoflagellates produces an organic resting cyst or dinocyst as an obligatory part of their lifecycle (Evitt, 1985). The organic resting cysts from autotrophic species have excellent preservation potential,
- are often highly oxidation-resistant (Zonneveld et al., 1997, 2019; Kodrans-Nsiah et al., 2008) and several ubiquitous extant genera and species, such as *Spiniferites* spp. and *Operculodinium centrocarpum*, have extremely long geological records (Fensome et al., 1996; Williams et al., 2004). The ecology and morphology of these long-ranging species seemingly remained unchanged for millions of years (Frieling and Sluijs, 2018). Most importantly, recent advances in methodology allow for analyses of species-specific single-cyst δ^{13} C (van Roij et al., 2017; Sluijs et al., 2018). This presents the opportunity to quantify
- 60 environmental controls on ε_p of individual dinoflagellate cysts and hence species to assess the potential to obtain more accurate paleo-*p*CO₂ estimates from sedimentary records.

Controlled growth experiments across a range of CO_2 levels representative for the last glacial (e.g. Barnola et al., 1987), modern and future carbon emission scenarios (Eberlein et al., 2016; Hoins et al., 2016a, 2016b, 2015; IPCC, 2014; Rost et al.,

2006; Van de Waal et al., 2013; Wilkes et al., 2017) showed species-specific CO₂-dependent ε_p for multiple dinoflagellate

- 65 species. From these, the species *Protoceratium reticulatum* and *Gonyaulax spinifera* (Hoins et al., 2015, 2016a, 2016b) are of particular interest as these produce the organic cyst species *Operculodinium centrocarpum* sensu Wall and Dale, (1966), hereafter referred to as *O. centrocarpum*, and *Spiniferites (cf.) ramosus*, hereafter referred to as *S. ramosus*, respectively (Head, 1996; Zonneveld et al., 2013). These cyst species have their first occurrences in the geological records around ~60 and 130 million years ago (Ma), for *O. centrocarpum* and *S. ramosus*, respectively (Williams et al., 2004), thus providing potential for
- 70 deep-time pCO₂ reconstructions.

Before ε_p values based on dinocysts can be used for reconstructing *p*CO₂, several fundamental questions need to be addressed. Although $\delta^{13}C_{DIC}$ exerts a major control on dinocyst $\delta^{13}C$ (Sluijs et al., 2018), it remains uncertain whether the CO₂ control on ε_p of motile cells from controlled growth experiments can be translated to their cysts formed in the natural environment. In addition, potential offsets in ε_p values between the motile cells and the cysts need to be established. This is especially important

as the cell-cyst relations in carbon isotope ratios are not necessarily straightforward, because bulk biomass such as cysts potentially deviates in δ¹³C values from the various cell components and potentially not by a constant offset (e.g. Freeman, 2001; Hayes, 2001; Pancost and Pagani, 2006; Schouten et al., 1998; Van de Waal et al., 2013; Wilkes et al., 2018).
We here present the first core-top data for single-species dinocysts to constrain the environmental controls on ε_p. We focus on

the species O. centrocarpum and compare this data, when possible, to several species of Spiniferites (S. ramosus, S. mirabilis,

80 *S. elongatus*) and *Impagidinium aculeatum*. The established environmental relations are subsequently evaluated using simple models converting carbon isotope fractionation in dinocysts into pCO_2 values for the surface waters.

2.1 Materials

- The primary dataset is based on 34 core-top samples (Table 1, Fig. 1), collected from the North Atlantic Ocean and Mediterranean Sea, with a secondary dataset based on pre-industrial sample material spanning $\sim 0 - 1500$ common era (CE) from ENAM9606 the North Atlantic (Richter et al., 2009). The core-top samples encompass a substantial natural *p*CO₂ (aq) gradient because the rate of cooling of the North Atlantic Current exceeds that of CO₂ uptake, whereas *p*CO₂ in the Mediterranean is close to or slightly above equilibrium with the atmosphere. Sample selection is further based on the dinocyst occurrence maps of Zonneveld et al. (2013), including only samples with an expected relative abundance of at least 10-20%
- 90 of the target species. Similarly, the coverage of environmental parameters such as sea surface temperature (SST) and pCO_2 and difference in environmental settings was maximized during sample selection. Existing ocean databases are used for obtaining the relevant environmental parameters (Table 1).

2.2 Methods

95

Using standard palynological techniques (see e.g. Brinkhuis et al., 2003), ca. 5–10 g freeze-dried sediment of the upper 1–2 cm of core material was processed for each sample. This involved dissolving carbonates and silicate components

using strong acids (HCl, 30% and HF, 38–40%). After acid steps, residues were pH-neutralized and sieved using an ultrasonic bath and 250 and 10µm nylon mesh sieve to remove large and small particles, respectively. Subsequently, samples were transferred to glass test tubes with demineralized water and centrifuged at 3200 rpm for 10 minutes to obtain an optimum concentrate of the sample material. Prior to dinocyst selection, samples were stored in 4 mL glass vials in demineralized water.

100

A micro-manipulator consisting of a Leica inverted microscope and a Narishige IM-9B microinjector connected to a strung-out pipette was used to manually select individual dinocysts from a water droplet on a glass petri dish. Dinocyst selection followed a strict protocol, in which cyst morphology (primarily cyst shape and process length) was kept constant and contribution of other organic particles minimized. Specimens with darker coloration or amorphous organic matter adhered to

- 105 the cyst or processes were avoided. In the case of *O. centrocarpum*, the morphological selection primarily involved selecting specimens of equal size and process length to avoid cysts that may be derived from different environments (e.g. Mertens et al., 2009). For *Spiniferites*, we were able to distinguish and separate three distinct morphological species in sufficient numbers; *S. ramosus*, *S. elongatus* and *S. mirabilis*. For all dinocyst species, the selected diameter excluding processes was in the order of ~30–40 µm, except for *S. mirabilis* (~60 µm), although constraining the exact size of each individual specimen was not feasible
- 110 within the current analytical procedures. Stable carbon isotope analyses for individual samples are based on replicating the analyses of single of dinocysts, with ~30 individual measurements being conducted to obtain a reasonably precise (~0.3–0.4‰) sample average (van Roij et al., 2017). Given the size of the dinocysts used here (~30–40 µm cyst diameter), 3–7 specimens were required for each measurement and hence ~150 cysts were required to obtain sample averages (Table 1).
- Dinocysts were placed on a 6 mm Ø nickel sample tray, after which an identical second tray is added on top and
 compressed to fixate the dinocysts. Before placement in the ablation chamber, approximately ~1 mm² of International Atomic Energy Agency CH-7 (IAEA-CH7) polyethylene standard (PE; certified δ¹³C value -32.15‰ ± 0.05‰; 1σ) was added to the sample tray. Stable carbon isotope analyses of the dinocysts followed the procedures described in previous work (van Roij et al., 2017; Sluijs et al., 2018), utilizing the recently developed Laser Ablation nano Combustion Gas Chromatography Isotope Ratio Mass Spectrometry (LA-nC-GC/IRMS) method. Fragments resulting from deep ultraviolet LA were carried
 using a continuous Helium flow in 0.32 mm capillaries and oxidized in a combustion oven at 940 °C. The resultant CO₂ was
- transported to a GC combustion interface, dried in a nafion tube using a He counterflow and subsequently into a ThermoFisher DeltaV Advantage IRMS for isotope analysis. Each analytical run included 5 standards with signal intensity above 4 Vs (ca. 40 ng C; δ^{13} C precision better than 0.5‰) to allow calibrating to the Vienna Peedee Belemnite (VPDB) scale. Direct visual monitoring of the ablation process was used as initial quality assessment of each individual measurement.
- 125 To calculate the fractionation factor ε_p of the dinocysts relative to dissolved inorganic carbon (DIC) from which the dinocyst was produced, we take the $\delta^{13}C_{DIC}$ from the modeled grid of Tagliabue and Bopp (2008). As many dinoflagellate species, including those that produce *O. centrocarpum* and *S. ramosus* cysts, are able to utilize both HCO₃⁻, which makes up the majority of DIC, and CO₂ for carbon fixation (Hoins et al., 2016a), we also compare the $\delta^{13}C_{DINO}$ data to $\delta^{13}C_{CO2}$ and with overall sea water carbon partitioning.

- 130 ϵ_{p-DIC} is calculated as: $\delta^{13}C_{DIC} \delta^{13}C_{DINO}$ and ϵ_{p-CO2} is calculated as $\delta^{13}C_{CO2} \delta^{13}C_{DINO}$. For the latter the $\delta^{13}C$ of dissolved CO₂ is calculated from $\delta^{13}C_{DIC}$ using the temperature-dependent fractionation between DIC and CO_{2(aq)} (Mook et al., 1974). To evaluate the dominant contributions to ¹³C-fractionation in dinocysts, we compare the ϵ_{p-DIC} and ϵ_{p-CO2} values to measured and interpolated physicochemical parameters. We test a suite of parameters, [NO₃⁻], [PO₄³⁻], [Si], alkalinity, *p*CO₂, SST and SSS, which are extracted using Ocean Data View (https://odv.awi.de/) from existing (gridded) datasets (Gouretski
- 135 and Koltermann, 2004; Takahashi et al., 2014, 2016) (Supplementary Data File). Where possible, data are averaged over a grid 4° longitude and latitude around the sample position. This is both to reduce errors introduced by data scarcity and to account for potential lateral transport of dinocysts during sinking (Nooteboom et al., 2019). Carbonate chemistry is calculated using the R-package seacarb (Gattuso et al., 2019), with alkalinity and pCO_2 as input variables to calculate the other relevant parameters of the carbonate system: the relative contributions of $CO_2(aq)$, HCO_3^- and CO_3^{2-} , i.e. carbon speciation.
- Ideally, all environmental parameters would be known for the different locations, as well as the time the dinoflagellates lived and encysted. This is, however, unfeasible because the dinocysts assemblage in core-top sediments (typically the upper 2 cm of sediment) integrates conditions over several centuries, assuming moderate to low average sediment accumulation rates (< 10 cm kyr⁻¹) that generally characterise open ocean settings such as examined here. We therefore apply a rough correction for pCO_2 , based on the assumption that local air-sea gas exchange has remained similar, that equals the
- 145 atmospheric pCO_2 rise between the sampling date and 1850 CE. The correction from actual measurements to 'pre-industrial' conditions for pCO_2 yields a substantial offset due to the ~90 ppmv atmospheric pCO_2 rise from 1850 CE to the average sampling date (*ca.* 2000 CE). As this correction is broadly similar for all sample localities, it has only a small impact on the overall pattern in the CO₂ data (Fig. 3). We employed a Monte Carlo simulation to assess the potential impact of the pCO_2 correction by propagating (1) the 5% analytical error on pCO_2 values and add +45 ± 15 ppm to reflect a normally distributed
- 150 mixture of modern and pre-industrial conditions and (2) a resampled uncertainty derived from the pCO_2 rise since pre-industrial times (1800 – 2000 CE). Both scenarios are set-up to represent worst-case scenarios; a single error drawn from the error distribution is imposed on a sample basis and not a resampled average of the number of $\delta^{13}C_{\text{DINO}}$ measurements within a sample, as that would reduce the error through averaging.
- Changes in SST, SSS and nutrient concentrations are also expected, partly also by anthropogenic activity, but offsets in these parameters are generally subtle and more local compared to the changes in pCO_2 and hence would require site-specific reconstructions. Still, recent wide-spread eutrophication and enhanced productivity may impact the carbon isotope results through increased DIC uptake in algal blooms (i.e. counteracting the impact of enhanced pCO_2). However, as eutrophication mainly affected coastal areas (Hallegraeff, 1993; Anderson et al., 2002), this is expected to play a minor factor at our, mostly open marine, sample localities (Fig. 1). Lastly, long-term natural changes in nutrients, SSS and SST also occur, and it is
- 160 currently not possible to fully filter out the various anthropogenic offsets. With the exception of pCO_2 , we hence assume all parameters (SST, SSS, nutrients) to have remained constant over the period the core top samples represent.

3. Results

3.1 Carbon yields from dinocyst analyses

- 165 Despite our pre-screening to include only samples with high relative abundances of the target species, some of the selected samples contained too few dinocysts or in too low abundance relative to other organic particles (amorphous organic matter, plant debris, pollen, non-dinocyst marine palynomorphs etc.), to be used in our study. Ultimately, out of the initial core-top sample set of 34 samples, 19 were found suitable for species specific dinocyst stable carbon isotope analyses (Table 1).
- Typically, ~150 individual cysts were picked and analyzed for a total of 20–50 measurements, amounting to 3–7 cysts per carbon isotope measurement. We calculate an average signal size of 0.2 Vs for a single cyst, which amounts to a carbon yield of ~6–7 ng C cyst⁻¹ (van Roij et al., 2017). Although the variability in signal intensity from individual measurements suggests there is substantial intra-sample (cyst-cyst) variability, no significant offsets in average carbon content per cyst were observed between samples, suggesting the average carbon content of the cysts within each of the analyzed populations is similar. *Spiniferites mirabilis* is the notable exception to this rule, as far fewer specimens of this species are needed for a single $\delta^{13}C_{cyst}$
- 175 measurement. Based on the signal intensity per specimen we estimate that this larger cyst species contains twice the amount of C compared to *S. ramosus*, *S. elongatus* and *O. centrocarpum*.

3.2 Carbon isotope data

3.2.1 Signal Intensity

- The 949 individual core-top analyses range in δ^{13} C from ~ -18.5 ‰ to -35.5 ‰ (all δ^{13} C are relative to Vienna Peedee Belemnite (VPDB)), and the 137 down-core analyses range from ~ -19.4 ‰ to -28.1 ‰. No relation was observed between δ^{13} C and signal size (Vs), except at the very low end (≤ 0.2 Vs) (Fig. 2), in line with earlier analyses (van Roij et al., 2017). In this low range, the median of the δ^{13} C values rises from -28 ‰ below 0.1 Vs to values between -22 and -25 ‰ above 0.2 Vs. In the ≤ 0.2 Vs range the δ^{13} C average of both the cysts and PE converge between -25 ‰ and -30 ‰, with substantial scatter. Poorer
- performance at such low C masses and signal intensities is expected, as these extremely small signal sizes and poor signal to noise ratio (below ~3:1) approach the limit of our method. Consequently, even a very minor contamination source would bias values and result in larger scatter, as also apparent in the PE standard at a similar signal intensity (van Roij et al., 2017; Fig. 2). Due to a worsening signal-noise ratio, we find a noticeable degree of δ¹³C biasing from a background C source within the system is likely to occur at signal intensities ≤0.5 Vs and are particularly pronounced ≤ 0.2 Vs, and values for the standard and
- 190 dinocysts converge around -27 ‰ in this range (Fig. 2A). A similar background δ^{13} C value was also obtained after liquid nitrogen trapping (van Roij et al., 2017). The source of this C remains elusive. It is unlikely to be related to the ablation (etching) of the nickel plate or associated with the water used to pick sample from, as measured blanks for those result in much lower signal intensities and neither source would affect the measurements of the PE standard. Lastly, a significant contribution of atmospheric CO₂ (δ^{13} C around -8‰) appears unlikely due to the δ^{13} C signature of the background signal (-27 ‰). Though
- 195 the origin of the background C remains unknown, we can use the trapping experiment to estimate the relevant background contribution (van Roij et al., 2017). We calculate the typical contribution is likely between 0.024 and 0.08 Vs, given a

background C flux of 0.0008 Vs per second (van Roij et al., 2017) and typical duration of measurements (30–50 s for $\delta^{13}C_{DINO}$ and up to ~100 s for PE standard).

Before comparing our data with environmental variables, we therefore assess the impact of a very minor, but consistent,

- 200 background contamination on the carbon isotope signal at low signal intensities (e.g. Fig. 2A). We find that a constant addition of *ca.* 0.04 Vs ($\leq 1 \text{ ng C}$) of a background C source with a δ^{13} C of -27 ‰ can explain the positive skewing in the standard PE δ^{13} C. Using a simple isotope endmember / mass balance mixing model to correct for skewing (Fig. 2B), we calculated an average deviation from the measured PE and dinocyst values for intensities below 0.2 Vs in the order of |2.6 ‰| and |1.3 ‰|, respectively. The standard deviation of the data increases approximately 3-fold (Fig. 2B) compared to the raw measurement
- 205 data below 0.2 Vs, but remains virtually unchanged above 0.2 Vs and the calculated deviation from the measured value is also much reduced above 0.2 Vs (|<0.3 ‰|).</p>

The data correction using our simple mixing model eliminates the skew towards -27 ‰ at low signal intensities, and removes signal size δ^{13} C-dependency below 0.2 Vs for both the isotopically homogeneous PE and the heterogeneous dinocyst data (Fig. 2A, B). This suggests our method of bias correction is warranted, but the increased variability at very low intensities and lack

of independent control on the exact size and δ^{13} C of the background contamination implies the data associated with the lowest signal intensities remain significantly less reliable. We therefore apply a conservative cut-off, and use only corrected data with a signal size above 0.2 Vs.

3.2.2 Skewed distributions and outlier omission

- The drift-corrected δ¹³C_{DINO} is non-normally distributed in many core-top samples and also in different species (Table 1, Fig. 4). Distributions tend to be tailed towards lower values, exaggerated by the presence of a small amount of outlier values. This is not due to analytical error or otherwise directly related to low signal intensity as we used a 0.2 Vs cut-off to eliminate samples with potentially unreliable signal-noise ratios (see above) and a minor correction for background C addition was sufficient to eliminate skewing at low signal intensities. The absence of such signals in the down-core samples (Supplementary Fig. 1) suggests that the outliers and skewing in the sampled core-top populations could represent a real signal.
- Based on typical deep ocean sedimentation rates in the range of centimetres per kyr, the core-top samples are expected to contain a mixed assemblage of dinocysts produced mostly within the last centuries to millennia but could also include cysts produced during the last few decades that are likely affected by anthropogenic influences. It is particularly relevant to consider because a steep δ¹³C decrease (~2‰ since 1850 CE of which >1.5‰ occurs after 1950 CE) (Francey et al., 1999; Keeling et al., 2017) accompanies the *p*CO₂ rise (>130 ppmv since 1850 CE, of which >100 ppmv after 1950 CE). So even if enhanced
- carbon isotope fractionation at higher pCO_2 (Freeman and Hayes, 1992; Hoins et al., 2015; Brandenburg et al., 2022) would not play a role, the most recent specimens are likely to be impacted by decreasing $\delta^{13}C_{DIC}$.

As age integration in the modern era may result in a mixture of cysts representing a range of environmental conditions, 230 especially with respect to CO₂ concentrations and $\delta^{13}C_{DIC}$, it is important to consider the potential age-distribution of dinocysts before comparing $\delta^{13}C_{DINO}$ and ε_p to environmental variables. In an ideal scenario, cysts produced after 1850 CE should be avoided in proxy-calibration efforts to eliminate a systematic bias towards the most recent times when atmospheric CO₂ was already elevated above pre-industrial Holocene background (~280 ppmv). Because an accurate age-correction for the Suesseffect is technically unfeasible because the age-distribution of $\delta^{13}C_{DINO}$ measurements cannot be constrained, we illustrate the

- influence of $\delta^{13}C_{\text{DINO}}$ data treatment (Fig. 3) and use Monte Carlo simulations of different error distributions to test and the potential impact of the *p*CO₂ correction. We also compared both measured *p*CO₂ and *p*CO₂ around 1850 CE (see section 2.2) to ε_p calculated using both our raw $\delta^{13}C_{\text{DINO}}$ data and the $\delta^{13}C_{\text{DINO}}$ data after drift-correction and removal of statistical outliers identified within the sample-specific single species populations. This final step of data-treatment removed positive and negative measurement outliers from the sample- and species-specific $\delta^{13}C$ population (outside ±2.5 IQR), after eliminating the extremely low-signal intensities (<0.2 Vs) and correcting for the drift induced by background C in the system.
- Altogether, out of a 949 core-top measurements, we omit 43 measurements with signals < 0.2 Vs and 24 statistical outliers, which leaves 882 individual $\delta^{13}C_{DINO}$ measurements, 560 for *O. centrocarpum*, 293 for *Spiniferites* (158 *S. ramosus*, 69 *S. elongatus* and 66 *S. mirabilis*) and 29 for *I. aculeatum* (Table 1). Most of the 67 omitted measurements have comparatively low $\delta^{13}C$ and the resulting $\delta^{13}C$ of the populations are close to statistically indistinguishable from a normal distribution
- 245 (Shapiro-Wilk p = 0.05-0.1) or representative of a normal distribution (Table 1). Although the data-treatment partly removed the negative skew on the $\delta^{13}C_{\text{DINO}}$ distribution (Table 1), the combined effects of drift-correction and outlier-removal on sample / species-mean $\delta^{13}C_{\text{DINO}}$ are generally small (Fig. 3). This is expected as drift-correction averages only ~0.25 ‰ and the negative and positive outliers represent only a small percentage of the total measurements (Table 1).
- Distinctly non-normally distributed δ^{13} C values were not previously observed in recent pollen and ancient dinocyst species analyzed with the same method (van Roij et al., 2016; Sluijs et al., 2018). The here presented down-core pre-industrial δ^{13} C_{DINO} show a similar mean, variance and data distribution to the nearby core-top samples (Supplementary Fig. 1), suggesting that, at least for these nearby localities, the analysed core-top specimens represent pre-industrial conditions. We find an influence of Suess-effect and increased *p*CO₂ impacts on the δ^{13} C_{DINO} data is the most likely factor to explain the appearance of a small number of predominantly ¹³C-depleted outliers and resulting (subtle) negative skewing of the δ^{13} C distributions (Fig. 4).
- We use the background and outlier-corrected $\delta^{13}C_{DINO}$ data and compare these data with CO₂ conditions prevalent around 1850 CE (Fig. 6) and explore the effects of age-integration by propagating different error distributions representative of the *p*CO₂ change since 1850 CE (Fig. 7B D). For practical purposes, we assume all $\delta^{13}C_{DINO}$ populations to be normally distributed for further statistical analyses. We then use the mean carbon isotope value ($\delta^{13}C_{DINO}$) and signal intensity in volt seconds (Vs) of each sample. The standard error of the mean ranges from ~0.2 to 0.7‰ and depends primarily on the number of measurements
- 260 in cases where n<30, in line with expected values based on replicate measurements of the PE standard (van Roij et al. 2017) (Fig. 5).

3.3 Environmental parameters and correlation

The range of measured $\delta^{13}C_{\text{DINO}}$ values (~5‰) far exceeds the variability in surface ocean $\delta^{13}C_{\text{DIC}}$ (~1‰) and $\delta^{13}C_{\text{CO2}}$ (~2.5‰),

265 implying the observed range likely reflects differences in fractionation related to changing uptake or leakage of different inorganic carbon phases (CO₂ and HCO₃⁻; Sharkey and Berry, 1985; Hoins et al., 2016a), and this hence determines most of the variability in the $\delta^{13}C_{\text{DINO}}$ data. Here, we quantitatively assess fractionation as a function of several environmental parameters.

The simple (non-weighted) linear regressions show poor correlations between ε_{p-DIC} and $\delta^{13}C_{DIC}$ for all environmental 270 parameters, and the correlations slightly improve when compared to ε_{p-CO2} (Table 2, 3). However, none of the tested parameters individually explain the majority of the observed variance in ε_{p-DIC} (maximum R² (~0.1) or ε_{p-CO2} (maximum R² with *p*CO₂

- (~0.38), despite high significance (low p-values) of the regressions. The explained variance increases when polynomial regressions are applied. Several controlled growth experiments indeed show a non-linear response of ε_p as a function of *p*CO₂ of the growth medium (Hoins et al., 2015) although the number of data points in such experiments limit full mathematical descriptions of the trends within the *p*CO₂ range of this field study. Here, a second-order polynomial (quadratic) regression
- achieves an R² of ~0.74 and ~0.79 for the non-weighted and weighted versions, respectively. It is conceivable that other environmental parameters also significantly contribute to ε_p variability (Fig. 6). For example, [PO₄³⁻], [NO₃], and *p*CO₂ contribute significantly to a (linear) multiple-regression model, which takes the form of $\varepsilon_{p-CO2} = c + xCO_2 + yPO_4 + zNO_3$, where c, x, y and z are numerical constants. The multiple regression model using these three parameters covers
- 280 ~58% of the variance in *O. centrocarpum* ε_{p-CO2} (not weighted), and 67% when weighted to number of measurements per sample. Including more parameters, such as SST, oxygen concentrations, or other carbonate system parameters, explains slightly more of the observed variance, but does not significantly improve the model. The residual mean standard error (RSME) of the CNP- ε_p multiple regression model is ~1.45 ‰ while a linear regression with only *p*CO₂ yields 1.7 ‰. Only weighted regressions are given here and reported ranges of the constants represent one standard error or equivalent percentiles in case
- 285 of Monte Carlo simulated errors. These models have the following optimal formats:

Equation 1 linear:

 $\epsilon_{p-CO2} = 6.6 \pm 2.1 + 0.031 \pm 0.008 \ pCO_2$

(Adjusted $R^2 = 0.48$, p = 0.001, RSME = 1.7 ‰) (Fig. 6B)

290

Equation 2a quadratic (data without error propagation and only suitable for use > 240 µatm) $\varepsilon_{p-CO2} = 40.8 \pm 7.2 - 0.23 \pm 0.055 \ pCO_2 + 4.88 \ x \ 10^{-4} \pm 1 \ x \ 10^{-4} \ pCO_2^{-2}$ (Adjusted R² = 0.79, p <0.001, RSME = 1.13 ‰) (Fig. 6B)

295 Equation 2b quadratic (Monte Carlo constrained errors – analytical for pCO_2 and ε_{p-CO2}) (Fig. 7B) $\varepsilon_{p-CO2} = 35.6^{+5.8}/_{-5.6} - 0.19^{+0.045}/_{-0.045} pCO_2 + 4.1^{+0.91}/_{-0.88} 10^{-4} pCO_2^{-2}$ Equation 2c quadratic (as 2b with additional $45 \pm 15 \text{ ppm } p\text{CO}_2 \text{ error}$) (Fig. 7C)

 $\epsilon_{p\text{-}CO2} = 39.3 \ ^{+11.5}\!/_{-8.8} - 0.19^{+\ 0.058}\!/_{-0.076} \ p\text{CO}_2 + 3.4 \ ^{+1.3}\!/_{-0.95} \ x \ 10^{-4} \ p\text{CO}_2 \ ^2$

300

Equation 2d quadratic (as 2b with resampled pCO_2 rise 1800 – 2000 CE) (Fig. 7D) $\epsilon_{p-CO2} = 29.8^{+11.0}/_{-8.0} - 0.13^{+0.061}/_{-0.084} pCO_2 + 2.6^{+1.5}/_{-1.1} \times 10^{-4} pCO_2^{-2}$

 $\varepsilon_{p-CO2} = 6.0 \pm 3.1 + 0.034 \pm 0.01 \ pCO_2 + 1.22 \pm 0.47 \ NO_3 - 10.85 \pm 3.7 \ PO_4^{3}$

Equation 3a multiple-regression linear (Fig. 6E):

(Adjusted
$$R^2 = 0.67$$
, p < 0.001, RSME = 1.45 ‰)

Equation 3b adjusted for application in the paleo-domain:

 $\varepsilon_{p-CO2} = 6.0 \pm 3.1 + 0.034 \pm 0.01 \ pCO_2 - 1.1 \pm 5.3 \ PO_4^{3-}$

310

Equation 4 multiple-regression linear:

 $\varepsilon_{p-DIC} = 18.4 \pm 3.1 + 0.025 \pm 0.01 \ pCO_2 + 1.45 \pm 0.47 \ NO_3 - 11.1 \pm 3.7 \ PO_4^{3-}$ (Adjusted R² = 0.52, p = 0.01, RSME = 1.44 ‰)

- The two linear multiple regression models are offset (Equations 3a and 4), primarily because of the carbon isotope fractionation between HCO₃⁻ and CO₂. The slope with respect to *p*CO₂ also varies slightly between the models for ε_{p-DIC} and ε_{p-CO2} due to the temperature dependent fractionation between HCO₃⁻ and CO₂, but the slopes with NO₃⁻ and PO₄³⁻ are indistinguishable. The quadratic regression seemingly better fits the variability observed in ε_{p-CO2} compared to other (multiple) linear regressions and removes any structure in the residuals, potentially signaling a non-linear response in ε_{p-CO2} to *p*CO₂. The quadratic regression also indicates insensitivity to *p*CO₂ ≤240 µatm and should not be used below this value (Fig. 6B). The Monte Carlo simulations of scenarios where an additional *p*CO₂ uncertainty is imposed as a normally distributed mixture of pre-industrial values and modern, offsetting *p*CO₂ by +45 ± 15 ppm (Fig. 7C, Eq. 2c), and a resampled uncertainty derived from the *p*CO₂
- rise since pre-industrial times (Fig. 7D, Eq. 2d) show that the parameters of the quadratic regression are fairly robust to these uncertainties (*i.e.* none of the parameters become insignificant at p > 0.05), although the absolute pCO_2 values and errors increase.

4. Discussion

4.1 Absolute values, comparison to marine organic matter

The recorded $\delta^{13}C_{DINO}$ range and absolute values (~ -18‰ to -35‰) correspond well with global $\delta^{13}C$ values previously

- 330 reported for marine particulate organic matter ($\delta^{13}C_{POC}$) (e.g. Freeman and Hayes, 1992; Goericke and Fry, 1994) and modelled phytoplankton biomass (e.g. Magozzi et al., 2017; Tagliabue and Bopp, 2008). Consequently, ε_{p-DIC} and ε_{p-CO2} are also within the expected range for general marine particulate organic matter. However, the intra-sample variance of $\delta^{13}C_{DINO}$ appears to be substantial, often spanning most of the full range (~10‰) observed for $\delta^{13}C_{POC}$. Some of the observed variability might be related to the limited analytical precision during measurements of the extremely small amounts of carbon of individual
- 335 dinocysts. Fully constraining the contribution of this analytical uncertainty to the observed variance is, however, not possible because of unresolvable micrometer-scale heterogeneity in the PE standard (van Roij et al., 2017; Sluijs et al., 2018). In most cases, the variance in $\delta^{13}C_{\text{DINO}}$ is similar to that of the standard. Still, it is likely that some of the seasonal $\delta^{13}C_{\text{DIC}}$ differences are also recorded in the $\delta^{13}C_{\text{DINO}}$, and that some additional inter-specimen $\delta^{13}C$ variance is present. This is to be expected since the $\delta^{13}C$ populations from our integrated core-top samples span seasons, decades and thus also considerable variability in
- seawater properties and population change. In addition, growth-induced randomness and changes in δ^{13} C and DIC in the cell's microenvironment likely contributed to inter-specimen variability. Note that in our data inter-specimen variability is still underestimated because we analyzed 3-7 specimens per ablation event, as single-cyst carbon yield (~7 ng C) from these cystsizes approached the limit for reliable measurements (van Roij et al., 2016). We minimized potential influence of differences in cell size or shape through manual selection. We thus analyzed a population where the pre-selection of similar-sized cysts
- 345 restricts the variance in cell surface area and volume, unlike biomarker-based proxies for which the cell size has to be reconstructed independently (Henderiks and Pagani, 2007; Stoll et al., 2019). This approach could reduce scatter in the relation of ε_p to environmental variables (Popp et al., 1998).

4.2 Cell – cyst offset

- 350 One of the striking differences between the here generated data and the existing culture experiments, is that carbon isotope fractionation of dinocysts in the natural environment appears to be much larger than that of motile cells from controlled growth (dilute batch) experiments (Hoins et al., 2015, 2016b). We find average ε_p values ranging between 13–20‰ and 23–29‰ with respect to CO₂ and DIC. Cultured cells of *O. centrocarpum* yielded not only a smaller overall ε_p , but also a smaller range (~8– 12 and 18.5–20‰) across a larger CO₂ gradient, implying the cysts have a much steeper fractionation slope with CO₂ compared
- to the motile cells. Despite these differences, the average ε_p for *Spiniferites* species (*S. ramosus, elongatus & mirabilis*) is often somewhat larger than for *O. centrocarpum* (Fig. 4). This is consistent with culture experiments that showed larger CO₂-dependency and overall slightly larger ε_p in the motile species *G. spinifera* compared to *P. reticulatum* (Hoins et al., 2015). While the cultured single strains and dinoflagellate populations in nature may behave somewhat differently, we do not expect that this alone underlies such a marked offset between the motile cultured cells and natural cysts. Natural cysts and cultured
- 360 cells seem consistently offset in δ^{13} C, although at present the exact amplitude of this offset cannot be determined. However, such an offset is in line with certain compounds in dinoflagellate cells being depleted in ¹³C relative to the bulk biomass

(Schouten et al., 1998; Wilkes et al., 2017). The organic-walled dinocysts consist of mostly aliphatic and aromatic compounds, forming a complex biopolymer referred to as dinosporin (de Leeuw et al., 2006; Versteegh et al., 2007, 2012). Depending on the biosynthetic pathway of the cyst-material and the derivation or degradation of the original compounds, this may result in

- 365 offsets in δ^{13} C values between cysts and the motile cells. A potential additional fractionation might be introduced during taphonomy and also later by the processing of sediments to concentrate the dinocysts from sediment samples. The sediment processing involves hydrochloric and hydrofluoric acids, which affects the more labile organic compounds. Last, it is conceivable that fractionation in the dilute batch experiments may be reduced by e.g., higher-than-natural growth rates. This may be supported by chemostat culture experiments on *Alexandrium tamarense* (Wilkes et al., 2017), which show a (much)
- 370 greater fractionation compared to the dilute batch experiments (Hoins et al., 2015). However, the enhanced fractionation recorded in chemostat experiments is likely an artifact of isotope equilibration times exceeding CO₂ uptake rates (Brandenburg et al., 2022; Zhang et al., 2022). The range of options cannot be narrowed down until cultured cysts are compared to their motile cells harvested from the same culture, and treated with similar techniques as used for the sediments. Until these data become available, inferences on the origin and amplitude of the offsets between the cells and cysts of *O. centrocarpum* and 375 *Spiniferites* remain speculative.

4.3 Environmental controls on carbon isotope fractionation

Carbon isotope fractionation is determined by RubisCO and several environmental parameters, dominantly *p*CO₂, but also cell size and shape, growth rates and nutrient or light regimes (e.g. Freeman and Hayes, 1992; Pagani, 2013; Popp et al., 1998;
Stoll et al., 2019 and many others). In most cases, fractionation is *p*CO₂ dependent, and consequently ε_p in various groups and genera has been used as a paleo-*p*CO₂ proxy (e.g., Freeman and Hayes, 1992; Pagani, 2013; Rae et al., 2021). We performed a broad-spectrum multiple regression analysis to identify environmental factors that contribute significantly to dinocyst ε_p. The most important parameter is *p*CO₂, in line with previous studies (Freeman and Hayes, 1992). A large part of the remaining variability can be attributed to growth rate and ultimately nutrient content, specifically nitrate and phosphate of the surface
waters, in line with previous studies on phytoplankton and dinoflagellate carbon isotope fractionation (Popp et al., 1998; Hoins et al., 2016b; Wilkes et al., 2017; Wilkes and Pearson, 2019). We find that *p*CO₂ and ε_p are positively correlated, while NO₃⁻

and PO₄³⁻ are negatively correlated with ε_p (Fig. 6C,D).

The inclusion of nutrient levels as environmental factors on the carbon isotope fractionation reduces the error in the fit between the measured and modelled fractionation. In the rare cases where paleo-nutrient concentrations are estimated, reconstructions

usually cover only $[PO_4^{3-}]$, implying the $[NO_3^{-}]$ in Eq. 3a and 4 has to be approximated from $[PO_4^{3-}]$. As $[PO_4^{3-}]$ and $[NO_3]$ are generally well-correlated (typically ~1:10 $[PO_4^{3-}]$: $[NO_3]$; here ~1:8) in ocean waters, this is relatively straightforward. For application in the paleo-domain however, nutrient ($[PO_4^{3-}]$) reconstructions may not be available or considered too unreliable to provide meaningful constraints. In specific cases where accurate paleo- $[PO_4]$ estimates or large changes are reconstructed, Eq. 3b may be applied, in which $[NO_3]$ is substituted for $[PO_4^{3-}]$ in a 1:8 ratio. However, unless there are clear changes in the 395 nutrient regime or sufficient proxy constraints on the nutrient concentrations, we prefer a calibration based exclusively on carbon isotope fractionation and carbonate system parameters, including sea surface temperature to calculate $\delta^{13}C_{CO2}$ from $\delta^{13}C_{DIC}$, to reconstruct *p*CO₂.

4.4 Influence of carbon concentrating mechanisms: CO₂ insensitivity

- 400 As many phototrophs, including dinoflagellates, have mechanisms for concentrating CO₂ near the cell membrane, the sensitivity of carbon isotope fractionation to ambient CO₂ is expected to diminish below a certain concentration (M.R. Badger, 2003, M.P.S. Badger, 2021; Stoll et al., 2019). This is particularly the case as dinoflagellates utilize type II RubisCO, which has generally poorer performance compared to type I RubisCO at low CO₂ concentrations (Giordano et al., 2005). Indeed, substantial activity of the carboxyl anhydrase (CA) enzyme, which facilitates the conversion of HCO₃⁻ to CO₂ inside the cell
- 405 or near the membrane, was observed in numerous dinoflagellate species, including *Lingulodinium* (Lapointe et al., 2008), *Symbiodinium* (Leggat et al., 1999) and the here analysed *Operculodinium* (Ratti et al., 2007). Also here we find a relatively low sensitivity at the lower end of the CO₂ scale. The lower CO₂ values correspond to the northern-most locations, with trends below 240 µatm becoming somewhat obscured, at a minimum ε_p -CO₂ around 13 ‰ (Fig. 6B). However, part of this levelling of the proxy-relationship may reflect the locally higher nutrient concentrations offsetting the higher CO₂. Though growth rates
- 410 have a clear influence on ε_p in algal groups (Burkhardt et al., 1999), including dinoflagellates (Wilkes et al., 2017), the dilute batch culturing experiments conducted with *P. reticulatum* showed no clear influence of growth rates on ε_p (see also Sec. 4.2). It is also conceivable that higher growth rates influence ε_p indirectly through, for example, seasonally enhanced CO₂ drawdown, resulting in higher δ^{13} C values in the remaining DIC. This effect may be enhanced by the relatively short growing season at the high latitudes. However, in culture experiments at low CO₂ concentrations with other dinoflagellate species, ¹³C-
- 415 fractionation was higher under nutrient limiting conditions than under replete conditions (Hoins et al., 2016b). Because of these confounding factors, the influence of carbon concentrating mechanisms on ε_p in *O. centrocarpum* is difficult to gauge with the presently available data, and would ideally be tested using high nutrient or very low CO₂ concentrations. Still, also in the relatively limited range the current ocean offers for testing *p*CO₂ proxies we have established a robust, albeit

not overly sensitive, relation between pCO_2 and dinocyst $\delta^{13}C$. Our cyst-based calibration yields more conservative and 420 arguably more realistic absolute CO₂ estimates and variability compared to available culture-based calibrations as it is based on the same compounds as will be analyzed in the paleo-domain. However, the low sensitivity at low CO₂ implies that, until better constraints become available, the proposed calibration is potentially less suitable for application across, for example, the Pleistocene glacial periods. Further, it is important to realise that the value of 240 µatm is based on the assumption that the ε_p -CO₂ – pCO_2 relation originated from cysts that have not been affected by the Suess-effect and thus represent a lower limit for

425 CO_2 (in)sensitivity. While our data does not preclude fractionation smaller than the here observed minimum (~13 ‰) during low *p*CO₂ periods, increased sensitivity at higher CO₂ suggests that CO₂ above (minimum) 240 µatm and CO₂ variability can be reconstructed with reasonable precision.

4.5 Challenges of age-control and potential caveats associated with anthropogenic carbon

- 430 A topic that warrants specific attention is the potential impact of anthropogenic carbon emissions on shaping the relation between $\delta^{13}C_{\text{DINO}}$ and ε_p . Here, we assume that our outlier analyses preferentially excluded samples that were significantly affected by anthropogenic CO₂ and that, on average, the remainder of our core-top $\delta^{13}C_{\text{DINO}}$ were not appreciably affected by either decreasing $\delta^{13}C_{\text{DIC}}$ or elevated *p*CO₂. The similarity in pre-industrial down-core $\delta^{13}C_{\text{DINO}}$ and that of three core-top localities in the North Atlantic corroborate the validity of this assumption locally but these observations cannot be extrapolated
- 435 to other regions. Unfortunately, sedimentation rates or other constraints for cyst production datums are not available. Further, if sedimentation rates were available for core-top localities, that would constitute an imperfect solution as it cannot provide the required dinocyst-specific age distribution needed to obtain an appropriate local pCO_2 correction to the datum of cyst production. This challenge may be unique to data such as presented here, as studies on other substrates with pCO_2 -proxy potential either could not generate data for individual single celled organisms or have avoided the issue through other means,
- 440 such as culture experiments (e.g. Pagani et al., 2002; Henehan et al., 2013), approaches that have other drawbacks. Although we have no constraints on the ages of the cysts analysed here, we can provide a meaningful test of the potential uncertainty added by our assumption that cysts are representative of pre-industrial conditions (Eq. 2a, b). The two scenarios that we have explored through Monte-Carlo simulations show that, depending on the error distribution imposed on the assumed *p*CO₂ used in the quadratic regressions (Eq. 2a,b), the fitted regression shifts towards higher values (Fig. 7C) and may be
- steeper (Fig. 7D). These error-propagation case studies illustrate that our proposed pre-industrial transfer function (Eq. 2a, b), if it indeed contains a substantial proportion of very recent dinocysts, is likely to lead to underestimated pCO_2 and perhaps pCO_2 variability when applied in the paleo-domain. Consequently, we recommend future studies target, for example, sediment trap collections and culture-derived dinocysts to validate the results obtained here.

450 5. Conclusions - Proxy potential, limitations and calibrations

Our new modern ocean single-species carbon isotope fractionation dataset shows promising trends with environmental variables, pCO_2 and nutrients. The selection of individual cysts allows control of cell size and species, which reduces uncertainty in proxy calibration and application compared to approaches based on organic substrates which inevitably integrate entire communities. Although this approach has clear benefits, it also poses a unique challenge as the impact of anthropogenic

455 carbon emissions on individual single celled organisms must be considered. Based on our analyses, we expect this to be a relatively minor factor. In a worst-case scenario, however, we find that, although a helpful simplification, the assumption that all dinocysts from the core-top samples represent pre-industrial conditions may lead to an underestimate of pCO_2 and perhaps also pCO_2 variability when applied as a proxy in the paleo-domain.

In addition, many of the challenges associated with other proxies based on organic substrates are encountered here as well. For 460 example, like in cultures (Hoins et al., 2016b), we observed an impact of nutrients on carbon isotope fractionation, possibly related to differences in growth rates. Similarly, at low pCO_2 values sensitivity is reduced, possibly because of carbon concentrating mechanisms involved in dinoflagellate C uptake, as observed in culturing experiments (e.g., Hoins et al., 2016a). Another remaining challenge is the observed difference between the cultured populations and cysts from the core top sediments. This is a pronounced difference, not only in the absolute isotope fractionation values but also in the slope of the

465 CO_2 sensitivity, which appears to be much larger for the cysts and requires attention in future culture studies. The offset in $\delta^{13}C$, combined with uncertainties in fractionation between the motile cells and dinocysts imply that CO_2 reconstructions using culture-based calibrations are more likely to overestimate past *p*CO₂. Furthermore, the large spread in our data (~5‰ between high and low CO₂) suggests that, due to this high sensitivity in the cysts, the method is also suited to study population dynamics.

470

Data availability

All newly generated data will be available via a permanent online repository (Mendeley data doi: 10.17632/z6285myxkm.2) upon publication.

475 Author contribution

AS & GJR designed the study, LvR, IK & JF processed samples, generated and analysed data, JF wrote the original draft, AS & GJR reviewed and edited the manuscript. AS acquired funding for this study.

Competing interests

480 The authors declare that they have no conflict of interest.

Acknowledgments

We thank A. van Dijk, M. Kienhuis and H. de Waard (Utrecht University) for technical and analytical assistance. AS acknowledges funding from Netherlands Organisation for Scientific Research (NWO) #ALWOP.223 and European Research

485 Council (ERC) Starting grant #259627. This work was carried out under the program of the Netherlands Earth System Science Centre (NESSC), financially supported by the Dutch Ministry of Education, Culture and Science. We thank two anonymous reviewers for their insightful and highly constructive comments and Steven Bouillon for editorial handling.

References

Anderson, D. M., Glibert, P. M. and Burkholder, J. M.: Harmful algal blooms and eutrophication: Nutrient sources, composition, and consequences, Estuaries, 25(4), 704–726, doi:10.1007/BF02804901, 2002.
Badger, M.: The roles of carbonic anhydrases in photosynthetic CO2 concentrating mechanisms, Photosynth. Res., 77(2–3), 83–94, doi:10.1023/A:1025821717773, 2003.

Badger, M. P. S.: Alkenone isotopes show evidence of active carbon concentrating mechanisms in coccolithophores as aqueous

495 carbon dioxide concentrations fall below 7 μmolL-1, Biogeosciences, 18(3), 1149–1160, doi:10.5194/bg-18-1149-2021, 2021.
Barnola, J. M., Raynaud, D., Korotkevich, Y. S. and Lorius, C.: Vostok ice core provides 160,000-year record of atmospheric CO2, Nature, 329(6138), 408–414, doi:10.1038/329408a0, 1987.
Bijl, P. K., Houben, A. J. P., Schouten, S., Bohaty, S. M., Sluijs, A., Reichart, G.-J., Damsté, J. S. S. and Brinkhuis, H.: Transient Middle Eocene Atmospheric CO2 and Temperature Variations, Science, 330(6005), 819–821,

500 doi:10.1126/science.1193654, 2010.

Boller, A. J., Thomas, P. J., Cavanaugh, C. M. and Scott, K. M.: Low stable carbon isotope fractionation by coccolithophore
RubisCO, Geochim. Cosmochim. Acta, 75(22), 7200–7207, doi:10.1016/j.gca.2011.08.031, 2011.
Boller, A. J., Thomas, P. J., Cavanaugh, C. M. and Scott, K. M.: Isotopic discrimination and kinetic parameters of RubisCO

from the marine bloom-forming diatom, Skeletonema costatum, Geobiology, 13(1), 33–43, doi:10.1111/gbi.12112, 2015.

- Brandenburg, K. M., Rost, B., Van De Waal, D. B., Hoins, M. and Sluijs, A.: Physiological control on carbon isotope fractionation in marine phytoplankton, Biogeosciences, 19(13), 3305–3315, doi:10.5194/bg-19-3305-2022, 2022.
 Brinkhuis, H., Sengers, S., Sluijs, A., Warnaar, J., Williams, G. L. and Exon, N. F.: Latest Creataceous-Earliest Oligocene and Quaternary Dinoflagellate Cysts, ODP Site 1172, East Tasman Plateau, Proc. Ocean Drill. Program, Sci. results, 189(October), 48, 2003.
- 510 Burkhardt, S., Riesebell, U. and Zondervan, I.: Effects of growth rate, CO 2 concentration, and cell size on the stable carbon isotope fractionation in marine phytoplankton, Geochim. Cosmochim. Acta, 63(22), 3729–3741, 1999.
 Eberlein, T., Van de Waal, D., Brandenburg, K., John, U., Voss, M., Achterberg, E. and Rost, B.: Interactive effects of ocean acidification and nitrogen limitation on two bloom-forming dinoflagellate species, Mar. Ecol. Prog. Ser., 543, 127–140, doi:10.3354/meps11568, 2016.
- 515 Evitt, W. R.: Sporopollenin dinoflagellate cysts: their morphology and interpretation, Amer Assn of Stratigraphic., 1985.
 Farquhar, G. D., Ehleringer, J. R. and Hubick, K. T.: Carbon Isotope Discrimination and Photosynthesis, Annu. Rev. Plant Physiol. Plant Mol. Biol., 40(1), 503–537, doi:10.1146/annurev.pp.40.060189.002443, 1989.
 Fensome, R. A., MacRae, R. A., Moldowan, J. M., Taylor, F. J. R. and Williams, G. L.: The early Mesozoic radiation of dinoflagellates, Paleobiology, 329–338, 1996.
- Francey, R. J., Allison, C. E., Etheridge, D. M., Trudinger, C. M., Enting, I. G., Leuenberger, M., Langenfelds, R. L., Michel,
 E. and Steele, L. P.: A 1000-year high precision record of δ13C in atmospheric CO2, Tellus, Ser. B Chem. Phys. Meteorol., 51(2), 170–193, doi:10.1034/j.1600-0889.1999.t01-1-00005.x, 1999.

Freeman, K. H.: Isotopic biogeochemistry of marine organic carbon, Rev. Mineral. Geochemistry, 43, 579–605, doi:10.2138/gsrmg.43.1.579, 2001.

525 Freeman, K. H. and Hayes, J. M.: Fractionation of carbon isotopes by phytoplankton and estimates of ancient CO2 levels, Global Biogeochem. Cycles, 6(2), 185–198, 1992.

Frieling, J. and Sluijs, A.: Towards quantitative environmental reconstructions from ancient non-analogue microfossil assemblages: Ecological preferences of Paleocene – Eocene dinoflagellates, Earth-Science Rev., 185(August), 956–973,

doi:10.1016/j.earscirev.2018.08.014, 2018.

- Gattuso, J. P., Epitalon, J. M., Lavigne, H. and Orr, J.: seacarb: Seawater carbonate chemistry. R package version 3.2. 12., n.d. Giordano, M., Beardall, J. and Raven, J. A.: CO2 Concentrating Mechanisms in Algae: Mechanisms, Environmental Modulation, and Evolution, Annu. Rev. Plant Biol., 56(1), 99–131, doi:10.1146/annurev.arplant.56.032604.144052, 2005. Goericke, R. and Fry, B.: Variations of marine plankton δ13C with latitude, temperature, and dissolved CO2 in the world ocean, Global Biogeochem. Cycles, 8(1), 85–90, doi:10.1029/93GB03272, 1994.
- 535 Gouretski, V. V. and Koltermann, K. P.: WOCE Global Hydrographic Climatology, Berichte des Bundesamtes fur Seeschifffahrt und Hydrogr., doi:10.5065/GS51-V170, 2004.
 Hallegraeff, G. M.: A review of harmful algal blooms and their apparent global increase, Phycologia, 32(2), 79–99, doi:10.2216/i0031-8884-32-2-79.1, 1993.
 Haves, J. M.: Fractionation of Carbon and Hydrogen Isotopes in Biosynthetic Processes, Rev. Mineral. Geochemistry, 43(1),

225–277, doi:10.2138/gsrmg.43.1.225, 2001.
Hayes, J. M., Strauss, H. and Kaufman, A. J.: The abundance of 13C in marine organic matter and isotopic fractionation in the global biogeochemical cycle of carbon during the past 800 Ma, Chem. Geol., 161(1), 103–125, 1999.
Head, M. J.: Modern dinoflagellate cysts and their biological affinities, Palynol. Princ. Appl., 3, 1197–1248, 1996.
Henderiks, J. and Pagani, M.: Refining ancient carbon dioxide estimates: Significance of coccolithophore cell size for

- alkenone-based pCO2 records, Paleoceanography, 22(3), doi:10.1029/2006PA001399, 2007.
 Henehan, M. J., Rae, J. W. B., Foster, G. L., Erez, J., Prentice, K. C., Kucera, M., Bostock, H. C., Martínez-Botí, M. A., Milton, J. A., Wilson, P. A., Marshall, B. J. and Elliott, T.: Calibration of the boron isotope proxy in the planktonic foraminifera Globigerinoides ruber for use in palaeo-CO2 reconstruction, Earth Planet. Sci. Lett., 364, 111–122, doi:10.1016/j.epsl.2012.12.029, 2013.
- 550 Hoins, M., Van de Waal, D. B., Eberlein, T., Reichart, G.-J., Rost, B. and Sluijs, A.: Stable carbon isotope fractionation of organic cyst-forming dinoflagellates: Evaluating the potential for a CO2 proxy, Geochim. Cosmochim. Acta, 160, 267–276, doi:10.1016/j.gca.2015.04.001, 2015.

Hoins, M., Eberlein, T., Van de Waal, D. B., Sluijs, A., Reichart, G.-J. and Rost, B.: CO2-dependent carbon isotope fractionation in dinoflagellates relates to their inorganic carbon fluxes, J. Exp. Mar. Bio. Ecol., 481, 9–14, doi:10.1016/j.jembe.2016.04.001, 2016a.

Hoins, M., Eberlein, T., Groβmann, C. H., Brandenburg, K., Reichart, G.-J., Rost, B., Sluijs, A. and Van de Waal, D. B.:
Combined Effects of Ocean Acidification and Light or Nitrogen Availabilities on 13C Fractionation in Marine Dinoflagellates,
PLoS One, 11(5), e0154370, doi:10.1371/journal.pone.0154370, 2016b.

IPCC: Climate Change 2014: Synthesis Report. Contribution of Working Groups I, II and III to the Fifth Assessment Report 560 of the Intergovernmental Panel on Climate Change, 2014.

Keeling, R. F., Graven, H. D., Welp, L. R., Resplandy, L., Bi, J., Piper, S. C., Sun, Y., Bollenbacher, A. and Meijer, H. A. J.: Atmospheric evidence for a global secular increase in carbon isotopic discrimination of land photosynthesis, Proc. Natl. Acad.

Sci. U. S. A., 114(39), 10361–10366, doi:10.1073/pnas.1619240114, 2017.

Kodrans-Nsiah, M., De Lange, G. J. and Zonneveld, K. A. F.: A natural exposure experiment on short-term species-selective aerobic degradation of dinoflagellate cysts, Rev. Palaeobot. Palynol., 152(1), 32–39, 2008.

- Lapointe, M., MacKenzie, T. D. B. and Morse, D.: An external δ-carbonic anhydrase in a free-living marine dinoflagellate may circumvent diffusion-limited carbon acquisition, Plant Physiol., 147(3), 1427–1436, doi:10.1104/pp.108.117077, 2008. de Leeuw, J. W., Versteegh, G. J. M. and van Bergen, P. F.: Biomacromolecules of Algae and Plants and their Fossil Analogues, Plant Ecol., 182(1–2), 209–233, doi:10.1007/s11258-005-9027-x, 2006.
- 570 Leggat, W., Badger, M. R. and Yellowlees, D.: Evidence for an inorganic carbon-concentrating mechanism in the symbiotic dinoflagellate Symbiodinium sp., Plant Physiol., 121(4), 1247–1255, doi:10.1104/pp.121.4.1247, 1999.
 Magozzi, S., Yool, A., Vander Zanden, H. B., Wunder, M. B. and Trueman, C. N.: Using ocean models to predict spatial and temporal variation in marine carbon isotopes, Ecosphere, 8(5), e01763, doi:10.1002/ecs2.1763, 2017.
 Mertens, K. N., Ribeiro, S., Bouimetarhan, I., Caner, H., Combourieu Nebout, N., Dale, B., De Vernal, A., Ellegaard, M.,
- 575 Filipova, M., Godhe, A., Goubert, E., Grøsfjeld, K., Holzwarth, U., Kotthoff, U., Leroy, S. A. G., Londeix, L., Marret, F., Matsuoka, K., Mudie, P. J., Naudts, L., Peña-Manjarrez, J. L., Persson, A., Popescu, S. M., Pospelova, V., Sangiorgi, F., van der Meer, M. T. J., Vink, A., Zonneveld, K. A. F., Vercauteren, D., Vlassenbroeck, J., Louwye, S., Nebout, N. C., Dale, B., De Vernal, A., Ellegaard, M., Filipova, M. and Godhe, A.: Process length variation in cysts of a dinoflagellate, Lingulodinium machaerophorum, in surface sediments: investigating its potential as salinity proxy, Mar. Micropaleontol., 70(1), 54–69,
- 580 doi:10.1016/j.marmicro.2008.10.004, 2009.

565

Mook, W. G., Bommerson, J. C. and Staverman, W. H.: Carbon isotope fractionation between dissolved bicarbonate and gaseous carbon dioxide, Earth Planet. Sci. Lett., 22(2), 169–176, doi:10.1016/0012-821X(74)90078-8, 1974.

Naafs, B. D. A., Castro, J. M., De Gea, G. A., Quijano, M. L., Schmidt, D. N. and Pancost, R. D.: Gradual and sustained carbon dioxide release during Aptian Oceanic Anoxic Event 1a, Nat. Geosci., 9(2), 135–139, doi:10.1038/ngeo2627, 2016.

585 Nooteboom, P. D., Bijl, P. K., van Sebille, E., von der Heydt, A. S. and Dijkstra, H. A.: Transport bias by ocean currents in sedimentary microplankton assemblages: Implications for paleoceanographic reconstructions, Paleoceanogr. Paleoclimatology, doi:10.1029/2019PA003606, 2019.

Pagani, M.: Biomarker-Based Inferences of Past Climate: The Alkenone pCO2 Proxy, 2nd ed., Elsevier Ltd., 2013. Pagani, M., Freeman, K. H., Ohkouchi, N. and Caldeira, K.: Comparison of water column [CO_{2aq}] with sedimentary alkenone-

based estimates: A test of the alkenone-CO₂ proxy, Paleoceanography, 17(4), 21-1-21–12, doi:10.1029/2002PA000756, 2002.
 Pagani, M., Liu, Z., LaRiviere, J. and Ravelo, A. C.: High Earth-system climate sensitivity determined from Pliocene carbon dioxide concentrations, Nat. Geosci., 3(1), 27–30, doi:10.1038/ngeo724, 2010.

Pagani, M., Huber, M., Liu, Z., Bohaty, S. M., Henderiks, J., Sijp, W. P., Krishnan, S. and DeConto, R. M.: The Role of Carbon Dioxide During the Onset of Antarctic Glaciation, Science (80-.)., 334, 1261–1265, doi:10.1126/science.1203909, 2011.

PALAEOSENS: Making sense of palaeoclimate sensitivity, Nature, 491(7426), 683–691, doi:10.1038/nature11574, 2012.

Pancost, R. D. and Pagani, M.: Controls on the carbon isotopic compositions of lipids in marine environments, Handb. Environ. Chem. Vol. 2 React. Process., 2 N(October 2005), 209–249, doi:10.1007/698_2_007, 2006.

Popp, B. N., Laws, E. A., Bidigare, R. R., Dore, J. E., Hanson, K. L. and Wakeham, S. G.: Effect of Phytoplankton Cell
Geometry on Carbon Isotopic Fractionation, Geochim. Cosmochim. Acta, 62(1), 69–77, doi:10.1016/S0016-7037(97)003335, 1998.

Rae, J. W. B., Zhang, Y. G., Liu, X., Foster, G. L., Stoll, H. M. and Whiteford, R. D. M.: Atmospheric CO2 over the Past 66 Million Years from Marine Archives, Annu. Rev. Earth Planet. Sci., 49(1), 599–631, doi:10.1146/annurev-earth-082420-063026, 2021.

- Ratti, S., Giordano, M. and Morse, D.: CO2-concentrating mechanisms of the potentially toxic dinoflagellate Protoceratium reticulatum (Dinophyceae, Gonyaulacales), J. Phycol., 43(4), 693–701, doi:10.1111/j.1529-8817.2007.00368.x, 2007.
 Richter, T. O., Peeters, F. J. C. and van Weering, T. C. E.: Late Holocene (0–2.4kaBP) surface water temperature and salinity variability, Feni Drift, NE Atlantic Ocean, Quat. Sci. Rev., 28(19–20), 1941–1955, doi:10.1016/j.quascirev.2009.04.008, 2009.
 Roeske, C. A. and O'Leary, M. H.: Carbon isotope effects on enzyme-catalyzed carboxylation of ribulose bisphosphate,
- Biochemistry, 23(25), 6275–6284, doi:10.1021/bi00320a058, 1984.
 van Roij, L., Sluijs, A., Laks, J. J. and Reichart, G.-J.: Stable carbon isotope analyses of ng quantities of particulate organic carbon (pollen) with laser ablation nano combustion gas chromatography isotope ratio mass spectrometry, Rapid Commun. Mass Spectrom., (October 2016), 47–58, doi:10.1002/rcm.7769, 2016.
 Rost, B., Richter, K.-U., Riesebell, U. and Hansen, P. J.: Inorganic carbon acquisition in red tide dinoflagellates, Plant, Cell
- Environ., 29(5), 810–822, doi:10.1111/j.1365-3040.2005.01450.x, 2006.
 Schoon, P. L., Sluijs, A., Sinninghe Damsté, J. S. and Schouten, S.: Stable carbon isotope patterns of marine biomarker lipids in the Arctic Ocean during Eocene Thermal Maximum 2, Paleoceanography, 26(3), doi:10.1029/2010PA002028, 2011.
 Schouten, S., Klein Breteler, W. C., Blokker, P., Schogt, N., Rijpstra, W. I. C., Grice, K., Baas, M. and Sinninghe Damsté, J. S.: Biosynthetic effects on the stable carbon isotopic compositions of algal lipids: implications for deciphering the carbon
- 620 isotopic biomarker record, Geochim. Cosmochim. Acta, 62(8), 1397–1406, doi:10.1016/S0016-7037(98)00076-3, 1998. Sharkey, T. D. and Berry, J. A.: Carbon isotope fractionation of algae as influenced by an inducible CO2 concentrating mechanism, in Inorganic carbon uptake by Aquatic Photosynthetic organisms, edited by W. J. Lucas and J. A. Berry, pp. 389– 401, the American Society of Plant Physiologists., 1985.

Sluijs, A., van Roij, L., Frieling, J., Laks, J. and Reichart, G.-J.: Single-species dinoflagellate cyst carbon isotope ecology across the Paleocene-Eocene Thermal Maximum, Geology, 46(1), 79–82, doi:10.1130/G39598.1, 2018.

Stoll, H. M., Guitian, J., Hernandez-Almeida, I., Mejia, L. M., Phelps, S., Polissar, P., Rosenthal, Y., Zhang, H. and Ziveri, P.:
Upregulation of phytoplankton carbon concentrating mechanisms during low CO2 glacial periods and implications for the phytoplankton pCO2 proxy, Quat. Sci. Rev., 208, 1–20, doi:10.1016/j.quascirev.2019.01.012, 2019.

Tagliabue, A. and Bopp, L.: Towards understanding global variability in ocean carbon-13, Global Biogeochem. Cycles, 22(1),

630 1–13, doi:10.1029/2007GB003037, 2008.

Takahashi, T., Sutherland, S. C., Chipman, D. W., Goddard, J. G., Ho, C., Newberger, T., Sweeney, C. and Munro, D. R.: Climatological distributions of pH, pCO2, total CO2, alkalinity, and CaCO3 saturation in the global surface ocean, and temporal changes at selected locations, Mar. Chem., 164, 95–125, doi:10.1016/j.marchem.2014.06.004, 2014. Takahashi, T., Sutherland, S. C. and Kozyr, A.: Global Ocean Surface Water Partial Pressure of CO2 Database: Measurements

Performed During 1957-2015 (Version 2015), Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory,
 U.S. Department of Energy, Oak Ridge, Tennessee, 2015.

Versteegh, G. J. M., Blokker, P., Marshall, C. and Pross, J.: Macromolecular composition of the dinoflagellate cyst Thalassiphora pelagica (Oligocene, SW Germany), Org. Geochem., 38(10), 1643–1656, doi:10.1016/j.orggeochem.2007.06.007, 2007.

640 Versteegh, G. J. M., Blokker, P., Bogus, K., Harding, I. C., Lewis, J., Oltmanns, S., Rochon, A. and Zonneveld, K. A. F.: Infra red spectroscopy, flash pyrolysis, thermally assisted hydrolysis and methylation (THM) in the presence of tetramethylammonium hydroxide (TMAH) of cultured and sediment-derived Lingulodinium polyedrum (Dinoflagellata) cyst walls, Org. Geochem., 43, 92–102, doi:10.1016/j.orggeochem.2011.10.007, 2012.

Van de Waal, D. B., John, U., Ziveri, P., Reichart, G.-J., Hoins, M., Sluijs, A. and Rost, B.: Ocean Acidification Reduces

- Growth and Calcification in a Marine Dinoflagellate, PLoS One, 8(6), doi:10.1371/journal.pone.0065987, 2013.
 Wall, D. and Dale, B.: "Living fossils" in Western Atlantic plankton, Nature, 211(5053), 1025–1026 [online] Available from: http://www.scopus.com/inward/record.url?eid=2-s2.0-0000259853&partnerID=40&md5=441e2fd50df53b93e4d31aa0ba9e6015, 1966.
- Wilkes, E. B. and Pearson, A.: A general model for carbon isotopes in red-lineage phytoplankton: Interplay between
 unidirectional processes and fractionation by RubisCO, Geochim. Cosmochim. Acta, 265, 163–181, doi:10.1016/j.gca.2019.08.043, 2019.

Wilkes, E. B., Carter, S. J. and Pearson, A.: CO 2 -dependent carbon isotope fractionation in the dinoflagellate Alexandrium tamarense, Geochim. Cosmochim. Acta, 212, 48–61, doi:10.1016/j.gca.2017.05.037, 2017.

Wilkes, E. B., Lee, R. B. Y., McClelland, H. L. O., Rickaby, R. E. M. and Pearson, A.: Carbon isotope ratios of coccolith–
associated polysaccharides of Emiliania huxleyi as a function of growth rate and CO2 concentration, Org. Geochem., 119, 1–
10, doi:10.1016/j.orggeochem.2018.02.006, 2018.

Williams, G. L., Brinkhuis, H., Pearce, M. A., Fensome, R. A. and Weegink, J. W.: Southern Ocean and global dinoflagellate cyst events compared: index events for the Late Cretaceous–Neogene, in Proceedings of the Ocean Drilling Program, Scientific Results, vol. 189, pp. 1–98., 2004.

Witkowski, C. R., Weijers, J. W. H., Blais, B., Schouten, S. and Sinninghe Damsté, J. S.: Molecular fossils from phytoplankton reveal secular PCO2 trend over the Phanerozoic, Sci. Adv., 4(11), eaat4556, doi:10.1126/sciadv.aat4556, 2018.
Zhang, H., Torres-Romero, I. and Stoll, H. M.: Re-examining extreme carbon isotope fractionation in the coccolithophore Ochrosphaera neapolitana, Nat. Commun., 13(1), 7606, doi:10.1038/s41467-022-35109-4, 2022.
Zonneveld, K. A. F., Versteegh, G. J. M. and De Lange, G. J.: Preservation of organic-walled dinoflagellate cysts in different

- 665 oxygen regimes: a 10,000 year natural experiment, Mar. Micropaleontol., 29(3), 393–405, 1997.
 - Zonneveld, K. A. F., Marret, F., Versteegh, G. J. M., Bogus, K., Bonnet, S., Bouimetarhan, I., Crouch, E., de Vernal, A.,
 Elshanawany, R., Edwards, L., Esper, O., Forke, S., Grøsfjeld, K., Henry, M., Holzwarth, U., Kielt, J.-F., Kim, S.-Y.,
 Ladouceur, S., Ledu, D., Chen, L., Limoges, A., Londeix, L., Lu, S.-H., Mahmoud, M. S., Marino, G., Matsouka, K.,
 Matthiessen, J., Mildenhal, D. C., Mudie, P., Neil, H. L., Pospelova, V., Qi, Y., Radi, T., Richerol, T., Rochon, A., Sangiorgi,
- F., Solignac, S., Turon, J.-L., Verleye, T., Wang, Y., Wang, Z. and Young, M.: Atlas of modern dinoflagellate cyst distribution based on 2405 data points, Rev. Palaeobot. Palynol., 191, 1–197, doi:10.1016/j.revpalbo.2012.08.003, 2013.
 Zonneveld, K. A. F., Gray, D. D., Kuhn, G. and Versteegh, G. J. m.: Postdepositional aerobic and anaerobic particulate organic matter degradation succession reflected by dinoflagellate cysts: The Madeira Abyssal Plain revisited, Mar. Geol., 408, 87–109, doi:10.1016/j.margeo.2018.11.010, 2019.

675

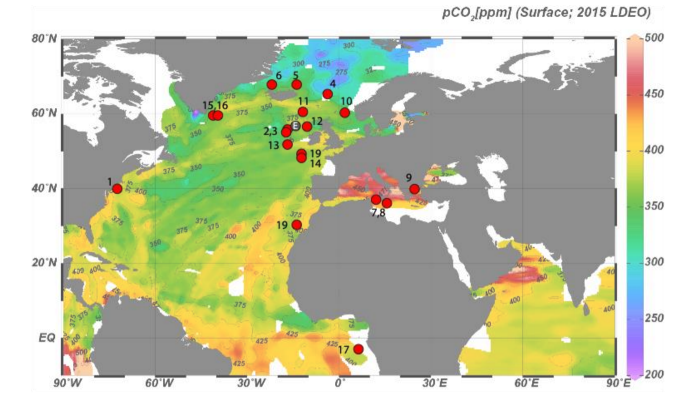


Figure 1. Locations of samples with sufficient *Operculodinium centrocarpum* and/or *Spiniferites* spp for dinoflagellate cyst
 δ¹³C analyses. Numbers correspond to localities listed in Table 1. Down-core data location (ENAM9606; 55.650 °N, -13.985 °E) is marked by a grey dot ("E") between core-tops #2, 3 and 12.

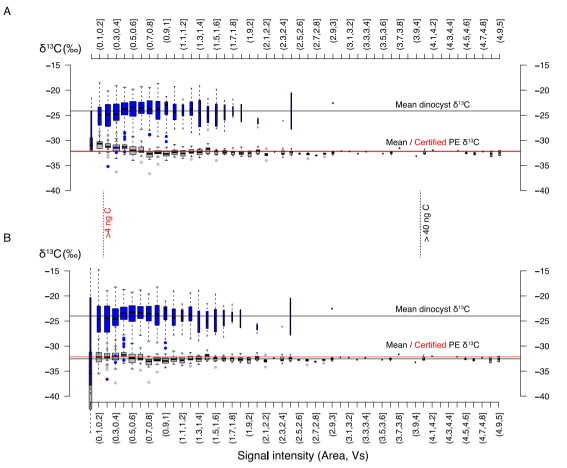


Figure 2. Signal intensity in Volt seconds (Vs) versus carbon isotope value distribution within bins of 0.1 Vs. A. The blue boxplots represent our new dinocyst δ^{13} C, and the gray boxplots represent the previously published data for the PE 685 standard (δ^{13} C value $-32.15\% \pm 0.05\%$; 1 σ). **B.** Same as A. but for background-corrected values (see §3.2). The vertical black and red line represent cut-off values for individual PE standard measurements (>4 Vs; ~42 ng C; 0.5‰ precision) and individual cyst measurements (>0.2 Vs; ~6 ng C), respectively. The width of each boxplot is square-root scaled with the number of measurements in the respective bins. Note that several bins at the high-end do not contain any data.

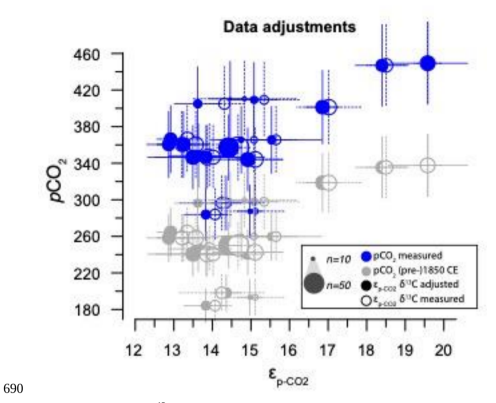


Figure 3. Effects of δ^{13} C and pCO₂ corrections (see also Figure 2 and §3.2).

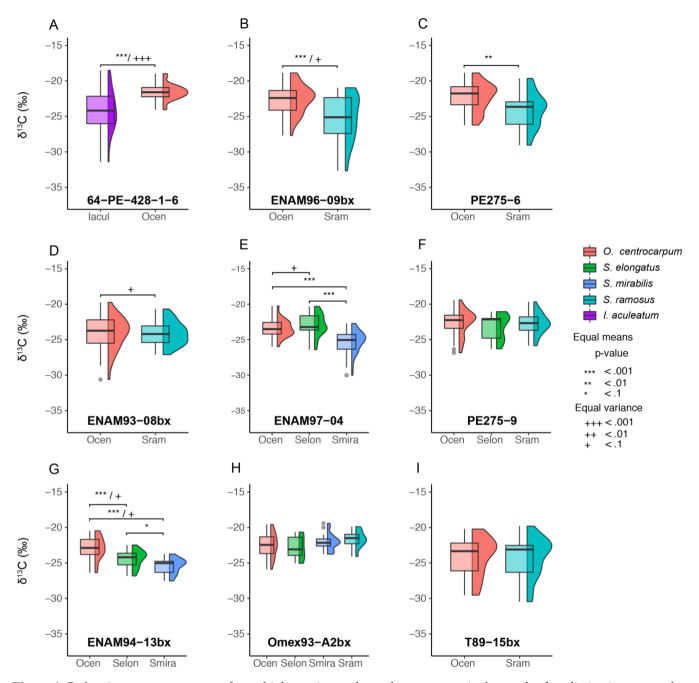


Figure 4. Carbon isotope measurements for multiple species; each panel represents a single sample after eliminating extremely small measurements sizes (<0.2 Vs), background correction and removing outliers (± 2.5 interquartile range) (paragraph §3.2).
 Each box-whisker and δ¹³C distribution plot represents a set of measurements for a single species at their respective locality; note that tailing towards negative δ¹³C is common. Brackets above species δ¹³C populations indicate significant differences in means and variance for different species within a single sample.

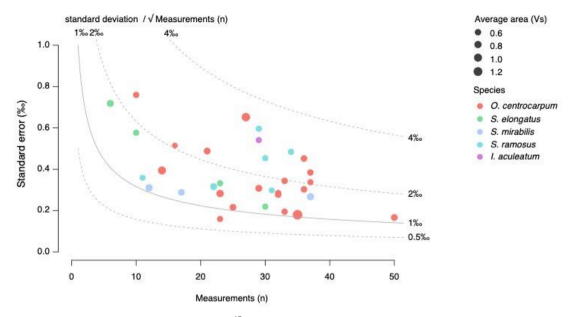


Figure 5. Relation of standard error of $\delta^{13}C_{DINO}$ (‰) with the number of measurements and signal intensity (area in Volt seconds (Vs)). Colors correspond to the various analyzed species.

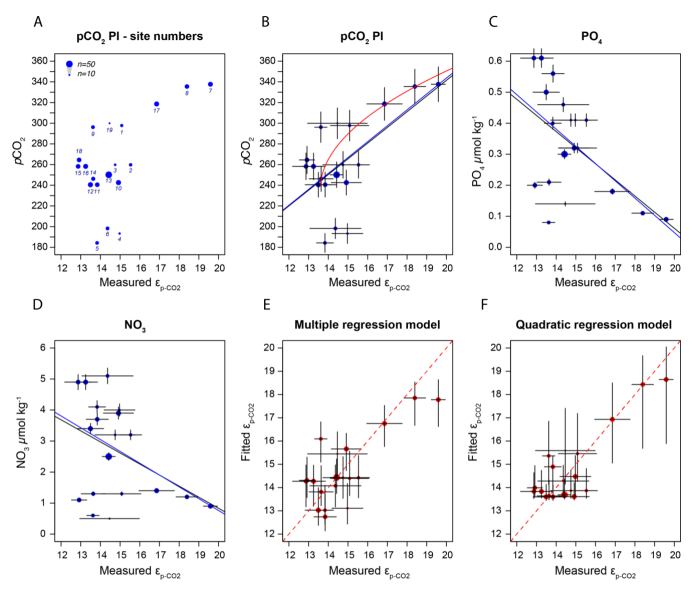


Figure 6. A. Sample – specific ε_{p-CO2} of *O. centrocarpum* relative to pre-industrial pCO₂, numbers correspond to localities listed in Table 1. **B.** Regression analyses for ε_{p-CO2} of *O. centrocarpum* relative to pCO_2 (measured, corrected to pre-industrial values); black line represents simple linear regression, blue lines represent weighted linear regressions and red line represents weighted quadratic regression. **C.** phosphate concentrations (PO₄³⁻), **D.** nitrate concentrations (NO₃⁻). **E.** Fitted values illustrating the multiple regression model performance using parameters a-c relative to measured ε_{p-CO2} . **F.** Fitted values using only pre-industrial *p*CO₂ but applying a quadratic regression (red curve in panel b). Errors in panels b-d represent 5% of the

705

measured value and errors on the fitted values in panels d and e represent propagated errors of both measurements and environmental variables (as shown in panel B – D) using Monte-Carlo simulations (n = 1000) for regression models. Symbol size (top left corner of panel A) represents the number of measurements within each sample.

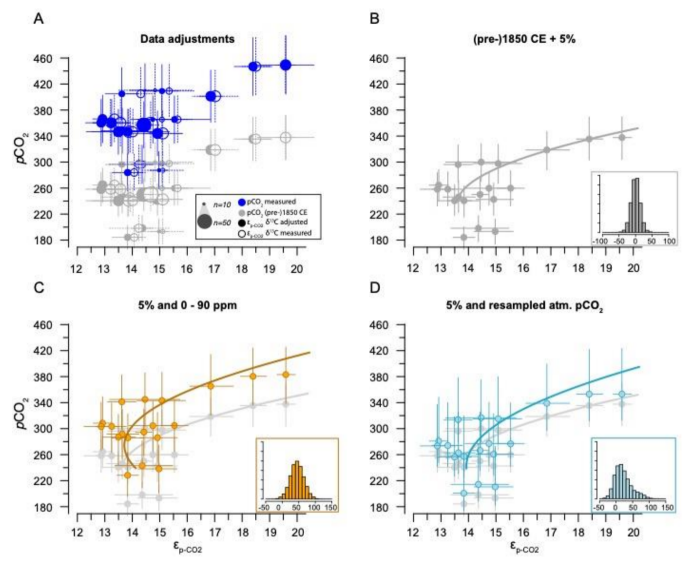


Figure 7. Sensitivity tests to potential effect of anthropogenic carbon emissions. A. Effects of data treatment on the difference between measured and adjusted *p*CO₂ and ε_{p-CO2} (same as Figure 3). Open symbols indicate measured δ¹³C, closed symbols represent data after eliminating small signals (<0.2 Vs) and outliers. Blue dots represent measured CO₂ values and grey dots indicate the CO₂ around 1850 CE. B. Quadratic regression (red line in Fig. 6B) with propagated analytical error on *p*CO₂ and δ¹³C only, using CO₂ values around 1850 CE (grey filled symbols in panel A. C. As in B but with addition of a +45 ± 15 ppm error to reflect potential impact of anthropogenic CO₂ in orange. Grey dots and curve of panel B are added as a comparison. D. As in B but with addition of a CO₂ increase relative to pre-industrial, sampled for the period 1800 – 2000
720 CE. Insets on the bottom right in panels B, C and D show the combined error distributions imposed on *p*CO₂. All error bars

in panels B – D on *p*CO₂ and ε_{p-CO2} are 2.5 – 97.5% percentile ranges from Monte Carlo simulations (n = 1000).

Site number	Core-ID	Latitude (°N)	Longitude (ºE)	Species	Measurements (n)	S-W normality
1	NF2012-091	(N) 37.977402	(° ⊏) -73.669403	O.centrocarpum	(11) 22 (21)	* ()
2	PE360-24	55.496231	-15.800755	O.centrocarpum	24 (23)	** ()
3	PE360-45	55.539398	-15.8453	O.centrocarpum	23 (16)	0
4	NA87-02	64.480003	-5.83	O.centrocarpum	20 (14)	*** ()
5	LCD13	67.501282	-15.069252	O.centrocarpum	25 (25)	
6	LCD10A	66.677437	-24.179598	O.centrocarpum	29 (27)	() ()
7	MedSea (MC-613)	35.8575	14.105556	O.centrocarpum	35 (35)	0
8	MedSea (MC-614)	35.8075	12.998056	O.centrocarpum	33 (32)	* ()
9	MedSea (MC-645)	40.2175	25.244167	O.centrocarpum	33 (23)	** ()
10	ENAM93-08bx	59.501667	3.69	O.centrocarpum	38 (37)	*()
10		39.301007	5.05	S.ramosus	33 (32)	
11	ENAM94-13bx	60.249997	-11.19	O.centrocarpum	36 (33)	() ()
11	LINAWI94-130X	00.249997	-11.19	S.elongatus	34 (30)	() *** ()
				S.mirabilis		0
12	ENAM96-09bx	57.159917	-10.26	O.centrocarpum	13 (12) 39 (37)	0
12		57.159917	-10.20	S.ramosus	30 (29)	0
13	ENAM97-04	52.410386	-14.94	O.centrocarpum	52 (50)	()
15	LINAW97-04	52.410500	-14.94	S.elongatus	23 (23)	
				S.mirabilis	38 (37)	() ** (*)
14	Omex93-A2 bx	49.483	-11.13	O.centrocarpum	30 (29)	
14	Omer93-Az br	49.405	-11.15	S.ramosus	13 (11)	() *()
				S.elongatus	6 (6)	0
				S.mirabilis	18 (17)	()
15	PE275-6	59.272369	-38.36	O.centrocarpum	34 (33)	() *()
10	1 22/0 0	00.272000	-00.00	S.ramosus	34 (30)	**()
16	PE275-9	59.272369	-38.36	O.centrocarpum	37 (36)	() *(*)
10	1 22/0 0	00.272000	-00.00	S.ramosus	23 (22)	() *(*)
				S.elongatus	12 (10)	() *(*)
17	T89-15bx	-4.199372	10.05	O.centrocarpum	36 (36)	* (*)
	100 100	4.100012	10.00	S.ramosus	35 (34)	* (*)
18	64PE428-1-1-6	47.079782	-10.197305	O.centrocarpum	35 (33)	()
10		41.010102	10.107000	I.aculeatum	30 (29)	()
19	64PE428-1-6-6	30.67917	-11.930478	0.centrocarpum	11 (10)	()
13		50.07317	11.000470	S.Com Ocarpulli	11(10)	0

Table 1. Core localities, analyzed species, number of measurements and normality of the carbon isotope data. Number of measurements total and in parentheses measurements used for environmental comparisons (see also results §3.4). Shapiro-Wilk (S-W) normality test on data: non-normal data distributions are indicated where p values are < 0.1 (*), <0.01 (**) and <0.001 (***), in parentheses the same for the data used for environmental comparisons. Site numbers correspond to those in Figs. 1 and 6A.

Table 2: Linear regression coefficients and significance for all samples where *O. centrocarpum* was analyzed (n = 19), with ε_{p-CO2} as dependent variable. Parameters with p-values <0.05 in bold.

	Coeff.	Std.err.	t	р	R^2
CO ₂ (mol/kg)	-2.506e+05	5.603e+05	-0.447	0.66039	0.01163
CO ₃ ²⁻ (mol/kg)	41263.663	15432.856	2.674	0.0160	0.296
HCO3 ⁻ (mol/kg)	809.88	7057.32	0.115	0.910	0.0007741
DIC(mol/kg)	5655.859	6266.998	0.902	0.379	0.04572
SST (°C)	0.16971	0.06043	2.809	0.0121	0.3169
SSS (psu)	0.6737	0.3604	1.869	0.0789	0.1705
PO4 ³⁻ (µmol /kg)	-5.5131	2.1102	-2.613	0.0182	0.2865
NO3 ⁻ (µmol /kg)	-0.4484	0.2493	-1.798	0.0899	0.1599
Si (µmol /kg)	-0.2553	0.3298	-0.774	0.449	0.03405
O ₂ (mL/L)	-1.274	0.433	-2.943	0.0091	0.3375
ALK (mol/kg)	7621.253	4712.178	1.617	0.124	0.1334
pCO ₂ ~1850	0.024945	0.007752	3.218	0.005050	0.3785

Table 3: As Table 2, but with ε_{p-DIC} as dependent variable.

	Coeff.	Std.err.	t	р	R^2
CO ₂ (mol/kg)	1.133e+05	4.760e+05	0.238	0.815	0.00332
CO ₃ ²⁻ (mol/kg)	19589.987	14818.493	1.322	0.204	0.09322
HCO3 ⁻ (mol/kg)	6547.21	5758.02	1.137	0.271	0.07068
DIC(mol/kg)	7803.835	5086.841	1.534	0.143	0.1216
SST (°C)	0.04959	0.06068	0.817	0.425	0.03781
SSS (psu)	0.4043	0.3201	1.263	0.224	0.08579
PO4 ³⁻ (µmol /kg)	-2.3993	2.0318	-1.181	0.254	0.07581
NO3 ⁻ (µmol /kg)	-0.05666	0.22969	-0.247	0.808	0.003567
Si (µmol /kg)	-0.01925	0.28388	-0.068	0.947	0.0002703
O ₂ (mL/L)	-0.4170	0.4385	-0.951	0.355	0.05051
ALK (mol/kg)	6754.843	3956.586	1.707	0.106	0.1464
pCO ₂ ~1850	0.010083	0.007952	1.268	0.222	0.0864

735