Coccolithophore productivity at the western Iberian Margin during the Middle Pleistocene (310–455 ka) – evidence from coccolith Sr/Ca data

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Abstract. Coccolithophores contribute significantly to marine primary productivity and play a unique role in ocean biogeochemistry by using carbon for photosynthesis (soft-tissue pump) and for calcification (carbonate counter pump). Despite the importance of including coccolithophores in Earth system models to allow better predictions of the climate system’s responses to planetary change, the reconstruction of coccolithophore productivity mostly relied on proxies dependent on accumulation and sedimentation rates and preservation conditions. In this study we used an independent proxy, based on the coccolith fraction (CF) Sr/Ca ratio, to reconstruct coccolithophore productivity. We studied the marine sediment core MD03-2699 from the western Iberian margin (IbM), concentrating on glacial–interglacial cycles of Marine Isotopic Stage (MIS) 12 to MIS 9. We found that IbM coccolithophore productivity was controlled by changes in the oceanographic conditions, such as in sea surface temperature (SST) and nutrient availability, and by competition with other phytoplankton groups. Long-term coccolithophore productivity was primarily affected by variations in the dominant surface water mass. Polar and subpolar surface waters during glacial substages were associated with decreased coccolithophore productivity, with the strongest productivity minima concomitant with Heinrich-type events (HtEs).

Subtropical, nutrient-poorer waters, increased terrigenous input, and moderate to strong upwelling during the deglaciation and early MIS11 are hypothesized to have attributed a competitive advantage to diatoms to the detriment of coccolithophores, resulting in intermediate coccolithophore productivity levels. During the progression towards full glacial conditions an increasing presence of nutrient-richer waters, related to the growing influence of transitional surface waters and/or intensified upwelling, probably stimulated coccolithophore productivity to maxima following the rapid depletion of silica by diatoms. We present conceptual models of the carbon and carbonate cycle components for the IbM in different time slices that might serve as a basis for further investigation and modelling experiments.

1 Introduction

Coccolithophores play a unique role in ocean biogeochemistry using carbon for both photosynthesis and calcification (e.g. Rost and Riebesell, 2004; Westbroek et al., 1993) and contributing up to 60 % to the total calcium carbonate in the ocean (Flores and Sierro, 2007). During the mid-Brunhes interval of Marine Isotope Stage (MIS) 14 to MIS 9, when the assemblages were vastly dominated by gephyrocapsids...
(Baumann and Freitag, 2004; Saavedra-Pellitero et al., 2017), coccolithophores provided > 80% of the total oceanic calcium carbonate. And, several studies hypothesized a relevant role of coccolithophores in glacial-interglacial dynamics (Duchamp-Alphonse et al., 2018; McClelland et al., 2016; Omta et al., 2013; Rickaby et al., 2007; Saavedra-Pellitero et al., 2017), but there are uncertainties arising from the complex interactions between coccolithophore productivity and growth rates, nutrient dynamics and competition with non-calcifiers, seasonality and the export of both carbonate and organic carbon, and the final accumulation/burial rates (Balch, 2018; Duchamp-Alphonse et al., 2018; McClelland et al., 2016; Omta et al., 2013; Rickaby et al., 2007; Ridgwell and Zeebe, 2005; Saavedra-Pellitero et al., 2017). These uncertainties make the inclusion of biogeochemical processes related to coccolithophores in Earth system models challenging. In order to provide useful information to modellers, it is thus fundamental to test the dynamic response of the various components of the system over different climatic scenarios through the use of multiproxy studies involving the reconstruction of coccolithophore growth rates and the export of both their organic and inorganic compounds, namely alkenones and nannofossil accumulation rate (AlkAR and NAR, respectively).

Since coccolithophores are sensitive to rapid fluctuations in temperature, salinity, nutrients, and turbidity of surface waters (Baumann et al., 2005; McIntyre and Bé, 1967), their calcareous remains, the coccoliths, retrieved from deep-sea sediments have been used extensively to reconstruct paleoenvironmental conditions (Amore et al., 2012; Baumann et al., 2005; Beaufort et al., 2001; Flores et al., 1997; Maieronzo et al., 2015; Marino et al., 2014; McIntyre and Molino, 1996; Saavedra-Pellitero et al., 2017). However, coccolithophore paleoproductivity reconstruction has been tentative and mostly relied on proxies dependent not only on the extent of the supply but also on dilution by mineral matter, changes in sedimentation or accumulation rates, and preservation conditions (Rullkötter, 2006). Beaufort et al. (1997) proposed a proxy to quantitatively reconstruct coccolithophore productivity, but its applicability is unfortunately limited to latitudes between 30°N and 30°S (Hernández-Almeida et al., 2019). A widely used alternative proxy is the coccolith fraction Sr/Ca (CF Sr/Ca) ratio that is independent of accumulation rate (Cavaleiro et al., 2018; Mejía et al., 2014; Saavedra-Pellitero et al., 2017; Tangunan et al., 2017). Coccolithophores construct coccoliths internally within their cell and several studies show the direct and proportional relationship between the Sr/Ca ratio of the coccolith and the coccolith calcification rate. The calcification rate is a function of growth rate (e.g. Daniels et al., 2018) and therefore of coccolithophore productivity (Balch et al., 1996; Rickaby et al., 2007; Stoll and Schrag, 2000). The faster coccolithophores grow, the faster they calcify and more Sr is incorporated into the calcite lattice of their coccoliths (Stoll et al., 2002b, a; Stoll and Schrag, 2000). With temperature and assemblage effects considered (see “Material and methods” section), the sea surface temperature (SST)-corrected CF Sr/Ca curve (or residual curve) can be expected to reflect coccolithophores’ growth rate and thus their productivity qualitatively (Müller et al., 2014; Stoll et al., 2002a).

Coccolithophores research has been gaining increased attention because of their potential relevant role in the carbon cycle (Ridgwell and Zeebe, 2005). The carbon cycle underwent major changes during the past 800 kyr as evidenced, for example, by the glacial-interglacial variations in atmospheric carbon dioxide (atmCO2) concentrations (Lüthi et al., 2008). Though the changing orbital parameters of Earth (i.e. the Milankovitch cycles) are undoubtedly related to the glacial-interglacial cycles (e.g. Berger, 1988), the mechanisms driving the almost 80 ppmv lower atmCO2 concentrations during glacial periods and the rapid rise during deglaciations are yet to be described and modelled accurately (Barker et al., 2006; Duchamp-Alphonse et al., 2018; Omta et al., 2013; Ridgwell and Zeebe, 2005). Since the ocean contains 60 times more carbon than the atmosphere, modest changes in its holding capacity could greatly affect atmCO2 (e.g. Sarmiento and Gruber, 2007). It is clear that atmCO2 changes are a consequence of the interplay of several factors and not a single mechanism at work (Archer et al., 2000; Sigman and Boyle, 2000). Yet, theories such as the “coral reef hypothesis” (Berger, 1982), “carbonate compensation” (Broecker and Peng, 1987), and “rain ratio hypothesis” (Archer and Maier-Reimer, 1994) give the marine carbonate cycle a deterministic role in atmCO2 changes on timescales shorter than 100 kyr (e.g. Ridgwell and Zeebe, 2005; Broecker, 1982).

Our studied period falls into the mid-Brunhes interval (Barker et al., 2006; Baumann and Freitag, 2004; Jansen et al., 1986) when, starting with MIS 11c, interglacial atmCO2 levels were comparable to the Holocene (Lüthi et al., 2008). The mid-Brunhes interval also has the advantage to cover the Gephyrocapsa caribbeanica acme (Baumann and Freitag, 2004), reducing the likelihood of a biasing effect of the CF Sr/Ca ratio from changing coccolith assemblages (see “Material and methods” section). It encompasses different climatic scenarios, from late MIS 12 to early MIS 9, straddling the harsh glacial of MIS 12 and the prolonged MIS 11c interglacial period, considered to be one of the best analogues for the current MIS 1 interglacial (Berger et al., 2015; Candy et al., 2014; Loutre and Berger, 2003; Oliveira et al., 2018). The deglaciation and termination (T) V is considered one of the longest deglaciations (Tzedakis et al., 2012; Vázquez Riveiros et al., 2013), and the highest glacial-interglacial amplitude change of the last 1 Myr (Lisiecki and Raymo, 2005) allowing a more detailed analysis due to the high sedimentation rates that characterize the western Iberian Margin (IbM).

The western IbM embodies the northern section of the Canary Current Eastern Boundary Upwelling System (Arístegui et al., 2009; Mason et al., 2006). The seasonal meridional shifts in the atmospheric highs revert the upwelling favour-
ing equatorward winds seasonally to become poleward in autumn and winter (Mason et al., 2006). Despite representing only 10% of the ocean, the Eastern Boundary Upwelling Systems contribute 80% to 90% to the oceanic new production (Hill et al., 1998, Ryther, 1969). These systems are sensitive to climate change by responding to shifts in processes that regulate ocean stratification, upwelling, wind stress, dust supply, and basin-wide circulation (Freon et al., 2009). However, to infer the role of climate modes in modulating air–sea CO₂ fluxes in such a complex environment is challenging although this knowledge is crucial to improve our ability to predict future evolution of atmCO₂ (Brady et al., 2019).

In this study we reconstruct coccolithophore productivity from the CF Sr/Ca ratio record at Site MD03-2699, an area under the influence of the Canary Current upwelling system, and particularly sensitive to global climate change (Hodell et al., 2015; Martrat et al., 2007; Oliveira et al., 2016; Rodrigues et al., 2017) and thus optimal to evaluate this phytoplankton group’s behaviour to varying conditions. Using a multiproxies approach, we compare the CF Sr/Ca data with the total alkenone concentration (i.e. the exported organic carbon produced by coccolithophores), as well as with the NARs in order to investigate the processes of production versus export and preservation of both organic and inorganic carbon. This approach allows us to gain a better understanding of (1) the coupling–decoupling between the different proxies used to reconstruct coccolithophores’ productivity and (2) the role of upwelling strength and terrigenous input in nutrient competition under different climate conditions, such as glacials, interglacials, deglaciations, and the transition from interglacial to glacial conditions.

2 Regional setting

2.1 Present hydrography of the IbM

The study area is located at the north-eastern edge of the subtropical gyre and influenced by a southward flowing branch of the North Atlantic Current forming the Portugal Current system (Fig. 1). The Portugal Current system is mainly characterized by a slow, southerly flow with intricate interaction between coastal and offshore currents (Peliz et al., 2005; Relvas et al., 2007), bottom topography, and water mass convergence (Bischof et al., 2003). One of the most important features of the IbM is the surface circulation’s seasonality as a response to the combined position of the Azores High and the Icelandic low-pressure systems (Barton, 2001; Haynes and Barton, 2018; Relvas et al., 2007). During summer, the migration of the Azores High to the central Atlantic exposes the IbM to northerly trade winds, strengthening the Portugal Current and forcing the upper layer (150 to 200 m) to flow towards the Equator. The induced offshore Ekman transport allows colder, less saline, and nutrient-rich subsurface water to rise to the surface (Alvarez et al., 2011; Fitsuza, 1983), which leads to high primary production (Figueiras et al., 2002; Fraga, 1981; Tenore et al., 2018). The upwelled water is the Eastern North Atlantic Central Water (ENACW), which has two different origins: a less saline, colder, nutrient-rich water mass of subpolar origin (ENACWsp) formed along with the Subpolar Mode Water in the Rockall Plateau region and a warmer, saltier, nutrient-poorer subtropical branch (ENACWst) formed along the Azores Front (Fiúza et al., 1984). Depending on the wind strength intensity either type can be upwelled (Fiúza et al., 1984), and in the current interglacial ENACWsp is mostly upwelled in the north-western coast of the Iberian peninsula, while ENACWst is mostly upwelled in the southern area of the western Iberian peninsula (Mason et al., 2006). The upwelling season typically lasts from late spring into summer (May to September), and upwelling filaments are mostly observed off the most prominent capes (Fiúza et al., 1982). Upwelling plumes and other mesoscale features can spread zonally offshore to distances of 200 km or more (Sousa and Bricaud, 1992). In autumn and winter the seasonal meridional shifts in the atmospheric highs revert the upwelling favouring equatorward winds to become poleward (Mason et al., 2006). The Azores High located further south and the Icelandic low intensification forces the wind regime to become more southerly, enabling the near-shore Iberian Poleward Current, which brings warmer and more oligotrophic waters from the Azores Current to the shelf and upper slope areas along the western IbM (Fiúza, 1983; Peliz et al., 2005; Vitorino et al., 2002). Downwelling can occur during this period but wintertime freshwater discharges from the major rivers are also a relevant feature, namely from the Tagus river that crosses the Iberian
Peninsula, carrying a major load of suspended organic-rich sediments onto the continental shelf off Lisbon (Cabeçadas and Brogueira, 1998) and the Estremadura spur where our sediment core is located.

2.2 Present coccolithophore productivity patterns in the IbM

Several studies on coccolithophore abundance and distribution (e.g., Abrantes and Moita, 1999; Cachão et al., 2000; Guerreiro et al., 2013; Moita, 2001) have been performed along the IbM, but only two were able to assess annual and seasonal productive cycles (Ausín et al., 2018; Silva, 2008; Silva et al., 2009). Maximum coccolithophores were found during spring and summer and associated with high irradiance levels (Ausín et al., 2017) and the convergence of warmer, more oligotrophic waters with the relaxation of northerly winds (Silva, 2008; Silva et al., 2009). Minimum absolute abundances, on the other hand, were observed during winter. The competitive dynamics between coccolithophores and diatoms are still not completely understood (Cermeño et al., 2011). Yet, these two main phytoplankton groups dominate the carbon dioxide mediation between the atmosphere and the ocean making them important contributors to the global carbon cycle. Diatoms are known to require more silica (Si) and iron (Fe) than coccolithophores (e.g. Merico et al., 2004), and Tyrrell and Young (2009) suggested that coccolithophore blooms might be associated with Si and Fe surface water depletion. Cermeño et al. (2011) suggested that coccolithophores could outcompete diatoms under steady-state nitrate limitation but under dynamical conditions (introducing nitrate pulses) diatoms outcompeted coccolithophores. They further found that the more frequent the pulses, the more rapidly diatoms outcompeted coccolithophores. The coastal upwelling conditions resemble these dynamical conditions, when pulses of nutrients are brought to the ocean surface. This could explain why Abrantes and Moita (1999) found a phytoplankton dominance of diatoms during upwelling events and a clear dominance of coccolithophores (∼90 %) during non-upwelling events. Also, Ausín et al. (2017) describe a decreasing abundance of diatoms with an increasing distance to shore, whereas coccolithophores showed higher abundances offshore than in coastal areas. This distribution pattern was maintained during the upwelling season (May to September), when coccolithophores are outcompeted by diatoms, in particular during the longer-lasting and more intense upwelling events (Moita, 2001; Silva et al., 2008).

Despite the general acceptance of coccolithophores as a single functional phytoplanktonic group associated with low-turbulence, low-nutrient, and high-irradiance environments, different species show varying life strategies. Several studies describe increased cell densities, mostly due to blooms of Emiliania huxleyi and Gephyrocapsa oceanica in the IbM (Moita, 2001; Guerreiro et al., 2013; Silva, 2008; Silva et al., 2009), together with other fast-growing opportunistic phytoplankton genera, such as the diatoms Chaetoceros s.l., Thalassiosira s.l., and Skeletonema s.l. This corroborates coccolithophores’ role among early successsion taxa, or at least some species, capable of rapid growth in a nutrient-rich environment and most likely explains why Abrantes and Moita (1999) found coccolithophores’ distribution in recent sediments to reflect their water column distribution during an upwelling situation and not the winter conditions.

3 Material and methods

3.1 Sediment sampling and coccolith fraction separation

In this study we used sediments of core MD03-2699 (39°02.20′ N, 10°39.63′ W; 1895 m water depth), retrieved from a sediment drift, located ca. 100 km offshore, on the Estremadura promontory (Fig. 1). Sediments were collected using a giant CALYPSO piston corer on board the R/V Marion Dufresne II (PICABIA Cruise, 2003). The sedimentary record is mainly composed of hemipelagic silty clays. For our research, we use the age model published by Voelker et al. (2010).

For the CF Sr/Ca ratio, a total of 183 samples was analysed. Samples were taken at 4 cm spacing from the ∼7 m long section (from 1190 to 1898 cm core depth) corresponding to MIS 12 to MIS 9 (∼455 to 310 ka). This resulted in a temporal resolution of ∼775 years, although the oldest part has lower resolution due to the lower sedimentation rates (Rodrigues et al., 2011). To obtain the coccolith fraction (CF) Sr/Ca record, ∼250 mg of freeze-dried sample was collected and suspended in 2 % ammonia (to avoid carbonate dissolution) and sieved through a 20 µm mesh. This sieving aimed to separate the coccoliths contained in the so-called coccolith fraction (< 20 µm) from mostly foraminifers and their fragments and other larger microfossils or sediment components. All sieving material was carefully washed with running tap water and rinsed with distilled water in between samples to avoid cross contamination.

3.2 Sample preparation and Sr/Ca analysis

We followed a three-step protocol to clean the sediment samples based on Stoll and Ziveri (2002):

1. the addition of 15 mL of MNX reagent (75 mg of hydroxylamine hydrochloride, 6 mL of concentrated ammonia, and 9 mL of ultrapure water) for a 12 h reaction in an automatic shaker; this step reduces Fe and Mn oxyhydroxides that scavenge metals from seawater and contain non-carbonate Sr;
2. the addition of 2 % ammonia to remove any non-carbonate Sr, e.g. from clays, by exchanging cations (Sr\(^2^+\)) with the excess of NH\(\_4^+\);
3. three ultrapure water rinses to extract the ammonia.
A weak buffered acid (6 g glacial acetic acid, 7 g ammonium acetate in 1 L of Milli-Q water) was then used to dissolve the coccoliths, minimizing the contribution of ions from non-carbonate phases. The samples were left in acid for 12 h and the obtained solution was centrifuged, extracted, and kept in acid-cleaned centrifuge tubes. A first ICP-AES (inductively coupled plasma–atomic emission spectrometry) measurement of Ca was performed by diluting 100 µL of the original sample (2 mL) into 2 mL of ultrapure Millipore water. The samples were subsequently diluted to Ca concentrations similar to the standard solutions. Calibration was conducted following the method described by de Villiers et al. (2002) using standards with constant Ca concentrations and different Sr concentrations to provide Sr/Ca ratios ranging from 0.75 to 4 mmol/mol. All measurements were conducted using the ICP-AES (Thermo ICAP DUO 6300) in the Geology Department at the University of Oviedo with a reproducibility better than 0.02 mmol/mol. To infer any possible contamination other metals, such as Fe and Mg, were also measured together with Sr.

3.3 Extraction of paleoproductivity record from the CF Sr/Ca ratios

Culture studies found a temperature dependence of coccolith Sr/Ca with a 0.03 mmol/mol increase per degree Celsius rise (Müller et al., 2014; Stoll et al., 2002a). This signal can, however, be removed from the Sr/Ca record using an independent SST reconstruction, so that the component of variation due to growth rate remains as residual (e.g. Cavaleiro et al., 2018; Mejía et al., 2014; (e.g. Cavaleiro et al., 2018; Mejía et al., 2014; Saavedra-Pellitero et al., 2017). The partitioning of Sr into the coccolith calcite varies among species (with larger and more heavily calcified coccoliths generally having a higher Sr content than smaller and lighter coccoliths; Fink et al., 2010; Stoll et al., 2007). Although it has been demonstrated that CF Sr/Ca ratios are not primarily controlled by variations in the coccolith assemblage in the modern ocean (Barker et al., 2006; Stoll and Schrag, 2000), the species composition, especially regarding the abundance of heavily calcified coccolithophores (e.g. *Calcidiscus leptoporus*, *Heliocapsa carteri*, *Coccolithus pelagicus*), should always be assessed. Our study period falls within the *Gephyrocapsa caribbeanica* acme (Baumann and Freitag, 2004; Bollmann et al., 1998), described between MIS 14/13 and MIS 8 when this species dominated (≥ 50 %) the coccolithophore community globally. In core MD03-2699, the gephyrocapsids contribute more than 92 % to the flora (on average 97 %; Fig. 2; Amore et al., 2012), which minimizes the bias of the CF Sr/Ca data due to significant coccolith assemblage changes. The sum of the remaining coccoliths, which may contain the larger and Sr-rich coccoliths *Calcidiscus leptoporus*, *Coccolithus pelagicus*, and *Heliocapsa carteri*, averaged 2.8 % (maximum 8.1 %, standard deviation 1.6 %). If we consider the abundance of the larger and Sr-rich coccoliths *Calcidiscus leptoporus*, *Coccolithus pelagicus*, and *Heliocapsa carteri*, the average and maximum abundances were, respectively, 1 % and 3.6 %, 0.1 % and 0.3 %, and 0.2 and 1.5 %. Their contribution to the total Sr of the coccolith fraction is, therefore, negligible when compared to the gephyrocapsids’ contribution.

With the assemblage changes considered we then moved to the temperature effect extraction. This was extracted from the CF Sr/Ca ratios record following Mejia et al. (2014) and Cavaleiro et al. (2018). We used the multi-species temperature dependence to calculate an SST-predicted Sr/Ca curve, and the extraction of the temperature effect consisted in subtracting the SST-predicted Sr/Ca values from the initial CF Sr/Ca ratios in Eq. (1):

\[
\frac{\text{Sr}}{\text{Ca}}_{\text{CF}} - \frac{\text{Sr}}{\text{Ca}}_{\text{predicted SST}} = \frac{\text{Sr}}{\text{Ca}}_{\text{residual}}. \tag{1}
\]

We calculated the temperature dependence using the \(U_{147}^K\)-based SST record of core MD03-2699 for the same period (Rodrigues et al., 2011). The alkenones present in the sediments are today mostly produced by a few species of the class Prynnesiophyceae, mainly by the coccolithophores *Emiliania huxleyi* and *Gephyrocapsa oceanica* (Prahl et al., 1988; Prahl and Wakeham, 1987), and the \(U_{147}^K\)-based SST is therefore the best estimator of coccolithophores’ habitat temperature. After extraction of the temperature influence, the resultant residual mainly represents coccolithophore calcification and growth rate and consequently coccolithophore long-term productivity fluctuations. We refer to the Sr/Ca residual as coccolithophore productivity because after correcting for the temperature changes, we expect the data to most reflect coccolithophore calcification and growth rate and thus their paleoproductivity qualitatively. This coccolithophore...
productivity record reflects relative productivity change, representing the productivity deviation around the average productivity of the time series.

An uncertainty envelope/confidence interval for the CF Sr/Ca residual estimation (shown as a grey envelope in Fig. 3) was calculated using Astrochron (Meyers, 2014) where the upper and lower limits correspond to the Monte Carlo 20 and 80 % confidence interval, respectively. One hundred Monte Carlo simulations were run for each of the 183 data points carrying out propagation of errors accounting for uncertainties measurement of: (1) temperature (with \( \sigma = 1.5^\circ C \)); (2) Sr concentration (with \( \sigma = 0.02 \text{mmol/mol} \)); and (3) the linear regression of temperature versus Sr (\( \sigma = 0.12 \text{mmol/mol} \)).

4 Results

4.1 Potential influence of detrital carbonate on the CF Sr/Ca data

When present, detrital carbonate could negatively bias any CF Sr/Ca ratio (and any coccolithophore productivity reconstruction) by increasing the background carbonate in relation to the coccolith carbonate. In our study, we considered potential sources of detrital carbonate biasing the CF Sr/Ca results: (1) ice-rafted debris (IRD) discharged by melting icebergs, (2) aeolian input, and (3) riverine discharge. Deposition of carbonate/dolomite-rich IRD is linked to extremely cold and lower-salinity conditions (Fig. 4), namely Heinrich-type events (HTEs; Andrews and Voelker, 2018; Hodell et al., 2008; Rodrigues et al., 2011; Salgueiro et al., 2010; Stein et al., 2009). Ice-rafted material, especially the fine-grained material, also contains reworked coccoliths, which can be Tertiary or Cretaceous in age (Marino et al., 2011, 2014; Rahman, 1995). Increased aridity and wind conditions enhance aeolian transport (e.g. Bozzano et al., 2002), which can transport detrital carbonate material, coming mostly from North Africa (e.g. Negral et al., 2012; Stumpf et al., 2010), to the IbM. On the other hand, increased moisture and precipitation intensifies rock weathering and riverine discharge, raising fluvial transport of detrital carbonate onto the IbM (Hodell et al., 2017; Hodell and Channell, 2016). However, the IbM documents that an increased detrital carbonate presence is prevalent during cold and arid rapid millennial-scale events related to low sea-level stands and IRD events (Lebreiro et al., 2009; Marino et al., 2011; Hodell et al., 2013, 2017; and Hodell and Channell, 2016).

In order to evaluate this potential biasing effect we match the CF Mg/Ca values in a cross-plot with the CF Sr/Ca results. We used CF Mg/Ca because Mg/Ca peaks can be attributed to the presence of increased detrital carbonate, such as dolomite (CaMg(CO\(_3\))\(_2\)). The Mg/Ca threshold was defined at 30 mmol/mol because from that point onwards both CF Mg/Ca and Sr/Ca show a clear negative relationship: i.e. CF Mg/Ca increases when CF Sr/Ca clearly decreases (Fig. 5a). Therefore the samples highlighted in Fig. 5b with a red circle were considered possibly contaminated by detrital carbonate. We estimate that (i) 16 % of the samples might have been biased by the presence of detrital carbonate and confirm that (ii) such samples are linked to rapid millennial-scale and low sea-level stands, namely during substages MIS 10a, MIS 12a, and the respective subsequent terminations V and IV (Fig. 4). Although a possible bias effect exists in such samples, it does not prove that coccolith Sr/Ca ratio and coccolithophore productivity were not indeed significantly lower during those intervals (see Discussion). We have considered the bias from reworked coccolith contribution to be negligible because reworked coccolith abundance was generally below 2 %, not only in our study site (Amore et al., 2012) but also at other IbM sites for the same time interval (Maiorano et al., 2015; Marino et al., 2014).

4.2 Coccolithophore productivity results

The CF Sr/Ca ratio results varied between 1.2 and 2.4 mmol/mol (sampling range of 1.16 mmol/mol) with 85 % of the samples falling between 1.8 and 2.3 mmol/mol. The Sr/Ca residual, from now on referred to as coccolithophore productivity (CP), generally mimics the CF Sr/Ca ratio record and shows minima and maxima at the same levels (Fig. 3). The high CF Sr/Ca ratios found in the record do not coincide with increased abundance of large/Sr-rich coccoliths (Fig. 2) or with higher temperatures (Fig. 3). This supports the statement that our CF Sr/Ca variation reliably records coccolithophore calcification rate and productivity changes. Both MIS 12 and MIS 10 are characterized by higher CP during the first half and lower productivity towards the second half. Large and rapid increases in CP characterize the deglaciations at the end of MIS 12 and MIS 10.

To better visualize and interpret the CF Sr/Ca results, the CP data were divided into three relative intervals. An “intermediate” interval is defined by the mean of the data plus and minus half of the standard deviation. The “high” and “low” CP levels are above and below the intermediate level of CP, respectively (highlighted by green and blue shading in figures, respectively). We would like to stress that our study focuses on the qualitative characteristics of the coccolithophore paleoproductivity record rather than quantitatively estimating the productivity of coccolithophores.

A preliminary visual inspection of the record already allows the identification of higher-amplitude events during glacial substages MIS 12b and 12a and MIS 10b and 10a in comparison to the less variable interglacial substages MIS 11c and MIS 9e. CP maxima happened during the first half of MIS 12 and MIS 10 and during the transition from interglacial to glacial substages (Fig. 3). Productivity minima, on the other hand, occurred during glacial maximum substages MIS 12a and MIS 10a and at the end of MIS 11a. Marine Isotope Stages 11c and 9e, commonly known as the interglacial substages (Past Interglacials Working Group
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5 Discussion

5.1 Causes for CP change

5.1.1 High CP levels during the transition from interglacial to glacial periods

CP increased steadily from mid-MIS 11c until the end of MIS 11b and remained generally high from late MIS 11c to MIS 10c, prior to the glacial substages within MIS 10 and MIS 12. We suggest this increasing CP is a consequence of a gradual rise in nutrient availability due to either enhanced northerly winds and upwelling strength or due to the decreasing influence of subtropical waters, substituted by increasing influence of transitional nutrient-richer surface waters (i.e. ENACWsp). During interglacial substages, such as MIS 11c and MIS 9e, the wind stress associated with the upwelling events has been suggested to be lower than during glacial substages in the IbM (e.g. Salgueiro et al., 2010). This is based on the observation that along with the expansion of the Northern Hemisphere continental ice sheets, atmospheric and oceanic circulation changed leading to a narrowed latitudinal temperature gradient and potentially enhanced northerly winds and upwelling intensity (Broccoli et al., 2006; Hostetler et al., 1999). Note, however, that for upwelling to be intensified during transitions from interglacial to glacial substages and during glacial substages, it would have required that the atmospheric circulation pattern governed by the relative position and gradients between high- and low-pressure centres have remained similar to today (i.e. interglacial substage). Based on differences in the planktonic oxygen isotope records of cores MD03-2699 and MD01-2446 (more offshore) (Fig. 1), Voelker et al. (2010) suggested that the higher variability in our core, MD03-2699, could reflect variations in the upwelling of deeper waters into the thermocline. Stronger winds would favour upwelling intensification, which could lead to the upwelling of the deeper, nutrient-richer ENACWsp on the IbM (Fiúza, 1984), as already suggested for some Late Pleistocene periods (Salgueiro et al., 2010, 2014). Still, the transition period is characterized by a decrease in SSTs most likely already reflecting the southward displacement of the North Atlantic frontal system and, consequently, increasing the influence of an ENACWsp origin, more nutrient rich than an ENACWst origin, in the surface water of western IbM. Increased nutrient availability would thus support an increase in productivity of the whole phytoplankton community (Álvarez-Salgado et al., 1997; Guerreiro et al., 2013), and after depletion of silica by diatoms, coccolithophore species capable of rapid growth, as Gephyrocapsa caribbeanica is thought to have been, given its cosmopolitan distribution and dominance in the sediments (Baumann and Freitag, 2004; Bollmann et al.,

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Figure 4. Coccolithophore paleoproductivity reconstruction with “high” and “low” coccolithophore productivity levels highlighted in green and blue shading, respectively (a), compared to (b) U$^{233}$-based reconstruction of sea surface temperature from core MD03-2699 (Rodrigues et al., 2011); (c) ice-rafted debris (IRD) abundance (Voelker et al., 2010) and heptatriatetraenone (C$^{37}$:4 alkenone) abundance (Rodrigues et al., 2011), both from core MD03-2699; (d) Neogloboquadrina pachyderma relative abundance and Gephyrocapsa margereli and G. muellerae group relative abundance, both from core MD01-2446 (Marino et al., 2014); (e) percentage IRD from ODP Site 980 (McManus et al., 1999) and dolomite/calcite ratio from IODP Site U1313 (Stein et al., 2009). Note that the MD03-2699 lithic fragments were counted in a coarser size fraction (> 315 µm) than the standard size fraction (> 150 µm; Hemming, 2004) and thus most likely only record major ice-rafting events at the IbM, as suggested by Marino et al. (2014). Chronology as in Fig. 3. Vertical bars: grey bars correspond to Heinrich-type events (HtEs) and blue bars to short-lived events of decreased coccolithophore productivity.

1998), could bloom. It is important to refer that such CP maxima would only be possible if bioavailable Si and Fe were not sufficiently high for diatoms to outcompete coccolithophores until complete nutrient depletion. Maybe seasonality could also have played an important role here (see intermediate CP level discussion below). As coccolithophores perform better in stable and continuously nutrient-replenished surface waters, in contrast with diatoms that thrive in dynamical (e.g. upwelling) conditions, (Cermeño et al., 2011), we believe that, during the transition to full glacial conditions, persistent and increasing nutrient replenishment on the IbM was more related to the advection of transitional waters of subpolar origin than to increased upwelling. Finally, our high CP supports the idea that the coccolithophore community was able to better perform under these transitional conditions, despite colder SST, and possibly more turbulent settings than during interglacial substages. Noteworthy is the fact that this maximal CP interval was frequently interrupted by abrupt cooling events (Fig. 6) when CP also abruptly decreased. Though detrital carbonate influence cannot be excluded (see methodology), such CP reductions might indicate that coccolithophore productivity was close to its lowest SST threshold and that
Figure 5. Detrital carbonate analysis: (a) coccolith fraction (CF) Sr/Ca results and CF Mg/Ca cross-plot (vertical red line delimits threshold at 30 mmol/mol of Mg/Ca); (b) CF Sr/Ca samples possibly contaminated with detrital carbonate (red makers) and notably associated millennial abrupt events.

Figure 6. Coccolithophore paleoproductivity reconstruction with “high” and “low” CP levels highlighted in green and blue shading, respectively (a), compared to (b) $U_{137}^{14}$-based reconstruction of sea surface temperature (Rodrigues et al., 2011), (c) nannofossil accumulation rate (NAR) (Amore et al., 2012), and (d) total alkenone (AlkAR) and n-alkane accumulation rates (Rodrigues et al., 2011). Chronology as in Fig. 3. Vertical bars: grey bars correspond to Heinrich-type events (HtEs) and blue bars to short-lived events of decreased CP.
abrupt drops in SST immediately resulted in abrupt CP declines.

5.1.2 Intermediate CP levels associated with deglaciations and interglacial periods

The rapid CP increase during deglaciation is contemporary with a rapid SST rise (T V and T IV; Fig. 3). The deglacial SST increase and the subsequent particularly stable SSTs during MIS 11c (and most likely MIS 9e) recorded in our MD03-2699 core are linked to the increasing influence and dominance of subtropical oligotrophic waters from the Azores and Iberian Poleward Currents (Rodrigues et al., 2011; Voelker et al., 2010). Several IbM sites support this with increased abundance of warm coccolith taxa (site MD03-2699; Amore et al., 2012 and Palumbo et al., 2013a; IODP Site U1385, Maiorano et al., 2015; site MD01-2446, Marino et al., 2014) and of tropical planktonic foraminifera species Globorotalia menardii and Sphaeroidinella dehiscens (site MD03-2699; Voelker et al., 2010; site MD01-2443; de Abreu et al., 2005) during the deglaciations and MIS 11c and MIS 9e. Despite the large amplitude change in the CP during the deglaciation, a comparably larger amplitude change is associated with glacial stage MIS 12a and the transitional stage MIS 11b, when SSTs were comparably lower than during MIS 11c (see previous subchapter). Thus, we suggest that during the deglaciation CP increased sharply because of the amelioration of the oceanographic conditions (mostly due to increasing SST) but their productivity was limited to intermediate levels of productivity because of either limited nutrient supply from the prevalent advected subtropical oligotrophic waters or upwelled waters (1) and/or increased competition for nutrients with diatoms (2), which mostly affected the coccolithophore species with opportunistic and fast-growing characteristics (r-selected species) such as gephyrocapsids. Such competition with diatoms could be related to upwelling intensity (Cermeño et al., 2011) during interglacial stages MIS 11c and MIS 9e. During the current interglacial, upwelling only happens in the IbM during the spring–summer months, when winds, mostly driven by trade winds, are strong and predominantly from the north–northwest (Fiúza et al., 1982; Fiúza, 1983; Mason et al., 2006). Given the current variability in the western IbM (Alvarez et al., 2008; Arístegui et al., 2009) and its dependency on changes in the regional atmospheric circulation, especially related to the latitudinal migration of the subtropical front and to the dynamics of the Azores anticyclonic cell (Arístegui et al., 2009; Fiúza et al., 1982; Mason et al., 2006), it is difficult to assert what has been the wind regime and upwelling seasonality during the deglaciations. However, when assessing the hypothesized upwelling strength between interglacial and glacial substages, and in the event of lower nutrient availability for the rest of the year, coccolithophores would be easily outcompeted by diatoms, as in the upwelling event of the current interglacial (e.g. Abrantes and Moita, 1999).

This evidence suggests that irrespective of the upwelling regime that influenced seasonal overall nutrient availability and phytoplanktonic productivity, the intermediate levels of CP could be explained by a competitive advantage of diatoms upon sufficient delivery of Si and Fe of terrigenous input, either by increased aeolian input or riverine discharge. Since changes in the SST and nutrient availability are not only promoted by fluctuations in the upwelling regime (1) and changes in the advection of the dominant water masses (2) in the IbM, we also considered changing seasonality, i.e. changes in precipitation (3) and arid plus windy (4) conditions to have played a role in determining CP. It is well known that the addition of Fe into the surface ocean positively affects the overall phytoplankton community (Blain et al., 2004; Martin et al., 1990), and diatoms require more Si and Fe (e.g. Merico et al., 2004), whereas coccolithophore blooms are thought to be linked with Si and Fe depletion (Merico et al., 2004; Tyrrell and Merico, 2004) giving coccolithophores a competitive advantage in an Si and Fe depletion scenario with plenty of other nutrients, such as nitrate and phosphate. Thus, under fluctuating nutrient availability and increased bioavailability of Si and Fe (Capellacci et al., 2013), diatoms could have outcompeted coccolithophores via nutrient competition, limiting coccolithophore growth to intermediate levels of productivity after diatom bloom. Periods of strengthened wind regime and aridity are generally limited to glacial substages and stadials (Desprat et al., 2009, 2017; Hodell et al., 2013b; Oliveira et al., 2016; Sánchez Goñi et al., 2016). But, Rodrigues et al. (2011) interpreted the synchrony between high terrigenous input (concentration of n-alkanes) and alkenone concentration in core MD03-2699 (Fig. 6d) with intensified dust export from North Africa (based on Fe counts at Site 958 off western North Africa; Helmke et al., 2008) as indicative of a strong wind regime in the IbM during the deglaciation and early phase of MIS 11c. However such high Fe counts evidencing strong winds are referred by Helmke et al. (2008) for the deglaciation but not for MIS 11d and early MIS 11c, where the same study suggests a humid period in North Africa associated with increased precipitation and a widespread vegetation cover. So during the deglaciation such strong winds might have been sufficient to replenish the surface waters with Si and Fe, allowing diatoms to bloom (Abrantes, 2000; Capellacci et al., 2013) and outcompeting coccolithophores. And, in the early MIS11c, Si and Fe could have also come into the IbM through riverine discharge.

Under full interglacial conditions, pollen records evidence the development of temperate and humid deciduous forest in north-western Iberia (Desprat et al., 2007), which during MIS 11c coincided with the North African humid period described by Helmke et al. (2008). Based on the increased abundance of Mediterranean forest taxa observed in south-west IbM IODP Site U1385, Oliveira et al. (2017) associated
climate during MIS 11c with the present-day SW Iberian climate, i.e. with pronounced seasonality between warm and dry summers and cool and wet winters. The warmer and drier summers, tied to the north-eastward expansion of the Azores subtropical High, associated with the descending branch of the Hadley cell (Lionello et al., 2006; Oliveira et al., 2017), could have enabled dust flux to the western IbM. During winters, on the other hand, nutrients in the surface waters could have been replenished by increased precipitation and associated river run-off, in agreement with the expansion of temperate humid forest. So, during interglacials, our theory that periods when CP seemed limited to intermediate levels are linked to conditions of increased terrigenous input, either aeolian or riverine, that allowed diatoms to out-compete coccolithophores is also valid.

5.1.3 Low CP associated with abrupt cooling events and glacial MIS 12a

CP shows consistent minima along the record that, with the exception of glacial maximum stage MIS 12a and the ∼10 kyr interval at the transition from MIS 11a to MIS 10c, are of short duration and related to abrupt climate change events. Thus, most of the minima coincide with the presence of colder and less saline surface waters that resulted from meltwater incursions onto the IbM evidenced either by the increased IRD content or higher percentages of C_{37:4} alkenones, concomitant with higher abundances of coccoliths belonging to *Gephyrocapsa margereli*–*G. muellerae* (> 4 µm), a cold/low-salinity water proxy in the IbM (Fig. 4; Marino et al., 2014; Rodrigues et al., 2011). The most extreme of these events are the HtEs (Stein et al., 2009; Fig. 4), with the ones during T IV and T V corresponding to the terminal stadial events of Hoddell et al. (2015). The associated cooling brought the SSTs nearly to local glacial levels (Fig. 4; Rodrigues et al., 2011). Similar to their counterparts during the last glacial cycle, the meltwater events resulted in an accentuated decrease or nearly the elimination of the Atlantic Meridional Overturning Circulation and in a southwards displacement of the North Atlantic frontal system with the subpolar/arctic front moving into IbM latitudes (Alonso-Garcia et al., 2011; Hoddell et al., 2008; McManus et al., 1999; Rodrigues et al., 2011, 2017). This led to the advection of polar and subpolar water masses onto the IbM evidenced by the increased percentage of polar planktonic foraminifer *Neogloboquadrina pachyderma* in the nearby core MD01-2446 (Fig. 4; Marino et al., 2014) and in cores off Galicia (Desprat et al., 2007, 2009).

The abrupt CP drops at 438 ka and at 340 ka prior to and during T V and T IV, respectively, are both related to HtEs (Fig. 4). As previously mentioned, during HtEs detrital carbonate might have negatively biased our CP. However, detrital carbonate might have acted as a low-productivity signal amplifier instead of attributing false low CP in such intervals. An extreme decline in CP during the TV HtEs/terminal stadial event was also observed at IODP Site U1313 in the central North Atlantic basin (Cavaleiro et al., 2018). During those events the abrupt SST decrease coupled with increased turbidity and blockage of sunlight (due to melting icebergs and/or sea ice cover) would have decreased the coccolithophores’ ability to survive. These conditions resemble those in the present-day polar domain of the Norwegian–Greenland Sea (Baumann et al., 2000). Here, a less diverse coccolithophore community was observed with a slightly diminished coccolithophore population, which could well support the abrupt decreases detected in our CP record.

Additional CP minima are detected at 389, 382, 371, 364 (the latter dropping to HtE levels; Fig. 4), 346, 334, 319, and 316 ka. Despite no significant concomitant IRD peaks in core MD03-2699, these decreases are associated with increases in the abundance of the *C_{37:4}* alkenone and colder SST (Fig. 4; Rodrigues et al., 2011) and the presence of *N. pachyderma* and *G. margereli–G. muellerae* (> 4 µm) in core MD01-2446 (Fig. 4; Marino et al., 2014), testifying to a rapid change to significantly colder and less saline conditions at this latitude. The four older drops in CP happened concomitantly with abrupt increases in *Neogloboquadrina pachyderma* abundance at ODP Site 980 (55°29′N, 14°42′W) in the eastern North Atlantic (Fig. 4; McManus et al., 1999; Oppo et al., 1998), placing them in a broader spatial scale of climate deterioration. Thus, we conclude that the IbM was subjected to drastic surface water changes that led to a coccolithophore community decrease in the IbM and generated abrupt decreases in our CP record.

Finally, the reduced productivity during MIS 12a at the location of core MD03-2699 is in stark contrast to the open-ocean record of IODP Site U1313 (Cavaleiro et al., 2018) and other past glacial periods in the IbM such as MIS 2 and MIS 6 (Salgueiro et al., 2010, 2014), all of which point to increased productivity during the glacial maxima. Furthermore, such prolonged low CP is also not observed during glacial maximum MIS 10a, when productivity at site MD03-2699 remained at intermediate levels between abrupt cold events (Fig. 4). However, the prolonged presence of (sub)polar waters in the eastern North Atlantic throughout MIS 12a is backed up by the IRD record and persistent high percentages of *N. pachyderma* (Oppo et al., 1998; McManus et al., 1999) at ODP Site 980 and in core MD01-2448 in the Bay of Biscay (Toucanne et al., 2009). It is thus likely that such low-temperature, and eventually salinity, water masses also affected the IbM, hampering upwelling (enhanced upwelling is seen as the cause for increased productivity at the IbM during glacial maxima; Abrantes et al., 2000; Salgueiro et al., 2010, 2014) and potentially leading to a low CP.
5.2 Evidence from different coccolithophore productivity proxies and conceptual models

Previous studies in the IbM used coccolith-derived proxies, such as coccolith assemblages, NARs, and total alkenone fluxes (produced mostly by coccolithophores) to reconstruct changes in coccolithophore paleoproductivity (Amore et al., 2012; Maiorano et al., 2015; Marino et al., 2014; Palumbo et al., 2013). These traditional proxies depend not only on the supply of coccoliths or organic compounds but also on dilution by minerals and other sediment constituents and on changing preservation conditions (Rullkötter, 2006). As the CP Sr/Ca ratio is an independent proxy, i.e. independent of sedimentation and accumulation rates, it offers a new perspective on CP dynamics and the opportunity to compare coccolithophore growth(calcification rate) with other components of the system. In a simple time series comparison of our CP data with the NAR or the AlkAR records we can already confirm that their relationship is not straightforward (Fig. 6). While in some intervals CP, NAR, and AlkAR show similar trends, in others they diverge.

As mentioned previously, several studies have hypothesized a relevant role of coccolithophores on the global carbon and carbonate cycles and on the glacial–interglacial dynamics (Duchamp-Alphonse et al., 2018; McClelland et al., 2016; Omta et al., 2013; Rickaby et al., 2007; Saavedra-Pellitero et al., 2017). Yet, the inclusion of biogeochemical processes related to coccolithophores in Earth system models is lacking, although they play a role in both the organic (soft tissue) and the carbonate counter pump (through calcification). Coccolith carbonate also has a ballasting effect, enhancing the Corg export efficiency (Armstrong et al., 2001; Francois et al., 2002), but the relative proportion of exported particulate organic carbon to inorganic carbon is difficult to predict and model (Barker et al., 2006; Ridgwell and Zeebe, 2005). Therefore, the respective and relative importance of these mechanisms is still poorly understood and the past and future net impact of calcification by coccolithophores on atmCO2 is still uncertain (Lawton et al., 2003). Among the difficulties in modelling coccolithophores’ responses are the complex interactions between coccolithophore productivity and growth rates and nutrient dynamics, non-calcifiers competition, seasonality, and their response to upwelling/non-upwelling conditions.

We suggest that the contemporaneous coupling or decoupling of the different coccolithophore productivity proxies reflect changes in the carbon and carbonate system and conceived conceptual models of the carbon and carbonate cycles in the IbM (Fig. 7).

During the MIS 12 deglaciation all coccolithophore proxies show similar trends. This deglaciation is marked by clear rapid increases in CP, NAR, and AlkAR, accompanied by a rapid, high-amplitude rise in SST, characteristic of terminations (e.g. Rodrigues et al., 2017). The increasing CP is linked to the growing influence and dominance of subtropical oligotrophic waters from the Azores and Iberian Polar Currents (SST evidence) combined with a moderate to strong wind regime and consequent upwelling (Voelker et al., 2010). Based on paleoclimatic evidence, deglacial hinterland conditions in western Iberia and NW Africa were arid and resulted in increased (Saharan) dust flux (Desprat et al., 2007; Helmke et al., 2008; Hodell et al., 2013a, 2015; Rodrigues et al., 2011), which would supply high amounts of Fe and Si to the IbM. We suggest that this increased nutrient combination fertilized the non-calcifying phytoplankton component, namely diatoms (Duchamp-Alphonse et al., 2018; Helmke et al., 2008; Meckler et al., 2013; Rodrigues et al., 2011; Thomson et al., 2000), attributing to them a competitive advantage over coccolithophores, outcompeting them (e.g. Abrantes and Moita, 1999; Balch, 2004; Guerreiro et al., 2013). The combined increased surface productivity with both diatoms and coccolithophores blooming would likely allow for an increase in the soft-tissue pump into the deep ocean. During the deglaciation and early interglacial phases, the augmented (though only to intermediate levels) coccolithophore productivity was associated with higher calcification rates, and thus CP, which resulted in a higher NAR. The higher CaCO3 production at the surface might have led to an increased CaCO3 and Corg export efficiency through the ballast effect of CaCO3 (e.g. Armstrong et al., 2001; Francois et al., 2002). This could explain the enhanced AlkAR as well as NAR at our study site for the deglaciations and early MIS 11c and MIS 9e, though with significantly different amplitudes.

During mid-interglacial MIS 11c and MIS 11b (410 to 390 kyr) and MIS 9e, all coccolithophore productivity proxies steadily increase, most likely associated with decreasing competition with diatoms for nutrients (see previous Discussion section). However the quite different AlkAR records for the deglaciation and early MIS 11c (428 to 410 kyr) in comparison with the rest of MIS 11c and MIS 11b (410 kyr to 390 kyr) suggest a change in the phytoplanktonic community, favouring coccolithophores to the detriment of diatoms. We suggest that while diatoms were receiving a surplus replenishment of Si and Fe (as discussed in Sect. 5.1.2), the soft-tissue pump became more efficient because more organic particulate matter and particulate inorganic matter was present to form fast-sinking aggregates (Armstrong et al., 2001; Francois et al., 2002) allowing also for an increasing amount of alkenones to be exported and accumulated. We hypothesize that the timing of the rapid decrease in AlkAR could actually coincide with the decrease in the surface particulate organic matter as a consequence of the decreasing diatom productivity and surplus replenishment of Si and Fe, which ceased by 410 ka, when AlkAR reach their minimum. This could have important implications for past CO2 fluxes from the atmosphere into the deep ocean, if indeed the soft-tissue pump increased its efficiency due to this combination of factors.
Figure 7. Conceptual models for the IbM of the different components of the carbon/carbonate cycle under different climatic scenarios: D/T, Tr, IGs, and Gs stand for deglaciation/termination, transition from interglacial to glacial substages, interglacial substages, and glacial substages, respectively; ENACWst, ENACWsp, NADW, and MOW stand for Eastern North Atlantic Central Water of subtropical or sub-polar origin, North Atlantic Deep Water, and Mediterranean Outflow Water, respectively. In the deglaciation scenario we depict increased surface productivity, associated with peaks of diatom and intermediate levels of coccolithophore productivity, and the largest marine snow export. Interglacial substages depict a decreased surplus of nutrients from terrigenous input, a decreased abundance of diatoms, and intermediate levels of coccolithophore productivity due to the prevalence of ENACWst origin and a large decrease export; Transitions show a general increase in phytoplankton but placing more emphasis on coccolithophores because of the increasing and persistent predominance of ENACWsp origin, with intermediate levels of export. Glacial substages show a general decline in the phytoplankton associated with export minima and the increased influence of MOW.

From mid-MIS 11c to the end of MIS 11b (410 ka and until 390 ka), all coccolithophore productivity proxies increase, and we suggest that this coupled signal reflects that the coccolithophore productivity had a diminished competition with diatoms for nutrients and an associated less efficient soft-tissue pump in comparison with the deglaciation and early MIS 11c.

Based on the evidence from IODP Site U1313 in the open midlatitudinal North Atlantic (Cavaleiro et al., 2018), increases in coccolithophore calcification rate and productivity do not necessarily have to be a consequence of enhanced upwelling but can just as likely be related to the southward migration of the high-productivity/high-coccolithophorid belt (McIntyre et al., 1972; Villanueva et al., 2001). This band, currently located between 45 and 55° N (Antoine et al., 1996; Henson et al., 2009; Longhurst et al., 1995), is associated with the convergence zone between the subpolar and subtropical gyres (McIntyre et al., 1972). Along with the southward expansion of the subpolar gyre during periods of slowdown of the North Atlantic’s thermohaline circulation (Alonso-Garcia et al., 2011; Rodrigues et al., 2017; Wright and Flower, 2002), the high-productivity band moved into the latitudes of the western IbM. So, the (gradual) displacement of this high-coccolithophorid belt could explain the rapid rise in CP productivity during MIS 11b and associated maxima during MIS 11a, MIS 10c and MIS 9c when all three records – CP, NAR, and AlkAR – show contemporary increases (Fig. 6). Another factor affecting the signal preservation during the glacial periods and abrupt cold events of MIS 11b to MIS 11a and of MIS 9d to MIS 9c could be shifts in the conditions prevailing at the sea floor. Whereas site MD03-2699 was bathed by NADW (North Atlantic Deep Water) when the North Atlantic’s thermohaline circulation was strong, i.e. during the interglacial substages and the interstadials of MIS 11a, the North Atlantic Deep Water was replaced by Mediterranean Outflow Water (MOW) during times of a weak thermohaline circulation (stadials of MIS 11b and MIS 11a; glacial MIS 12a and most of MIS 10) (Veelker and Lebreiro, 2010). Periods of MOW presence were associated with increased bottom current activity at site MD03-2699, especially during MIS 12a, when the winnowing of finer particles could have contributed to the low NAR and AlkAR (Fig. 6). On the other hand, the less oxygenated MOW would induce less Corg remineralization, which could have played a role in the AlkAR maxima occurring during late MIS 11a.
5.3 Further research suggestions

The CP reconstruction for the mid-Brunhes interval at site MD03-2699 yielded important findings but also raised issues that should be explored in the future. Extending this research into other interglacial substages could help disentangle the phytoplankton community response to upwelling intensity, namely if both diatoms and coccolithophores increase or if one decreases to the detriment of the other with changing upwelling intensities. This would offer further insights into the role of particulate organic vs. inorganic production in the surface ocean in an upwelling area such as the IbM. Further attention should also be given to the role of seasonality in CP and diatom productivity dynamics in the IbM, which could potentially be disentangled by using a direct land–sea correlation, namely evidence from pollen records. Given the discrepancy between our and other surface productivity and export estimations for glacial periods on the IbM (Abrantes, 1992; Salgueiro et al., 2010, 2014), it would be interesting to assess coccolithophore productivity in Late or early Pleistocene glacial stages to assess if our observation of low CP is restricted to mid-Brunhes conditions.

Additional research, specially incorporating the information gained into carbon and carbonate cycle models, could help to better predict the role of the IbM and other (eastern boundary) upwelling systems in glacial–interglacial atmCO₂ changes. Did the coupled high productivity of non-calcifiers and calcifiers during deglaciations operate as sink of carbon or did the carbonate counter pump act to balance or even counteract the amplified soft-tissue pump?

Finally, Omta et al. (2013) presented a potential role of coccolithophores in the glacial–interglacial cycles of atmCO₂ and based their experiments on prey–predator models with the characteristic spiking predator (like coccolithophores) and a sawtooth-shaped curve of the prey behaviour (like alkalinity, which is positively and directly correlated to [CO₃²⁻]). Despite the undeniable direct link between calcifying phytoplankton and ocean alkalinity (e.g. Ridgwell and Zeebe, 2005), their model with the spikes of enhanced productivity at deglaciations/terminations actually resembles the non-calcifier behaviour in the IbM and off Africa, namely diatoms (Meckler et al., 2013; Thomson et al., 2000), more than that of coccolithophores, as our record shows. Coccolithophore productivity in the IbM margin seems to rise abruptly during the deglaciations, followed by a constant level or slight increase during the interglacial substages, and then steadily increases further during the progressive global cooling towards the glacial maximum only to then abruptly decrease under full glacial conditions (Fig. 7).

We hope that this information could be included in future modelling experiments, with both coccolithophore and diatom components of the dynamic phytoplanktonic community, to better describe the role of each functional group in the carbon and carbonate cycles in this upwelling system and with possible implications for other eastern boundary upwelling systems. We urge climate scientists to research the combined responses of the different components of the phytoplanktonic community, namely calcifiers and non-calcifiers. Much research has been conducted exploring only the productivity changes of one of these functional phytoplanktonic groups. Since the global ocean phytoplanktonic community with a potentially greater impact on atmCO₂ fluxes is mostly composed of diatoms and coccolithophores in either the open ocean or upwelling areas, it is important to understand how the phytoplankton community dynamics have changed in the past.

6 Conclusions

In this study we applied the CF Sr/Ca ratio to reconstruct CP during the mid-Brunhes interval of MIS 9 to MIS 12 and discussed the different factors controlling CP in the western IbM. We found that the long-term coccolithophore productivity was primarily controlled by changes in the paleo-
oceanographic surface conditions, namely the origin of the predominant surface waters and their nutrient content. CP maxima were found during the transition from interglacial to glacial substages and associated with increasing presence of nutrient-richer transitional surface waters. The CP maxima during this transition were, however, frequently interrupted by abrupt decreases in CP linked to abrupt cooling events (HtEs) and a growing predominance of polar and subpolar surface waters, such as during glacial maximum substages. The deglaciations during early MIS 11c were marked by intermediate CP linked to the increased influence of subtropical, nutrient-poorer waters during winter, combined with increased terrigenous input of nutrients and moderate to strong upwelling during summer, conditions that are hypothesized to have an attributed to a competitive advantage of diatoms over coccolithophores.

Using a multiproxy approach and gathering CP (surface production signal) and AlkAR and NAR (bottom export and accumulation signals), we infer conceptual models of the carbon and carbonates cycles in the IbM during different periods and suggest future research questions to be tested through modelling. We hope that our new data can encourage Earth system modellers to look deeper into constraining the role of coccolithophores in the regional and global carbon and carbonate cycles, especially in the eastern boundary upwelling systems of the world.

**Data availability.** The data used will be available from the Pangaea data repository at https://doi.org/10.1594/PANGAEA.908192

**Author contributions.** CC and AV designed the study. CC prepared the samples and analysed them under the supervision of HS. CC drafted the paper, and all authors contributed to the discussion and to the final version.

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