Environmental boundary conditions of cold-water coral mound growth over the last 3 million years in the Porcupine Seabight, Northeast Atlantic

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IODP Expedition 307 made it for the first time possible to investigate the entire body of a cold-water coral carbonate mound. Here we provide new insights into the long-term history of Challenger Mound on the European continental margin off Ireland. This study is based on age determinations ($^{230}$Th/U, $^{87}$Sr/$^{86}$Sr) and geochemical signals (Mg/Li and Ba/Ca) measured in the scleractinian cold-water coral \textit{Lophelia pertusa} from IODP Site 1317 in the Porcupine Seabight. The paleoceanographic reconstructions reveal that coral growth in the Porcupine Seabight was restricted to specific oceanographic conditions such as enhanced export of primary production and Bottom-Water Temperatures (BWT) between $\sim 8$ and $10^\circ C$, related to the water mass stratification of the Mediterranean Outflow Water (MOW) and Eastern North Atlantic Water (ENAW). The geochemical signals from the coral skeletons can be explained by the close interaction between cold-water coral growth, sea-surface productivity and the surrounding water masses – the boundary layer between MOW and ENAW. Enhanced sea-surface productivity and the build-up of a stable water mass stratification between ENAW and MOW caused enhanced nutrient supply at intermediate water depths and facilitated a steady mound growth between $\sim 3.0$ and $2.1$ Ma. With the decrease in sea-surface productivity and related reduced export productivity the food supply was insufficient for rapid coral mound growth between $\sim 1.7$ and $1$ Ma. During the late Pleistocene (over the last $\sim 0.5$ Myr) mound growth was restricted to interglacial periods. During glacial the water mass boundary between ENAW/MOW probably was below the mound summit and hence food supply was not sufficient for corals to grow.

1. Introduction

1.1. Background

Cold-water coral carbonate mounds are common features in certain parts of the worlds oceans (Roberts et al., 2006). Over the last 20 years these unique ecosystems have attracted increasing interest in their origin, growth and demise. The environmental boundary conditions necessary for their initiation are still debated. The continental slopes along the European margins contain large provinces of cold-water coral mounds and reefs (Freiwald et al., 2004; Roberts et al., 2006). Here mounds are mainly built by the scleractinian cold-water corals \textit{Madrepora oculata} and \textit{Lophelia pertusa}. The appearance and distribution of these heterotrophic and filter-feeding corals is controlled by several parameters. Settlement of coral larvae only occurs on hard substrata preferentially on continental slopes, seamounts or oceanic ridges. These areas are often associated with high sea-surface productivity and enhanced current strength (e.g. Guinotte et al., 2006; White et al., 2005). In particular, \textit{L. pertusa} tolerates a wide range of temperatures (4–14 $^\circ$C) and salinities of 32–39, but mostly occurs in salinities between 35 and 37. It appears that the seawater density plays an important role in the distribution of living \textit{Lophelia}-reefs and mounds on the European continental margin. The study of Dullo et al. (2008) highlighted that vivid \textit{Lophelia}-reefs along the SW Irish and Norwegian margin tend to occur within a narrow density envelope between sigma theta ($\sigma_t$) = 27.35–27.65 (Dullo et al., 2008). However, this finding appears not to be valid for each

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Lophelia – reef as exceptions can be found for example in submarine canyons in the Bay of Biscay and in the Mediterranean Sea (e.g. Freiwald et al., 2009; Huvenne et al., 2011). Living and fossil reefs occur from northern Norway in the Barents Sea (70°N, Lindberg et al., 2007) to NW Africa off Mauritania (16°N, Colman et al., 2005). On the Norwegian margin large flourishing reefs developed after the retreat of the glaciers at the end of the last glacial, forming the largest known living cold-water coral reefs in the worlds oceans (Freiwald et al., 2004; Fossà et al., 2005). The margin southwest of Ireland represents a major region of abundant cold-water coral carbonate mounds that tend to cluster in provinces (De Mol et al., 2007; Freiwald et al., 2004) and vary in height from a few metres up to >380 m (Wheeler et al., 2007). Especially, the Porcupine Seabight is characterised by different mound provinces some of which contain over 1000 buried and exposed mounds (Huvenne et al., 2007). The Belgica Mound province is one of five mound provinces (De Mol et al., 2002; Foubert et al., 2005; Wheeler et al., 2011) and the Challenger Mound is one of 66 mounds in the Belgica Mound Province. Here, the coral mounds are elongated, subconical structures and occur at depths between 600 and 1000 m, corresponding to the transition zone of Eastern North Atlantic Water (ENAW) and the underlying Mediterranean Outflow Water (MOW) with temperatures of ~9.5 °C (White, 2007; Fig. 1). Due to the large differences in density between the ambient intermediate water masses a pycnocline forms at around ~800 m (Dickson and McCave, 1986; Dullo et al., 2008; White and Dorschel, 2010). Here, corals benefit from organic matter that settles from the sea surface through vertical migration. The resuspension of organic matter by internal waves might be one reason for a general steady supply of nutrients into the Porcupine Seabight mound regions (White, 2007; Kiriakoulakis et al., 2007). Purser et al. (2010) demonstrated that L. pertusa polyps capture more efficiently zooplankton under low flow velocities of about 5 cm s⁻¹. These specific hydrodynamic conditions facilitate dense cold-water coral growth and mound build up SW of Ireland (De Mol et al., 2002; Kano et al., 2007).

1.2. Paleoceanographic reconstruction with scleractinian cold-water corals

Deep-sea or cold-water corals thrive in dark, cold and nutrient rich waters. Similar to their tropical counterparts their skeleton is composed of aragonite and hence can be used as an archive for paleoceanographic reconstructions (e.g. Adkins et al., 1998.). Recent developments highlighted the use of Mg/Li (Li/Mg) and Li/Ca ratios in scleractinian corals as a proxy for temperature (Case et al., 2010; Hathorne et al. 2013). In this study we use the Mg/Li
et al. (2013) to calculate the Bottom-Water-Temperature (BWT) as it combines all available Li/Mg temperature calibration including Ba/Ca ratios were used to calculate Ba/Casw ratios according to and hence records variations in nutrient supply. In this study, cold-water coral L. pertusa served as a proxy for the reconstruction of Ba/Caseawater ratio measured in the scleractinian cold-water coral Desmophyllum dianthus serves as a proxy for paleoproductivity (e.g. Dymond et al., 1992; Nürnberg and Tiedemann, 2004). The Ba/Ca ratios measured in marine carbonates have been used to reconstruct changes in ocean circulation, upwelling and nutrients (e.g. Lea and Boyle, 1991). Subsequently, Ba/Ca ratios were introduced as a nutrient tracer in cold-water corals (Montagna et al., 2005; Lea et al., 1989: Lea and Boyle, 1991). Consequently, Ba/Ca ratios were used as a nutrient tracer in cold-water corals (Montagna et al., 2005; Anagnostou et al., 2011; LaVigne et al., 2011) as they are not related to seawater temperature (Anagnostou et al., 2011). In particular, Anagnostou et al. (2011) demonstrated that the Ba/Ca ratio measured in the scleractinian cold-water coral Desmophyllum dianthus serves as a proxy for the reconstruction of Ba/Ca seawater and hence records variations in nutrient supply. In this study, Ba/Ca ratios were used to calculate Ba/Ca_{sw} ratios according to Anagnostou et al. (2011):

\[ \text{Ba/Ca}_{\text{coral}} = 1.4(\pm0.3) \times \text{Ba/Ca}_{\text{sw}} + 0(\pm2) \]

We are aware of the use of D. dianthus for this calibration (Anagnostou et al., 2011), however we use the same calibration on L. pertusa to reconstruct trends and not absolute values.

1.3. The Challenger Mound (Site U1317)

In 2005 IODP Expedition 307 drilled for the first time through the entire sediment body of a cold-water coral mound, and obtained complete records of 155 m high Challenger Mound, located in the Porcupine Seabight (Site U1317, 52°23'N, 11°43'W; ~800 m below sea level, Fig. 1, Williams et al., 2006). The recovered sediment cores contained mainly the scleractinian cold-water coral L. pertusa and only to a minor degree M. oculata. The mound is situated on top of an unconformity with a Miocene age of up to 16.58 Ma (Louwye et al., 2007; Kano et al., 2007). The sedimentary record of the Challenger Mound can be subdivided into two units, respectively M1 between 155 and 23.6 m below seafloor (mbsf) and M2 between 23.6 and 0 mbsf for Hole 1317 E (Expedition 307 Scientists, 2006; Thierens et al., 2010; Titschack et al. 2009). Challenger Mound stratigraphic work carried out by Kano et al. (2007), (Strontium Isotope Stratigraphy (SIS)) shows that mound growth initiated at ~2.6 Ma, whereas ages based on magnetostratigraphy point to an earlier onset at ~2.74 Ma (Foubert and Henriet, 2009). Nevertheless, both studies recognised a significant hiatus at around 23.6 mbsf in Site U1317 Hole E. This major hiatus separates the mound record into a period of fast (~15 cm/kyr: 2.6–1.7 Ma=M1) and slower mound growth (~5 cm/kyr: 1.0–0.5 Ma=M2).

Here we compare the oceanographic settings in times of rapid Challenger Mound initiation versus mound decline. Based on new age constraints (combined U/Th and \(^{87}\text{Sr/86}\text{Sr}\)) and different geochemical signals from coral skeletons (Mg/Li and Ba/Ca) this study contributes to the understanding of environmental and oceanographic boundary conditions required for cold-water coral mound growth in the Northeast Atlantic and adds to the debate if intermediate water masses had an impact on these deep-sea organisms and ecosystems throughout the last 3 Myr.

2. Material and methods

Coral fragments from Site U1317 recovered during IODP Expedition 307 with R/V Joides Resolution in 2005 (Expedition 307 Scientists, 2006) were investigated in this study. In order to obtain undamaged half core sections, the drill cores were frozen before splitting (Dorschel et al., 2005; Foubert et al., 2007). IODP Site 1317 consists of 5 different holes A, B, C, D and E. In this study we used samples from Holes B, C, D and E. For example at 147.95 mbsf sediments from Hole U1317C are characterised by an...
unconformity, marked by a sharp colour change from the greenish-grey underlying unit to the grey, coral bearing sediments. Clean and well-preserved coral samples were taken from the base of Holes C, D and E (148–115 mbsf) and from 0–30 mbsf of Holes B and C for age determination and elemental ratio analyses. In addition, also one modern (in-situ, live collected, M61/3-551) L. pertusa sample from the nearby Galway Mound (837 m water depth, 9.5°C) was also analysed for elemental/Ca ratios. All samples were drilled according to Rüggeberg et al. (2008) avoiding the centres of calcification, since these parts of the skeleton have different isotopic and elemental compositions compared to the rest of the skeleton. X-ray diffraction did exclude potential early diagenesis as all coral samples retain their initial aragonitic skeleton (>98% aragonite, below detection limit). Before analyses corals were cleaned according to Cheng et al. (2000a), but leaving out the reductively cleaning step after Shen and Boyle (1988). Briefly, corals were physically cleaned with a Proxxon dremel tool and all evidence (very rare) for endolithic organisms and potential bio-erosion were drilled out as far as possible. Subsequently, corals were ultrasonically cleaned several times (2–3) according to samples cleanness. Corals were then submerged into a 50/50 mixture 1 M NaOH and 30% H2O2 for 15 min. with ultrasonification. This step was repeated several times (>2) times. Finally corals were rinsed in a 50/50 mixture of 30% peroxide and 1% HClO4 for maximum of 1 min.

2.1. Strontium isotope stratigraphy on coral fragments

Strontium isotope ratios were determined by Thermal Ionisation Mass Spectrometry (TIMS, TRITON, ThermoFisher Scientific) after chemical separation via cation exchange chromatography using a Sr-specific resin (Eichrom). For Sr isotope measurements about 500 ng of Sr was used. All isotope ratios were internally normalised to an 86Sr/88Sr ratio of 0.1194. Repeated analysis of the standard NIST SRM 987 in the course of this study yielded an average value of 0.710245 ± 0.000012 (2σ, n=15). For comparison to literature values all 87Sr/86Sr were normalised to a value of 0.710248 for the NIST SRM 987. Ages were obtained by comparison to the seawater evolution curve (Lookup Table Version 4B-08-04, McArthur and Howarth, 2004, Age data: GTS 2004, Gradstein et al., 2004).

2.2. Thorium/Uranium age determinations on coral fragments

The Thorium and Uranium isotope measurements were performed on a VG Elemental AXIOM MC-ICP-MS at GEOMAR applying the approach of Fietzke et al. (2005). For isotope dilution measurements a combined 233/234U/229Th-spike was used, with stock solutions calibrated for concentration using NIST-SRM 3164 (U) and NIST-SRM 3159 (Th) as combi-spike calibrated against CRM-145 uranium standard solution (also known as NBL-112A) for U-isotope composition, and against a secular equilibrium standard (HU-1, uranium ore solution) for determination of 230Th/234U activity ratio. Whole procedure blank values of this sample set were around 0.0006 pg for 230Th, 2 pg for 232Th and around 2 pg for U, which are in the typical range of this method and laboratory. The element separation procedure was based on Eichrom-UTEVA resin. Calculation of geochronological data and activity ratios is based on the decay constants given by Cheng et al. (2000b).

2.3. Elemental ratio determinations on coral fragments

Selected samples of the cold-water coral powders used for 87Sr/86Sr and 230Th/U age determinations measurements were split for additional ICP-MS measurements (Agilent 7500 CS). The Elemental/Ca ratios were calculated from the raw counts using the method of Rosenthal et al. (1999). Calcium concentrations were measured in a first step and samples diluted to a Ca concentration of ∼10 ppm. Six aliquots of Porites sp. coral powder reference material JCp-1 (Okai et al., 2002) were treated like samples and the average values obtained during the course of this study (n=8, including repeated measurements) for Mg/Ca 4.17 ± 0.03 mmol/mol, Li/Ca 6.17 ± 0.18 μmol/mol, Ba/Ca 7.19 ± 0.36 μmol/mol resulting in an average value for Mg/Li values of 0.67 ± 0.02 mol/mmol. The reproducibility (2 SD) of the measurements was 2.64% for Mg/Ca, 10.1% for Ba/Ca and 5.42% for Li resulting in 7.13% for the Mg/Li ratio for the JCp-1 (2 SD). The absolute concentrations are within the uncertainty of the recommended values (Okai et al., 2004).

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Fig. 3. Age constraints of the Challenger Mound showing the strontium isotope based ages for Site U1317 (Holes B, C, D, and E) and the published Sr-Ages of Kano et al. (2007) for Hole E. The SIS reveals that the decline of mound growth occurred at least in Holes B, C, and E.
3. Results

3.1. Strontium isotope measurements

Radiogenic strontium isotope measurements were carried out on corals covering the upper (0–30 mbsf) and lower mound sequence (115–148 mbsf). Overall the $^{87}$Sr/$^{86}$Sr isotope ratios vary from 0.709061 to 0.709174 and show an increasing trend from the lower to the upper mound part. In particular, $^{87}$Sr/$^{86}$Sr isotope ratios in the interval between 110 and 148 mbsf of M1 are similar and vary only between 0.70906 and 0.70908. In contrast, in the interval between 0 and 30 mbsf the $^{87}$Sr/$^{86}$Sr isotope ratios increase from 0.70910 to 0.70917. Corresponding ages were calculated after the Lookup Table Version 4B-08-04, McArthur and Howarth, 2004, Age data: GTS 2004 (Gradstein et al., 2004). Calculated Sr-ages reveal that the mound generally becomes younger from the base to the top. In particular the mound base shows ages between 3.02 and 2.01 Ma (SIS mean age), whereas the upper mound interval displays ages between 0.008 and 1.67 Ma (Fig. 3).

3.2. Th/U age determinations

Calculated U-series ages of 16 corals vary from 1.2 to 513 ka (Supplements). The measured $^{232}$Th concentrations are small < 2 ng g$^{-1}$ and hence negligible due to the high activity ratio of $^{230}$Th/$^{232}$Th of > 1000. However, calculated initial $\delta^{234}$U$_{0}$ values vary between 15.6 and 156.84 (Supplements).

Four samples are within the range of 149.6 ± 10‰ (compare Stirling et al., 1998; Robinson et al., 2004) and show $^{230}$Th/U ages similar to the Sr-ages (Fig. 4). The $^{230}$Th/U dated coral growth occurs generally in interglacial periods at 1.2 ka (MIS 1), 104.7 ± 0.7 ka (MIS 5) and 513.9 ± 35.2 (MIS 13). The coral based ages at 460.2 (MIS 12) might also be assigned to an interglacial period, due to the large uncertainty of ± 63 ka.

The $\delta^{234}$U$_{0}$ scatter for the older samples is larger, compared to the younger samples probably due to more open system behaviour (Thompson et al., 2003). The study of Titschack et al. (2009) reported many hiatuses in the upper mound M2 interval, therefore we assume that corals, indicating open system behaviour, could have been exposed at the sediment/seawater interface for extended periods and may have been affected by pore waters. Future studies will focus on more detailed analysis of the U-series data including uncertainties of potential seawater U-isotopic variations, diagenetic alteration and U-series system opening, but this is beyond the scope of this study and will be presented elsewhere.

For further geochemical investigations corals indicating open-system behaviour were not used for paleoceanographic reconstruction. However, corals from the lower mound sequence were not measured for Thorium/Uranium. Here, the quality of coral samples is only based on XRD measurements, which imply the rather good preservation of the studied coral fragments from the lower mound sequence M1.

3.3. Elemental ratios of coral skeletons

Downcore Mg/Ca ratios vary from 2.32 to 3.89 mmol/mol and the corresponding Li/Ca values range from 7.57 to 15.75 μmol/mol. The calculated Mg/Li ratios presented here vary from 0.23 to 0.32 mol/mmol with a mean of 0.28 mol/mmol in the entire mound sequence. In general the Mg/Li ratios display an increasing trend from the mound base to the top mound from 0.23 to 0.32 mol/mmol (Fig. 5). The same pattern is observed in the Li/Ca record. Overall, the Mg/Ca and Li/Ca ratios are in phase with the observed modern values of scleractinian cold-water corals (Case et al., 2010) and display the expected positive relationship indicating that the corresponding Mg/Li ratios are not disturbed by any potential diagenetic overprint (Fig. 6). We used the temperature relationship of Mg/Li (Li/Mg) ratios (Hathorne et al., 2013) in L. pertusa to calculate the BWT of the ambient seawater during times of coral growth (Fig. 7). Reconstructed BWT show an increasing trend from the mound base to the top mound from ~79 to 9.6°C (Fig. 7).

Recent studies have highlighted that both Mg/Li and Li/Ca ratios may serve as a potential paleotemperature proxy in scleractinian corals (Case et al., 2010; Montero-Serrano et al., 2013; Hathorne et al., 2013). However, Mg/Li ratios, introduced by the study of Bryan and Marchitto (2008), are suggested to be more robust since they show less heterogeneity within the coral skeleton compared to Li/Ca (Case et al., 2010; Hathorne et al., 2013).

As we aim to reconstruct paleotemperatures over a relatively long period of time (~3 Myrs) the elemental and isotopic composition of seawater has to be considered as this may result in over and/or underestimated paleotemperatures. In particular, the
Mg/Li is calculated from the Mg/Ca and Li/Ca ratio. For Magnesium and Lithium it was suggested that the seawater value has been increased similar by a factor of 2 over the last 3 Myrs (Kano et al., 2007; Foubert and Henriet, 2009). Our radiogenic Sr-based stratigraphy is similar to those findings indicating that coral mound growth initiated at the Pliocene/Pleistocene boundary (Fig. 3). Mound initiation in the Porcupine Seabight has been earlier related to the late Pliocene reintroduction of the Mediterranean Outflow Water into the Porcupine Seabight (De Mol et al., 2002). This is in good accordance with the study of Khéli et al. (2009) showing that the MOW intensified between 3 and 3.5 Ma in the Porcupine Seabight (DSDP Site 548, Fig. 1). Therefore, intermittent water mass dynamics are crucial for the build-up and development of cold-water coral carbonate mounds at the Porcupine Seabight margins (Raddatz et al., 2011).

The early development of Challenger Mound in M1 was fast and rather continuous (Mound Booster Stage; Henriet et al., 2002) and hence indicates that initial environmental conditions until ∼2.0–2.1 Ma were favourable for cold-water coral growth (Fig. 3). This period of fast initial mound development in the early Pleistocene was characterised by weak sea-level fluctuations triggered by the orbital low-amplitude 41-ka cycle (Lisiecki and Raymo, 2005). The growth patterns for Challenger Mound have been described by several studies (Kano et al., 2007; Foubert and Henriet, 2009; Hvenne et al., 2009; Thierens et al., 2010). Mound growth started to cease at ∼2 Ma and at ∼1 Ma. This growth behaviour was already identified by other studies (Kano et al., 2007; Foubert and Henriet, 2009; Sakai et al., 2009; Titschack et al., 2009). During this growth decline Challenger Mound was only partly covered by corals (Kano et al., 2007). These findings are supported by the $^{87}$Sr/$^{86}$Sr ages presented here showing that coral mound growth in this crucial period was reduced but did not entirely cease (Fig. 3).

For the top mound part we applied a combined age determination using $^{230}$Th/U and $^{87}$Sr/$^{86}$Sr ages. Kano et al. (2007) documented Sr ages until ∼0.6 Ma at 4 mbsf in Hole E, whereas our $^{87}$Sr/$^{86}$Sr ages of Hole C (0.1 mbsf) and B (8.37 mbsf) reveal younger ages (Fig. 3). In particular, from 13 mbsf to the top of the Challenger Mound Sr ages generally become younger again and show ages between 0.5 Ma and 0. However, mound growth in the upper mound part M2 appears not to be as fast as in the lower mound part M1. Early
mound development (M1) of Challenger Mound between 3.0 and 2.1 Ma was characterized by an initial period of fast mound growth with depositional rates of ∼15 cm/ka and a second period with lower growth at rates of < 1 cm/ka. The depositional rate of the upper mound M2 is remarkably lower with ∼2 cm/ka (12.0–0.001 Ma, Kano et al., 2007; Tittschack et al., 2009).

Furthermore, our results indicate that both age chronometers are within uncertainty = good accordance with each other (Fig. 4). The 230Th/U ages presented here reveal that Challenger Mound growth in the late Pleistocene occurred only in interglacial periods, except for MIS 12 (Fig. 4). A significant hiatus is identified at ∼4 mbsf revealing a period of no deposition and/or erosion of ∼0.3 Ma between MIS 5 and MIS 12. This interpretation is consistent with the findings of Foubert et al. (2007) indicating the same hiatus in gravity cores on Challenger Mound, and Tittschack et al. (2009) who highlighted that the sedimentary record might be disturbed by several hiatuses. Other studies have shown that carbonate mound growth on the Irish margin in the latest Pleistocene was restricted to interglacial periods (Dorschel et al., 2005; Eisele et al., 2008; Frank et al., 2005, 2009, 2011; Rüggeberg et al., 2007) without only a few exceptions. Further south in the Bay of Biscay cold-water coral growth tends to have occurred during interstadials (Schröder-Ritzrau et al., 2005). In the Gulf of Cadiz (Wienberg et al., 2009, 2011) as well as off Mauritania (Eisele et al., 2011; Frank et al., 2011) growth is observed during glaciars. Focusing on the very top of the Challenger Mound 230Th/U age determinations show that cold-water coral growth was indeed only active during interglacials, namely MIS 1 and MIS 5. Since the Mid-Pleistocene-Transition interglacial/glacial cycles are dominated by a 100 ka periodicity, characterised by changes in sea level of ∼120 m (Lisiecki and Raymo, 2005).

We therefore suggest that similar to other cold-water coral carbonate mounds on the Irish margin (Dorschel et al., 2005; Eisele et al., 2008; Frank et al., 2005, 2009, 2011; Rüggeberg et al., 2007) pronounced glacial-interglacial changes during these periods had an important impact on mound growth. Overall, the growth of the Challenger Mound seems to have been connected to major global climatic changes, such as the intensification of the Mediterranean Outflow Water (Khelifi et al., 2009), the Northern Hemisphere Glaciation and the Mid-Pleistocene Transition (Mudelsee and Schulz, 1997). Moreover, from our observation it appears that the restriction of carbonate mound growth to interglacial periods at around 50 N on the Irish Margin is at least valid for the last ∼0.5 Myr, but does not apply for the early growth stages of Challenger Mound (Raddatz et al., 2011).

4.2. Environmental boundary conditions of cold-water coral mound growth

Environmental control mechanisms of cold-water coral reef growth and mound build-up on the European continental margin between 20° and 70° N are currently the subject of ongoing paleoceanographic research. Recent studies have shown that late Pleistocene distribution of flourishing coral-reefs and mounds oscillated from the NE Atlantic down to the Gulf of Cadiz and off Mauritania in correspondence to Interglacial/Glacial cycles induced by the expansion and retreat of the Northern-Hemisphere ice-sheets (Dorschel et al., 2005; Eisele et al., 2008; Frank et al., 2005, 2009, 2011; Rüggeberg et al., 2007; Wienberg et al., 2009, 2011; Thieren et al., 2010). Their biogeographical limitation is caused by a combination of changes in sea-surface productivity (Eisele et al., 2011; Wienberg et al., 2011), upwelling (Wienberg et al., 2011) and bottom currents (Rüggeberg et al., 2005; Huvenne et al., 2009), which in turn are affected by changes in (intermediate) water mass variability and dynamics. Strong bottom currents, enhanced sea-surface productivity, temperatures above 4°C, and a prominent strong density gradient between the upper and intermediate water masses favoured coral growth during the last interglacial periods on the Irish Margin (Dorschel et al., 2005; Dullo et al., 2008; Eisele et al., 2008; Frank et al., 2005; Rüggeberg et al., 2007).

Our Mg/Li-BWT/Lophelia reconstructions vary between −8 and 10°C and are consistent with the optimal temperature range of L. pertusa and the modern BWT in the Belgica Mound Province (∼9.5°C, Fig. 7; White, 2007). Additionally, our results are also in good accordance with the Mg/Ca-based foraminalifer paleotemperatures from south of the Porcupine Seabight (DSDP Site 548) by Khelifi et al. (2009) and δ18O-based foraminalifer paleotemperatures from IODP Site 1317C (Raddatz et al., 2011). Hence, we propose from our reconstructions that coral mound growth mainly occurred within the optimal BWT envelope between 8 and 10°C. We relate this to the inflow of the warm and saline MOW causing a specific temperature setting at intermediate water depth in the Porcupine Seabight. In general cold-water corals are known to thrive in cold and nutrient-rich water that lack large seasonal variability (Roberts et al., 2006) and active mound growth is restricted to specific environmental conditions (Rüggeberg et al., 2007). In the Porcupine Seabight a pycnocline is formed between the inflowing MOW and ENAW. This density gradient favours the settlement of organic matter providing sufficient nutrients for

![Fig. 7. Paleooceanographic reconstruction based on 230Th/U and 87Sr/86Sr dated corals. Elemental ratios were measured in scleractinian cold-water coral L. pertusa. The Mg/Li ratios were transferred into BWT using the equation of Hathorne et al. (2013). Results show that corals thrived in a temperature envelope between 8 and 10°C comparable to the modern BWT (9.5°C dashed line; White, 2007). The Ba/Ca/Lophelia were transferred into Ba/Ca of seawater based on the study of Anagnostou et al. (2011) showing a clear shift towards lower values from the lower to the upper mound. The dashed line illustrates the recent Ba/Ca/Lophelia values from the nearby Galway Mound. The Ca/MAR based reconstructions of export productivity from ODP Site 982 are taken from Bolton et al. (2011). Also plotted is the benthic LR04 stack of Lisiecki and Raymo (2005) showing the increase of amplitudes in interglacial/glacial cycles over the last 3 Myr.](http://doc.rero.ch)
active cold-water coral growth, similar to recent observation at living coral reefs (Dullo et al., 2008). Today the majority of the living cold-water reefs and mounds in the Northeast Atlantic occur within a narrow density envelope and an associated permanent thermocline (Dullo et al., 2008; White and Dorschel, 2010). Corals benefit from organic matter that settles down from the sea-surface (Guinotte et al., 2006) and concentrates around this specific density envelope. White et al. (2005) investigated the productivity over the Porcupine Bank and concluded that coral growth depends on high sea-surface productivity and high utilisation of organic matter resulting in a dense nutrient rich water mass. In our records we observe a gradual trend in the reconstructed Ba/Ca$_{sys}$ values from $-29$ to $-10$ μmol/mol (Fig. 7). We attribute this trend to a decrease in nutrient supply (Figs. 2 and 7) due to changes in sea-surface productivity and to a shifting water mass boundary between the ENAW and MOW (Figs. 2 and 3). A deepening of the MOW would lead to a stronger influence of the warmer nutrient depleted ENAW at the core summit (Fig. 7). Such a trend can be observed in our Mg/Li$_{Lophelia}$ temperature record revealing a temperature increase of $-2$ °C (Fig. 7). Such an increase of 2 °C cannot be explained by the growth of the coral mound (155 m) through the water column (< 0.5 °C). However, the modern oceanography of Porcupine Seabight reveals that the top of the dead Challenger Mound is situated well above the boundary layer between the ENAW and the MOW, whereas other active mounds appear to occur deeper (Huvenne et al., 2003; Foubert et al., 2005). This implies that corals on the top of the Challenger Mound are not sufficiently provided by nutrients and food, leading to the recently observed decline or sparse growth of Challenger Mound.

This study indicates that in the past sea-surface productivity was also one of the major controlling parameters of cold-water coral growth. Enhanced primary sea-surface productivity in the North Atlantic can be associated with changes of the polar front at the Pliocene–Pleistocene boundary (Naafs et al., 2010; Bolton et al., 2011) and decreased sea-surface productivity during the Mid-Pleistocene Transition (MPT, Stein et al., 2009) to an enhanced impact of the British ice-sheets into the Porcupine Seabight. This crucial period corresponds to the MPT initiated as early as 3 Ma following an intensification of the Mediterranean Outflow Water (MOW). Bottom-Water-Temperature (BWT) reconstructions by Mg/Li$_{Lophelia}$ ratios and nutrient reconstructions based on Ba/Ca$_{Lophelia}$ Ratios reveal that coral growth was restricted to specific oceanographic settings. In particular coral and mound growth mainly occurred within a narrow temperature range between 8 and 10 °C. The MOW initiated the recent water mass stratification in the Porcupine Seabight that in turn established the necessary density gradient for enhanced nutrient and food supply between the Eastern North Atlantic Water (ENAW) and MOW at the mound summit. Between 1.7 and 1 Ma mound development was reduced or interrupted coinciding with the Mid-Pleistocene-Transition, a period of global climate change. High amplitude interglacial/glacial cycles (100 ka) initiated and glacial conditions became unfavourable for corals in the Porcupine Seabight.

From our reconstructions we propose that coral growth benefited from a complex oceanographic setting that supported an enhanced nutrient and food supply. Finally, this study underlines that cold-water coral communities are restricted to specific environmental conditions (specific water mass stratification at intermediate water depths in combination with enhanced palaeoproductivity) and hence climate change may cause a dramatic decline in cold-water coral mound growth on the European continental margins.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.dsr2.2013.06.009.

References


