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Key Points:

- We produced high-resolution records of benthic foraminiferal elemental ratios, δ¹⁸O, and δ¹³C from two cores from the northern Indian Ocean
- Evaluation of past IWT from the Mg/Li ratio and the δ¹⁸O since last deglaciation exhibit millennial-scale changes
- Variations of the IW Δ [CO₃^{2–}] calculated from *H. elegans* Sr/Ca are related to atmospheric CO₂ changes on glacial-interglacial timescales

Supporting Information:

- Supporting Information S1
- Data Set S1
- Data Set S2

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North Indian Ocean Circulation Since the Last Deglaciation as Inferred From New Elemental Ratio Records for Benthic Foraminifera *Hoeglundina elegans*

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Abstract The evolution of intermediate circulation in the northern Indian Ocean since the last deglaciation has been reconstructed from two marine cores located at intermediate depths off the southern tip of India (MD77-191) and in the northern Bay of Bengal (BoB) (MD77-176). Benthic for a for a seawater carbonate ion concentration ($[CO_3^{2-}]$) estimated from the Sr/Ca, and paleotemperature reconstructed on the basis of the Mg/Li of aragonite benthic species Hoeglundina elegans were used to trace the evolution of past intermediate-deepwater masses and to constrain ocean-atmosphere exchanges during the two-stage increase in atmospheric CO₂ across the last deglaciation. The intermediate water [CO3²⁻] was mainly affected by changes in the ocean alkalinity inventory, associated with the modulation of atmospheric CO₂ on glacial-interglacial timescales. Higher benthic foraminiferal δ^{13} C, depleted [CO₃²⁻], and decreased benthic-planktonic ¹⁴C age offsets at intermediate water depths suggest a release of deep-sea CO₂ to the atmosphere through the Antarctic Intermediate Water (AAIW) in the Southern Ocean during the 17-15.2 and 12.6-10.5 cal kyr BP time intervals. In addition, the decreased H. elegans Mg/Li record seems to reflect an increased contribution of cold water mass during the 17-15.2 and 12.6-11.9 cal kyr BP intervals and throughout the Holocene. In contrast, two warm events occurred in the 15–13.3 and 11–10.3 cal kyr BP time intervals. During the late Holocene, a decrease in the intermediate water $[CO_3^{2-}]$ indicates a contribution to atmospheric CO₂ rise since 8 cal kyr BP, due to the depleted global ocean alkalinity and/or the variations in surface productivity (at least for MD77-191).

1. Introduction

Atmospheric CO_2 records from ice cores in Antarctica exhibit variations at glacial-interglacial and millennial timescales over the last 800 kyr that are correlated to global climate changes (Lisiecki & Raymo, 2005; Lüthi et al., 2008). This indicates that the climate system and, in particular, the Earth's heat budget and temperature are highly sensitive to variations in CO_2 greenhouse gas (IPCC, 2013). Past variations in the global carbon cycle are, therefore, a key element to understand the processes controlling atmospheric CO_2 and its relationships to climate changes. As the ocean is the largest external carbon reservoir that can exchange rapidly with the atmosphere, changes in the oceanic carbon pool are strongly linked to past atmospheric CO_2 variations (Broecker & Peng, 1982), through complex feedback relationships. A decrease in atmospheric CO_2 to a net decrease of ~15 ppm in atmospheric CO_2 during the glacial period (Brovkin et al., 2007; Sigman & Boyle, 2000; Yu, Anderson, & Rohling, 2014). At a shorter timescale, abrupt millennial events, such as during the last deglaciation, require more complex ocean processes, involving changes in ocean biogeochemistry and circulation.

Exchanges of carbon through the ocean and to/from the atmosphere occur through changes in the dissolved inorganic carbon (DIC) system (e.g., Anderson et al., 2009; Ridgwell & Zeebe, 2005; Sigman & Boyle, 2000). Among the DIC species, the carbonate ion concentration ($[CO_3^{2^-}]$) is a particularly important one as it determines the seawater carbonate saturation state with respect to the calcium carbonates in the water column as well as at the seafloor. Variations in seawater $[CO_3^{2^-}]$, by driving changes in the

preservation/dissolution of sedimentary carbonates, make this fraction an active player of the ocean and global carbon cycle. Thus, reconstructing past seawater $[CO_3^{2-}]$ variations is mandatory to better constrain the processes and mechanisms controlling past global carbon cycle.

As a means of reconstructing bottom water parameters over a geological timescale, benthic foraminifera elemental/Ca ratios have received increasing attention over recent decades (e.g., Boyle et al., 1995; Boyle & Keigwin, 1985; Marchitto et al., 2018; Raitzsch et al., 2011; Yu et al., 2008). For instance, the Mg/Ca in benthic foraminifera has been widely used to reconstruct bottom water temperature changes (e.g., Barrientos et al., 2018; Lear et al., 2002; Marchitto et al., 2007; Martin et al., 2002; Rosenthal et al., 1997; Yu & Broecker, 2010). However, this proxy is also sensitive to bottom water carbonate ion saturation $(\Delta[CO_3^{2-}])$, defined as the difference between $[CO_3^{2-}]$ and $[CO_3^{2-}]_{sat}$ (e.g., Bryan & Marchitto, 2008; Marchitto et al., 2018; Rosenthal et al., 2006; Yu & Elderfield, 2008). Other elemental/Ca ratios, such as the benthic foraminifera Li/Ca and Sr/Ca, have been shown to covary with both past bottom temperatures and Δ [CO₃²⁻] (e.g., Bryan & Marchitto, 2008; Doss et al., 2018; Hall & Chan, 2004; Marchitto et al., 2018; Rosenthal et al., 2006). More recent studies have demonstrated that the benthic foraminiferal U/Ca, Li/Ca, and Sr/Ca are also promising proxies for the marine dissolved carbon/carbonate system (Bryan & Marchitto, 2008; Keul et al., 2013; Raitzsch et al., 2011; Yu, Elderfield, Jin, Tomascak, & Rohling, 2014); they can thus increase our ability to fully reconstruct past changes in the intermediate and deepwater chemistry linked to the global carbon cycle (Came et al., 2008; Makou et al., 2010; Poggemann et al., 2017; Yu et al., 2008).

Most of the studies referred to above were devoted to calcitic foraminifera. Only a few studies have focused on benthic foraminifera *Hoeglundina elegans*, since this aragonite species is not widely found in the sediment due to its higher sensitivity to dissolution than calcitic foraminifera (Marchitto et al., 2018; Rosenthal et al., 2006). Previous works suggest that Mg/Ca, Li/Ca, and Sr/Ca in *H. elegans* could also be used as paleoproxies for both carbonate ion concentration and temperature (e.g., Bryan & Marchitto, 2008; Hall & Chan, 2004; Marchitto et al., 2018; Rosenthal et al., 2006).

Furthermore, most of these studies that apply elemental ratios have focused on the Atlantic Ocean (e.g., Makou et al., 2010; Poggemann et al., 2017; Yu et al., 2008). As the northern Indian Ocean plays an important role in global ocean circulation in terms of deepwater ventilation and stratification (e.g., Ahmad et al., 2008, 2012; Bryan et al., 2010; Raza et al., 2014), it has received increasing attention over recent decades (e.g., Bryan et al., 2010; Ma et al., 2019; Sijinkumar et al., 2016; Yu et al., 2018). However, no continuous, high-resolution benthic foraminiferal elemental ratio records have been produced for this area.

In this study, we investigated benchic and planktonic foraminiferal δ^{13} C, δ^{18} O, and elemental ratios from two cores located at intermediate water depths off the southern tip of India and in the northeastern Bay of Bengal (BoB); our aim is to better constrain the temporal evolution of the source and ventilation of intermediate water mass in the northern Indian Ocean. We explored the potential of the Mg/Li and Sr/Ca ratios of *H. elegans* as proxies to reconstruct past changes in the temperature and $[CO_3^{2-}]$, respectively. Reconstructing past variations in $[CO_3^{2-}]$ and comparing our results with other areas as well as with atmospheric CO₂ records allow us to better understand the relationships between past changes in intermediate ocean circulation and variations in the global carbon cycle.

2. Material and Hydrological Settings

Cores MD77-191 and MD77-176 were collected off the southern tip of India on the edge of the Arabian Sea (07°30'N–76°43'E, 1,254 m water depth) and in the northeastern BoB (14°30'5 N–93°07'6E, 1,375 m water depth), respectively, during the OSIRIS III cruise of the N.0 *Marion Dufresne* in 1977 (Figure 1). These two cores cover a similar range of $[CO_3^{2-}]$ (75 µmol/kg for MD77-191 and 82 µmol/kg for MD77-176) in the modern ocean (supporting information Figure S1). The modern $[CO_3^{2-}]$ values are calculated using CO₂sys.xls (Ver. 12) based on the data collection from the Global Alkanity & TCO₂ data sets (Goyet et al., 2000; Pelletier et al., 2005).

Today, the surface water masses above 150 m in the Arabian Sea correspond to the Arabian Sea High Salinity Water (ASHS; salinity 36.5 psu) (Talley et al., 2011). In contrast, in the BoB, the surface water shallower than 100 m, that is, Bay of Bengal Surface Water (BoBSW), is much fresher (salinity ~31 psu) because of the





Figure 1. (a) Geographical setting, locations of MD77-191 (red star) in the Arabian Sea, MD77-176 (red star) in the Bay of Bengal, and Reference Site SK237 GC04 (red circle, Naik et al., 2017). The black arrows indicate the general direction of surface circulation in the northern Indian Ocean drifted by Southwest Monsoon (Schott & McCreary, 2001). (b and c) Salinity (psu, colored shading) depth-latitude section using Ocean Data View (ODV) software (Schlitzer, 2015) and vertical distribution of water masses in the Arabian Sea and Bay of Bengal (N-S cross section). AABW = Antarctic Bottom Water; ASHS = Arabian Sea High Salinity Water; BoBLS = Bay of Bengal Low Salinity Water; IDW = Indian Deep Water; NIIW = North Indian Intermediate Water; RSOW = Red Sea Overflow Water.

considerable input from rivers (Talley et al., 2011). The surface mixed layer and the boundary currents in the Arabian Sea and in the BoB are directly controlled by wind forcing and, therefore, display changes related to the seasonal reversal of the Indian monsoon wind system (Figure 1). The surface waters at the site of MD77-191 and MD77-176 are a mix of high-salinity ASHS and low-salinity BoBSW, respectively (Talley et al., 2011; Tomczak & Godfrey, 2003). MD77-191 is also located in the summer upwelling area of the Arabian Sea, and the average depth of the permanent thermocline is around 100 m (Ravichandran et al., 2012).

The main intermediate water masses in the Indian Ocean are Antarctic Intermediate Water (AAIW), Red Sea Overflow Water (RSOW), and Indonesian Intermediate Water (IIW) (Talley et al., 2011; You, 1998). The northward extension of AAIW in the Indian Ocean reaches only as far as 10°S in the modern Indian



Ocean (Lynch-Stieglitz et al., 1994). Red Sea Water is a high-salinity intermediate water, and it fills the Arabian Sea (between 400 and 1,400 m) (Beal et al., 2000; Talley et al., 2011). RSOW spreads eastward to the eastern boundary at 5°N and flows through the much stronger western boundary toward the Agulhas (Beal et al., 2000; Talley et al., 2011; You, 1998). Thus, RSOW could impact upon the MD77-191 core site at the southern tip of India but should make only a small contribution at Core Site MD77-176 in the BoB. IIW originates from the Pacific Central Water and enters the BoB via the Indonesian through flow. IIW flows clockwise at thermocline levels in the BoB (You, 1998).

The deepwater masses of the northern Indian Ocean are characterized by a mixture of Atlantic and Antarctic water masses (Reid, 2003; Talley et al., 2011). Between 1,200 and 3,800 m, the dominant deep water is Indian Deep Water (IDW), originating from the Circumpolar Deep Water and NADW (Tomczak & Godfrey, 2003; You & Tomczak, 1993). Below 3,800 m, the abyssal basins are bathed chiefly by the Antarctic Bottom Water (AABW) (Reid, 2003; Tomczak & Godfrey, 2003). However, AABW only reaches the southern part of the BoB as most of the bay is shallower than 4,000 m (Tomczak & Godfrey, 2003). In the BoB, the bottom water upwells when it moves northward (Talley et al., 2011); thus, changes in bottom waters are transferred upward to shallower water masses.

3. Methods

3.1. $\delta^{18}O$ and $\delta^{13}C$ Analyses From Core MD77-191

Stable oxygen (δ^{18} O) and carbon (δ^{13} C) isotope records for MD77-176 have already been published in Ma et al. (2019). For Core MD77-191, measurements were performed on well-preserved (clean and intact) samples of the planktonic foraminifera *Globigerinoides ruber* (250–315 µm fraction) and the endobenthic foraminifera *Uvigerina peregrina* (250–315 µm fraction). As *U. peregrina* is an infaunal benthic foraminiferal species, the δ^{13} C values may reflect pore water isotopic compositions rather than deepwater signature. To obtain a high-resolution benthic δ^{13} C record that would more accurately reflect bottom water changes, epibenthic foraminifera *Cibicidoides wuellerstorfi* and *Cibicidoides pachyderma* were also analyzed.

These stable isotope analyses were performed on approximately 4 to 8 clean and well-preserved specimens per sample using a Finnigan MAT 251 mass spectrometer at the Laboratoire des Sciences du Climat et de l'Environnement (LSCE, France). The δ^{18} O and δ^{13} C values are presented relative to the Pee Dee Belemnite (PDB) scale through calibrations performed with standards from the National Bureau of Standards (NBS). The mean external reproducibility of carbonate standards is $\pm 0.05\%$ for δ^{18} O and $\pm 0.03\%$ for δ^{13} C.

3.2. Analytical Protocols for Elemental Ratios

Mg/Ca, Sr/Ca, U/Ca, and Li/Ca ratios were measured in shells of the epifaunal benthic foraminifera *H. elegans* from MD77-191. Only the Mg/Ca and Sr/Ca results are available for MD77-176 due to the paucity of material and the difficulty in measuring the Li/Ca and U/Ca ratios on very small samples. *H. elegans* is an epifaunal species that grows on the sediment-seawater interface, thus minimizing the influence of pore water composition on the elemental ratios (Lutze & Thiel, 1989). Each sample contained approximately 8–15 individual foraminifera larger than 250 μ m. The foraminiferal test samples were crushed open between two glass plates. The resulting fragments were ultrasonically cleaned to remove clays and then subjected to oxidative and reductive protocols to remove organic matter and oxides following methods described by Boyle and Keigwin (1985) and Barker et al. (2003). Samples were dissolved in 0.075 N HNO₃ and analyzed using a single collector sector field high-resolution inductively coupled plasma mass spectrometer (HR-ICP-MS) Thermo Element XR hosted at the Laboratory GEOPS (University Paris-Sud, France).

Instrumental settings used for trace element analyses are listed in Data Set S1. The instrument sensitivity was optimized daily using 100 ppt tuning standard solutions. Given the limited amount of solution obtained after dissolution of the foraminifera samples, a high efficiency PFA MicroFlow Nebulizer (ES-2000-3503-080) was used and resulted in an uptake rate of 50 μ l/min. Aliquots of samples were first analyzed for Ca concentrations, and after, the remaining solutions were diluted to reach a Ca concentration of ~50 ppm for the second analyses of Mg, Sr, Cd, Li, B, Ba, and U (Yu et al., 2005). One measurement requires 1 ml solution.



Table 1

Calibrated AMS ¹⁴C Age Determined on Planktonic (10 Data Taken From Bassinot et al., 2011)

Depth (cm)	Calendar age (yr BP)	Taxa	¹⁴ C age (yr)	1 sigma error (±yr)
28	1,526	G. bulloides	1,970	60
76	2,226	G. bulloides	2,560	70
127	2,797	G. bulloides	3,020	60
175	3,567	G. bulloides	3,660	60
222	4,236	G. bulloides	4,160	60
271	5,070	G. bulloides	4,790	60
373	6,582	G. bulloides	6,150	80
425	8,757	G. bulloides	8,230	90
482	9,648	G. bulloides	8,970	80
550	11,294	G. ruber	10,300	50
580	13,285	G. ruber	11,810	50
594	14,258	Pteropods sp.	12,630	190
710	16,142	G. ruber	13,820	60

Note. 14 C ages were converted into calendar years (cal. yr BP, BP = AD 1950) USING THE CALIB REV. 7.1 SOFTWARE (STUIVER & BRAZIUNAS, 1993).

The procedures here are similar to the method described by Yu et al. (2005). Two series of mother standard solutions were prepared for elemental ratio analyses. The first six standard solutions were Mn, Al, and Fe mixtures with linear Ca concentration (0.5–7 ppm) to get more precise Ca concentration dilutions for the second run. Then, seven multielement standard stock solutions (J0–J6) were prepared gravimetrically by spiking a 1,000 μ g/ml Ca standard with appropriate amounts of Mg, Sr, Cd, Li, B, Ba, and U monoelemental 1,000 μ g/ml certified ICP-MS-grade stock solutions. The ratios of stock solutions were spaced linearly to contain the natural ratio ranges expected in both planktonic and benthic for-aminifera. Working mother standards were made by diluting the stock standard solutions with 0.1 N HNO₃ to get calcium concentrations of 50 ppm.

Sample solutions were systematically adjusted at 50 ppm Ca through dilution using 0.1 N HNO_3 . A blank consisting of the same 0.1 N HNO_3 used to dilute the standards and samples was also analyzed. The elemental blanks are better than 0.14% for Mg, 0.02% for Sr, 5.3% for Li, and 0.05% for U. All raw intensities (including standards) were corrected by removing the specific blank intensity values. Drift corrections were performed during daily

analyses by bracketing and linear regression estimated using standards interspersed every four samples. Standard curves were used to calculate elemental/Ca ratios, and coefficients of determination (r^2) were always >0.9999 for all elemental ratios. The mean reproducibility is 2% for Mg/Ca, 0.9% for Sr/Ca, 0.2% for Li/Ca, and 3.5% for U/Ca.

4. Chronological Framework of the Studied Cores

The age model of Core MD77-191 was established using nine accelerator mass spectrometry (AMS)¹⁴C dates from monospecific planktonic foraminifer *Globigerina bulloides* (Bassinot et al., 2011), one ¹⁴C date from pteropods (Mléneck, 1997), and three ¹⁴C dates obtained from planktonic foraminifera *G. ruber* measured with the *ECHo*MICADAS at the LSCE, France (see the analytical method in detail in Ma et al., 2019) (Table 1). The ¹⁴C dates were converted to calendar dates using CALIB Rev. 7.1 software (Stuiver & Braziunas, 1993) and the marine calibration data set (Reimer et al., 2013) and corrected for a surface marine reservoir of around 400 yr. MD77-191 provides a continuous record spanning the last 17 kyr BP with an average sedimentation rate of about 53 and up to 90 cm/kyr for the Holocene.

The age model of Core MD77-176 was previously established using 31 planktonic foraminifera (*G. ruber*) AMS ¹⁴C dates combined with the MD77-176 oxygen isotope record obtained on planktonic foraminifera *G. ruber*, which was correlated to the GISP2 Greenland ice core record (Marzin et al., 2013). MD77-176 displays high accumulation rates (average ~25 and up to 40 cm/kyr for the Holocene) and provides a high-resolution record for the period since 40 cal kyr BP.

5. Results

Results obtained from Cores MD77-191 and MD77-176 reveal millennial-scale events that punctuated the last deglaciation. However, we will not discuss the significance of these variations at the core sites in terms of Heinrich Stadial 1, Younger Dryas or Bølling-Allerød events, because the timing of these events, especially from where they have been described in the northern Atlantic Ocean, may be not strictly synchronous with observations from the northern Indian Ocean, as already observed in previous works from the area (e.g., Ma et al., 2019). Thus, we prefer to use the labeling of the time intervals rather than interpreting them in terms of deglacial climate events.

5.1. Stable Isotope Results From Core MD77-191

Stable isotope analyses were performed on planktonic foraminifera *G. ruber* and on benthic foraminifera *C. pachyderma*, *C. wuellerstorfi*, and *U. peregrina* from MD77-191 to produce the most complete records. Benthic δ^{18} O values were obtained from *U. peregrina*, which is known to record the calcite δ^{18} O at





Figure 2. (a) δ^{18} O records of the planktonic foraminifera *G. ruber*, (b) benthic δ^{18} O, and (c) δ^{13} C records for MD77-191. The black inverted triangles represent the ¹⁴C calculated calendar age points (Bassinot et al., 2011). The gray-shaded intervals mark the two-step increase in atmospheric CO₂ (Monnin et al., 2001), and the yellow-shaded interval marks the 15–13.3 cal kyr BP interval.

equilibrium (e.g., Shackleton, 1974). The δ^{13} C obtained from *C. wuellerstorfi* is known in the literature to record the bottom water δ^{13} C at equilibrium (Duplessy et al., 1984; Zahn et al., 1986), without any microhabitat effect (except in some special areas, like Southern Ocean, Mackensen et al., 1993). Because of the lack of *C. wuellerstorfi*, and in order to get a higher resolution, we completed the δ^{13} C record with results from *C. pachyderma*. We corrected the values from *C. pachyderma* to *C. wuellerstorfi* by using the average difference of 0.27‰ we measured between both species, obtained from 19 pair measurements of samples where the two species coexisted (Data Set S2 and Figure S2).

The δ^{18} O of *G. ruber* was enriched during the last deglaciation compared with the Holocene (Figure 2). A marked maximum in *G. ruber* δ^{18} O is evident during the early deglaciation, reaching -0.7% at 16 cal kyr BP. After that, the δ^{18} O displays a general decrease and reaches a more depleted level during the Holocene with an average of $\sim -2.58\%$. In addition, the general trend is punctuated by higher values of $\sim -1.53\%$ in the interval between 12.6 and 10.5 cal kyr BP (the late deglaciation).

Down-core benthic δ^{18} O values range between 2.36% and 4.25% (Figure 2). The most enriched δ^{18} O value (4.25%) was found during the last deglaciation period (16 cal kyr BP), whereas the most depleted (2.36%) occurred in the late Holocene (2.36 cal kyr BP). In the late part of the deglaciation, the δ^{18} O shows a shift toward higher values of around ~3% in the 12.6–10.5 cal kyr BP. This interval is followed by a decrease of δ^{18} O from 10 cal kyr BP to the core top (mean value of 2.75%).

The *Cibicidoides* δ^{13} C values of MD77-191 range from -0.04% to 0.62% (Figure 2). Although the record seems quite noisy, an increase (~0.11‰) in the δ^{13} C from the last deglaciation to the Holocene could be



observed. In addition, shorter-scale events occurred during the last deglaciation; the *Cibicidoides* δ^{13} C values increased during the time intervals 17–15.2 and 12.6–10.5 cal kyr BP and decreased (~0.03‰) at around 15–13.3 cal kyr BP. From 10.3 to about 9 cal kyr BP, the *Cibicidoides* δ^{13} C record suggests that considerable oscillations occurred with high peak values (0.52–0.62‰) at around 9.8–9.1 cal kyr BP. Then, the δ^{13} C records display a decrease during the middle and late Holocene, with a mean value of ~0.31‰ for *Cibicidoides* δ^{13} C.

5.2. Elemental Ratios in H. elegans From Cores MD77-191 and MD77-176

To check the robustness of our results and the potential diagenetic imprint (i.e., oxides), the Mn/Ca ratio was systematically measured. Results range between 6.5 and 10 μ mol/mol, which is much lower than the 100 μ mol/mol threshold proposed by Boyle (1983) to sort out contaminated samples and eliminate trace element results that chiefly reflect the presence of oxides. Barker et al. (2003) also suggested that potential clay contamination should be checked for using an Al/Ca < 0.5 mmol/mol. All Al/Ca results are below 0.2 mmol/mol, thus indicating that the samples were well cleaned and free of sedimentary clay contamination.

Mg/Ca, Sr/Ca, Li/Ca, and U/Ca ratios for MD77-191 range from 0.5–1.5 mmol/mol, 0.8–2.1 mmol/mol, 2.7– 6.4 µmol/mol, and 7.4–34.1 nmol/mol, respectively (Figures 3a–3d). All of these elemental ratios display a minimum during the 17–15.2 cal kyr BP time interval, with average values of ~0.8 mmol/mol for Mg/Ca, ~1.4 mmol/mol for Sr/Ca, ~3.9 µmol/mol for Li/Ca, and ~15.1 nmol/mol for U/Ca. Elemental ratios display higher values in the 15–13.3 cal kyr BP interval, and the most enriched values of Mg/Ca (1.5 mmol/mol), Sr/Ca (2.1 mmol/mol), Li/Ca (6.4 µmol/mol), and U/Ca (34.1 nmol/mol) occur during the same time interval. Thereafter, the Mg/Ca and Sr/Ca records generally show a continuous decrease throughout the deglaciation until the late Holocene. However, the continuous decrease is interrupted by an increase during the early Holocene (10.3–9 cal kyr BP). The lowest values are found in the late Holocene with mean values of Mg/Ca and Sr/Ca reaching 0.6 and 1.0 mmol/mol, respectively. The Li/Ca and U/Ca values also display a decreasing trend during the 12.6–10.5 cal kyr BP. U/Ca reaches its most depleted value (7.4 nmol/mol) at 10.8 cal kyr BP and then displays a significant increase during the early Holocene (10.3–9 cal kyr BP). Thereafter, lower values of Li/Ca and U/Ca (mean value of 3.5 µmol/mol and 17.3 nmol/mol, respectively) occurred during the 8.8–1.3 cal kyr BP time interval.

For Core MD77-176 from the BoB, Mg/Ca and Sr/Ca values for *H. elegans* range between 0.35 and 0.82 mmol/mol and between 0.55 and 1.32 mmol/mol, respectively. The *H. elegans* Mg/Ca and Sr/Ca records from MD77-176 show strong covariations (Figures 3e and 3f). The mean values of Mg/Ca (0.65 mmol/mol) and Sr/Ca (0.85 mmol/mol) during the LGM (22–17 cal kyr BP) are slightly higher compared with the Holocene (average values for Mg/Ca of 0.47 mmol/mol and for Sr/Ca of 0.72 mmol/mol). From 17 to about 12.5 cal kyr BP, the Mg/Ca and Sr/Ca records exhibit a succession of oscillations with a first minimum (0.39 and 0.7 mmol/mol, respectively) occurring between 17 and 15.4 cal kyr BP and a second shorter low (0.54 and 0.68 mmol/mol, respectively) between 13.4 and 12.5 cal kyr BP. In addition, during the last deglaciation period, the records display higher values (0.59 mmol/mol for Mg/Ca and 1.03 mmol/mol for Sr/Ca) at around 14.6 cal kyr BP, and the most enriched Mg/Ca (0.82 mmol/mol) and Sr/Ca (1.32 mmol/mol) values occur at the end of the last deglaciation (11.7–11.5 cal kyr BP). Then, during the Holocene, the values of Mg/Ca and Sr/Ca decrease continuously to reach a minimum (Mg/Ca for 0.35 mmol/mol and Sr/Ca for 0.55 mmol/mol) at the end of Holocene.

6. Discussion

Stable isotopes and elemental ratio records obtained for benthic foraminifera from Cores MD77-176 and MD77-191 show significant changes from the last deglaciation to the Holocene, as well as millennial-scale events. To better understand these variations, we compare our records with those from other areas and use the geochemical proxies to calculate different intermediate water mass parameters such as temperature and $[CO_3^{2-}]$. Moreover, since all proxies seem to follow the same trends, it is quite challenging to identify the main environmental control on each of them; thus, the limitations of these proxies are also discussed.

6.1. Assessing Past Variations in Intermediate Water Temperature

On the evidence of previous works, the environmental controls on benthic foraminiferal elemental ratios are not always well understood. For instance, the U/Ca of both calcitic benthic and planktonic foraminifera shows a negative correlation with $[CO_3^{2-}]$ (e.g., Keul et al., 2013; Raitzsch et al., 2011; Russell





Figure 3. Comparison of benthic foraminifera *H. elegans* elemental ratios. (a–d) MD77-191 Mg/Ca, Sr/Ca, Li/Ca, and U/Ca; (e and f) MD77-176 Mg/Ca and Sr/Ca. The color-shaded intervals are the same as in Figure 2.

et al., 2004), whereas Chen et al. (2017) indicated that there is no significant correlation between calcitic benthic U/Ca and carbonate system parameters. Furthermore, as *H. elegans* is an aragonitic species, we cannot discard the influence of temperature on this proxy (Chen et al., 2017; Keul et al., 2013; Raitzsch et al., 2011). Thus, due to the difficulty to interpret properly this proxy, we preferred not to include the results in the discussion.

For the other elemental ratios, based on several studies carried out on aragonite benthic species *H. elegans*, Rosenthal et al. (2006) proposed that the Mg/Ca and Sr/Ca of *H. elegans* may reflect a combination effect of $[CO_3^{2-}]$ and bottom water temperature when the degree of $\Delta[CO_3^{2-}]$ is below 15 µmol/mol. The Li/Ca ratio from both calcite and aragonite benthic species also seems to reflect changes in the $[CO_3^{2-}]$ saturation state and bottom paleotemperature (Hall & Chan, 2004; Lear et al., 2010; Lear & Rosenthal, 2006; Marchitto





Figure 4. (a) Benthic $\delta^{18}O_{ivc}$ obtained from MD77-176 (Bay of Bengal; red line) and MD77-191 (Arabian Sea; black line), respectively. The benthic $\delta^{18}O_{ivc}$ is corrected by using the global ice volume $\delta^{18}O$. (b) *H. elegans* Mg/Li ratios obtained from MD77-191. (c) IWT record based on the Mg/Ca_{Uvigerina} obtained from M78/1-235-1 in the tropical West Atlantic Ocean (Poggemann et al., 2018); the blue line is the smoothed curves using a 10-point average. The color-shaded intervals are the same as in Figure 2.

et al., 2004). However, as already observed in previous studies (Bryan & Marchitto, 2008; Marchitto et al., 2018), and in contrast to the Mg/Ca and Li/Ca ratios of *H. elegans*, the Mg/Li ratio shows a better correlation to bottom temperature. Thus, we decided to calculate the Mg/Li ratio from MD77-191 as a proxy for past intermediate water temperature (IWT). The Mg/Li values range between 0.14 and 0.28 mol/mmol (Figure 4b). However, according to the calibration published by Marchitto et al. (2018), this range of values corresponds to the coldest part of the equation, for which the Mg/Li ratio is less sensitive compared to the higher values. We observe that the Mg/Li values we calculated are in the lower range of the published results (Marchitto et al., 2018), but when we compare the Mg/Ca and Li/Ca of MD77-191 to the values from the same species *H. elegans* found in the literature (Mg/Ca and Li/Ca range from 0.5–1.7 mmol/mol and 1.4–9.3 μ mol/mol, respectively; Bryan & Marchitto, 2008; Marchitto et al., 2018), results are in agreement. Thus, this observation gives us confidence to the Mg/Li ratios we calculated, despite a low range of values, maybe indicating small variations in the coldest range of the observed values. Therefore, rather than trying to calculate IWT values, it seems that the Mg/Li raw data are a reasonable reflection of variations in the IWT from MD77-191.

The *H. elegans* Mg/Li record shows decreasing trends during the 17–15.2 and 12.6–11.9 cal kyr BP time intervals; however, it displays higher values in the 15–13.3 cal kyr BP period. This may indicate colder IWT at the beginning and the end of the last deglaciation, compared to the 15–13.3 cal kyr BP (Figure 4b). Another significant increase in the Mg/Li ratio is observed at around 11–10.3 cal kyr BP, followed by stable and low

values during the Holocene. During the last deglaciation, previous studies have indicated an enhanced northward flow of AAIW in both the Atlantic and Indian Oceans (e.g., Bryan et al., 2010; Ma et al., 2019; Pahnke et al., 2008; Poggemann et al., 2017, 2018), suggesting that the source of intermediate water masses could at least partly be the same in both oceans. To test this hypothesis, we can compare the *H. elegans* Mg/Li results from MD77-191 with another IWT record obtained from the tropical West Atlantic Ocean, calculated using *Uvigerina* spp. Mg/Ca ratios (Core M78/1-235-1, 11°36.53'N–60°57.86'W, 852 m water depth; Poggemann et al., 2018; Figures 4b and 4c). Despite a higher resolution for M78/1-235-1 compared to MD77-191, both records seem to exhibit some similar trends, at least for the last 15.2 cal kyr BP. To better highlight these patterns, we compare the smoothed curve of M78/1-235-1 in the tropical West Atlantic Ocean with the MD77-19 record. Both records display an increasing trend centered at 14 cal kyr BP, followed by decreased values at around 13.3–11.9 cal kyr BP, and then a rise from about 11 to 10.3 kyr. The same trend in decreasing values can be observed over the Holocene (Figure 4).

In order to assess the validity of this reconstruction, benthic foraminiferal δ^{18} O can be used, since it records changes in global ice volume, local temperature, and, potentially, salinity. We used the ice volume-related, global seawater δ^{18} O estimate provided by Bintanja and van de Wal (2008) to correct the benthic δ^{18} O record for ice volume changes for both Cores MD77-191 and MD77-176. Thus, the resulting corrected δ^{18} O record sindicate variations in both deepwater paleotemperature and/or salinity. In order to constrain past changes in the salinity, we can compare both core sites in the modern ocean. Indeed, the RSOW flows through a much stronger western boundary toward the Agulhas compared with eastward flows to the eastern boundary at 5°N (Beal et al., 2000; Talley et al., 2011; You, 1998). Thus, the site of MD77-191 (located in the eastern Arabian Sea) may be influenced by the RSOW, which is characterized by high salinity (Beal et al., 2000; Talley et al., 2011). However, the site of MD77-176 in the northeastern BoB should show little or no influence from the RSOW (Talley et al., 2011). Thus, the comparison between the δ^{18} O_{ivc} records obtained from MD77-176 and MD77-191 could reflect the influence of high-salinity RSOW (Figure 4a).

At the site of MD77-176, past IWT is more difficult to assess, as the benthic $\delta^{18}O_{ivc}$ was the only proxy used to reconstruct the past IWT. However, some similar trends are observed between the benthic $\delta^{18}O_{ivc}$ from MD77-176 and MD77-191. For the benthic $\delta^{18}O_{ivc}$ records obtained from these two cores, more enriched benthic $\delta^{18}O_{ivc}$ values (3.34‰ and 2.88‰, respectively) occurred during the 17–15.2 and 12.6–11.9 cal kyr BP time intervals, corresponding to colder and/or saltier deep water, and more depleted values (1.98‰ and 2.7‰, respectively) related to warmer and/or fresher deep water in the 15–13.3 cal kyr BP interval (Figure 4a). These trends match well with the MD77-191 *H. elegans* Mg/Li record at these time intervals, which may indicate cold IWT. By contrast, an increased trend in the Mg/Li ratio occurred at around 11–10.3 cal kyr BP, in opposition to the benthic $\delta^{18}O_{ivc}$ (Figures 4a and 4b). Differences are also observed during the Holocene, which could be related to past changes in the local salinity recorded in the benthic $\delta^{18}O_{ivc}$.

Thus, past IWT seem to have changed from the last deglaciation to the Holocene in the Arabian Sea, in the BoB, and in the tropical Atlantic, with the incursion of cold water masses at 17–15.2 and 12.6–11.9 cal kyr BP and two warmer events at 15–13.3 and 11–10.3 cal kyr BP. Furthermore, the $\delta^{18}O_{ivc}$ records obtained from MD77-176 and MD77-191 all display depleted values associated with warm and/or freshwater masses at 15–13.3 cal kyr BP during the last deglaciation, suggesting limited/no contribution of RSOW at the studied sites.

Then, a global trend to colder IWT is observed from the Mg/Li ratios across the Holocene, whereas both $\delta^{18}O_{ivc}$ records exhibit an opposite trend that may instead reflect the influence of fresher water masses. During the Holocene, increased benthic $\delta^{13}C$ and B-P age offsets obtained from MD77-176 in the BoB are synchronous with depleted ϵ_{Nd} records, suggesting the increased influence of NADW (Ma et al., 2019; Yu et al., 2018). All these records seem to indicate a limited influence of RSOW at both core sites during the Holocene, rather indicating an increased contribution of NADW (Ma et al., 2019; Yu et al., 2018).

However, during last deglaciation, colder water masses at 17–15.2 and 12.6–11.9 cal kyr BP in the North Indian Ocean seem to be inconsistent with previous studies suggesting a warming of AAIW during Heinrich Stadial 1 and Younger Dryas (Poggemann et al., 2018). This appearing discrepancy may relate to a smaller range of IWT variations in the northern Indian Ocean compared to other areas and/or to differences associated to local effects on the two areas that are very different. Thus, after having explored the past variations of IWT to track past intermediate water circulation changes, we investigate another important parameter, $[CO_3^{2-}]$.





Figure 5. Core-top *H. elegans* Sr/Ca versus deepwater $\Delta [CO_3^{2^-}]_{aragonite}$ (Rosenthal et al., 2006; Yu, Elderfield, et al., 2014).

6.2. Reconstructing Past Changes in the Intermediate Water $[CO_3^{2-}]$ 6.2.1. Calculation of Past $[CO_3^{2-}]$

According to previous studies by Rosenthal et al. (2006) and, more recently, by Yu, Elderfield, et al. (2014), the *H. elegans* Sr/Ca values seem to covary with the bottom water $\Delta[CO_3^{2-}]$. The $\Delta[CO_3^{2-}]$ is defined as $\Delta[\text{CO}_3^{2-}] = [\text{CO}_3^{2-}] - [\text{CO}_3^{2-}]_{\text{sat}}, \text{ where } [\text{CO}_3^{2-}] \text{ is carbonate ion and } [\text{CO}_3^{2-}]_{\text{sat}} \text{ is aragonite saturation}$ concentration. The calculation of $[CO_3^{2-}]_{sat}$ at the studied sites was carried out using CO_2 sys.xls (Ver. 12) (Pelletier et al., 2005). However, Rosenthal et al. (2006) have suggested that the H. elegans Sr/Ca is less sensitive to the Δ [CO₃²⁻]_{aragonite} variations when values are above 15 µmol/mol. Yu, Elderfield, et al. (2014) summarized the published Sr/Ca calibration data for C. wuellerstorfi, Cibicidoides mundulus, Uvigerina spp., and H. elegans. Although not yet tested thoroughly, they proposed that the benthic Sr/Ca may be used as an auxiliary proxy for seawater Δ [CO₃^{2–}]. To complete this approach, we collected the published core-top data for $\Delta [CO_3^{2-}]_{aragonite}$ paired with *H. elegans* Sr/Ca at intermediate water depth (500–1,500 m) (Rosenthal et al., 2006; Yu, Elderfield, et al., 2014) and plotted all of these Sr/Ca values together. Our database shows a positive relationship between Sr/Ca and bottom water Δ [CO₃²⁻]_{aragonite} defined as follows: $Sr/Ca = -0.0002 \pm 0.00003 \Delta [CO_3^{2-}]^2_{aragonite} + 0.0292 \pm 0.00207 \Delta [CO_3^{2-}]_{aragonite} + 1.3187 \pm 0.03404$ $(r^2 = 0.85)$ (Figure 5). The standard error of Δ [CO₃²⁻] estimation from the Sr/Ca according to the determined equation is $\pm 1 \,\mu$ mol/kg at $-18.5 \,\mu$ mol/kg and $\pm 7 \,\mu$ mol/kg at 74 μ mol/kg. Thus, we calculated the intermediate water Δ [CO₃²⁻] from the *H. elegans* Sr/Ca by using this equation. The seawater [CO₃²⁻]_{sat} is mainly affected by salinity, temperature, and water depth (pressure), for which variations on glacial-interglacial timescales are about $\pm 0.5 \,\mu$ mol/kg, remaining roughly unchanged in the deep ocean (Yu et al., 2008; Yu, Anderson, et al., 2014). Therefore, it is possible to get the paleo- $[CO_3^{2-}]$ of the intermediate water at our core sites, thanks to the equation defined above and assuming little changes in the seawater $[CO_3^{2-}]_{sat}$

The reconstructed intermediate water $[CO_3^{2-}]$ at the site of MD77-191 displays a decreasing trend during the 17–15.2 and 12.6–10.5 cal kyr BP time intervals, ranging between 97 and 77 µmol/kg and between 95 and 84 µmol/kg, respectively. The $[CO_3^{2-}]$ concentrations reach maximum values (~119 µmol/kg) in the 15–13.3 cal kyr BP. During the early Holocene (10.3–9 cal kyr BP), mean values are about 87 µmol/kg, and then we observed lower values of ~67 µmol/kg during the late Holocene (8.8–1.3 cal kyr BP).

The record of intermediate water $[CO_3^{2-}]$ at the site of MD77-176 shows slightly higher values during the LGM (~71 µmol/kg) than during the Holocene (~68 µmol/kg). For the period of the last deglaciation, the intermediate water $[CO_3^{2-}]$ record displays a decreasing trend to around ~67 µmol/kg at the 17–15.2 cal kyr BP interval and reaches a value of about ~77 µmol/kg at 14.6 cal kyr BP. An increasing trend is



observed from 13.4 to 11.7–11.5 cal kyr BP, with two extreme high values of 84–86 μ mol/kg. Then, a global decrease in the [CO₃^{2–}] is recorded during the Holocene.

6.2.2. Possible Controls on $[CO_3^{2-}]$ Records

On the basis of approximation, $[CO_3^{2-}] \approx 0.6 \times (ALK - DIC)$ (Yu et al., 2016), and thus, carbonate ion concentration in the seawater reflects the balance between mechanisms that affect changes in alkalinity (ALK) and total DIC. These mechanisms are varied, and the most important ones involve biological regeneration, changes in the water mass sources, and/or mixing of different water masses and air-sea exchanges (Yu et al., 2008). Deep water $[CO_3^{2-}]$ affects the dissolution/preservation of CaCO₃ on the seafloor by determining the seawater carbonate saturation and therefore acts as a control on the global oceanic ALK inventory (Yu, Anderson, et al., 2014).

Biological regeneration is linked to the sinking of organic matter and CaCO₃. The remineralization of sinking organic matter releases CO₂ in the deep ocean (Holligan & Robertson, 1996; Yu et al., 2008). The released CO₂ will consume $[CO_3^{2-}]$ to produce bicarbonate ions (HCO_3^{-}) , resulting in a decrease in carbonate ions (Holligan & Robertson, 1996). Thus, the ratio of organic to inorganic carbon (the so-called rain ratio) has an important effect on the $[CO_3^{2-}]$ concentration; an increasing rain ratio leads to the reduction of the $[CO_3^{2-}]$ concentration of the deep ocean (Berger & Keir, 1984; Holligan & Robertson, 1996; Yu et al., 2008).

At the location of MD77-191, paleoproductivity is controlled by summer monsoon upwelling activity. An abundance of *G. bulloides* identified in Cores MD77-191 and SK237 GC04 (10°58.65'N–74°59.96'E, 1,245 m water depth) indicates a progressive increase of surface productivity during the Holocene, with a significant increase since ~6 cal kyr BP (Bassinot et al., 2011; Naik et al., 2017; Figure 6). In addition, the total organic carbon (C_{org}) can be used as an indicator of paleoproductivity (e.g., Calvert et al., 1995; Naidu et al., 1992). The C_{org} record obtained from SK237 GC04 also shows an increasing trend during the late Holocene and reaches a maximum value at the end of the Holocene (Figure 6). An increasing rain ratio (more organic matter sinking) caused a decreased [CO_3^{2-}] concentration in the deep ocean (Berger & Keir, 1984; Holligan & Robertson, 1996; Yu et al., 2008; Yu et al., 2019). In addition, a decrease in the [CO_3^{2-}] concentration is observed, which is consistent with high productivity during the Holocene (8.8–1.3 cal kyr BP). This hypothesis is also supported by the benthic Cd/Ca record from MD77-191 as a proxy to reconstruct the paleonutrient concentration of the intermediate water (unpublished data). Thus, we suggest that at the site of Core MD77-191, changes in the paleoproductivity resulting from the progressive increase of upwelling activity have involved an increase in organic matter rain and a depletion of $[CO_3^{2-}]$.

However, at the site of Core MD77-176 in the northern BoB, primary productivity is much lower than in the Arabian Sea in modern times and may not have a strong influence on the intermediate water $[CO_3^{2^-}]$. Indeed, the modern distribution of chlorophyll in the surface water of the northeastern BoB indicates a slight increase in productivity during the winter monsoon (O'Malley, 2017; Thushara & Vinayachandran, 2016), and to our knowledge, there is no paleorecord of primary productivity in this area. Moreover, the paleosurface productivity in the BoB is much lower compared to the Arabian Sea (Prasanna Kumar et al., 2001). Thus, assuming that changes in primary productivity since the last deglaciation may not have been as strong as in the Arabian Sea, indicating less organic matter sinking from the surface, we suggest that paleoproductivity does not appear to be a major control in affecting the variation of $[CO_3^{2^-}]$ concentrations in the northeastern BoB.

In addition, in order to examine the variations at a millennial timescale since the last deglaciation at intermediate and deepwater depth from a global view, we compared the seawater $[CO_3^{2-}]$ records from MD77-191 and MD77-176 with $[CO_3^{2-}]$ results reconstructed from the benthic foraminiferal B/Ca at different water depths from the Atlantic Ocean (BOFS 17K, 58.0°N–16.5°E, 1,150 m water depth) and the Indian Ocean (WIND 28K, 10.2°S–51.8°E, 4,147 m water depth) (Yu et al., 2008; Yu, Broecker, et al., 2010) (Figure 6). Similar trends are observed during the Holocene, with lower values recorded throughout these locations (Figure 6). This global decrease is likely to result from a depletion in the oceanic ALK (Yu, Anderson, et al., 2014), since we know that reducing the whole ocean ALK inventory may decrease the CO_2 solubility, thus driving up the atmospheric CO_2 over the last 8 cal kyr BP (Menviel & Joos, 2012; Ridgwell et al., 2003; Yu, Anderson, et al., 2014). In addition, the decreased $[CO_3^{2-}]$ during the Holocene could also be associated with increased coral reef construction due to the higher sea level, which could





Figure 6. (a) The total organic carbon percentage (%C_{org}) and (b) *G. bulloides* percentage from Core SK237 GC04 (1,245 m, Naik et al., 2017). (c) The relative abundance of *G. bulloides* and (d–g) comparison between $[CO_3^{2-}]$ records obtained from studied Cores MD77-191 (Arabian Sea) and MD77-176 (Bay of Bengal) with other $[CO_3^{2-}]$ records at different depths in the Indian Ocean and Atlantic Ocean (Yu et al., 2008; Yu, Broecker, et al., 2010). The color-shaded intervals are the same as in Figure 2.

remove the global ALK and induce a decline in $[CO_3^{2-}]$ over the whole ocean (Opdyke & Walker, 1992; Yu, Anderson, et al., 2014).

During the last deglaciation, the $[CO_3^{2-}]$ records at intermediate water depth from the Arabian Sea and the Atlantic Ocean display low values during the 17-15.2 and 12.6-10.5 kyr intervals. The record of intermediate $[CO_3^{2-}]$ obtained from MD77-176 closely matches previous records for the 17–15.2 cal kyr BP interval but, in contrast, displays maximum values (~84-86 µmol/kg) at 11.7-11.5 cal kyr BP, corresponding to a marked decrease trend of the benthic $\delta^{18}O(2.51\%)$ and benthic $\delta^{13}C(-0.19\%)$ at 11.5 cal kyr BP. Variations during this specific time interval thus seem to be associated with a local effect at the site of MD77-176, combining changes in the water mass salinity and/or temperature and $[CO_3^{2-}]$. Furthermore, the decrease of benthic δ^{13} C may correspond to the influenced discharge of terrigenous organic matter into the ocean, in a good agreement with the marked depleted benthic δ^{18} O, which associated with warm and/or fresh water. Thus, this significant variation seems to be associated to the local large input of fresh water in the BoB. At the same time, in the deep Atlantic, equatorial Pacific, and Indian Oceans, deepwater $[CO_3^{2-}]$ concentrations rose by about ~10 µmol/kg (Yu et al., 2008; Yu, Anderson, et al., 2014; Yu, Broecker, et al., 2010; Yu, Foster, et al., 2010). The increased deepwater $[CO_3^{2-}]$ would have promoted deep-sea carbonate preservation and hence would have depleted the oceanic ALK inventory. Such an increase in deepwater $[CO_3^{2-}]$ concentration is coeval with the two-step deglacial increase of atmospheric CO_2 (Figure 7). This probably indicates that the deepwater carbonate ion increases results from a drop in DIC associated with the transfer of CO2 from the deep ocean to the upper water depth and, ultimately, to the atmosphere. The release of deep-sea CO_2 may have induced a $[CO_3^{2-}]$ decline at intermediate depths (Yu, Anderson, et al., 2014). Such a hypothesis is consistent with the variations in intermediate $[CO_3^{2-}]$ concentration records from the northern BoB, Arabian Sea, and northern Atlantic Ocean (Figure 6).

6.3. Significance With Regard to Paleoclimate and Atmospheric CO₂ Changes

Comparing the paleo- $[CO_3^{2-}]$ records with other proxies, the decrease in $[CO_3^{2-}]$ records corresponds to an increase in benthic δ^{13} C during the last deglaciation. Such an increase in benthic δ^{13} C values has also been observed in previous studies from the northern Indian Ocean (e.g., Curry et al., 1988; Duplessy et al., 1984; Jung et al., 2009; Ma et al., 2019; Naqvi et al., 1994). This increase of benthic δ^{13} C in the Holocene may reflect more invigorated circulation associated with better-ventilated waters at intermediate depth in the northern Indian Ocean (Duplessy et al., 1984; Ma et al., 2019; Waelbroeck et al., 2006). The similarity of the benthic δ^{13} C increases reflects the northward expansion of AAIW during time intervals 17-15.2 and 12.6-10.5 cal kyr BP in the western Arabian Sea, Pacific Ocean, and BoB (Figure S3) (Jung et al., 2009; Ma et al., 2019; Pahnke & Zahn, 2005). However, as AAIW is formed from the deep waters of the AABW, air-sea exchange could also affect the values of δ^{13} C, as the intermediate benthic δ^{13} C in the Southern Ocean is dominated by the influence of air-sea exchange (Lynch-Stieglitz et al., 1994). Thus, values for intermediate benthic $\delta^{13}C$ could increase relatively via stronger upwelling during the formation of AAIW. In addition, the transition in the ϵ_{Nd} and Δ^{14} C records during the deglaciation also indicates a strong northward penetration of AAIW within the North Atlantic and BoB (e.g., Cao et al., 2007; Ma et al., 2019; Pahnke et al., 2008; Pena et al., 2013; Yu et al., 2018). During the last deglaciation, MD77-176 B-P age offsets display a decreasing trend in the 17–15.2 and 12.6–10.5 cal kyr BP intervals, corresponding to increases in benthic δ^{13} C values (Figure 7). The variations in the B-P age offsets obtained from MD77-176 (northern BoB), Arabian Sea, and Pacific Ocean all indicate stronger upwelling and an enhanced northern flow of AAIW from the Southern Ocean during these two periods.

Therefore, during the 17–15.2 and 12.6–10.5 cal kyr BP intervals, changes in the benthic δ^{13} C, B-P age offsets and ε_{Nd} records obtained from MD77-191 and MD77-176, associated to an increased atmospheric CO₂, could strongly support the hypothesis that enhanced vertical ventilation in the Southern Ocean could have led to an increased production of intermediate water masses (AAIW). As a consequence, enhanced Southern Ocean ventilation could have played an important role in the CO₂ increase during the last deglaciation (Anderson et al., 2009; Poggemann et al., 2017, 2018; Skinner et al., 2014).

In addition, as mentioned before, during the last deglaciation, the synchronicity of depleted intermediate water $[CO_3^{2-}]$ in the Indian and Atlantic Oceans increased deepwater $[CO_3^{2-}]$ in the global ocean (Yu





Figure 7. (a) Ice core atmospheric CO₂ from Antarctic Dome C (Monnin et al., 2001), (b) the intermediate water B-P ¹⁴C age offset of MD77-176 in the BoB (Ma et al., 2019), (c and d) benthic δ^{13} C from studied Cores MD77-176 and MD77-191, respectively, and (e and f) compilation of [CO₃²⁻] records from Cores MD77-191 and MD77-176. The color-shaded intervals are the same as in Figure 2.

et al., 2008; Yu, Foster, et al., 2010; Yu, Broecker, et al., 2010; Yu, Anderson, et al., 2014), associated with the transfer of CO_2 from the deep ocean to the atmosphere (Yu, Anderson, et al., 2014). Thus, the depleted intermediate water $[CO_3^{2-}]$ could strongly support that variations of the Southern Ocean circulation contribute to these increases of atmospheric CO_2 by transferring the deep ocean carbon to the upper



ocean and atmosphere during the last deglaciation, by enhancing the upwelling and increasing the northward penetration of the AAIW in all ocean basins (e.g., Anderson et al., 2009; Jung et al., 2009; Ma et al., 2019; Pahnke & Zahn, 2005) at 17–15.2 and 12.6–10.5 cal kyr BP. In addition, the variations in intermediate-deepwater $[CO_3^{2-}]$ are associated with the carbonate compensation and/or reef buildup due to increased sea level; these could confirm the influence of whole ocean ALK inventory in carbon reorganization in the Earth's biosphere system (Yu, Anderson, et al., 2014). Furthermore, for MD77-191 located in the south tip of India, the depleted intermediate water $[CO_3^{2-}]$ during the Holocene may also be influenced by increased local paleoproductivity.

7. Conclusions

Benthic foraminifera δ^{13} C, δ^{18} O, and elemental ratios of *H. elegans* have been analyzed on Core MD77-191 (1,254 m water depth) located at the southern tip of India, as well as Core MD77-176 (1,375 m water depth) obtained from the northern BoB, to reconstruct the evolution of intermediate water masses in the northern Indian Ocean since the last deglaciation.

The past IWT in the northern Indian Ocean were mainly reconstructed using the *H. elegans* Mg/Li record obtained from MD77-191 in the Arabian Sea. It seems to indicate the influence of cold water mass during the last deglaciation (17–15.2 and 12.6–11.9 cal kyr BP time intervals) and a global trend toward colder IWT during the Holocene. In addition, two warmer events occurred at around 15–13.3 and 11–10.3 cal kyr BP time intervals.

We reconstructed seawater $[CO_3^{2-}]$ concentration by converting *H. elegans* Sr/Ca to intermediate water $[CO_3^{2-}]$ using a modern, core top-based, empirical relationship with the standard error of $\Delta[CO_3^{2-}] \pm 1 \,\mu$ mol/kg at -18.5 μ mol/kg and $\pm 7 \,\mu$ mol/kg at 74 μ mol/kg. During the last deglaciation, increased benthic δ^{13} C, depletion in $[CO_3^{2-}]$, and decreased B-P age offsets occurred in the 17–15.2 and 12.6–10.5 cal kyr BP intervals. All of these results suggest a strong linkage between Southern Ocean enhanced ventilation via AAIW and release of CO_2 during the last deglaciation. The decreased $[CO_3^{2-}]$ in intermediate water masses also provides evidence for the important influence of variations in the global ALK inventory during the late Holocene, corresponding to the rise in atmospheric CO_2 since 8 kyr BP and/or increased productivity throughout the Holocene (at least for MD77-191).

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