The ²²⁶Ra-Ba relationship in the North Atlantic during GEOTRACES-GA01

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Abstract. We report detailed sections of radium-226 (226 Ra, $T_{1/2} = 1602$ y) activities and barium (Ba) concentrations determined in the North Atlantic (Portugal-Greenland-Canada) in the framework of the

international GEOTRACES program (GA01 section-GEOVIDE project, May-July 2014). Dissolved 23 ²²⁶Ra and Ba are strongly correlated along the section, a pattern that may reflect their similar chemical 24 behavior. Because ²²⁶Ra and Ba have been widely used as tracers of water masses and ocean mixing, we 25 investigated more thoroughly their behavior in this crucial region for thermohaline circulation taking 26 advantage of the contrasting biogeochemical patterns existing along the GA01 section. We used an 27 Optimum Multiparameter (OMP) analysis to distinguish the relative importance of physical transport 28 (water mass mixing) and non-conservative processes (sedimentary, river, or hydrothermal inputs; 29 uptake by particles, and dissolved-particulate dynamics) on the ²²⁶Ra and Ba distributions in the North 30 Atlantic. Results show that the measured ²²⁶Ra and Ba concentrations can be explained by conservative 31 mixing for 58 and 65 % of the samples respectively, notably at intermediate depth, away from the ocean 32 interfaces. ²²⁶Ra and Ba can thus be considered as conservative tracers of water mass transport in the 33 ocean interior at the space scales considered here, namely, of the order of few thousand km. However, 34 regions where ²²⁶Ra and Ba displayed non-conservative behaviors and in some cases decoupled 35 behaviors were also identified, mostly at the ocean boundaries (seafloor, continental margins, and 36

surface waters). Elevated ²²⁶Ra and Ba concentrations found in deep waters of the West European Basin
 suggest that lower North East Atlantic Deep Water (NEADWI) accumulates ²²⁶Ra and Ba from

sediment diffusion and/or particle dissolution during transport. In the upper 1500 m of the West 1 European Basin, deficiencies in ²²⁶Ra and Ba are likely explained by their incorporation in planktonic 2 calcareous and siliceous shells, or in barite (BaSO₄) by substitution or adsorption mechanisms. Finally, 3 because Ba and ²²⁶Ra display different source terms (mostly deep-sea sediments for ²²⁶Ra and rivers for 4 Ba), strong decoupling between ²²⁶Ra and Ba were observed at the land-ocean boundaries. This is 5 especially true in the shallow stations near the coasts of Greenland and Newfoundland where high 6 ²²⁶Ra/Ba ratios at depth reflect the diffusion of ²²⁶Ra from sediment and low ²²⁶Ra/Ba ratios in the upper 7 water column reflect the input of Ba associated with meteoric waters. 8

9 1. Introduction

The primary source of radium-226 (226 Ra, $T_{1/2} = 1602$ y) to the ocean was found to be the diffusion 10 from deep-sea sediments following the decay of its parent isotope, ²³⁰Th (Koczy, 1958; Kröll, 1953). 11 12 This mode of introduction led Koczy to use radium data to derive vertical eddy diffusivities and velocities in the deep sea (Koczy, 1958). Since then, ²²⁶Ra has been widely used to study the ocean 13 circulation and mixing at a global scale (Chung and Craig, 1980; Ku et al., 1980). In the framework of 14 the Geochemical Ocean Sections Study (GEOSECS) program, special attention was given to ²²⁶Ra as its 15 solubility and half-life made it an ideal candidate as a chronometer of the global thermohaline 16 circulation. In particular, its 1602 y half-life is more adapted than the longer half-life of carbon-14 (¹⁴C, 17 $T_{1/2} = 5730$ y) that had also been used for that purpose. Therefore, the global oceanic distribution of 18 ²²⁶Ra in seawater was extensively documented throughout the major ocean basins and a unique database 19 was generated during the GEOSECS program (Broecker et al., 1970, 1967; Chung et al., 1974; Ku et 20 21 al., 1970; Ku and Lin, 1976).

Barium (Ba) is an alkaline earth element like ²²⁶Ra. Therefore they share a similar geochemical 22 behavior in the ocean (Chan et al., 1976; Fanning et al., 1988; Mathieu and Wolgemuth, 1973). As such, 23 Ba was proposed as a stable analog of ²²⁶Ra in order to use the ²²⁶Ra/Ba ratio as a clock in a similar 24 manner as the ¹⁴C/¹²C ratio. However, the recognition that ²²⁶Ra and Ba participate in upper ocean 25 biological cycles (Ku and Luo, 1994) introduced additional complications for the use of the ²²⁶Ra/Ba 26 ratio as a time tracer for deep water ventilation. Both ²²⁶Ra and Ba indeed increase with increasing 27 depth, thus reflecting uptake due to biological processes in surface waters, particles scavenging and 28 subsequent release at depth through the dissolution of the settling particles (Broecker et al., 1967; Ku et 29 al., 1970; Ku and Luo, 1994). ²²⁶Ra and Ba are thus not only controlled by physical processes, but 30 31 appear to be incorporated in settling particles such as calcareous and siliceous shells, or in barite (BaSO₄) that precipitates in the mesopelagic zone (Bishop, 1988; Chan et al., 1976; Dehairs et al., 1980; 32 Lea and Boyle, 1989, 1989). Hence, despite different principal sources to the ocean, rivers in the case of 33

Ba and marine sediment diffusion for ²²⁶Ra, their distributions are affected by similar processes in the 1 water column. Barium displays a linear correlation with ²²⁶Ra in the global ocean, resulting in a fairly 2 constant ²²⁶Ra/Ba ratio of 2.2 ± 0.2 dpm μ mol⁻¹ (dpm, disintegrations per minute) (Chan et al., 1976; 3 Foster et al., 2004; Ku et al., 1980; Li et al., 1973; Östlund et al., 1987). Similarly, strong correlations 4 were also found between Ba-Si (silicate) and ²²⁶Ra-Si although no obvious process links ²²⁶Ra or Ba 5 with Si. These latter relationships appeared to be more surprising because Si is not a chemical analog of 6 Ra and Ba. It was first proposed that diatom frustules exported from the upper water column could 7 8 adsorb Ra and Ba, these elements being then released at depth following the dissolution of the siliceous 9 tests (Bishop, 1988; Chung, 1980; Dileep Kumar and Li, 1996). More recent studies showed that the 10 similar behaviors of Ba and Si (and alkalinity) reflect similar dissolved-particulate interactions (Jeandel 11 et al., 1996; McManus et al., 1999; Rubin et al., 2003). Indeed, Ba is not mechanistically coupled with alkalinity or silicate. Rather, the observed relationships may result from the spatial coherence of 12 different carriers (barite, opal and carbonate) overprinted by hydrodynamics. The formation of biogenic 13 silica, CaCO₃ and barite in surface water and their subsequent dissolution in the deeper water column 14 15 may thus generate parallel oceanic distributions. While barite has been shown to be the main carrier that controls the Ba water column distribution, the relationship between Ba-Ra remains unclear. 16

While the global GEOSECS program provided valuable information on the coupling between biogeochemical cycles of ²²⁶Ra and Ba in the ocean, several unknowns still remain. In this work, we take advantage of a new worldwide program, GEOTRACES, to provide new information on the distribution of ²²⁶Ra and Ba and their relationship in the ocean. GEOTRACES program aims to characterize the distribution of trace elements and their isotopes (TEIs) (sources, sinks, internal cycling) in the ocean through a global survey consisting of ocean sections and regional process studies.

In the present study, we report dissolved ²²⁶Ra activities and dissolved Ba concentrations in the 23 North Atlantic Ocean and Labrador Sea (GEOVIDE project, GA01 section). The North Atlantic region 24 hosts a variety of globally significant water masses with complex circulation patterns (García-Ibáñez et 25 al., this issue; Lherminier et al., 2010). This area is crucial for the thermohaline circulation, and thus for 26 27 global climate, through its important role in the ventilation of the deep layer of the global ocean (Seager et al., 2002). As part of this process, the Meridional Overturning Circulation (MOC) includes the 28 northward transport of warm subtropical waters. These surface waters are then cooled and transformed 29 30 into subpolar waters, and may reach the Labrador and Irminger Seas where deep-water formation and deep convection take place (Bennett et al., 1985; Pickart and Spall, 2007; Yashayaev et al., 2007). We 31 propose to study the relationship between ²²⁶Ra and Ba and to test the conservative behavior of these 32 tracers in this specific region. We further document the Ra-Ba-Si relationship along the GA01 section, 33

as it was done in previous sections conducted during the GEOSECS program and more recently along
 the GEOTRACES GA03 section.

3 2. Materials and Methods

4 2.1. Study area; the GEOVIDE project

5 The GEOTRACES GA01 section (GEOVIDE project; PIs : Géraldine Sarthou, LEMAR, France and Pascale Lherminier, LOPS, France) was conducted in the North Atlantic Ocean between Lisbon, 6 7 Portugal, and St John's, Canada (15 May 2014-30 June 2014; Fig.1). The water samples described here were collected on board the R/V Pourquoi Pas?. The section crossed different topographic features and 8 9 regions with contrasting biogeochemical patterns. It complemented the sections GA03 (U.S.-GEOTRACES) and GA02 (Dutch GEOTRACES) also conducted in the Atlantic Ocean in the 10 framework of the GEOTRACES program. Seventy-eight stations were visited during the GEOVIDE 11 12 project.

13 2.2. Sample collection

14 At 15 of the 78 stations completed during the GA01 cruise, up to 22 discrete 10-L seawater samples were collected through the water column from Niskin bottles. The seawater samples were passed by 15 gravity through 10 g of acrylic fibers impregnated with MnO₂ (so called "Mn fibers"), which 16 quantitatively adsorb radium isotopes (assumed to scavenge 100 % of Ra; van Beek et al., 2010; Moore 17 and Reid, 1973). High-resolution vertical profiles of ²²⁶Ra were thus built to provide a detailed ²²⁶Ra 18 section. The samples were unfiltered since particulate ²²⁶Ra activities are typically two orders of 19 magnitude lower than the dissolved ²²⁶Ra activities (van Beek et al., 2007, 2009). From the same Niskin 20 bottles, 15 mL was collected to determine the Ba concentration, so that Ba and ²²⁶Ra analyses were 21 conducted from the same initial sample, which allows us to investigate the ²²⁶Ra/Ba ratio in the 22 samples. The Ba samples were collected in pre-cleaned polypropylene bottles (rinsed three times with 23 the same seawater sample), acidified with 15 µL of HCl (10 M, Merck, Suprapur) and kept at room 24 temperature for later analysis. 25

26 2.3. Analysis of dissolved ²²⁶Ra activities *via* ²²²Rn emanation

Radium-226 was determined *via* its daughter, radon-222 (222 Rn; T_{1/2} = 3.8 days) using a radon extraction system followed by alpha scintillation counting (Key et al., 1979). The Mn Fiber samples were placed into gas-tight PVC cartridges (Peterson et al., 2009) that were flushed with helium (He) for 5 min at 250 mL min⁻¹. The cartridges were sealed and held for approximately 2 weeks (minimum of 5

days) to allow for ²²²Rn ingrowth from ²²⁶Ra decay. The ²²²Rn was then flushed out from the cartridges 1 using He and cryo-trapped in copper tubing using liquid nitrogen. The copper trap was heated to allow 2 3 the ²²²Rn to be transferred to an evacuated "Lucas cells" via a stream of He. The "Lucas cells" are airtight chambers with inner walls coated with silver-activated zinc sulfide that emits photons when struck 4 by alpha decay particles (Key et al., 1979; Lucas, 1957; Peterson et al., 2009). The cells were held 3 5 hours to reach the secular equilibrium of all ²²²Rn decay chain daughters. After 3 hours, the samples 6 were counted overnight on a radon counting system (Model AC/DC-DRC-MK 10-2). The counting 7 uncertainties (1SD, Standard Deviation) were within the range of 2-5% for 10 L volume samples. All 8 9 samples were appropriately ingrowth and decay corrected. The combined Lucas cell and detector background was \sim 7 % of the typical total measured sample activity. The method was standardized using 10 NIST (U.S. National Institute of Standards & Technology)²²⁶Ra (20 dpm) sorbed onto MnO₂ fiber and 11 analyzed in the same manner as the samples, with uncertainties (1SD) of 5 % (Charette et al., 2015; 12 Henderson et al., 2013). Vertical profiles of ²²⁶Ra from the GEOTRACES GA01 (this study) and GA03 13 14 (Charette et al., 2015) sections that were located in close proximity off Portugal (Fig.1) were compared, and showed a good agreement with increasing activities with increasing depth (Fig.S1). 15

16 2.4. Analysis of dissolved Ba concentrations

17 Barium concentrations were measured using an isotope dilution (ID) method (Freydier et al., 1995; 18 Klinkhammer and Chan, 1990) by high resolution-inductively coupled plasma-mass spectrometry (HR-19 ICP-MS). This method was adapted to a Thermo Finnigan Element XR instrument (MIO, Marseille). 20 The Ba measurements presented here are the sum of dissolved Ba and a very small fraction of 21 particulate Ba (generally <1 % of total Ba, along GEOVIDE up to 1.3 % at the bottom of station 32 due to presence of a nepheloid layer; Lemaitre et al., this issue) released from the samples as a result of the 22 acidification step. Hence, while the measurements reported herein are total Ba, they are within 23 analytical uncertainty representative of the dissolved Ba pool. The samples (0.5 mL) were spiked with 24 300 µL of a ¹³⁵Ba-enriched solution (93 % ¹³⁵Ba; 95 nmol kg⁻¹) and diluted with 15 mL of acidified 25 (2 % HNO₃, 14 M, Optima grade) Milli-Q grade water (Millipore). The amounts of sample, spike and 26 dilution water were assessed by weighing. The reproducibility of this method is about 1.5 % (1 RSD, 27 28 Relative Standard Deviation), as tested on repeated preparations of the reference solution SLRS-5 29 (NRC-CNRC river water reference material for trace metals). Average Ba values obtained for SLRS-5 were $13.48 \pm 0.20 \ \mu g \ L^{-1}(1 \ \sigma)$ with RSD of 1.5 %, which is in good agreement with the certified values 30 31 (SLRS-5 13.4 \pm 0.6 µg L⁻¹). The limit of detection calculated as three times the standard deviation of the procedural blank was 0.09 nmol L^{-1} . 32

1 2.5. Multiparameter mixing model

2 An Optimum MultiParameter (OMP) analysis was used to distinguish the relative importance of 3 physical transport (i.e., water mass mixing) from non-conservative processes (input from the sediments, rivers or hydrothermal vents, dissolution of particles; uptake by particles and dissolved-particulate 4 dynamics) on the ²²⁶Ra and Ba distributions in the North Atlantic. We used the OMP analysis computed 5 for the GA01 section by Garcia Ibanez et al. (this issue) with 12 source water types (SWTs). Based on 6 historical data reported from the North Atlantic, we defined ²²⁶Ra and Ba endmember concentrations 7 associated with each SWT (Table S1). The characteristics of SWTs (potential temperature, salinity, and 8 9 geographical location) reported by Garcia Ibanez et al. (this issue) were used to determine the SWTs endmembers for ²²⁶Ra and Ba,). In some cases, data from the GA01 section were used for the SWT 10 endmember (Table S1). These ²²⁶Ra and Ba SWT endmembers were then used to calculate the ²²⁶Ra 11 and Ba concentrations that strictly result from mixing of the different water masses. In this way, we 12 estimated the conservative component of ²²⁶Ra and Ba, which can in turn be compared to the *in situ* 13 concentrations to generate the non-conservative component of ²²⁶Ra and Ba along the GA01 section. 14

The uncertainties for the SWT endmembers were considered to be ± 0.6 dpm 100 L⁻¹ for ²²⁶Ra and 15 ± 1.5 nmol L⁻¹ for Ba, based on typical measurement uncertainties and spatial variability. A Monte 16 Carlo method (250 random perturbations) was used to propagate the SWT endmember uncertainties to 17 the ²²⁶Ra and Ba results of the OMP analysis. This allowed us to estimate conservative component 18 uncertainties of \pm 0.9 dpm 100 L⁻¹ for ²²⁶Ra and \pm 4.4 nmol L⁻¹ for Ba. When the measured ²²⁶Ra and Ba 19 values were different from the conservative component values (taking into account the uncertainties on 20 these values), ²²⁶Ra and Ba were considered as non-conservative. These non-conservative values can 21 either be positive (representative of a net addition of ²²⁶Ra and Ba) or negative (representative of a net 22 removal of ²²⁶Ra and Ba). 23

Note that the OMP analysis was not solved where non-conservative behavior of temperature and salinity is expected (that is, for waters above 100 m and for waters with salinities lower than 34.7). In these latter cases, changes in water mass properties may indeed be due to air-sea interaction or inputs of fresh waters (e.g., near Greenland shelf; Daniault et al., 2011).

28 3. Results

29 3.1. Hydrodynamic context

The OMP analysis was used to identify the different water masses (Table 1) crossing the GA01 section. The potential temperature-salinity diagram for all the GA01 stations along with the different SWT endmembers used in the OMP analysis are represented in Fig.2. The salinity section is shown in
 Fig.3. The different water masses present along the GA01 section are described below.

3 3.1.1. Upper waters

4 Three main water masses were found in upper waters (<1000 m) in the investigated area (Fig.3). First, the Central Waters occupied the upper eastern part of the GA01 section from the Iberian Peninsula 5 to the Rockall Through (stations 1 to 26). Their distribution was associated with the circulation of the 6 North Atlantic Current (NAC). The NAC transports warm and saline waters northward, connecting the 7 subtropical and the subpolar latitudes, and is part of the upper layer of the Atlantic Meridional 8 9 Overturning Circulation (AMOC) in the North Atlantic subpolar gyre. The NAC flows eastward from 10 the Grand Banks of Newfoundland, splitting into four branches west of the Mid-Atlantic Ridge (MAR), while incorporating local water masses (Fig.1). East of the MAR, the two northern branches of the NAC 11 12 flow northward into the Icelandic Basin, the Rockall Plateau and the Rockall Trough, while the two southern branches flow southward into the West European Basin. The Central Waters can be identified 13 by the highest potential temperature of the entire GA01 section and are represented by two endmembers 14 called East North Atlantic Central Waters (ENACW₁₆ and ENACW₁₂). The ENACW₁₆ is warmer 15 16 (16 °C) than the ENACW₁₂ that can be identified with a potential temperature of 12.3 °C (Fig.2).

Part of the Central Waters carried by the NAC recirculates toward the Iceland Basin and the Irminger Sea, leading to the formation of subpolar mode waters by mixing and cooling in the subpolar gyre (Lacan and Jeandel, 2004; McCartney, 1992). Iceland Subpolar Mode Water (IcSPMW) is formed in the Icelandic Basin and is located, along GA01, over the Reykjanes Ridges (stations 32 and 38) (Fig.3). The IcSPMW is described by two endmembers, the SPMW₇ and the SPMW₈, which are distinguished by their potential temperature of 7.0 °C and 8.0 °C, respectively (Fig.2). Once formed, the IcSPMW follows the Irminger Current.

Finally, the Irminger Subpolar Mode Water (IrSPMW) is the result of the transformation of the Central Waters and the IcSPMW, and is formed northwest of the Irminger Sea (Krauss, 1995). The IrSPMW is located near Greenland (stations 53, 57 and 60) (Fig.3) (García-Ibáñez et al., 2015; Lacan and Jeandel, 2004; Read, 2000).

28 3.1.2. Intermediate waters

The Subarctic Intermediate Water (SAIW) originates in the Labrador Current (Read, 2000). The SAIW is associated with the advection of waters from the Labrador Sea within the NAC; it subducts below the Central Waters at approximately 600 m. Low salinities (34.8 and 34.7) and potential temperatures of 4.5 °C and 6 °C are representative of the two SAIWs, SAIW₄ and SAIW₆, respectively
(Fig.2).

Around the Rockall Plateau, the SAIW overlies the Mediterranean Water (MW). The MW enters the North Atlantic through the Gibraltar Strait and flows northward while extending westward. The MW can be identified in the West European Basin at approximately 1200 m (stations 1 and 13 in Fig.3) with a salinity of 36.5 (Fig.2; García-Ibáñez et al., 2015).

7 The Labrador Sea water (LSW) is found in multiple locations and at different water depths along the GA01 section (Fig.3). The LSW is formed by progressive cooling and freshening in winter due to 8 9 deep convection. The LSW can be characterized by its minimum salinity (34.87) and its minimum 10 potential temperature (3 °C) (Fig.2). The LSW contributes to the stratification of the interior of the North Atlantic and its boundary currents and spreads at intermediate depths in three different basins 11 intersected by the GA01 section (Fig.1). The three independent pathways are: (i) northward into the 12 Irminger Sea (station 44), (ii) eastward across the MAR, through the Charlie-Gibbs fracture zone, then 13 14 northward into the Iceland Basin (station 32) and eastward into the West European Basin (stations 21 and 26), and (iii) equatorward as a major component of the North Atlantic Deep Water in the Deep 15 Western Boundary Current (DWBC), which constitutes the lower limb of the AMOC. Along these 16 paths, the LSW mixes with both the overlying and underlying water masses and becomes warmer and 17 18 saltier (Lazier, 1973).

The Polar Intermediate Water (PIW) is characterized by very low salinity (34.9) and potential temperature (less than 2 °C) (Fig.2) and is defined as a separate upper core on the Greenland slope. The PIW is episodically injected into the Irminger Sea and originates from either the Arctic Ocean or the Greenland shelf (Falina et al., 2012; Jenkins et al., 2015; Rudels et al., 2002).

23 **3.1.3.** Overflow waters and deep waters

The Iceland-Scotland Overflow Water (ISOW) originates at the Iceland-Scotland sill, and entrains the overlying warm saline atlantic waters (SPMW and LSW). ISOW identification features are a potential temperature of 2.6 °C and a salinity of 34.98 (Fig.2; van Aken and Becker, 1996). ISOW was found at stations located on the Eastern flank of the Reykjanes Ridge (stations 32 and 38) and near Greenland (stations 60 and 64) at great depth (2000-3500 m) (Fig.3).

Overflow waters coming from the Denmark Strait mix with both the SPMW and the LSW during descent into the Irminger Sea to form the Denmark Strait Overflow Water (DSOW) (Fig.1) (Read, 2000; Yashayaev and Dickson, 2008). DSOW is located at the northern end of the Irminger Sea (station 44) and occupies the deepest part of the Greenland continental slope (stations 69 and 77) 1 (Fig.3). At bottom depth, DSOW is easily identified by a minimum potential temperature of 1.3 °C
2 (Fig.2).

In the Southern Ocean, the Antarctic Bottom water (AABW) is formed by deep winter convection of surface waters. AABW flows to the north along the eastern side of the Atlantic and contributes to the formation of the lower North East Atlantic Deep Water (NEADWl) once this water penetrates the Iberian Abyssal Plain by crossing the Mid-Atlantic Ridge (Fig.1). The NEADWl is laying at the bottom of the West European Basin (stations 1 to 26 in Fig.3) with a mean salinity of 34.895 and a potential temperature of 1.98 °C (Fig.2). Then, the NEADWl recirculates into the Rockall Trough and meets ISOW in the Iceland Basin (van Aken, 2000; McCartney, 1992; Schmitz and McCartney, 1993).

10 **3.2.** Distribution of ²²⁶Ra and Ba along the GA01 section

The ²²⁶Ra distribution for the GA01 section is presented in Fig.4 with Ba concentrations and ²²⁶Ra/Ba ratios. The ²²⁶Ra activities and Ba concentrations in the water column range from 7 to 25 dpm 100 L⁻¹ and from 33.6 to 81.5 nmol L⁻¹, respectively. These data are in good agreement with Atlantic data from the GEOSECS program, which range from 6.8 to 23.4 dpm 100 L⁻¹ for ²²⁶Ra and from 35 to 105 nmol L⁻¹ for Ba (Broecker et al., 1976).

For both ²²⁶Ra and Ba, the vertical gradient is stronger in the eastern part of the section (West 16 European Basin) than on the western part of the section (from Reykjanes Ridge to Newfoundland). Both 17 are particularly high near the seafloor in the West European Basin (²²⁶Ra: 14-25 dpm 100 L⁻¹: Ba: 18 63-82 nmol L⁻¹) and are in agreement with data previously reported for this region (Broecker et al., 19 1976; Charette et al., 2015). At intermediate depths, Ba concentrations range from 40 to 50 nmol L⁻¹ in 20 the West European Basin (stations 1 and 21) and ²²⁶Ra activities range from 9.5 to 10.9 dpm 100 L⁻¹, 21 also in good agreement with literature data (Charette et al., 2015; Schmidt and Reyss, 1996). Low ²²⁶Ra 22 and Ba are found in upper waters of the West European Basin and the Iceland Basin 23 (8.1-8.9 dpm 100 L⁻¹ and 35-43 nmol L⁻¹, respectively). Intermediate ²²⁶Ra activities and Ba 24 concentrations (9 dpm 100 L⁻¹ and 42 nmol L⁻¹, respectively) are observed in bottom waters in Irminger 25 and Labrador Seas. Between the Reykjanes Ridge and Newfoundland, ²²⁶Ra activities range between 7-26 10 dpm 100 L⁻¹ in surface and intermediate waters. Similar to ²²⁶Ra, Ba concentrations are relatively 27 low in this area, ranging from 39-50 nmol L⁻¹. The distributions in ²²⁶Ra and Ba are to a first order 28 29 explained by the different water masses present in the region, as discussed below.

1 4. Discussion

2 4.1. ²²⁶Ra-Ba and ²²⁶Ra-Ba-Si relationships

A linear correlation between ²²⁶Ra and Ba is observed for all data collected along the GA01 section 3 (Fig.5). The slope of the ²²⁶Ra-Ba linear regression obtained by this study in the North Atlantic is 4 2.5 ± 0.1 (2SD) dpm μ mol⁻¹ which agrees with the slope of the ²²⁶Ra-Ba linear regression of 5 2.3 dpm µmol⁻¹ reported during the GEOSECS program for all the oceanic basins (Chan et al., 1976). 6 The intercept on the horizontal Ba axis is 4.4 nmol L^{-1} for the GA01 section, which is in agreement with 7 GEOSECS data (Chan et al., 1976; Li et al., 1973). This positive intercept may be the result of a larger 8 riverine Ba input relative to ²²⁶Ra (Ku and Luo, 1994). The slope of the ²²⁶Ra-Ba linear regression 9 reported from the GEOSECS program is similar from one oceanic basin to the other. The ²²⁶Ra/Ba ratio 10 (slightly different from the slope) is also fairly constant throughout the global ocean 11 $(2.2 \pm 0.2 \text{ dpm }\mu\text{mol}^{-1}; \text{ Östlund et al., 1987})$. This pattern indicates that ²²⁶Ra and Ba may behave 12 similarly in the ocean. Since ²²⁶Ra and Ba are incorporated in settling particles such as calcareous and 13 siliceous shells or barite (BaSO₄) by substitution or adsorption mechanisms (Bishop, 1988; Dehairs et 14 al., 1980; Lea and Boyle, 1989, 1990) and are then released at depth following the dissolution of these 15 particles, the constant ²²⁶Ra/Ba ratio suggests that fractionation between ²²⁶Ra and Ba during these 16 processes is small. 17

Investigations conducted during the GEOSECS program further concluded that ²²⁶Ra and Ba were 18 tightly correlated to orthosilicic acid (Si(OH)₄) (Chan et al., 1976; Chung, 1980; Ku et al., 1970; Ku and 19 Lin, 1976) despite the fact that ²²⁶Ra, Ba, and Si(OH)₄ exhibit different chemical behavior in the water 20 21 column and different source terms. A Ra-Ba-Si relationship is also observed along the GA01 section (Fig.5). Si(OH)₄ concentrations generally increase with increasing depth, with a steeper gradient in the 22 West European Basin (Introduction Paper, 2017; This issue), as it was also the case for ²²⁶Ra and Ba 23 (Fig.S2). The link between ²²⁶Ra, Ba and Si has been shown to reflect parallel dissolved-particulate 24 25 interactions between barite and biogenic silica (Bishop, 1988; Chung, 1980; Jacquet et al., 2005, 2007; Jeandel et al., 1996); the main carrier of ²²⁶Ra in the ocean, however, remains an open question. The 26 oceanic Ba-Si and ²²⁶Ra-Si relationships may thus be the result of the interaction between ocean 27 biogeochemistry and the water mass transport. 28

In contrast to the ²²⁶Ra-Ba relationship, the slope of the ²²⁶Ra-Si(OH)₄ relationship observed during GEOSECS exhibited significant spatial variability from one oceanic basin to the other (Li et al., 1973). First, ²²⁶Ra and Si are not chemical analogues, as it is the case for ²²⁶Ra and Ba. Second, the variability observed in the ²²⁶Ra-Si(OH)₄ relationship may be related to the large variability of the Si(OH)₄ distribution which is mostly governed by the preformed nutrient concentrations of waters feeding into

the main thermocline from surface waters of the Southern Ocean (Sarmiento et al., 2007). In the case of 1 GA01, the ²²⁶Ra-Si(OH)₄ linear regression slope is 2.4 ± 0.9 (2SD) 10^3 dpm mol⁻¹. As a comparison, the 2 ²²⁶Ra-Si(OH)₄ slope reported for the GEOTRACES-GA03 section conducted south of the GA01 section 3 in the Atlantic Ocean was 2.1 10³ dpm mol⁻¹ (Charette et al., 2015). The slope of the ²²⁶Ra-Si(OH)₄ 4 linear regression is 34.3 10³ dpm mol⁻¹ in the Pacific Ocean and 14.5 10³ dpm mol⁻¹ in the Antarctic 5 Ocean. The ²²⁶Ra-Si(OH)₄ relationship has an intercept with the vertical axis of 7.3 ± 0.1 dpm 100 L⁻¹. 6 which represents the residual ²²⁶Ra resulting from the total usage of Si in surface waters (Ku et al., 7 1970). According to (Shannon and Cherry, 1971), the removal of ²²⁶Ra in the upper waters is limited by 8 Si. For both the ²²⁶Ra-Ba and ²²⁶Ra-Si(OH)₄ relationships, several values are clearly outside of the linear 9 regression trend (Fig.5), a pattern that indicates deviation from the relationship usually observed 10 between ²²⁶Ra and Ba (or Si(OH)₄). Such deviations may result from non-conservative processes. 11

12 4.2. ²²⁶Ra and Ba distributions and their relationship with hydrography

A striking feature of the GA01 section is that the ²²⁶Ra activities and Ba concentrations are 13 particularly high in the West European Basin below 2000 m (Fig.4), in the NEADWI. This pattern can 14 also be observed in the GA03 section conducted south of the GA01 section (Charette et al., 2015), the 15 16 two sections being separated by only ca. 500 km in that basin. The NEADWI is mainly formed from waters with a southern origin (Read, 2000). South of the Antarctic Convergence, the surface waters 17 contain high ²²⁶Ra activities from the upwelling of deep waters enriched in ²²⁶Ra associated with the 18 circumpolar current (Ku and Lin, 1976). The convection of these surface waters leads to the formation 19 of the ²²⁶Ra-rich AABW that circulates northward into the Atlantic Ocean. However, the high ²²⁶Ra 20 activities of the NEADWI cannot be solely explained by the high ²²⁶Ra activities of these waters of 21 southern origin. This will be discussed in section 4.3.1. 22

In contrast, the lowest ²²⁶Ra activities and Ba concentrations reported on the GA01 section are associated with the Central Waters (upper waters of the West European Basin; Fig.4). Central Waters are derived from the NAC and mix with the SAIW and the SPMW. Along their path, Central Waters remain in the upper water column, and therefore are not affected by the deep sedimentary source of ²²⁶Ra. West of the Iceland Basin between 200 and 400 m (stations 32 and 38), an increase in the ²²⁶Ra activities and Ba concentrations is associated with the IcSPMW.

A slight increase in ²²⁶Ra is observed between 1000-1600 m in the West European Basin (Fig.4; Stations 1 and 13) where a salinity maximum is identified. This pattern is associated with the MW. This is corroborated by the slightly higher Ba concentrations and lower ²²⁶Ra/Ba ratios, as observed in the Western Mediterranean Sea (van Beek et al., 2009), these waters spreading westward into the Atlantic Ocean. At these stations, between 30 and 79 % of the water found at 1000-1600 m is of Mediterranean
 origin (MW), according to the OMP analysis (Garcia Ibanez et al., 2017; this issue).

Relatively uniform and low ²²⁶Ra activities and Ba concentrations are found between the surface 3 and 2500 m in the Labrador Sea, up to 2000 m in the Iceland Basin and deeper in the Irminger Basin 4 5 (Fig.4). These distributions can be related to the LSW which is formed by winter convection in the Labrador Sea (Fröb et al., 2016; Pickart et al., 2003; Yashayaev and Loder, 2016). When formed, the 6 LSW transports the characteristics of surface waters to the deep ocean (i.e., low ²²⁶Ra activities and low 7 Ba concentrations). The LSW then spreads into the Irminger and the Iceland Basin while conserving its 8 low ²²⁶Ra and Ba signatures. Relatively low ²²⁶Ra activities and Ba concentrations are found at bottom 9 depths in the Irminger and Labrador Seas and may be associated with DSOW, which is also a recently 10 ventilated water mass (Lazier, 1973). 11

Finally, according to the OMP analysis, ISOW is present at several stations along the GA01 section (Garcia-Ibanez et al., this issue). First, on the eastern flank of the Reykjanes Ridge (station 32), 68 % of the water between 2700 and 3000 m is considered as ISOW. Then, in the Labrador Sea (stations 69 and 77), an average of 58 % of the water between 2100 and 3000 m is identified as ISOW. The slight increase in ²²⁶Ra activities and Ba concentrations observed at these locations may thus be related to the ISOW.

18 4.3. Conservative versus non-conservative behavior of ²²⁶Ra and Ba

19 The use of an Optimum Multiparameter (OMP) analysis allowed us to distinguish the relative importance of physical transport (i.e., mixing of water masses) from non-conservative processes on the 20 ²²⁶Ra, Ba and ²²⁶Ra/Ba ratios distributions in the North Atlantic (Fig.6). The comparison between the 21 vertical profiles of ²²⁶Ra and Ba determined along the GA01 section, and those derived from OMP 22 analysis (Fig.7) clearly indicates deviations from the conservative behavior and reflects either an input 23 of ²²⁶Ra or Ba (positive anomalies highlighted in red; same color code as in Fig.6) or a removal of ²²⁶Ra 24 25 or Ba (negative anomalies highlighted in blue; same color code as in Fig.6). This comparison reveals that for 58 % of the samples ²²⁶Ra can be considered as conservative (activities due to mixing and 26 transport) along the GA01 section (i.e., 58 % of the 226Ra anomalies are within the [-0.9 and 27 $0.9 \text{ dpm } 100 \text{ L}^{-1}$ interval), whereas for 65 % of samples Ba can be considered as conservative (i.e., 28 65 % of the Ba anomalies are within the [-4.4 and 4.4 nmol L^{-1}] interval). A major finding of this study 29 is that ²²⁶Ra and Ba are predominantly conservative at intermediate depths: mostly between 500 m to 30 2000 m, but slightly deeper in the center of deep basins such as at stations 21, 44 and 69 (Fig.6). These 31 locations correspond to the depths where the waters are far from the main sources and sinks of ²²⁶Ra 32 and Ba. The non-conservative ²²⁶Ra (42 % of the ²²⁶Ra) is mostly distributed close to the interfaces such 33

as surface/subsurface waters and bottom waters (both in the deep West European Basin and the
 Labrador Sea), near the main sources (seafloor or shallow sediments deposited onto the margins). The
 non-conservative Ba is mostly distributed in the upper 1500 m and in the deep West European Basin
 (Fig.6).

5 The ²²⁶Ra/Ba ratios are also reported for all samples in Fig.7. The mean ratio determined along the 6 GA01 section is identical to the ratio determined during the GEOSECS program ($2.2 \pm 0.2 \text{ dpm }\mu\text{mol}^{-1}$; 7 Östlund et al., 1987). 77 % of the ²²⁶Ra/Ba ratios determined along the GA01 section are within the 8 confidence interval [2.0-2.4 dpm μmol^{-1}], indicating little deviation from the mean ratio, a pattern that is 9 likely related to the similar chemical behavior between ²²⁶Ra and Ba.

10 4.3.1. ²²⁶Ra inputs and their relationship with Ba

Deep waters of the West European Basin display positive ²²⁶Ra and Ba anomalies (stations 1 to 11 26; Fig.7). The ²²⁶Ra anomalies are initiated at shallower depths (ca. 300-2000 m) than the Ba 12 anomalies (ca. 1000-2000m) (Fig.7). As mentioned above, the NEADWI-which is the main water 13 mass of the deep West European Basin-is mainly formed from waters with a southern origin (mainly 14 AABW) that are characterized by high ²²⁶Ra and Ba concentrations. However, these southern waters 15 experience a very specific history along their northward transport to the GA01 section that suggests that 16 the high ²²⁶Ra activities (and Ba) of the NEADWl cannot be solely explained by the high ²²⁶Ra 17 activities (and Ba) of these waters of southern origin. In order to explain the positive ²²⁶Ra and Ba 18 anomalies in the deep waters of the West European Basin, we thus need to investigate the fate of ²²⁶Ra 19 and Ba in the waters of southern origin that travel northward and reach section GA01. Figure 8 was 20 computed combining GEOSECS and TTO data (²²⁶Ra, Si(OH)₄, salinity and temperature) associated 21 with the AABW (Spencer, 1972) that travels northward between 60 °S and 40 °N in the West Atlantic 22 Basin. The same data (²²⁶Ra, Si(OH)₄, salinity and temperature) associated with the NEADWl in the 23 East Atlantic Basin and along GA01 are also reported. 24

Between 60 °S and the equator, the high ²²⁶Ra activities of the AABW are associated with 25 relatively low salinity, low temperature, and high Si(OH)₄ (Fig.8). Then, while crossing the Mid-26 Atlantic Ridge at the equator and at 11 °N, the AABW undergoes several important transformations: 27 ²²⁶Ra activities and Si(OH)₄ concentrations decrease while salinity and temperature tend to increase 28 (Fig.8). The ²²⁶Ra and Ba endmembers for the NEADWl were chosen at this specific location to 29 coincide both geographically and with the characteristics (salinity, temperature and Si(OH)₄) of the 30 31 NEADWl endmembers used for the OMP analysis (Fig.8; Fig.S3). This endmember has been chosen far from the GA01 section in the OMP analysis (García-Ibáñez et al., this issue), because between 11 °N 32 and the GA01 section (Fig.8), salinity, temperature, and Si(OH)₄ concentrations display relatively 33

constant trends indicating no major modifications. In contrast, the ²²⁶Ra activities display a significant 1 spatial variability north of 11 °N, and clearly increase towards the north (Fig.8). This ²²⁶Ra increase is 2 thus decoupled from salinity, temperature, and $Si(OH)_4$), and likely explains the positive anomalies 3 deduced from the OMP analysis in the deep West European Basin (Fig.7). The specific history of these 4 waters of southern origin (waters initially with a high ²²⁶Ra activity; decrease in the ²²⁶Ra activity at the 5 equator and at 11 °N; new increase of ²²⁶Ra activity north of 11 °N) suggest that the ²²⁶Ra anomalies 6 observed in the West European Basin are explained by inputs of ²²⁶Ra along the northward transport of 7 8 these waters.

The positive anomalies result from the input of ²²⁶Ra (and Ba) following either i) 9 dissolution/remineralization of settling particles that incorporated ²²⁶Ra and Ba in the upper water 10 11 column (this includes the dissolution of barite, since the waters of the Atlantic Ocean are undersaturated with respect to barite; Monnin et al., 1999) and/or ii) diffusion of ²²⁶Ra and Ba from deep-sea sediments 12 (Cochran and Krishnaswami, 1980) (see 4.4). Of special note are stations in the West European Basin, 13 which are especially deep (down to 5500 m). Deep sediments generally display elevated ²³⁰Th activities 14 due to scavenging of ²³⁰Th from the entire water column (Bacon and Anderson, 1982; Nozaki, 1984). 15 The highest dissolved ²³⁰Th activities reported along the GA01 section were thus observed in the deep 16 17 waters of the West European Basin (Deng et al., 2017, this issue). Consequently, because ²²⁶Ra is produced by the decay of ²³⁰Th in the sediment, the ²²⁶Ra diffusive flux in this area is expected to be 18 19 especially high.

The input of ²²⁶Ra in the West European Margin is accompanied by a Ba input since i) positive Ba 20 anomalies are also observed in the deep waters and ii) the ²²⁶Ra/Ba ratios do not significantly deviate 21 from the mean GEOSECS ²²⁶Ra/Ba ratio (Fig.7a). One exception is found at station 21 in the West 22 European Basin, which displays high ²²⁶Ra/Ba at approximately 4000 m (up to 3.17 dpm µmol⁻¹). At 23 several stations (21, 26, 32, 38, 44, 60, 64 and 77), lower beam transmission values near the seafloor 24 25 indicate presence of suspended sediments likely associated with a nepheloid layer. Nepheloid layers are turbid layers formed episodically by strong and intense abyssal currents that are transported along 26 isopycnals away from the site of resuspension of bottom sediments (McCave, 1986). The presence of a 27 benthic nepheloid layer is also indicated by high particulate iron concentrations at these stations 28 (Gourain et al., 2017; this issue). Such process may thus contribute to release ²²⁶Ra (and potentially Ba) 29 to the deep water column, following desorption or dissolution of the particles. Similar ²²⁶Ra maxima 30 have been observed in the deep waters of the Northeast Pacific suggesting that the ²²⁶Ra flux is not 31 32 uniform over the ocean bottom even on a regional scale (Chung, 1976). Suspended particle dissolution may also play a role here, notably for Ba. This will be discussed in more detailed in section 4.4. 33

Positive ²²⁶Ra anomalies are also found in deep waters at several other stations located in 1 relatively deep basins (> 1200 m) along the GA01 section (e.g. stations 32, 38, 44, 60, 64, 69 and 77). 2 Most of these anomalies are associated with ²²⁶Ra/Ba ratios higher than 2.4 dpm µmol⁻¹. The ²²⁶Ra 3 positive anomalies observed at the stations mentioned above are thus best explained by the diffusion of 4 ²²⁶Ra from the sediment. However, these latter stations do not exhibit a positive Ba anomaly and Ba 5 tends to be conservative. Consequently, the ²²⁶Ra/Ba ratios in the deep waters of these stations are often 6 significantly higher than the mean GEOSECS value (stations 21, 32, 38, 60, 64; Fig.7). This pattern is 7 8 different to that observed in the West European Basin, a discrepancy that may be explained by the different sediment composition in the two regions, by the different residence time of deep waters in 9 10 contact with deep-sea sediments (Chung, 1976) and/or different role played by suspended particles 11 dissolution.

A strong ²²⁶Ra positive anomaly is observed in the deepest sample collected at station 38 above 12 the Reykjanes Ridge. It cannot be completely excluded that this positive anomaly is attributed to 13 hydrothermal vent since hydrothermal systems are known in the area (Fig.1). High particulate iron and 14 15 aluminum concentrations were also observed at these stations (Gourain et al., 2017; Menzel et al., 2017 : this issue). Enrichment in ²²⁶Ra have indeed been observed in hydrothermal systems plume at 16 mid-ocean Ridges (Kadko, 1996; Kadko and Moore, 1988; Kipp et al., 2017; Rudnicki and Elderfield, 17 1992). Moreover, the ²²⁶Ra enrichments are accompanied by slight Ba enrichments, which may support 18 19 the hydrothermal origin hypothesis, since hydrothermal venting at mid-ocean Ridge constitutes the second major external source of Ba to the ocean (Edmond et al., 1979). 20

Finally, high ²²⁶Ra/Ba ratios variations are also observed in shallow coastal waters (Fig.7c). At 21 stations 53 and 61, high ²²⁶Ra/Ba ratios are found close to the bottom, in agreement with the input of 22 ²²⁶Ra from the sediment, whereas low ²²⁶Ra/Ba ratios are found in subsurface at stations 57, 61 and 78, 23 in association with low salinities (Fig.S2). The low ²²⁶Ra/Ba ratios are thus explained by the input of 24 meteoritic waters in coastal areas, since such waters are known to be the predominant source of Ba to 25 the ocean (Martin and Meybeck, 1979; Wolgemuth and Broecker, 1970). At these shallow stations, the 26 different source terms between ²²⁶Ra and Ba therefore leads to important variations of the ²²⁶Ra/Ba 27 ratios (Fig.7c.; stations 53, 57, 61 and 78). These observations clearly indicate that ²²⁶Ra may 28 sometimes be decoupled from Ba. 29

30 4.3.2. ²²⁶Ra removal and its relationship with Ba

Relatively few ²²⁶Ra negative anomalies were found along the GA01 section. At the deep openocean stations, the negative anomalies are mostly observed in the upper 1000 m (Fig.7; stations 13, 21, 32, 32, 38, 44 and 77), but can also be found as deep as 2000 m (i.e., stations 64 and 69). In most cases, the negative ²²⁶Ra anomalies are associated with significant negative Ba anomalies (stations 13,21, 26, 38, 44, 64 and 69). Such features are likely explained by biological mediated processes including incorporation of ²²⁶Ra and Ba in planktonic as calcareous and siliceous shells (Bishop, 1988), in acantharian skeletons made of celestite (SrSO₄) or in barite (BaSO₄) crystals (van Beek et al., 2007; Chow and Goldberg, 1960; Shannon and Cherry, 1971; Szabo, 1967; Wolgemuth and Broecker, 1970).

Particularly low dissolved ²²⁶Ra/Ba ratios (<2 dpm µmol⁻¹) are found in the upper 50 m at 6 stations 21, 32, 64, 69 and 77, a pattern that was also observed in the upper 150 m of the Sargasso Sea, 7 where van Beek et al., (2007) reported similar low dissolved ²²⁶Ra/Ba ratios that were accompanied by 8 high ²²⁶Ra/Ba ratios in suspended particles. This latter pattern was attributed to the incorporation of 9 ²²⁶Ra and Ba in acantharian skeletons. The low dissolved ²²⁶Ra/Ba ratios (e.g. 1.7 dpm µmol⁻¹, 10 11 station 77) observed in the upper 200 m along the GA01 section may thus be attributed to acantharians, which have skeletons that incorporate ²²⁶Ra preferentially to Ba (van Beek et al., 2007, 2009; Bernstein 12 et al., 1998). The presence of Acantharians was not studied during GEOVIDE. However, previous 13 14 studies reported presence of acantharians in this area, like for example in the Iceland Basin and in the 15 East Greenland Sea (Antia et al., 1993; Barnard et al., 2004; Martin et al., 2010).

16 Several phytoplankton blooms were observed along the GA01 section, as indicated by the 17 chlorophyll a concentrations (Chl-a) (Fig.S4). The largest bloom was observed in the Labrador Sea in May 2014. Diatoms were the dominant species in the Irminger and Labrador Seas and on the Greenland 18 19 and Newfoundland margins during GA01 (up to 55 % of the total Chl-a concentration; Tonnard et al., in prep). In the West European Basin, Chl-a was lower in May and June 2014 and coccolithophorids were 20 the dominant species in that area (Tonnard et al., in prep). In these two regions, diatom frustules and 21 coccolithophorids may thus contribute to the removal of ²²⁶Ra and Ba (Bishop, 1988; Dehairs et al., 22 1980), from the water column in these areas that were characterized by noticeable negative anomalies. 23

24 Additionally, because the Labrador Sea was sampled in June, during the decline of the bloom, 25 barite that is presumably formed following the decay of settling organic matter may also contribute to the removal of Ba (and ²²⁶Ra). High particulate excess Ba (Ba_{xs}) concentrations were indeed observed at 26 stations displaying significant Ba negative anomalies: Baxs concentrations reached a maximum at 400 m 27 at station 13, 21 and 26) and between 400 and 800 m near Greenland, at stations 44, 64 and 69 28 (Lemaitre et al., 2017, Ba paper, this issue). These Baxs profiles can be related to the relatively high 29 30 particulate organic carbon (POC) export flux determined at these stations (e.g. at station 69, Lemaitre et al., 2017, Export paper; this issue). This POC flux would promote barite formation in subsurface 31 32 (Dehairs et al., 1980; Legeleux and Reyss, 1996) but also deeper in the water column (van Beek et al., 2007). Similarly, Jullion et al., (2017)-by using a parametric OMP analyses as applied in the 33 34 Mediterranean Sea-also reported quantification of the non-conservative component of the Ba signal 1 and suggested that the Ba negative anomalies potentially reflected Ba subtraction during barite 2 formation occurring during POC remineralization. The winter deep convection in the Labrador Sea may 3 also potentially explain this relatively deep Ba anomalies by transporting negative Ba and ²²⁶Ra 4 anomalies waters toward the deep-sea (Jullion et al., 2017). With the exception of the acantharian 5 skeletons that may impact the dissolved ²²⁶Ra/Ba ratios in the upper 200 m, the removal of ²²⁶Ra and Ba 6 that takes place deeper in the water column or that involves other processes (e.g. barite precipitation) 7 does not seem to affect significantly the dissolved ²²⁶Ra/Ba ratios.

In the shallow coastal stations, lower ²²⁶Ra/Ba ratios are observed (Fig.7). These low ratios may 8 also result from the removal of ²²⁶Ra and Ba by planktonic shells and/or barite or scavenging by 9 10 lithogenic particles. However, because these stations are coastal stations, various processes are at play in a relatively shallow water column (i.e. diffusion of ²²⁶Ra from the sediments; input of Ba from 11 meteoritic water; removal of Ba and ²²⁶Ra by shells and barite) thus complicating the interpretation of 12 the vertical profiles. We note that the low ²²⁶Ra/Ba ratios observed in surface of shallow stations near 13 the coast of Greenland (stations 57 and 61) and Newfoundland (station 78) are associated with low 14 salinities (Fig.7c). This decoupling between ²²⁶Ra and Ba may be explained by input of freshwater into 15 16 the coastal zone.

Finally, at several stations, a decrease in the ²²⁶Ra activities is observed near the seafloor 17 (stations 13, 21, 44, 60, 64 and 77; Fig.7). Similar decreasing trends near the seafloor have been 18 reported in the Southwest Atlantic and North Pacific for ²³⁰Th (Deng et al., 2014; Okubo et al., 2012), a 19 reactive element that strongly adsorbs onto suspended particles. This trend for ²³⁰Th was explained by 20 nuclide scavenging at the seafloor (Deng et al., 2014; Okubo et al., 2012). Radium-226-although it is 21 much less particle-reactive than ²³⁰Th—and Ba may also be scavenged by resuspended particles near 22 the seafloor and may adsorb onto the surfaces of Mn oxides (Moore and Reid, 1973). High particulate 23 trace element concentrations were also observed at stations 26, 38, and 69 and may be related to 24 nepheloid layers that impact the deep water column up to 200-300 m above the seafloor (Gourain et al., 25 2017; Menzel et al., 2017; this issue). 26

27 4.4. Estimation of ²²⁶Ra and Ba input fluxes into the West European Basin

A strong ²²⁶Ra positive anomaly is observed in the NEADWl between stations 1 and 21 and below 3500 m. On average, it is 3.3 dpm 100 L⁻¹ over this vertical section. This anomaly reflects a concentration difference between the ²²⁶Ra measured along GA01 and the ²²⁶Ra due to water mass mixing. This concentration difference is associated to the northward transport of the NEADWl, estimated to be 0.9 ± 0.3 Sv (10⁶ m³ s⁻¹) at 45 °N (GA01 section) (Daniault et al., 2016; McCartney, 1 1992). Therefore, the positive concentration anomaly can be converted into a 226 Ra flux that has to be 2 added to this water mass, $F_{Input-Ra}$, calculated as follows:

3

 $F_{Input-Ra} = A \times T_{NEADWl} (1)$

4 where *A* is the mean positive anomaly of 226 Ra (in dpm m⁻³) determined by the OMP analysis; 5 T_{NEADW1} is the transport associated with the NEADW1 (in m³ s⁻¹).

6 This ²²⁶Ra input may be associated with a sedimentary source and/or may result from the 7 dissolution of suspended particles. In a first place, we will study the hypothesis of the sedimentary 8 source; the suspended particle source will be discussed in a second place.

9 The NEADWl at 45 °N is made of up to 92 % of the 11 °N NEADWl endmember. Therefore, the 10 sedimentary input along the northward transport of the NEADWl is calculated across a sediment area 11 between 11 °N and 45 °N (Fig.S3). We consider the distance of 4209 km between 11 °N and the GA01 12 section and the distance of 1475 km between stations 1 and 21. This leads to a horizontal area of 6.21 13 10⁶ km² (assuming a constant bathymetry), across which the sedimentary input is assumed to take place. 14 The ²²⁶Ra flux diffusing out of bottom sediments, *F_{Sed-Ra}* (in dpm cm⁻² y⁻¹) can be calculated using 15 Eq. (2), assuming that the anomaly is entirely explained by the sediment source:

$$F_{Sed-Ra} = \frac{F_{Input-Ra}}{S}$$
(2)

16 where S is the surface area described above (in cm²) and $F_{Input-Ra}$ is 1.67 10⁸ dpm s⁻¹.

17

The calculated F_{Sed-Ra} is $14.8 \pm 3.1 \ 10^{-3} \text{ dpm cm}^{-2} \text{ y}^{-1}$, which is within the range of fluxes 18 reported in the literature. For example, Cochran (1980) reported F_{Sed-Ra} in the range of 19 $1.5 \ 10^{-3} \text{ dpm cm}^{-2} \text{ y}^{-1}$ for the Atlantic Ocean to 2.1 $10^{-1} \text{ dpm cm}^{-2} \text{ y}^{-1}$ in the Pacific Ocean (Fig.8). Li et 20 al. (1973) estimated ²²⁶Ra fluxes diffusing out of the sediment in the Southern Ocean and on the 21 Antarctic shelf of 6.2 10^{-4} dpm cm⁻² y⁻¹ and 1.6 10^{-3} dpm cm⁻² y⁻¹, respectively. The F_{Sed-Ra} calculated here 22 is thus slightly higher than the ²²⁶Ra sedimentary fluxes reported in the Atlantic Ocean by Cochran 23 (1980). Note, however, that the ²²⁶Ra fluxes released from the sediments vary locally as a function of 24 the sedimentary ²³⁰Th activity, bioturbation rates, but also as a function of the sediment type and 25 composition (Chung, 1976; Cochran, 1980). The ²²⁶Ra fluxes reported in the Atlantic Ocean by Cochran 26 (1980), which are the lowest of all basins, are only available for calcareous ooze type sediment 27 (Cochran, 1980). The NEADWI may cross different types of sediments along its northward path in the 28 Atlantic Ocean. This includes calcareous oozes, fine-grained calcareous sediments and clay (Dutkiewicz 29 et al., 2015). In particular, ²²⁶Ra diffusion is expected to be higher in these two latter sediment types 30 31 (Cochran, 1980).

As for Ba is concerned, the mean positive anomalies deduced from the OMP analysis is 7.0 nmol L⁻¹ leading to a $F_{Input-Ba}$ of 69.5 mol s⁻¹. In the same way as ²²⁶Ra, a Ba sedimentary flux F_{Sed-Ba} 1 of 3.19 ± 1.4 nmol cm⁻² y⁻¹ would be required to explain the Ba anomalies observed in the West 2 European Basin. This flux is on the low end of the Ba sedimentary fluxes reported in different ocean 3 basins, which range from 1.0 to 30 nmol cm² y⁻¹ (Chan et al., 1977; Falkner et al., 1993; McManus et 4 al., 1999; Paytan and Kastner, 1996).

Alternatively, the dissolution of settling particles could also contribute to the ²²⁶Ra and Ba 5 anomalies observed in the deep waters of the West European Basin. Assuming steady state, we may 6 undertake a mass balance calculation for particulate ²²⁶Ra and Ba in the same box as described above 7 (i.e., box defined between 11 °N and the GA01 section, between stations 1 and 21 -1475 km-and 8 between 3500 m depth and the seafloor; Fig.S5). Particles enter the box from above as settling particles, 9 10 but also horizontally, carried within the water masses at 11 °N that travel northward. Particles leave the box by different processes (accumulation in the sediment or northward transport by the water masses) or 11 alternatively may dissolve while settling in the box. In the absence of precise information about the 12 particulate ²²⁶Ra and Ba fluxes entering and exiting the box horizontally (i.e. the particulate ²²⁶Ra and 13 Ba concentrations at 11 °N and at the GA01 section), we assume that they are of equal importance and 14 therefore that they cancel each other in the mass balance calculation. 15

16 The vertical particulate flux entering the box, from above, can be calculated as follows:

$$F_{Part-x} = Cp_{3500} \times Vs \times S (3)$$

17

18 where x is either ²²⁶Ra or Ba, *Cp3500* is either the particulate ²²⁶Ra activities or the particulate Ba 19 concentrations at 3500 m, *Vs* is the settling speed for suspended particles and *S* is the horizontal surface 20 area described above (6.21 10^6 km²).

We use the value of 0.007 dpm 100 L^{-1} for the mean ²²⁶Ra particulate activity at 3500 m, a value 21 that was reported for the Atlantic Ocean, Sargasso Sea (van Beek et al., 2007) and the value of 22 0.087 nmol L⁻¹ for the mean Ba particulate concentration at 3500 m, a value that was determined along 23 24 the GA01 section (Lemaitre et al., this issue). We use the settling speeds (Vs) reported for suspended particles in the literature and that typically range from 100 to 1000 m y⁻¹ (Bacon and Anderson, 1982; 25 Krishnaswami et al., 1976; Roy-Barman et al., 2002). The F_{Part} thus obtained range from 1.4 10⁶ to 26 13.8 10^6 dpm s⁻¹ for ²²⁶Ra, while the F_{Part} range from 1.7 and 17.2 mol s⁻¹ for Ba. Of this total F_{Part} , some 27 fraction may dissolve while settling, while the remainder will accumulate in the sediment. This 28 dissolution flux is named F_{dissolution-x}, where x is either ²²⁶Ra or Ba. We use the sediment Ba 29 accumulation rates reported by Gingele and Dahmke (1994) in the Atlantic Ocean to calculate the 30 particulate Ba flux that exits the box ($F_{Accumulation-Ba}$; 2.0 to 13.4 mol s⁻¹); hence, by difference the 31 $F_{dissolution-Ba}$ is 0-15.2 mol s⁻¹ (Fig.10). This value is of the same order of magnitude of the $F_{Input-Ba}$ 32

needed to explain the Ba anomalies (6.28 mol s^{-1}). Therefore, in the case of Ba, the dissolution of 1 settling particles may entirely explain the OMPA-derived anomalies. The sediment ²²⁶Ra accumulation 2 rates can be calculated from the Ba accumulation rates estimated above, using the ²²⁶Ra/Ba ratio 3 determined in sinking particles collected in the Sargasso Sea near the seafloor (i.e., 1.5 dpm µmol⁻¹; van 4 Beek et al., 2007). The sediment ²²⁶Ra accumulation flux thus calculated, F_{Accumulation-Ra}, is 2.9 10⁶-5 19.6 10⁶ dpm s⁻¹, leading to F_{dissolution-Ra} of 0-10.9 10⁶ dpm s⁻¹ (Fig.10). Therefore, F_{dissolution-Ra} cannot 6 account for more than 37 % of the required 226 Ra flux ($F_{Input-Ra}$). This implies that even if the settling 7 speed is high (1000 m y^{-1} ; high turnover of the particles), the particle dissolution cannot account for the 8 entire $F_{Input-Ra}$. The remaining (minimum of 63 %) therefore has to be sustained by ²²⁶Ra diffusion from 9 10 the sediments.

11 While the above calculations have to be taken with caution given the numerous assumptions in the mass balance model, overall they suggest that the ²²⁶Ra positive anomalies observed in the West 12 European Basin may be explained entirely by ²²⁶Ra that diffuses out of the sediments. However, it 13 cannot be excluded that the dissolution of settling particles also contributes to the ²²⁶Ra enrichment. In 14 contrast, the Ba positive anomalies may be explained either by the diffusion of Ba from sediment or by 15 the dissolution of settling particles or by a combination of these two processes. These conclusions are in 16 line with the current knowledge about ²²⁶Ra and Ba sources in the deep ocean (Broecker et al., 1970; 17 Chan et al., 1976, 1977; Ku et al., 1980). 18

19 5. Conclusion

We investigated the distribution of dissolved ²²⁶Ra activities and Ba concentrations in the North 20 Atlantic Ocean along the GA01 section. To a first order, the ²²⁶Ra and Ba patterns appear to be 21 correlated to the water masses (e.g. high ²²⁶Ra and Ba in NEADWl in the West European Basin; low 22 ²²⁶Ra and Ba in Central Waters; slight increase of ²²⁶Ra in the MW). Using a mixing model, we show 23 that the measured ²²⁶Ra and Ba concentrations can be explained by conservative mixing for 58 % and 24 65 % of the samples respectively, notably at intermediate depth (mostly between 1000 m and 2000 m) 25 and slightly deeper in the middle of deep basins, away from the ocean interfaces. These locations 26 correspond to the depths where the waters are away from the main sources of ²²⁶Ra and Ba. ²²⁶Ra and 27 Ba can thus be considered as conservative tracers of water mass transport in the ocean interior at the 28 29 space scales considered here, namely, of the order of few thousand km.

Our study also highlighted several regions where significant input or loss of ²²⁶Ra and Ba takes place. In the West European Basin, the deep waters (NEADWI) accumulate both ²²⁶Ra and Ba while flowing from 11 °N to the GA01 section. Mass balance calculations suggest that these enrichments are predominantly explained by sediment diffusion for ²²⁶Ra, with a possible contribution from suspended

1 particle dissolution, while both the sediment and suspended particle dissolution could significantly 2 contribute to the Ba enrichments. This pattern contrasts with that observed in the deep waters collected elsewhere along the section that do not display Ba enrichments associated to the ²²⁶Ra enrichments. 3 Bottom nepheloid layers may also contribute to the release of ²²⁶Ra (and Ba) to the deep water column 4 at several stations. Interestingly, nepheloid layer processes seem to also act as potential removal of 5 ²²⁶Ra (and Ba) at several other stations. Significant input of Ba-likely associated with meteoritic 6 waters-is found in the upper water column near Greenland. Finally, ²²⁶Ra and Ba are removed from 7 the upper water column, likely primarily due to biological mediated processes such as incorporation of 8 ²²⁶Ra and Ba into barite (BaSO₄) that are presumably formed following the decay of settling organic 9 matter and/or adsorption onto diatom frustules, a mechanism that would explain the ²²⁶Ra-Ba 10 relationship reported here. Similarly, strong correlations were also found between Ba-Si and ²²⁶Ra-Si 11 although no obvious process links ²²⁶Ra or Ba with Si, except maybe for the adsorption of Ba and 12 (²²⁶Ra) onto diatom frustules. It cannot be excluded, however, that the observed Ba-Si and ²²⁶Ra-Si 13 relationships may result from the spatial coherence of different carriers overprinted by hydrodynamics. 14

Our study also provides evidence of significant decoupling between ²²⁶Ra and Ba. In the upper 200 m, the ²²⁶Ra/Ba ratios reach low values ($<2 \text{ dpm }\mu\text{mol}^{-1}$), a pattern that has been observed in other regions and was related to acantharian skeletons that incorporate ²²⁶Ra preferentially to Ba (van Beek et al., 2007; Bernstein et al., 1998). Finally, deviations from the mean GEOSECS ²²⁶Ra/Ba ratios were observed in the shallow coastal waters of Greenland and Newfoundland: the predominant input of Ba due to input of meteoritic waters leads to lower ²²⁶Ra/Ba ratios whereas near the seafloor, the input of sedimentary ²²⁶Ra leads to higher ²²⁶Ra/Ba ratios.

The absence of a stable isotope for radium led geochemists to consider Ba as a stable analog for 22 ²²⁶Ra because ²²⁶Ra and Ba display a similar chemical behavior, with the aim of using the ²²⁶Ra/Ba ratio 23 as a chronometer for the thermohaline circulation. This study confirms that ²²⁶Ra and Ba behave 24 similarly in the ocean interior away from external sources, both elements being predominantly 25 conservative in the studied area over distances of the order of a few thousands of km. However, this 26 study also highlights regions where ²²⁶Ra and Ba deviate from a conservative behavior, an important 27 consideration when considering the balance between the large-scale oceanic circulation and biological 28 activity over long time scales. Decoupling between ²²⁶Ra and Ba has been observed, in most cases at the 29 ocean boundaries as the result of dissolved ²²⁶Ra and Ba external sources. In addition, suspended 30 particle dissolution may differently impact the dissolved ²²⁶Ra and Ba content of intermediate and deep 31 waters (as shown for the NEADWI); such process would therefore potentially modify their ²²⁶Ra/Ba 32 ratios and would complicate the use of this ratio as a chronometer. Inclusion of the different sources and 33 34 sinks and particle/dissolved interactions in global ocean models should help to refine the use of the 1 ²²⁶Ra/Ba ratio as a clock to chronometer the thermohaline circulation, as was proposed several decades

- 2 ago during the GEOSECS program.
- 3

4 **Figure Caption**

Figure 1: Station locations of the GA01 section between Portugal and Newfoundland in the North Atlantic (black and blue dots). Stations investigated for ²²⁶Ra and Ba are marked as blue dots. The main currents and water masses in the North Atlantic are also represented. The major hydrothermal vents located near the GA01 section are indicated by black triangles. Stations investigated during the US-GEOTRACES-GA03 section, also conducted in the Atlantic Ocean, are reported on the lower panel (red dots).

Figure 2: Potential temperature-salinity diagram—including a zoom for bottom waters—of the water samples (colored dots) from the GA01 section. The properties of the source water types (based on García-Ibáñez et al., 2017; This issue) used in the Optimum Multiparameter (OMP) analysis are reported with white circles. Isopycnals are also plotted (potential density referenced to 0 dbar).

Figure 3: Distribution of salinity (CTD data) along the GA01 section. The different water masses are also reported, following García-Ibáñez et al. (2017; this issue). The station numbers are found on top of the figure. The sampling depths for ²²⁶Ra are shown for each vertical profile (black dots).

Figure 4: Distribution of (a) dissolved ²²⁶Ra activities (dpm 100 L⁻¹), (b) dissolved Ba concentrations (nmol L⁻¹) and ²²⁶Ra/Ba ratio (dpm μ mol⁻¹) along the GA01 section. Station numbers are found on top of the panels. The sampling depths are shown for each vertical profile (black dots).

Figure 5: Relationships between ²²⁶Ra and Ba (red dots) and between ²²⁶Ra and Si(OH)₄ (blue dots) along the GA01 section in the North Atlantic. The best linear fit for the two plots is also reported (R=0.93 for the two plots). The slopes of the relationships between ²²⁶Ra-Ba and between ²²⁶Ra-Si(OH)₄ are expressed in 10^{-2} dpm nmol⁻¹ and in 10^{-2} dpm µmol⁻¹, respectively.

Figure 6: Difference between the measured concentrations and those calculated by the OMP analysis, for ²²⁶Ra (a), Ba (b) and (c) ²²⁶Ra/Ba ratio along the GA01 section. Positive anomalies reflect recent tracer addition, while negative ones reflect recent tracer removal. Station numbers are found on top of the panels.

Figure 7: Vertical profiles of dissolved ²²⁶Ra activities and Ba concentrations determined along the GA01 section: (a) West European Basin; (b) Iceland Basin and Irminger Basin, (c) the Greenland and Newfoundland margins, and (d) Labrador Basin. As a comparison, the conservative ²²⁶Ra and Ba vertical profiles derived from the OMP analysis are also reported in solid grey lines. The discrepancy between the two vertical profiles indicates deviation from the conservative behavior and reflects either an input of ²²⁶Ra or Ba (positive anomalies highlighted in red; same color code as Fig.6) or a removal of ²²⁶Ra or Ba (negative anomalies highlighted in blue; same color code as Fig.6). The OMP analysis has not been solved for the shallow coastal stations 53 and 78. The ²²⁶Ra/Ba ratios are also reported, together with the mean GEOSECS ²²⁶Ra/Ba ratio (2.2 ± 0.2 dpm µmol⁻¹; black dashed line) together with its one standard deviation (grey shaded areas). Note that the scale may be different from one station to the other and the vertical axis was cut to 1000 m. The seafloor is represented by the bottom axis.

Figure 8: Geographical variation of 226 Ra activities (red dots), salinity (blue dots), temperature (yellow dots) and Si(OH)₄ concentrations (green dots) in AABW and the NEADWI between 60 °S and 45 °N (GA01 section) in the Atlantic Ocean based on data from the GEOSECS and TTO programs. The 226 Ra activities, salinity, temperature and Si(OH)₄ concentrations from GA01 are represented with open circles. The values used as endmembers for the OMP analysis are also identified by the black circles. The shaded area represents the region where transformation of the AABW into NEADWI takes place.

Figure 9: 226 Ra fluxes diffusing out of the sediment in relationship with bottom water 226 Ra activities determined in different oceanic basins (P=Pacific Ocean, A=Atlantic Ocean, I=Indian Ocean and AA=Southern Ocean) by Cochran (1980). The 226 Ra flux calculated in this study to explain the positive anomalies in the West European Basin is also reported (red dot).

Figure 10: ²²⁶Ra and Ba flux estimations: $F_{Part - x}$ is the particulate flux entering the box and $F_{Accumulation-x}$ is the sediment accumulation flux and $F_{Dissolution-x}$ is the flux of particle dissolution assuming that all the settling particles dissolve. x is either ²²⁶Ra or Ba. Both maximum and minimum values are shown for $F_{Part - x}$ and $F_{Accumulation-x}$. Max $F_{Dissolution-x}$ represents a maximum value since it is calculated by subtracting the minimum value of $F_{Accumulation-x}$ from the maximum value of $F_{Part - x}$. $F_{Dissolution-x}$ is thus comprised between zero (if all F_{part-x} accumulates in the sediment) and this latter value.

46 **Figure S1:** Comparison of the vertical profiles of dissolved ²²⁶Ra at stations 1 and 13 of the GA01 section (black and red dots, respectively) and station 1 of the GA03 section (U.S.-GEOTRACES; blue dots) off Portugal.

Figure S2: Vertical profiles of dissolved ²²⁶Ra activities and dissolved Ba concentrations with the conservative ²²⁶Ra and Ba vertical profiles derived from the OMP analysis, ²²⁶Ra/Ba ratios, Si(OH)₄ concentrations, salinity (black line) and potential temperature (red line) for (a) the Iberian margin and the West European Basin, (b) the Iceland Basin and the Irminger Sea, (c) the Greenland margin, and (d) the

- Labrador Sea and the Newfoundland margin. Note that the scale may be different from one station to the other and the vertical axis was cut to 1000 m. The bottom is represented by the bottom axis.
- Figure S3: Location of each endmember source water types (SWTs) used for the OMP analysis (black circles). The surface of the basin,
 S, used to calculate the fluxes is represented by the grey hatched area.
- 5

Figure S4: Satellite Chlorophyll-a concentrations (MODIS Aqua from http://oceancolor.gsfc.nasa.gov), in mg m⁻³ during the GA01 cruise
 in (a) May 2014 and (b) June 2014. The dashed line indicates the location of the GA01 section. Stations investigated in this work are
 indicated by dots. White dots indicate the stations investigated during the corresponding month.

Figure S5: Schematic box model used to calculate the input fluxes in the West European Basin: F_{Sed-x} is the flux diffusing out of bottom sediments, F_{Part-x} is the vertical flux of particles entering the box from above, $F_{Accumulation-x}$ is the flux of particles accumulating in the sediment and F_{H-In-x} and $F_{H-Out-x}$ represent horizontal fluxes of dissolved species or particles coming in and out of the box due to transport, respectively. *x* is either ²²⁶Ra or Ba.

- 13 Table S1: Characteristics and location of each endmember source water types (SWTs).
- **Table S2:** ²²⁶Ra activities, Ba concentrations, ²²⁶Ra/Ba ratios, potential temperature and salinity at the different stations of the GA01 section.
- 16

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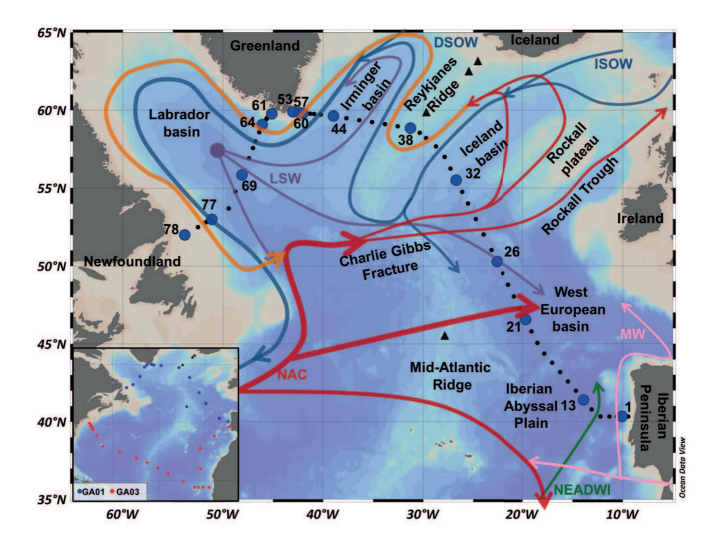
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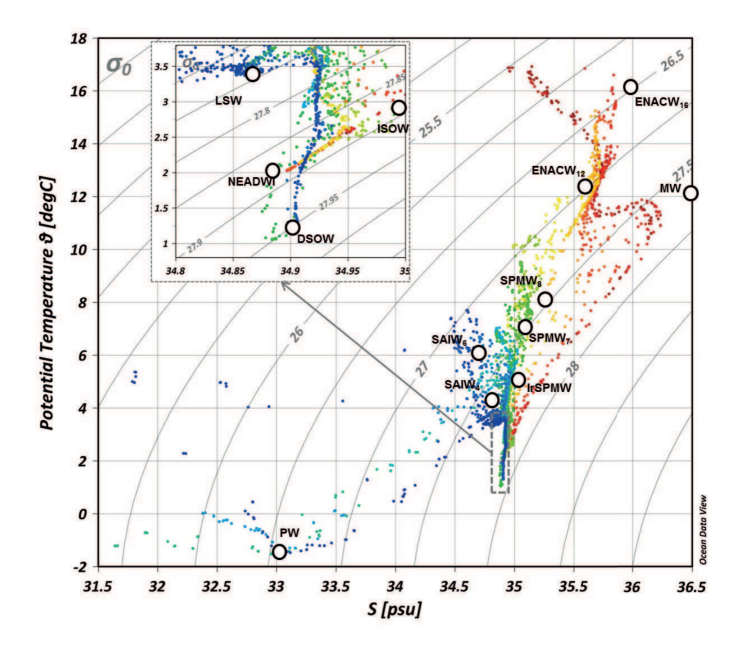
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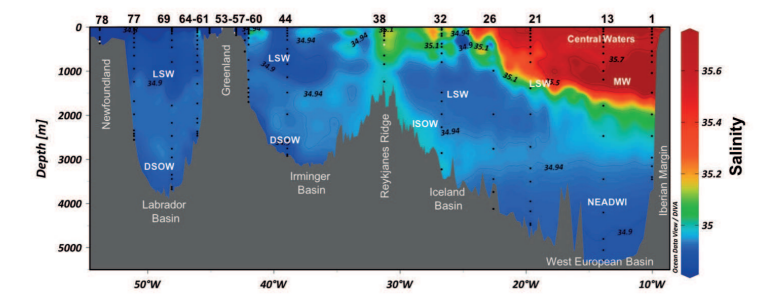
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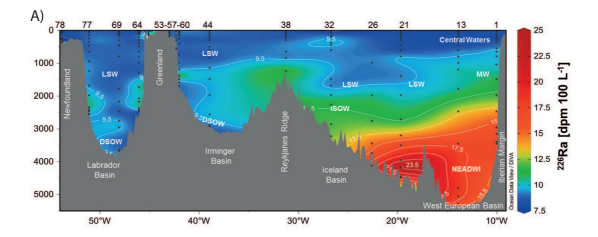
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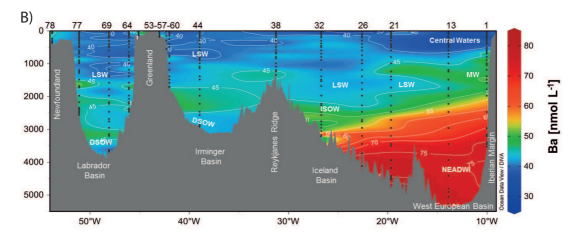
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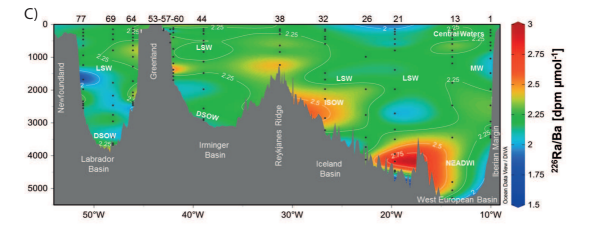


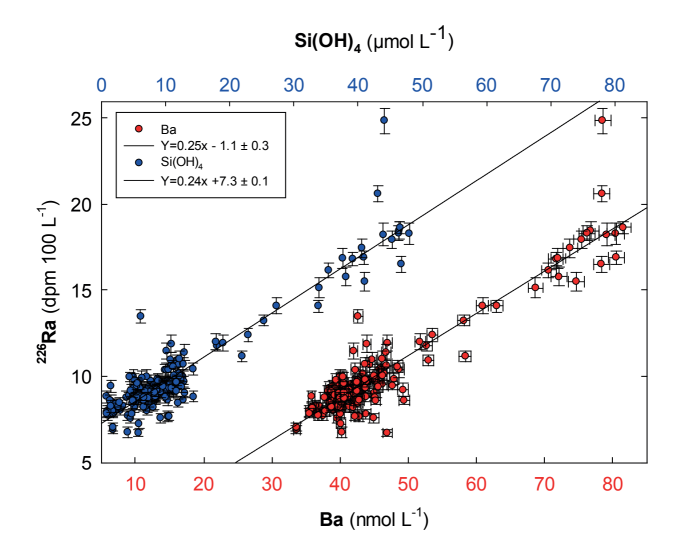


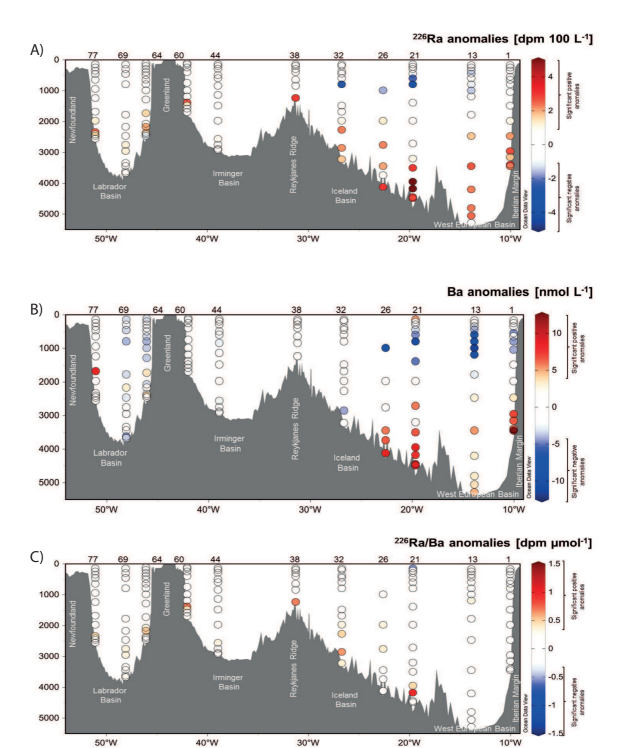




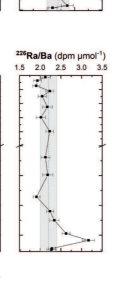








A) West European Basin



226Ra/Ba (dpm µmol⁻¹)

2.5

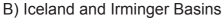
2.5

3.0

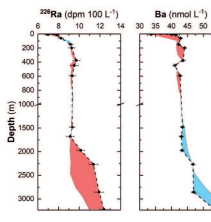
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2.0

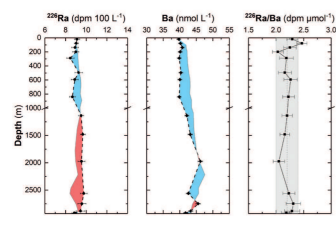
55 1.5

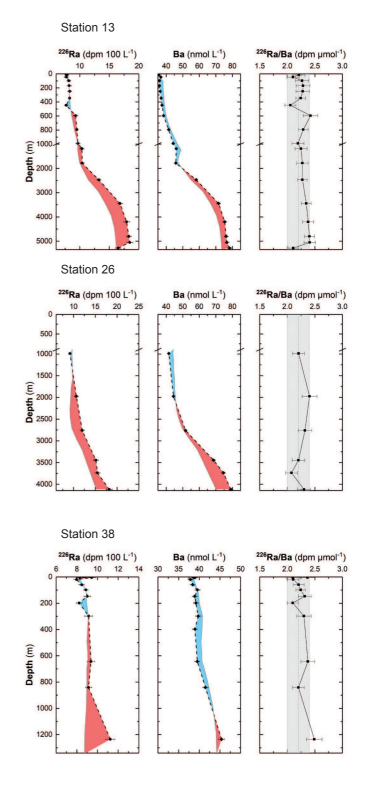


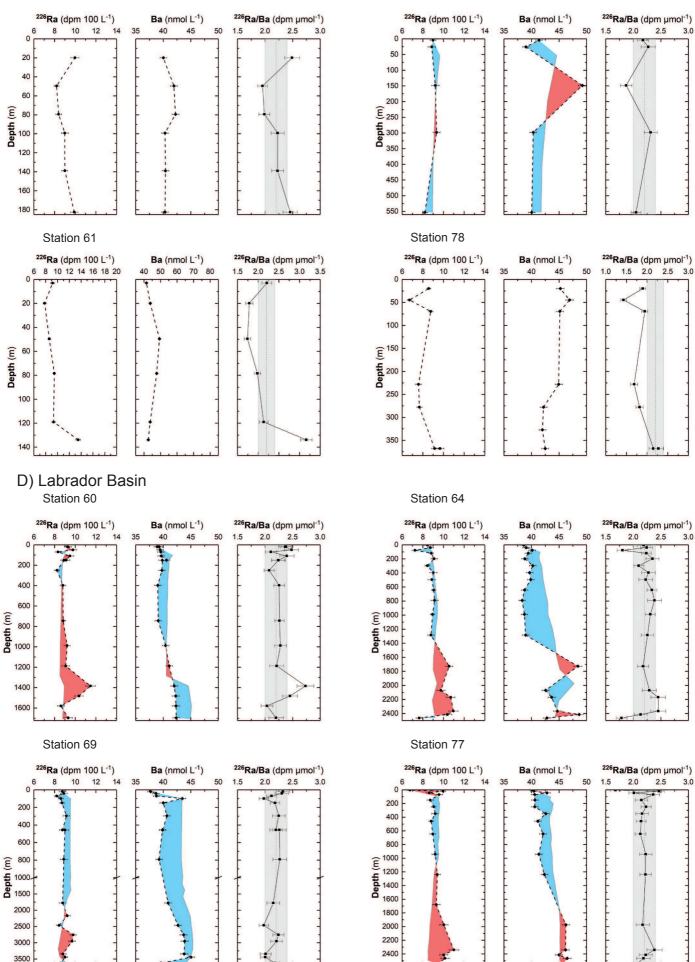
Station 32



Station 44







C) Greenland and Newfoundland Margins Station 53

Station 57

