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Upper ocean mixing controls the seasonality of planktonic foraminifer fluxes and associated strength of the carbonate pump in the oligotrophic North Atlantic

K. H. Salmon1, P. Anand1, P. F. Sexton1, and M. Conte2

1Environment, Earth and Ecosystems, The Open University, UK
2Bermuda Institute of Ocean Sciences, St George’s GE01, Bermuda

Correspondence to: K. H. Salmon (kate.salmon@open.ac.uk)

Received: 23 July 2014 – Published in Biogeosciences Discuss.: 12 August 2014
Revised: 5 December 2014 – Accepted: 11 December 2014 – Published: 13 January 2015

Abstract. Oligotrophic regions represent up to 75 % of Earth’s open-ocean environments. They are thus areas of major importance in understanding the plankton community dynamics and biogeochemical fluxes. Here we present fluxes of total planktonic foraminifera and 11 planktonic foraminifer species measured at the Oceanic Flux Program (OFP) time series site in the oligotrophic Sargasso Sea, subtropical western North Atlantic Ocean. Foraminifera flux was measured at 1500 m water depth, over two ∼2.5-year intervals: 1998–2000 and 2007–2010. We find that foraminifera flux was closely correlated with total mass flux, carbonate and organic carbon fluxes. We show that the planktonic foraminifera flux increases approximately 5-fold during the winter–spring, contributing up to ∼40 % of the total carbonate flux. This was primarily driven by increased fluxes of deeper-dwelling globorotaliid species, which contributed up to 90 % of the foraminiferal-derived carbonate during late winter–early spring. Interannual variability in total foraminifera flux, and in particular fluxes of the deep-dwelling species (Globorotalia truncatulinoides, Globorotalia hirsuta and Globorotalia inflata), was related to differences in seasonal mixed layer dynamics affecting the strength of the spring phytoplankton bloom and export flux, and by the passage of mesoscale eddies. As these heavily calcified, dense carbonate tests of deeper-dwelling species (3 times denser than surface dwellers) have greater sinking rates, this implies a high seasonality of the biological carbonate pump in oligotrophic oceanic regions. Our data suggest that climate cycles, such as the North Atlantic Oscillation, which modulates nutrient supply into the euphotic zone and the strength of the spring bloom, may also in turn modulate the production and flux of these heavily calcified deep-dwelling foraminifera by increasing their food supply, thereby intensifying the biological carbonate pump.

1 Introduction

Planktonic foraminifera (PF) comprise 23–56 % of the total open marine calcite flux and thus exert an important control on global carbon cycling (Schiebel, 2002). They are used extensively in palaeoceanographic and palaeoclimatic reconstructions via utilisation of their species abundance and assemblage composition (e.g. Lutz, 2011; Sexton and Norris, 2011), geochemical signatures (e.g. Zeebe et al., 2008), shell mass (e.g. Barker and Elderfield, 2002) and in evolutionary and biogeographic studies (e.g. Sexton and Norris, 2008). However, gaps remain in our understanding of the controls on their spatial and temporal distribution in the upper water column. Following the early 1980s when sea surface temperatures (SSTs) were thought to dominantly control PF distributions and abundance (CLIMAP project members, 1994), a number of other environmental parameters have also been shown to exert influence on the distribution and abundance of PF, such as salinity (Kuroyanagi and Kawahata, 2004), productivity, nutrient availability (Schiebel, 2002; Northcote and Neil, 2005; Žarić et al., 2005; Storz et al., 2009; Sexton and Norris, 2011) and water column stability (Hemleben et al., 1989; Lohmann and Schweitzer, 1990; King and Howard, 2003). It is thus imperative to better understand the
environmental factors controlling modern-day PF abundance in order to produce accurate interpretations of palaeorecords based on PF assemblages.

The response of PF flux and species composition to environmental and/or oceanographic factors have been studied using plankton tow materials which can give information about living populations’ species distribution and depth habitats within the upper ocean (Tolderlund and Be; 1971, Fairbanks et al., 1980; Schiebel, 2002). However, temporal resolution is often limited when using plankton tows. The continuous time series records provided by sediment traps allow a more complete understanding of the seasonal and interannual changes in PF flux and can aid in integrating living assemblages with the sedimentary record.

Earlier studies of planktonic foraminifer flux off Bermuda at the Seasonal Changes in Foraminifera Flux (SCIFF) site (Fig. 1) (Deuser et al., 1981; Hemleben et al., 1985; Deuser, 1987; Deuser and Ross, 1989) were based on a bi-monthly sampling interval and provide a general description of foraminifera species, composition and seasonality. These studies found that PF > 125 µm comprise on average 22 % of the total calcium carbonate flux in the Sargasso Sea (Deuser and Ross, 1989), although this average underestimates the importance of the PF flux contribution during different seasons. Here we utilise a higher resolution bi-weekly sediment trap time series from the Oceanic Flux Program (OFP), ideal for studying the detailed response of PF species flux to physical oceanographic changes because PF species lifespan is approximately 2–3 weeks (Spero, 1998; Erez et al., 1991). These samples also benefit from the availability of upper ocean hydrographic and biogeochemical data collected at the nearby Bermuda Atlantic Time Series (BATS) site, as well as remote sensing data, which allows us to evaluate the environmental factors that control the total foraminifer flux as well as the response of individual species flux. Furthermore, we assess the relative contribution of PF flux to regional carbonate export and explore the implications of our findings for carbonate cycling in the oligotrophic North Atlantic.

2 Oceanographic setting

The Sargasso Sea is located within the North Atlantic gyre, which is characterised by high temperatures and salinities, and weak, variable surface currents (Lomas et al., 2013, and references therein). The OFP and BATS sites are situated in a transition region between the northern eutrophic waters and the relatively oligotrophic subtropical convergence zone in the south (Steinberg et al., 2001, and references therein). Subtropical Mode Water (STMW) forms on the fringes, north of the gyre, owing to convective deep winter mixing and entrainment of nutrients and is characterized by temperatures of 17.8–18.4 °C and salinities of ~36.5 ± 0.05 % (Bates et al., 2002), typically occurring between ~250 and 400 m water depth (Bates, 2007).

The hydrography and biogeochemistry of the area have been summarised by Michaels and Knap (1996), Steinberg et al. (2001), Lomas et al. (2013) and references therein. In the absence of large changes in salinity, the 10 °C seasonal change in surface temperatures driven by solar insolation controls the shoaling and erosion of the mixed layer, which reaches a maximum of 250–400 m in late winter, increasing vertical mixing and entraining nutrient-rich waters. The depth of mixing determines the strength of seasonal particulate flux, nutrient concentrations and primary production during the subsequent spring bloom (Michaels and Knap, 1996; Steinberg et al., 2001). With the onset of seasonal stratification in late February–March, a spring bloom develops when phytoplankton biomass and particulate organic carbon standing stocks are maximal. As seasonal stratification intensifies, a nutrient-depleted, shallow surface mixed layer develops which is underlain by a subsurface chlorophyll maximum at approximately 80–100 m depth. Strong stratification in summer and autumn results in low vertical mixing that limits nutrient availability and primary production. Seasonal cooling in late autumn results in erosion and gradual deepening of the mixed layer, with renewed nutrient entrainment into the euphotic zone and an increase in primary production. Mesoscale physical variability in this area is the dominant method of nutrient transport (McGillicuddy et al., 1998). In particular, passage of cyclonic and mode water eddies may lead to nutrient entrainment which generates short-lived phytoplankton blooms and community restructuring (Wiebe and Joyce, 1992; Olaizola et al., 1993; McNeil et al., 1999; Letelier et al., 2000; Seki et al., 2001; Sweeney et al., 2003) which could, in turn, impact higher trophic levels such as planktonic foraminifers. In addition, these blooms often result in short-lived, episodic periods of enhanced export fluxes of

3 Materials and methods

3.1 The OFP sediment trap time series

The OFP mooring is located at 31°50’N, 64°10’W, about 55 km southeast of Bermuda at 4200 m water depth (Fig. 1). Three Mark VII Parflux sediment traps (McLane Labs, Falmouth, MA) are deployed at depths of 500, 1500 and 3200 m. The traps (0.5 m² surface area) are programmed to collect a continuous bi-weekly time series of the particle flux. Collected samples were processed according to Conte et al. (2001) and split into <125, 125–500, 500–1000 and > 1000 µm size fractions. We analysed foraminifera in the 125–500 and 500–1000 µm size fractions of 1500 m trap samples collected during two time periods: 1998–2000 and 2008–2010 (109 samples total). We selected the two equivalent 2.5-year intervals a decade apart to generate a bi-weekly resolved time series which would enable assessment of seasonality as well as interannual variability. Our analyses focused on 11 species that fall within three general groupings: (i) surface-dwelling species living within the upper 50 m water column (Globigerinoides ruber var. white/pink Globigerinella siphonifera, Globigerinoides sacculifer), (ii) intermediate-dwelling species living in the ~50–200 m depth range (Orbulina universa, Globigerinoides conglobatus, Neogloboquadrina dutertrei, Palleniata in obtuiloculata) and (iii) deep-dwelling species (or species that are thought to calcify over a large depth range) living in the ~100–800 m depth range (Globorotalia inflata, Globorotalia crassaformis, Globorotalia truncatulinoides, Globorotalia hirsuta). Our assignments of the depth habitats were based on measured species depth distributions and/or inferred distributions based on oxygen isotopic composition (Fairbanks et al., 1980; Anand et al., 2003). The temporal offset between the foraminiferan species fluxes reaching the trap at 1500 m depth versus the timing of these species’ growth in overlying waters will vary depending on habitat depths and individual species’ sinking rates (Takahashi and Bé, 1984). A surface-dwelling G. ruber living at 25 m depth may sink at ~198 m day⁻¹, taking ~7 days to reach the 1500 m trap, whereas a more heavily calcified deeper-dwelling species such as G. inflata may sink ~504 m day⁻¹, taking only ~3 days to reach the 1500 m trap. These fast sinking rates are much shorter than the typical lifespans of PF and are thus not anticipated to cause any offset between the hydrographic and sediment trap flux data (Honjo and Manganini, 1993).

On average, ~440 tests were counted in each sample fraction. To generate the flux data, counts of total and individual foraminifera species in the sample aliquots for each size fraction were converted to total counts per sample fraction and then the totals for the two fractions were combined (i.e. total planktonic foraminifera between 125 and 1000 µm in size). Total counts were then scaled for the processing split (60 %) and converted to flux (tests m⁻² d⁻¹). PF flux data is available through www.pangaea.de.

3.2 BATS and remote sensing data

The BATS site (31°40’N, 64°10’W) is located just south of the OFP mooring (Fig. 1). Monthly hydrographic and geochemical data collected by the BATS time series were obtained from the BATS website (http://bats.bios.edu). Mixed layer depth (MLD) was available from Lomas et al. (2013) and was calculated from CTD profiles using the variable sigma-t criterion equivalent to a 0.2°C temperature change (Sprintall and Tomczak, 1992). The mesoscale eddy field was assessed using interpolated data on sea surface anomaly available from the CCAR Global Historical Gridded SSH Data Viewer (http://eddy.colorado.edu/ccar/ssh/hist_global_grid_viewer).

4 Total planktonic foraminiferal fluxes

4.1 In relation to other mass fluxes

The seasonal cycle and interannual variability of the PF flux at 1500 m depth is highly correlated with that of the total mass, carbonate and organic carbon fluxes. All fluxes are strongly characterized by an abrupt spring maximum during February–April, which varies significantly on an interannual basis (Fig. 2). For example, the spring PF flux peak ranged from a low of 400 tests m⁻² d⁻¹ in 2008, coinciding with minimal spring mass fluxes, to a high of 900 tests m⁻² d⁻¹ in 2009, coinciding with an extreme peak in spring mass fluxes. All fluxes typically drop to a minimum over the summer months (May–August) and remain low until the following spring bloom. During these minima, the PF flux generally amounts to <200 tests m⁻² d⁻¹. In some years (e.g. 2009 and, to a lesser extent, 2008), the PF flux displays a smaller, but distinct second peak in the months September–October. This secondary autumn peak can also be seen in the mass flux and carbonate flux in 2009 but is absent in the organic carbon flux. Over the entire record, the correlation between PF flux and mass, carbonate and organic carbon flux is 0.65, 0.64 and 0.55, respectively.

4.2 Relative to upper ocean hydrography

In Fig. 3 we compare interannual variations in bi-weekly resolved total PF flux to ~monthly resolved changes in key upper ocean hydrographic parameters, measured at the BATS site. PF flux exhibits an inverse relationship with seasonal variations in SSTs and reaches a maximum when SST is coolest in January–March (Fig. 3a). Of note, is the particularly large and prolonged PF bloom in 2010, which coincided
with a cyclonic eddy that passed through the area causing the lowest SSTs on record for this site $\sim 18.9^\circ\text{C}$ (Fig. 3a–b).

Sea level anomaly (SLA) provides information about eddies passing through the area (Fig. 3b). A negative anomaly is associated with cyclonic eddies and a positive anomaly associated with anticyclonic and mode water eddies. The SLA data show that the particularly high and prolonged PF fluxes, total mass flux and organic carbon flux in spring 2009 and 2010 coincided with the passage of cold, cyclonic eddies (Fig. 2), which enhance nutrient upwelling into the euphotic zone.

The annual and interannual PF flux is in phase with the deepening and shoaling of the mixed layer depth (MLD) (Fig. 3c) and with chlorophyll $a$ concentrations (Fig. 3d). The seasonal PF flux maximum coincides with the chlorophyll $a$ maximum (which is used here as a proxy for the spring phytoplankton bloom) and the organic carbon flux from 200 m, which represents organic carbon export from surface productivity (Fig. 3e), and the deepest MLD during February–March. During April–May, the MLD shoals back towards the surface coinciding with decreasing chlorophyll $a$ concentrations and PF flux. The strong correlation between the seasonality in PF flux and that of primary production and export is demonstrated by the regressions between total PF flux and chlorophyll $a$ concentration (Fig. 4a) and the 1500 m mass flux (Fig. 4b). During the winter–spring period the magnitude of PF flux generally follows the evolution in MLD and is maximal when the MLD is maximal (Fig. 4c). However, when the mixed layer depth shoals to <80 m during the low productivity period in late spring and summer, this correlation is not significant (Fig. 4d).

### 4.3 Planktonic foraminifera species fluxes

In general, all planktonic foraminifera, and especially deeper-dwelling species, show strong, consistent seasonal variance (Figs. 5–7). Our results demonstrate a clear depth progression towards more pronounced seasonality in the deeper species, compared to a larger intra-seasonal variability in the surface and intermediate dwellers. In addition, the deep-dwelling PF species exhibit repeatable species successions throughout the winter and early spring (Fig. 8, Table 1). Figure 8 shows that *Globorotalia truncatulinoides* dominates the flux of deeper dwellers, and thrives each December, reaching a maximum during January. *G. truncatulinoides* is then followed by *G. hirsuta*, *G. crassaformis*, and *G. inflata*, which all peak between March and April. *G. truncatulinoides* displays large interannual variability (Table 1), ranging from lows of $\sim 4000$ tests m$^{-2}$ year$^{-1}$ in 2009–2010 to highs of up to $\sim 14$ 000 tests m$^{-2}$ year$^{-1}$ in 1999–2000 (Fig. 6). The remaining deeper dwellers (*Globorotalia hirsuta*, *G. crassaformis*) also...
Thus, the environmental factors controlling PF flux should be closely aligned with the factors controlling phytoplankton productivity and export flux.

Figure 4. Correlation between total planktonic foraminifera flux in the 1500 m OFP trap (thin, black line) with environmental parameters measured at the BATS site. (a) Chlorophyll a concentration at 0–25 m. The correlation excludes an anomalous peak in chlorophyll a concentration observed in 2010. (b) Regression with mass flux (c) MLDs > 80 m, excluding the extremely deep MLD observed in 2010. (d) MLDs < 80 m.

Figure 5. Temporal changes in surface-dwelling planktonic foraminifera fluxes in the 1500 m trap with changes in sea surface temperature (0–25 m) shown in the dashed black line for reference. The approximate depth habitat (Anand et al., 2003) is shown in figures.

5 Discussion

The controls on PF flux in the Sargasso Sea was first introduced by Bé (1960) and later developed by Tolderlund and Bé (1971), who suggested that PF flux is dominantly controlled by the availability of their food phytoplankton. Thus, the environmental factors controlling PF flux should be closely aligned with the factors controlling phytoplankton productivity and export flux.

5.1 Environmental controls on PF fluxes

5.1.1 Depth of the mixed layer

Previous studies suggest that increased chlorophyll concentrations and larger phytoplankton abundances occur when the MLD deepens (Townsend et al., 1994; Waniek, 2003; Nelson et al., 2004) and the amplitude and timing of MLD deepening determines the size of the following spring bloom (Menzel and Ryther, 1961; Michaels et al., 1994). Here, we also observe a simultaneous seasonal peak in chlorophyll a and maximum depth of the MLD, as observed by previous studies at BATS (Steinberg et al., 2001; Cianca et al., 2012), the timing and amplitude of which coincides with the maximum PF flux (Fig. 3c, d). Similarly, seasonal changes in mixed layer depth are closely associated with changes in foraminifera production (Thunell and Reynolds, 1984; Sautter and Thunell, 1989; Pujol and Vergnaud Grazzini, 1995; Schmuker and Schiebel, 2002) and chlorophyll a concentrations (King and Howard, 2003, 2005) in other ocean basins. Siegel et al. (2002) proposed that, south of 40° N, the initiation and extent of the spring bloom is dominantly limited by nutrients, and this is supported by the simultaneous increase in phytoplankton concentrations with mixing depth at BATS (Treusch et al., 2012). Vertical mixing in late winter and spring distributes nutrients into the euphotic zone to support the spring phytoplankton bloom, causing the consequent seasonal peak in export fluxes of organic carbon, to fuel symbiont-barren foraminifera production (Fig. 2d). In contrast, no correlation exists between PF flux and MLD during the late spring to autumn when the mixed layer fails to penetrate the minimum depth of the deep chlorophyll maximum layer (∼80 m), where many species of planktonic foraminifera reside in association with other zooplankton and
Table 1. Annual fluxes for planktonic foraminifera species at 1500 m depth in 1998–1999, 1999–2000, 2008–2009 and 2009–2010 and the 4-year averages. Fluxes were calculated from the sum of bi-weekly averages between July and June for each year and converted to tests m$^{-2}$ yr$^{-1}$. Species are listed according to their estimated depth habitats.

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<tbody>
<tr>
<td><strong>Surface dwellers:</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>G. ruber</em> (pink)</td>
<td>Oct$^1$, Mar$^2$</td>
<td>256</td>
<td>292</td>
<td>1007</td>
<td>348</td>
<td>1903</td>
<td>425</td>
</tr>
<tr>
<td><em>G. ruber</em> (white)</td>
<td>Oct$^1$, Mar$^2$</td>
<td>16 197</td>
<td>19 633</td>
<td>13 917</td>
<td>18 719</td>
<td>17 117</td>
<td>3200 m avg (1978–1984)$^3$</td>
</tr>
<tr>
<td><em>G. sacculifer</em></td>
<td>Oct$^1$, Mar$^2$</td>
<td>256</td>
<td>292</td>
<td>1007</td>
<td>348</td>
<td>1903</td>
<td>425</td>
</tr>
<tr>
<td>Surface totals</td>
<td></td>
<td>18 977</td>
<td>21 903</td>
<td>16 500</td>
<td>21 189</td>
<td>17 346</td>
<td></td>
</tr>
<tr>
<td><strong>Intermediate dwellers:</strong></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td><em>G. siphonifera</em></td>
<td>Apr–May$^1$, Oct–Nov$^2$</td>
<td>1429</td>
<td>694</td>
<td>1056</td>
<td>2250</td>
<td>1357</td>
<td></td>
</tr>
<tr>
<td><em>O. universa</em></td>
<td>Nov</td>
<td>277</td>
<td>180</td>
<td>0</td>
<td>4</td>
<td>115</td>
<td>300</td>
</tr>
<tr>
<td><em>N. dutertrei</em></td>
<td>Mar–Apr$^1$, Nov–Dec$^2$</td>
<td>1290</td>
<td>185</td>
<td>471</td>
<td>839</td>
<td>696</td>
<td>876</td>
</tr>
<tr>
<td><em>P. obliquiloculata</em></td>
<td>Dec–Mar</td>
<td>398</td>
<td>205</td>
<td>708</td>
<td>352</td>
<td>416</td>
<td>762</td>
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<tr>
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<td></td>
<td>9495</td>
<td>4446</td>
<td>4466</td>
<td>6278</td>
<td>6171</td>
<td></td>
</tr>
<tr>
<td><strong>Deep dwellers:</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>G. truncatulinoides</em></td>
<td>Jan–Feb</td>
<td>5248</td>
<td>13 796</td>
<td>9517</td>
<td>4031</td>
<td>8148</td>
<td>3420</td>
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<tr>
<td><em>G. hirsuta</em></td>
<td>Feb–Mar</td>
<td>1784</td>
<td>9888</td>
<td>3859</td>
<td>2770</td>
<td>4575</td>
<td>1520</td>
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<tr>
<td><em>G. crassaformis</em></td>
<td>Feb–Mar</td>
<td>26</td>
<td>100</td>
<td>122</td>
<td>139</td>
<td>97</td>
<td>192</td>
</tr>
<tr>
<td><em>G. inflata</em></td>
<td>Mar–Apr</td>
<td>844</td>
<td>995</td>
<td>1652</td>
<td>1869</td>
<td>1340</td>
<td>1270</td>
</tr>
<tr>
<td>Deep totals</td>
<td></td>
<td>7902</td>
<td>24 779</td>
<td>15 150</td>
<td>8809</td>
<td>14 160</td>
<td>5402</td>
</tr>
<tr>
<td><strong>Other species</strong></td>
<td></td>
<td>51 442</td>
<td>43 704</td>
<td>43 172</td>
<td>70 446</td>
<td>51 191</td>
<td></td>
</tr>
<tr>
<td><strong>Totals</strong></td>
<td></td>
<td>87 816</td>
<td>94 831</td>
<td>79 289</td>
<td>106 722</td>
<td>92 165</td>
<td></td>
</tr>
</tbody>
</table>

$^1$ Primary peak. $^2$ Secondary peak. $^3$ Averages from Deuser and Ross (1989). $^4$ This species has low seasonality.

algal cells (Fairbanks and Wiebe, 1980) (Fig. 4d). This is also the depth of the nitricline where nitrate concentrations are greater than 0.1 µmol kg$^{-1}$ (Schiebel et al., 2001).

The majority of the increased PF flux in the winter–spring is driven by increased fluxes of deeper-dwelling species, in particular *G. truncatulinoides* and *G. hirsuta* (Fig. 9b). These species are symbiont-barren and rely on the flux of phytodetritus and other labile organic carbon as a food source from the spring phytoplankton bloom (Hemleben et al., 1989). The discrepancy in timing of peaks between the deeper-dwelling species (Fig. 8) is likely due to subtle changes in phytoplankton succession related to the species’ diets (Deuser and Ross, 1989; Hemleben et al., 1989). Overall, the seasonal PF species succession is broadly similar to previous observations from 1959 to 1963 and 1978 to 1984 (Tolderlund and Bé, 1971; Deuser, 1987; Deuser and Ross, 1989) which suggests that, despite long-term environmental change, species seasonality have remained consistent over the past 50 years.

The correlation observed here between the seasonality in the PF flux, chlorophyll $a$ concentration and mass flux at 1500 m (Fig. 4a and b) clearly demonstrates that the seasonality of non-symbiont-bearing foraminifera, such as the globorotaliids, is controlled by phytoplankton production and the export flux of phytodetritus to depth. As these globorotaliids are up to 3 times denser than surface species (unpublished data), their sinking rates are significantly higher than those of other species. Thus, increased production by these species can accelerate the transfer of carbonate from surface to deep ocean, thereby strengthening the carbonate pump.

In contrast, the surface-dwelling symbiont-bearing foraminifera have lifecycles which strongly benefit from stratified surface waters and shallow mixed layers in order to photosynthesise – allowing them to succeed in low-nutrient conditions (Hemleben et al., 1989). Surface dwellers generally calcify in late summer when sea surface temperatures are at a maximum and dinoflagellates are abundant (Tolderlund and Bé, 1971). We thus conclude that the depth and structure of the mixed layer plays an important role in regulating PF species flux by controlling the abundance and timing of their food availability throughout the seasonal cycle.

5.1.2 MLD deepening and shoaling rates

Current models based on the light-limited higher latitudes (Waniek, 2003; Mao, Y., personal communication, 2013), suggest that if the MLD shoals early and slowly, the
The ratio of the rate of deepening to the rate of shoaling, and computed a mixed layer dynamics index, \( \frac{r^2}{S_r} \), which is the ratio of the rate of deepening to the rate of shoaling, and compared this to the integrated PF flux (Table 2). The \( \frac{D_r}{S_r} \) ratio never exceeds 1, indicating that the shoaling rate always exceeds the deepening rate. For all the years studied, there is a strong inverse relationship between the integrated PF flux over the duration of spring bloom, and the \( \frac{D_r}{S_r} \) ratio (Fig. 10b, \( r^2 = 0.93 \)). This relationship is also present in the maximum in chlorophyll a concentration and the \( \frac{D_r}{S_t} \) ratio (Figure 10c, \( r^2 = 0.76 \)). This correlation indicates that when the MLD shoals more quickly during spring stratification (lower \( \frac{D_r}{S_t} \) ratio), the chlorophyll a concentrations and PF flux are higher, as supported by a strong correlation (\( r^2 = 0.87 \)) between shoaling rate and integrated PF flux (Fig. 10d).

To test whether the rates of mixed layer deepening in early winter and of shoaling in spring affect the PF flux, we computed a mixed layer dynamics index, \( \frac{D_r}{S_t} \), which is the ratio of the rate of deepening to the rate of shoaling, and

Figure 6. Temporal changes in intermediate-dwelling planktonic foraminifera fluxes in the 1500 m trap with changes in sea surface temperature (0–25 m) for reference. The approximate depth habitat (Anand et al., 2003) is shown in figures. Graphs are ordered according to seasonal succession.

Figure 7. Temporal changes in deeper-dwelling planktonic foraminifera fluxes in the 1500 m trap with changes in sea surface temperature (0–25 m) for reference. The approximate depth habitat (Anand et al., 2003) is shown in figures. Graphs are ordered according to seasonal succession.

Table 2. Mixed layer depth and mean rates of mixed layer (ML) deepening and shoaling. The $D_t / S_t$ ratio is a derived value calculated from the rate of ML deepening divided by the rate of ML shoaling (see text). The winter–spring PF flux represents the PF flux integrated over the whole bloom, which varied interannually in length but ranged from December to May. Bold faced years indicate years when a cyclonic eddy was present during the spring bloom period.

<table>
<thead>
<tr>
<th>Year</th>
<th>MLD max (m)</th>
<th>ML deepening rate (m day$^{-1}$)</th>
<th>ML shoaling rate (m day$^{-1}$)</th>
<th>$D_t / S_t$ ratio (m)</th>
<th>Maximum PF flux (tests m$^{-2}$ day$^{-1}$)</th>
<th>Integrated winter–spring PF flux (tests m$^{-2}$ day$^{-1}$)</th>
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<tr>
<td>1997–1998</td>
<td>235</td>
<td>0.93</td>
<td>1.91</td>
<td>0.49</td>
<td>641</td>
<td>28</td>
</tr>
<tr>
<td>1998–1999</td>
<td>222</td>
<td>0.78</td>
<td>7.78</td>
<td>0.10</td>
<td>816</td>
<td>41</td>
</tr>
<tr>
<td>1999–2000</td>
<td>197</td>
<td>0.63</td>
<td>Data missing</td>
<td>–</td>
<td>761</td>
<td>30</td>
</tr>
<tr>
<td>2007–2008</td>
<td>130</td>
<td>0.55</td>
<td>0.75</td>
<td>0.73</td>
<td>385</td>
<td>17</td>
</tr>
<tr>
<td>2008–2009</td>
<td>198</td>
<td>0.95</td>
<td>2.21</td>
<td>0.43</td>
<td>946</td>
<td>28</td>
</tr>
<tr>
<td>2009–2010</td>
<td>464</td>
<td>1.76</td>
<td>3.82</td>
<td>0.46</td>
<td>815</td>
<td>32</td>
</tr>
</tbody>
</table>

Years where the shoaling rate is twice as quick as the deepening rate (e.g. winters 1997, 2008, and 2009) have average $D_t / S_t$ ratios, average-length blooms and PF flux (∼ 30 tests m$^{-2}$ day$^{-1}$, Table 2). Years with comparatively equal rates of shoaling and deepening (e.g. winter 2007) have larger $D_t / S_t$ ratios, longer and slower blooms with shallower MLDs and small PF fluxes. Years when the shoaling rate is much quicker than deepening rate (e.g. winter 1999) have the smallest $D_t / S_t$ ratios and shorter, sharper blooms with greater numbers of intermediate thermocline-dwelling species such as N. dutertrei, P. obliquiloculata, G. siphonifera, suggesting that when the rate of shoaling is higher the seasonal thermocline is nearer to the surface for longer, which is beneficial for these symbiont-bearing and symbiont-facultative species. The PF fluxes were large (and prolonged) respectively in winter 2008–2009 and 2009–2010 despite having average $D_t / S_t$ ratios but were probably enhanced by additional factors discussed in the next section.

5.1.3 Eddies

The negative sea level anomalies in spring of 2009 and 2010 indicate that the large (and in 2010 prolonged) PF fluxes in these years were clearly associated with the passage of cyclonic eddies (Fig. 3b). Eddy pumping of nitrate into the euphotic zone has been shown to significantly increase new production (Oschlies and Garçon, 1998; Oschlies, 2002). Cianca et al. (2007) estimate that eddy pumping contributes ∼ 50% of the nutrient input into the euphotic zone in the Sargasso Sea. Studies at the BATS site have demonstrated the influence of cyclonic and mode water eddies in promoting phytoplankton blooms and increased secondary production (Eden et al., 2009; McGillicuddy et al., 2007, 1999; Goldthwait and Steinberg, 2008; Sweeney et al., 2003; Lomas et al., 2013; Cianca et al., 2012) and therefore affecting PF food availability and quality (Schmuker and Schiebel, 2002). Previous studies have found higher fluxes of certain PF species such as Globigerinoides glutinata associated with cyclonic eddy structures in the Caribbean Sea (Schmuker and Schiebel, 2002) and North Atlantic (Beckman et al., 1987), also in conjunction with upwelling frontal regions in the Mexican Pacific (Machain-Castillo et al., 2008) and deep mixed layers during winter in the Mediterranean (Pujo and Vergnaud Grazzini, 1995). Here we observe a similar response during the passage of a cyclonic eddy in spring 2009, particularly for deeper-dwelling species.
In fact, the largest PF flux observed over the entire record was associated with this eddy passage, even though the maximum MLD and $D_t / S_r$ were modest (Table 2). Similarly, the mass and organic carbon flux measured during the passage of this eddy (Fig. 2b–d) were the highest fluxes measured over the last 25 years of the OFP time series, indicating that the conditions in this eddy promoted an extremely large export flux to fuel the production of deep-dwelling foraminifera species such as *G. truncatulinoides*, *G. hirsuta*, and especially *G. inflata* which all experienced higher seasonal fluxes in 2009 (Fig. 7).

This observation is consistent with an exceptionally large increase in the flux of *G. truncatulinoides* (>600 tests m$^{-2}$ day$^{-1}$) seen at the OFP traps during the spring of 2007, which was also influenced by the passage of a productive cyclonic eddy (Fang et al., 2010; Conte and Weber, 2014). Both the 2007 and 2009 eddies occurred between January and March during the seasonal flux of the deeper dwellers (Fig. 7), underscoring the importance of the timing of eddy passage in enhancing PF flux. The influence of eddies here is similar to observations from the eastern Mediterranean where increased numbers of grazing species, such as *G. truncatulinoides* and *G. inflata*, have been found in association with eddy structures and deep mixed layers (Pujol and Vergnaud Grazzini, 1995). These findings suggest that productive cyclonic eddies, when co-occurring with deep MLDs, act to enhance the existing seasonal abundance of deeper-dwelling species through mixing of the water column, which aids their annual reproductive migration in addition to increasing food supply.

Along with the timing of the eddy passage, our observations also suggest that the PF flux response is dependent on whether the eddy is intensifying or weakening. For instance, both cyclonic eddies in 2009 and 2010 intensified over the spring bloom (Fig. 3b) eliciting a large biological response indicated by elevated subsurface Chl $a$ concentrations and increased PF flux. In contrast, the cyclonic eddy in winter 2007–2008 was weakening over the spring bloom and therefore elicited no PF flux response.

Recent studies have found that eddies which are a minimum of 1–2 months in duration are more likely to induce a larger biological response (Mourino-Carballido and McGillicuddy, 2006, Sweeny et al., 2003). Our observations also suggest that eddies need to be present for at least a month to elicit responses in the flux of PF which have minimum lifecycles of 2 weeks. For instance, in winter 1998–1999 a
cycloonic eddy passed over the sediment trap site in only 1 month and elicited no biological response, compared to cycloonic eddies in 2009 and 2010, which both remained over the site for a minimum of 2–3 months and elicited large biological responses (Fig. 3b). These findings suggest that cycloonic eddies which modify the spring bloom and last for 1–3 months can elicit a significant biological response and increased PF flux.

6 Implications

Our results show that environmental factors and mesoscale eddy variability play an important role in regulating the planktonic foraminifera fluxes, by regulating the MLD and consequent magnitude of the spring bloom and biological export flux.

An overarching climatological variable affecting this region especially is the North Atlantic Oscillation (NAO), which exerts a strong influence on air temperature, storminess, heat loss, winter mixed layer depth, and, therefore, nutrient injection into the upper ocean during the winter months (Bates, 2012; Bates and Hansell, 2004; Rodwell et al., 1999). Modelling studies have shown that when the NAO is in its low phase, i.e. negative NAO (e.g. winter 2010), there is increased heat loss that intensifies convective mixing and results in enhanced nutrient upwelling into the euphotic zone to support primary production (Oschlies, 2001). The NAO influence on upper ocean productivity and biogeochemical fluxes is demonstrated by the inverse correlation between the wintertime (NDJF) NAO index and the deep particulate nitrogen flux in the OFP traps over a 30-year period (Conte and Weber, 2014) and increased primary productivity in negative wintertime NAO phases (Lomas et al., 2010). If convective mixing and nutrient entrainment into the euphotic zone is stronger during negative NAO years, this could serve to modulate PF flux, and therefore carbonate flux, on decadal timescales. When we compare PF fluxes covering a range of NAO indexes, from this study using the 1500 m trap to the 3200 m trap between 1978 and 1984 (Deuser and Ross, 1989; Deuser, 1987), we find a weak inverse correlation between total PF flux and (DJFM) NAO index in-phase (not significant), but we do find a significant inverse correlation with a (DJF) NAO with a 1-year lag ($p < 0.005$) (Fig. 11). Cianca et al. (2012) showed that their correlation between winter NAO and total chlorophyll $a$ at BATS improved when applying a + 1-year time lag, but still remained insignificant. They attributed this to variability in the subtropical mode water, which can laterally advect nutrients on interannual timescales (Palter et al., 2005; Patara et al., 2011). We acknowledge that additional longer-term data are needed to test the mechanism behind this correlation, but our results suggest that changes in NAO status and/or mesoscale eddy frequency could significantly modulate planktonic foraminifera flux and export flux from the surface ocean on interannual timescales.

This study shows that the productivity of the dominant deep-dwelling species $G$. truncatulinoides and $G$. hirsuta is especially responsive to interannual variability in overlying surface water conditions and especially to the transient high-production/flux events that are associated with the passage of productive cycloonic eddies that coincide with their seasonal spring production peak. Our data show that deeper-dwelling species can account for up to ~90% of the total PF carbonate flux, representing up to ~40% of the total carbonate flux during winter–spring at the OFP site. Changes in NAO status, which modulates nutrient supply into the euphotic zone and the strength of the spring bloom, also may in turn modulate the production and flux of these heavily calcified deep-dwelling foraminifera by increasing their food supply, thereby intensifying the carbonate pump.

7 Conclusions

Our study demonstrates that the interannual variability in planktonic foraminifera flux can be linked to the MLD and the rate of deepening/shoaling of the mixed layer associated with nutrient injection into the euphotic zone. We find that higher PF fluxes coincide with deeper MLDs, especially when combined with cycloonic eddy-induced nutrient upwelling. In particular, the production of the dominant deep-dwelling species $G$. truncatulinoides and $G$. hirsuta is shown to be particularly responsive to interannual variability in overlying surface water conditions and especially to the transient high-production/flux events that are associated with productive cycloonic eddies. These species dominate the major late winter–early spring pulses of foraminifera and have higher sinking rates than surface-dwelling species because they are up to 3 times denser (unpublished results). We suggest deeper-dwelling species strengthen the carbonate pump by accelerating the transfer of carbonate from surface to deep ocean and contribute up to 40% of the contemporaneous peak in total carbonate export fluxes. It follows that any increase in fluxes of these deep dwellers arising from climate-induced changes in winter–spring mixed layer dynamics will also increase the average sinking rate of foraminiferal carbonate and intensify the overall carbonate pump. Our findings suggest that the North Atlantic Oscillation, via its influence on mixed layer depth, nutrient upwelling, phytoplankton production and export flux, may also serve to modulate the foraminiferal component of the carbonate pump in the subtropical North Atlantic.
Acknowledgements. We would like to thank two anonymous reviewers for their time and constructive comments that helped improve the manuscript. This research was funded through the U.K. Ocean Acidification Research Program by Natural Environment Research Council grant to P. Anand and P. Sexton (grant NE/I019891/1). We acknowledge the National Science Foundation for its support of the Oceanic Flux Program time series (most recently by grant OCE-1234292) and the Bermuda Atlantic Time Series (most recently by grant OCE-0801991). We thank Mike Lomas for providing MLD data and Yolanda Mao for providing insights and useful discussion on the data. P. Anand is also thankful to Werner Deuser for communication regarding published data.

Edited by: J. Bijma

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