Silicate:nitrate ratios of upwelled waters control the phytoplankton community sustained by mesoscale eddies in sub-tropical North Atlantic and Pacific

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Abstract. Mesoscale eddies in sub-tropical gyres physically perturb the water column and can introduce macronutrients to the euphotic zone, stimulating a biological response in which phytoplankton communities can become dominated by large phytoplankton. Mesoscale eddies may therefore be important in driving export in oligotrophic regions of the modern ocean. However, the character and magnitude of the biological response sustained by eddies is variable. Here we present data from mesoscale eddies in the Sargasso Sea (Atlantic) and the waters off Hawai‘i (Pacific), alongside mesoscale events that affected the Bermuda Atlantic Time-Series Study (BATS) over the past decade. From this analysis, we suggest that the phytoplankton community structure sustained by mesoscale eddies is predetermined by the relative abundance of silicate over nitrate (Si\textsuperscript{∗}) in the upwelled waters. We present data that demonstrate that mode-water eddies (MWE) in the Sargasso Sea upwell locally formed waters with relatively high Si\textsuperscript{∗} to the euphotic zone, and that cyclonic eddies in the Sargasso Sea introduce waters with relatively low Si\textsuperscript{∗}, a signature that originated in the iron-limited Southern Ocean. We propose that this phenomenon can explain the observed dominance of the phytoplankton community by large-diatom species in MWE and by small prokaryotic phytoplankton in cyclonic features. In contrast to the Atlantic, North Pacific Intermediate Water (NPIW) with high Si\textsuperscript{∗} may influence the cyclonic eddies in waters off Hawai‘i, which also appear capable of sustaining diatom populations. These observations suggest that the structure of phytoplankton communities sustained by eddies may be related to the chemical composition of the upwelled waters in addition to the physical nature of the eddy.

1 Introduction

Coupling between physical and biological processes in oceanic systems sets constraints on primary production in the upper ocean that control both the energy entering the ecosystem and the potential amount of carbon exported from the upper ocean via the biological carbon pump (Ducklow, 2001; Volk and Hoffert, 1985). Mesoscale eddies provide a mechanism of physical-biological coupling (McGillicuddy et al., 1999; Oschlies, 2002; Oschlies and Garcon, 1998) that may be particularly significant in the large oligotrophic gyres such as the North Atlantic Sub-Tropical Gyre (NASO), where estimates of export flux made locally are lower than geochemical estimates (Jenkins et al., 1988). Three main types of mesoscale eddy have been studied in the Sargasso Sea: mode-water eddies (MWE) and cyclonic eddies that cause an upward displacement of the seasonal-thermocline, and anticyclonic eddies that cause a downward displacement of the seasonal-thermocline. MWE and cyclonic mesoscale eddies can lift macronutrients into the euphotic zone and stimulate a biological response in the phytoplankton community, leading to enhanced export (McGillicuddy et al., 2007).

The episodic introduction of macronutrients from mesoscale eddies is thought to both stimulate phytoplankton growth and induce a shift from smaller to larger phytoplankton, which may subsequently export carbon from the euphotic zone more efficiently (Bibby et al., 2008; Brown et al., 2008; Mourino-Carballido, 2009; Vaillancourt et al., 2010).
2003). Such changes in community composition, enhanced production and export have been reported from a wide variety of mesoscale eddies. However, the character and magnitude of biological responses are variable, probably owing to the transient nature of eddies (Benitez-Nelson et al., 2007; Falkowski et al., 1991; Martin and Pondaven, 2003; McGillicuddy et al., 2007; Oschlies, 2002; Oschlies and Garçon, 1998). An interesting observation from recent studies is the difference in phytoplankton community composition that is maintained in both MWE and cyclonic eddies in the Sargasso Sea region of the NASG. MWE can sustain large-diatom communities dominated by species such as Chaetoceros spp., which are rare for the region, while cyclonic eddies are dominated by small prokaryotic phytoplankton that are more typical of the region (Bibby et al., 2008, 2009; Mouriño-Carballido, 2009; Ewart et al., 2008). In addition, measures of biogenic silicate production (0.9–1.1 mmol Si m$^{-2}$ d$^{-1}$) and accumulation (15–20 mmol Si m$^{-2}$) in a MWE in the Sargasso Sea have been reported that are six times greater than those recorded for the nearby Bermuda Atlantic Time-Series Study (BATS) site (Krause et al., 2010). Correspondingly, the estimated biogenic organic SiO$_2$ export rate of 1.27 mmol Si m$^{-2}$ d$^{-1}$ in this MWE was 20 times higher than the BATS average, suggesting that MWE can maintain substantial diatom growth and export (Krause et al., 2010). Two studies over 3-yr (Sweeney et al., 2003) and 9-yr (Mouriño-Carballido, 2009) periods observed a total of 8 MWE, 13 cyclonic eddies and 6 anti-cyclonic eddies that affected the BATS station. In these studies, only MWE features were shown to sustain significant diatom communities; for example, Mouriño-Carballido reported a statistically significant difference in the contribution of diatom pigments to the total chlorophyll-α concentration integrated over the top 160 m of the water column in MWE compared with cyclonic eddies that had affected the BATS site between 1993 and 2002 (Mouriño-Carballido, 2009). While the age of eddy features may have an important role in determining the phytoplankton community composition (Rii et al., 2008; Sweeney et al., 2003), the species differences between MWE and cyclonic eddies in the Sargasso Sea appear to be robust. A potential mechanism that has been proposed to account for the difference in overall magnitude of biological response between MWE and cyclonic eddy communities in the Sargasso Sea involves the effects of wind stress. Wind stress causes enhanced upwelling at the centre of MWE and downwelling in cyclonic eddies. This can persist throughout the lifetime of the eddy and thus sustain larger biological signals in MWE than cyclonic eddies (McGillicuddy et al., 2007; Martin and Pondaven, 2003). However, in contrast to the situation in the Atlantic, cyclonic eddies studied in the North Pacific Sub-Tropical Gyre (NPSG) have been shown to sustain high numbers of large diatoms for long periods and may even be active mechanisms of silica export in this region (Benitez-Nelson et al., 2007).

Recently, Si$^*$, the difference between available silicate [Si(OH)$_4$] and nitrate [NO$_3^-$], has been used as a tracer of the return path of deep waters upwelled in the Southern Ocean into the thermoclines of ocean systems (Sarmiento et al., 2004). The global distribution of Si$^*$ indicates that Sub-Antarctic Mode Water (SAMW) acts as a significant source of nutrients to the thermocline in the Southern Hemisphere and the North Atlantic (Sarmiento et al., 2004; Palter et al., 2010). In contrast, North Pacific Intermediate Waters (NPIW) play a more important role in thermocline nutrient supply in the Pacific (Sarmiento et al., 2004; Palter et al., 2010). In addition to acting as a tracer, we propose that the distinct Si$^*$ signatures of these water masses, negative in SAMW and positive in NPIW, may also play a role in dictating phytoplankton community responses to euphotic zone nutrient input. Si$^*$ can be considered an important indicator of nutrient availability related to the requirement of diatoms (Ragueneau et al., 2000; Brzezinski et al., 2003). When sufficient light and nutrients (including iron) are available, diatoms accumulate biomass with silicate and nitrate at a molar ratio of ~1:1 (Brzezinski et al., 2003, 2005). Consequently, for a diatom population to efficiently utilize all available nitrate, it might be expected that the ambient Si$^*$ would have to be >0µM (Ragueneau et al., 2000). Here, we investigated the possibility that the nature of the biological response instigated by upwelling mesoscale eddies, and the observed discrepancy in the contribution of large diatoms to phytoplankton communities between MWE and cyclonic eddies in the Atlantic and between cyclonic eddies in the North Atlantic and Pacific sub-tropical gyres, are related to the chemical composition of the upwelled waters impacting the euphotic zone.

2 Methods

2.1 General field work

The EDDIES project consisted of four cruises on the R/V Oceana over two summer field seasons (OC404-1 11 June–3 July 2004; OC404-4 25 July–12 August 2004; OC415-1 20 June–15 July 2005; and OC415-3 7–25 August 2005) in the Bermuda region of the Sargasso Sea (sub-tropical North Atlantic). E-Flux I (4–22 November 2004) and E-Flux III (10–28 March 2005) were conducted in the lee of the Hawaiian Islands aboard the R/V Kaimikai-O-Kanaloa and R/V Wecoma, respectively. Prior to each cruise, candidate eddies in the appropriate location were detected and tracked using satellite data, including the geostationary operational environmental satellites (GOES-10 and AVHRR) to detect sea surface temperature (SST), MODIS imagery to monitor surface chlorophyll-α, and satellite altimetry data from the CCAR (http://cordc.ucsd.edu/projects/satellite/altimetry/ccar/) (Dickey et al., 2008; Leben et al., 2002; Seki et al., 2001; Siegel et al., 1999). Each eddy was
sampled by CTD hydrocasts at several sampling stations. These stations were strategically chosen to sample sections from the outside of each eddy and through the eddy centre of each feature. The locations of these stations were based on ongoing ADCP measurements and, in the EDDIES project, were supported by along-track XBT profiles (0–700 m) to quantify the main thermocline signal.

2.2 Analytical measurements

Discrete water samples and in situ vertical profiles of temperature and salinity were collected using CTD systems, which consisted of a rosette of Niskin bottles attached to a frame containing a Seabird S9+CTD. Discrete samples were taken using this system to measure parameters such as phytoplankton pigmentation and inorganic nutrient analysis. In addition to sampling standard depths throughout the euphotic zone, the phytoplankton community at the deep chlorophyll maximum (DCM) was specifically targeted with reference to the real-time output from the fluorometer on the CTD rosette. In the EDDIES project, nutrient measurements were collected using a highly sensitive nanomolar technique with long-path-length spectrophotometry (Li and Hansell, 2008), and in the E-Flux project by a continuous segmented flow system consisting of components of both a Technicon Autoanalyser II™ and an Alpkem RFA 300™ (Rii et al., 2008). Phytoplankton pigment data were determined by HPLC analysis of the total phytoplankton community, as described previously (McGillicuddy et al., 2007; Rii et al., 2008). Discrete measurements from this sampling are now publicly available (through http://science.whoi.edu/users/olga/eddies/EDDIES_Project.html and http://www.soest.hawaii.edu/oceanography/). In this analysis, total chlorophyll-α concentration was used to estimate the relative phytoplankton abundance, while total fucoxanthin concentration was used to estimate the relative diatom abundance (Rii et al., 2008), with the ratio of fucoxanthin:chlorophyll-α providing an index of the proportional contribution of diatoms to total community biomass. One-way analysis of variance (one-way ANOVA) was used to compare differences in variables between different eddy types.

2.3 Retrospective analysis of BATS data

Eddy features affecting the BATS site in the period 1993–2002 were therefore used to study the distributions of nitrate and silicate in the present analysis. Details of the BATS sampling scheme, analytical methods, data quality control and inter-calibration procedures appear in the BATS Methods Manual (Knap et al., 1993). Data are available from the BATS website at http://bats.bios.edu/.

2.4 Whole Atlantic sections

Meridional Atlantic sections of Si* throughout the Atlantic were constructed using the GLODAP data collection (Key et al., 2004; Sabine et al., 2005).

3 Results

Here we present results of nutrient profiles and phytoplankton pigment composition of mesoscale eddy features specifically targeted in research cruises to the Sargasso Sea region of the NASG (EDDIES program 2004–2005, McGillicuddy et al., 2007) and from mesoscale eddies that form in the lee of the Hawaiian islands (NPSG) (E-Flux project 2004, Benitez-Nelson et al., 2007). These in situ measurements have been complemented by analysis of mesoscale eddy features that affected the BATS location in the Sargasso Sea (Mouriño-Carballido, 2009). We used Si* = [Si(OH)4]- [NO3] (Sarmiento et al., 2004) both as a tracer of water masses and as an indicator of the relative availability (or deficiency) of both nutrients for the phytoplankton community.

During the EDDIES and E-Flux programs a total of six cyclonic and two MWE were specifically targeted, and certain features were sampled on multiple occasions over time scales up to 2 months to assess the temporal development of the features. Three representative features of these programs were: (1) cyclonic eddy Opal (E-Flux program March 2005, Lat 19.522° N Long −157.082° W, NPSG); (2) cyclonic eddy C5 (EDDIES program June 2005, Lat 30.982° N Long −66.180° W, NASG); and (3) MWE A4 (EDDIES program June–August 2005, Lat 30.181° N Long −86.159° W, NASG). The locations of the eddies were determined remotely by satellite altimetry (Dickey et al., 2008; Siegel et al., 2008) and, more precisely, through in situ measurements collected at sea (Benitez-Nelson et al., 2007; McGillicuddy et al., 2007).

Figure 1 shows contour maps of vertical depth profiles of temperature (Fig. 1a–c), total chlorophyll-α concentration on a log scale (Fig. 1d–f), total dissolved nitrate concentration (Fig. 1g–i) and Si* (Fig. 1j–l) from approximately 150-km transects through the physical centres of the three representative mesoscale features. The physical perturbation associated with the “centre” of each eddy system can be seen from the temperature profiles; this is slightly off-centre in the transect of cyclonic eddy C5. The distribution of chlorophyll throughout each feature was dominated by a DCM (Fig. 1d–f). The DCM in the centres of cyclone Opal and MWE A4 is

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a dominant feature, whereas in cyclonic eddy C5 chlorophyll is less enhanced with respect to the DCM community at the edge of the eddy. DCM form in well-stratified water columns throughout oligotrophic ocean systems and are normally located at the top of the nutractline (Fig. 1g–i). The relative abundance of the macronutrient silicate $[\text{Si(OH)}_4]$ over nitrate $[\text{NO}_3^-]$, both measured in $\mu$mol l$^{-1}$, can be seen from the contour plot of $\text{Si}^+$ (Fig. 1j–l). Both cyclonic eddy Opal and MWE A4 have water with positive $\text{Si}^+$ in the eddy centre at depths that are close to the base of the DCM and so may influence the phytoplankton community. Cyclonic eddy C5, however, has water with negative $\text{Si}^+$ at the eddy centre close to the DCM.

Temperature-salinity plots of two representative profiles of the central stations measured in MWE A4 and cyclonic eddy C5 demonstrate that the central stations of MWE A4 and cyclonic eddy C5 in the Sargasso Sea contained similar water masses (Fig. 2a). However, the depth distribution of these water masses is clearly very different (Fig. 2b). Specifically, for MWE A4, the mode water layer between $\sim$100 and 500 m effectively isolates the DCM from the deeper thermocline waters that are found in close proximity to the base of the thermocline within C5 (Fig. 2b).

All three of the example eddy features have previously been shown to support chlorophyll biomasses in excess of what is normally measured at the Hawaiian Ocean Time Series (HOTS) and BATS at similar times of year (McGillicuddy et al., 2007). The highest chlorophyll concentration measured in the NASG MWE A4 was three orders of magnitude higher than anything previously measured at BATS (McGillicuddy et al., 2007). Microscopic analysis revealed that cyclonic eddy Opal supported diatom communities dominated by the large chain-forming diatoms Chaetoceros and Rhizosolenia spp. (Brown et al., 2008), whereas diatom communities supported by MWE A4 were dominated by Chaetoceros spp only (Bibby et al., 2008). Previous studies have shown that cyclonic eddy C5 did not support diatom communities and was dominated by the cyanobacteria Synechococcus at levels higher than those seen in the background BATS community (Ewart et al., 2008).

In Fig. 3 we show the average phytoplankton community composition associated with the DCM at the eddy centre of a total of eight eddies studied as part of the ED-DIES and E-Flux programmes. Due to nutrient exhaustion...
The biological community sustained at the deep-chlorophyll maximum (DCM) at the central station of the three eddy types described in the paper, two from the Sargasso Sea (NASG) (MWE, \( n = 2 \), and cyclonic eddy, \( C, n = 4 \)) and one from the Hawai’i region of the NPSG (cyclonic eddy, \( C, n = 2 \)). The DCM at the eddy centre station from each feature was sampled multiple times; averages are shown for each eddy type. The y-axis represents the ratio of fucoxanthin:chlorophyll-\( \alpha \) (a), \( \text{Si}^* \) at the DCM (\( \mu \text{mol} \text{l}^{-1} \)) (b), and \( \text{Si}^* \) at 300 m (\( \mu \text{mol} \text{l}^{-1} \)) (c). Standard error bars are shown.

![Graph](image)

**Fig. 3.** The biological community sustained at the deep-chlorophyll maximum (DCM) at the central station of the three eddy types described in the paper, two from the Sargasso Sea (NASG) (MWE, \( n = 2 \), and cyclonic eddy, \( C, n = 4 \)) and one from the Hawai’i region of the NPSG (cyclonic eddy, \( C, n = 2 \)). The DCM at the eddy centre station from each feature was sampled multiple times; averages are shown for each eddy type. The y-axis represents the ratio of fucoxanthin:chlorophyll-\( \alpha \) (a), \( \text{Si}^* \) at the DCM (\( \mu \text{mol} \text{l}^{-1} \)) (b), and \( \text{Si}^* \) at 300 m (\( \mu \text{mol} \text{l}^{-1} \)) (c). Standard error bars are shown.

above and light limitation below (Cullen, 1982), the DCM-associated communities are those most affected by the perturbed nutrient profiles associated with mesoscale eddy features. Hence, any changes in community composition resulting from eddy activity are most likely to be observed here. For each eddy type (Cyclonic NPSG, Cyclonic NASG and MWE NASG), three parameters are shown: the ratio of fucoxanthin:chlorophyll-\( \alpha \), an index of the proportional contribution of diatoms to total community biomass (Fig. 3a); \( \text{Si}^* \) at the DCM, a measure of the relative availability of silicate over nitrate at the DCM (Fig. 3b); and \( \text{Si}^* \) at 300 m, a measure of the relative availability of silicate over nitrate at a depth well below the euphotic zone but indicative of the potential ratio of nitrate and silicate supply to the euphotic zone (Fig. 3c). Each mesoscale feature was sampled on multiple occasions and the averages of each eddy type are reported (Cyclonic NPSG \( n = 2 \), Cyclonic NASG \( n = 4 \), and MWE NASG \( n = 2 \)). Absolute concentrations of the diatom marker pigment fucoxanthin and ratios of fucoxanthin:chlorophyll-\( \alpha \) indicated that the eddy types support different phytoplankton communities, with NPSG cyclones and NASG MWE, but not NASG cyclones, supporting significant populations of diatoms (Fig. 3a). Moreover, \( \text{Si}^* \) was lower in NASG cyclonic features at both the DCM and 300 m, which is in clear contrast to the higher values found in the other two systems. One-way ANOVA revealed significant differences in \( \text{Si}^* \) at both the DCM and at 300 m (DCM, \( p = 0.0027 \), \( F = 7.76 \); 300 m, \( p < 0.0000001 \), \( F = 33.66 \)) and the ratio of fucoxanthin:chlorophyll-\( \alpha \) (\( p < 0.0131 \), \( F = 5.11 \)) between eddy types. Subsequent Tukey-Kramer means comparison tests confirmed that both \( \text{Si}^* \) (at both the DCM and 300 m) and fucoxanthin:chlorophyll-\( \alpha \) were significantly (\( \alpha = 0.05 \)) lower in cyclonic eddies within the NASG than both NASG mode-water and NPSG cyclonic features, the latter two systems being indistinguishable at the 95% confidence level. Anti-cyclonic eddies, although not considered in this paper, generally support low biomass phytoplankton communities dominated by *Prochlorococcus* ecotypes (Ewart et al., 2008).

We also considered the mesoscale eddies that have affected the BATS site in the Sargasso Sea (NASG). These features have been identified through previous studies (Mourño-Carballido, 2009; Sweeney et al., 2003) and indicate that while MWE support diatom populations, cyclonic eddies in the Sargasso Sea typically support enhanced *Synechococcus* populations. In Fig. 4, we present the averaged vertical profiles from 0–400 m (Fig. 4a–c) and from 0–1000 m (Fig. 4d–f) of the relative abundance of silicate over nitrate, \( \text{Si}^* \), as well as both silicate concentration [\( \text{Si(OH)}_4 \text{]} \) and nitrate concentration [\( \text{NO}_3^- \)]. These results show that the base of the euphotic zone at the BATS site is affected by different water masses with a different chemical composition during cyclonic and MWE events (Fig. 4d–f). Despite these data averaging numerous features that affected BATS throughout the year, and that the influence of eddies at the BATS site is inferred both remotely from analysis of sea level and from in situ hydrographic sampling, the trend observed in mesoscale eddies studied in situ in spring/summer seasons during the EDDIES program is maintained. \( \text{Si}^* \) is always positive at depths from 0 to the inferred nutrcline (where nitrate becomes detectable using standard colorimetric techniques) (Fig. 4a and c), presumably due to surface non-siliceous phytoplankton growth stripping all the available nitrate from the system and leaving residual silicate (Fig. 4b). At depths below 100 m, \( \text{Si}^* \) becomes negative (\( \text{Si}^* = 0 \mu \text{mol} \text{l}^{-1} \) is shown in Fig. 4a and d as a vertical dotted line), and from this point to \(~500\) m \( \text{Si}^* \) is more negative in cyclonic features than MWE features. This indicates that
the trend observed in the EDDIES program (Fig. 1) is consistent with mesoscale features sampled over the past decade at the BATS site.

4 Discussion

In this paper we identify the tracer Si* (the relative abundance of silicate over nitrate) as a key parameter in determining the character of the phytoplankton community sustained by upwelling mesoscale eddies in the NASG and NPSG. In the Sargasso Sea, mesoscale eddies have been proposed to be an important mechanism by which new nutrients enter the euphotic zone and stimulate primary production (Jenkins et al., 1988; McGillicuddy et al., 1999; Oschlies, 2002). While this is clearly the case, there has emerged a clear and consistent trend relating to the nature of the biological response, namely that MWE support a phytoplankton community a significant fraction of which are diatoms and cyclonic eddies support prokaryote-dominated populations (Bibby et al., 2008; Mouriño-Carballido, 2009; Sweeney et al., 2003). This has clear implications for the capacity for export within each type of feature and should be considered when deriving mesoscale-resolving models (McGillicuddy et al., 1999; Oschlies, 2002; Oschlies and Garcon, 1998). However, both features can maintain enhanced phytoplankton populations for significant periods (> 6 weeks) and the mechanism for the marked community difference has remained enigmatic (Bibby et al., 2008; McGillicuddy et al., 1999). A mechanism of enhanced nutrient supply in MWE involving the interaction of wind stress on anti-cyclonic features has been suggested to be responsible for the overall biological response (McGillicuddy et al., 2007). However, this may not explain why there is a difference in the phytoplankton community structure or why cyclonic eddies in the NPSG are capable of maintaining communities of large diatoms (Benitez-Nelson et al., 2007).

The results presented here suggest that mesoscale eddy features in the Sargasso Sea are chemically preconditioned to support different types of phytoplankton community with implications for the potential for export from different features. Nitrate supplied to the euphotic zone is completely utilized within the DCM, resulting in extremely low concentrations above this feature (Figs. 1 and 4). As a result, the rate of nutrient supply will likely be a key determinant of the overall magnitude of the biological response (McGillicuddy et al., 2007). However, the ratio of supplied nutrients will be important in determining the structure of the phytoplankton community. Importantly, this influence on community structure (but not necessarily overall biomass) can occur even...
if the silicate concentration and/or supply within the waters influencing the base of the euphotic zone are identical. Indeed, the difference in \( \text{Si}^* \) between MWE and cyclones in the NASG is principally determined by differences in nitrate below the euphotic zone rather than silicate. For example, the silicate concentration in both MWE and cyclones at 300 m is about 2 µmol l\(^{-1} \) (±0.5 µmol l\(^{-1} \)) during the EDDIES project.

The residual silicate concentration above the DCM is similar in both cyclonic and MWE features (~1 µmol l\(^{-1} \), Fig. 4), potentially indicative of the concentration at which diatoms are out-competed for any remaining nitrate (Egge and Akssnes, 1992). Assuming that diatoms utilize silicate down to ~1 µmol l\(^{-1} \) alongside an equimolar consumption of nitrate (Brzezinski et al., 2003) and that non-siliceous phytoplankton utilize any remaining nitrate, a simple model of the expected community structure response can be formulated (Fig. 5a). At any given silicate concentration for supply waters, the resultant community structure (i.e. the contribution of diatoms to the total standing stock) is determined by \( \text{Si}^* \) (or effectively the excess of nitrate available after complete silicate removal). Although highly simplistic, for example neglecting any differential loss terms for different components of the community, this simple calculation at least qualitatively captures the observed relationship between MWE and cyclonic eddies (Fig. 5b). In MWE, where silicate and nitrate are both supplied and utilized at the DCM at relatively high \( \text{Si}^* \), diatoms become a significant fraction of the community. However, in cyclonic eddies where silicate and nitrate are supplied at lower \( \text{Si}^* \), and hence can only both be fully utilized at the DCM at a lower overall silicate/nitrate ratio, diatoms comprise a smaller fraction of the community.

Again we emphasize that, although complete nutrient depletion within the DCM drives the community structure to be dependent on the ratio of silicate to nitrate supply, the physical rate of supply will then determine the overall magnitude of biomass sustained in each eddy. Although this rate can vary during the lifetime of a particular feature, it appears to be lower in cyclonic eddies than MWE (McGillicuddy et al., 2007; Ledwell et al., 2008). Consequently, the overall high phytoplankton biomass within MWE may be driven by physical factors, while we suggest that the greater proportion of diatoms is a consequence of the higher \( \text{Si}^* \) at the base of the euphotic zone. In contrast cyclonic eddies in the Sargasso Sea are characterized by lower phytoplankton biomass dominated by smaller prokaryotes. Moreover, cyclonic eddies in the NPSG have a higher \( \text{Si}^* \) than cyclonic eddies in the Sargasso Sea and hence appear to be able to support diatom populations even with a potentially low rate of nutrient supply (Benitez-Nelson et al., 2007). Therefore, we suggest that it is only within NASG MWE (and NPSG cyclonic eddies) that nutrient inputs at relative high \( \text{Si}^* \) can result in higher diatom standing stocks within the DCM.

The observation that cyclonic eddies in the NPSG support diatom populations (Benitez-Nelson et al., 2007; Bibby et al., 2008) clearly indicates that the mechanism responsible for governing the dominance or not of diatoms in oligotrophic eddy communities is not a cyclone-specific physical mechanism. The nature of the biological community response to eddy activity in oligotrophic systems therefore appears to be predetermined by the characteristic waters influencing the DCM within each eddy feature. In Fig. 6, we provide a simplified schematic of the potential sources for these waters in cyclonic and MWE features in the Atlantic.
Fig. 6. Proposed model of the source of nutrients that affect the thermocline in the North Atlantic Sub-tropical Gyre (NASG). The figure shows a stylized section of Si* in both the Southern Ocean and the NASG. Sub-Antarctic Mode-Water (SAMW), where a negative Si* is formed in Antarctica from upwelling of nutrient-rich deep waters followed by subsequent northward Ekman transport and concurrent preferential silicate removal (a). Water with a Si* signal derived from SAMW is transported north and can reach the thermocline in the NASG through the action of mesoscale cyclonic eddies (b), but is downwelled by the action of mesoscale mode-water eddies (c) and uncoupled from the thermocline. Winter mixing of SAMW and surface water (d) on the northern boundary of the NASG leads to the formation of sub-tropical mode water (STMW) with positive Si* that impacts the base of the euphotic zone (EHZ, dashed horizontal line) in the NASG through the action of mesoscale mode-water eddies (e). This mechanism determines whether the phytoplankton community stimulated by mesoscale eddies can sustain significant diatom populations. (e) Latitudinal section of Si* through the Atlantic from GLODAP data.

Alongside a section of Si* within the upper 3000 m, which is clearly suggestive of the southern source of the low Si* signal within the thermocline (Sarmiento et al., 2004). The sub-DCM waters within cyclonic eddies are more influenced by SAMW with low Si*, as used as a tracer of this water mass (Sarmiento et al., 2004). The low Si* in SAMW is the result of the selective silicate pump in the Sub-Antarctic, whereby iron-limited phytoplankton populations, dominated by diatoms, are heavily silicified (have a Si:N ratio > 1), resulting in relatively more silicate than nitrate being exported, leaving a low Si* signature (Sarmiento et al., 2004; Boyd et al., 2001; Hutchins and Bruland, 1998; Takeda, 1998). We suggest that this mechanism caused by iron-limitation in the Southern Ocean predetermines the biological effects of cyclonic eddies in the Sargasso Sea some 6000 miles and 50–100 yr away (Zuo et al., 2009). However, the base of the DCM in MWE is more influenced by Sub-Tropical Mode-Water (STMW) that is formed during winter mixing in the northern boundary of the NASG (Palter et al., 2005, 2010; McCartney, 1982; Talley and Raymer, 1982). Surface waters in the NASG have a high Si* owing to the preferential removal of nitrate over silicate by non-silicifying phytoplankton over the summer. During winter in the STMW formation region, these surface waters will mix with deeper thermocline waters containing the lower Si* signature of SAMW. STMW thus forms with a higher Si* than these lower thermocline waters. Consequently, when a MWE lens impacts the euphotic zone it is has the capacity to maintain a significantly enhanced contribution of diatoms to overall community biomass. The combination of our proposed mechanism with an overall enhanced nutrient flux also likely explains the observed increase in biogenic silicate production at BATS when mode waters influence the site (Krause et al., 2010) and suggest that the extent of winter mixing to the north could have a direct effect on the Si* of STMW and therefore the biological response to MWE in this region. We believe this is the first demonstration of how mechanisms such as the selective silicate pump in the Southern Ocean, which are temporally and spatially separate from the Sargasso Sea, can affect biology and export in this region. The
study further illustrates how large-scale circulation patterns can govern differences in biogeochemical responses between the North Atlantic and Pacific sub-tropical gyres.

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