

1 **Abstract**

2 Despite slow nutrient supply to the subtropical surface ocean, its rates of annual inorganic carbon
3 drawdown and net oxygen production are similar to those of nutrient-rich high latitude waters.
4 This surprisingly rapid carbon drawdown, if due to the production and export of marine biomass,
5 cannot be explained in terms of known nutrient supply mechanisms. Moreover, carbon budgets
6 have failed to detect the export of this organic matter. One possible explanation is the export of
7 nutrient-poor organic matter with a composition that avoids detection as sinking particles. We
8 describe three signs of the decomposition of such organic matter in the shallow Sargasso Sea
9 subsurface. First, summertime oxygen consumption at 80-400 m occurs without the rate of
10 nitrate and phosphate production expected from the remineralization of marine biomass,
11 satisfying the observed summertime mixed layer inorganic carbon drawdown. Second, a seasonal
12 change in the $^{18}\text{O}/^{16}\text{O}$ of subsurface nitrate suggests summertime heterotrophic bacterial nitrate
13 assimilation down to ~400 m, as may be required for the remineralization of nutrient-poor
14 organic matter. Third, incubation of subsurface seawater leads to nitrate drawdown and
15 heterotrophic bacterial growth, supporting the thermocline nitrate $^{18}\text{O}/^{16}\text{O}$ evidence for
16 heterotrophic nitrate assimilation. These three pieces of evidence suggest the export of nutrient-
17 poor organic matter from the surface at a rate adequate to explain net community production in
18 the Sargasso Sea. We propose that transparent exopolymer particles or related compounds,
19 generated by a nutrient-limited upper ocean ecosystem, comprise this nutrient-poor export, and
20 that its properties cause its flux out of the euphotic zone to be underestimated by sediment traps.
21 Such nutrient-poor organic matter would contribute little to fisheries, deep ocean carbon dioxide
22 storage, or organic carbon burial, so that it may change our view of the significance of net
23 community production in the subtropical ocean.

1 **Introduction**

2 The net production of organic matter by upper ocean ecosystems is a central characteristic of the
3 global ocean. It underpins the ocean's "biological pump," whereby organic matter is exported
4 from surface waters prior to its remineralization back to carbon dioxide (CO₂). This lowers
5 atmospheric CO₂, and portions of the organic matter sustain upper trophic levels. Moreover, a
6 small fraction of the organic matter export is buried, thus contributing to the maintenance of
7 diatomic oxygen (O₂) in the atmosphere.

8

9 The biological productivity of much of the open ocean is limited by the supply of the major
10 nutrients nitrogen (N) and phosphorus (P). In particular, the sunlit upper waters of the subtropical
11 gyres receive the major nutrients at a lower rate than the polar, subpolar, and equatorial
12 upwelling regions (Williams and Follows, 2003). By many metrics, such as surface ocean
13 chlorophyll concentrations and the flux of organic matter that reaches 2000 m, the productivity
14 of the subtropical gyres appears to be appropriately depressed (Honjo et al., 2008; Lomas et al.,
15 2013; Yoder et al., 1993). However, a different picture is suggested by upper ocean budgets of
16 dissolved inorganic carbon (DIC) and dissolved O₂ (Emerson, 2014, and references therein).

17

18 DIC and dissolved O₂ have been used to study both the euphotic zone (the sunlit upper ocean
19 down to the depth of the 1% light level, typically ~100 m in the subtropical North Atlantic near
20 Bermuda) and the surface wind-mixed layer, which is typically <30 m near Bermuda during the
21 summer but can deepen to ~200 m for brief periods in the winter (Lomas et al., 2013). For both
22 the euphotic zone and the surface mixed layer, the warm-season (spring-summer-early fall)
23 decline in the concentration of DIC provides a first order measure of net community production

1 (NCP) (Michaels et al., 1994), the net production of organic matter in the sunlit upper water
2 column that should be equivalent to organic carbon export on time scales of months and longer.
3 Gas exchange and other terms are also significant for seasonal changes in the upper ocean DIC
4 concentration but can be addressed, for example, with the use of carbon isotopes (Gruber et al.,
5 1998). For dissolved O₂, gas exchange plays a much greater role, such that the mixed layer O₂
6 concentration ([O₂]) during the summer approximately reflects a steady state between NCP
7 (which produces O₂) and evasion of O₂ to the atmosphere (Jenkins and Goldman, 1985;
8 Emerson, 1987; Hendricks et al., 2004). Due to the importance of gas exchange, approaches for
9 estimating NCP from O₂ depend on whether measurements are from the surface mixed layer
10 (Kaiser et al., 2005; Emerson et al., 2008; Stanley et al., 2010; Nicholson et al., 2015), the
11 euphotic zone (Jenkins and Goldman, 1985; Spitzer and Jenkins, 1989; Nicholson et al., 2008,
12 Riser and Johnson, 2008; Howard et al., 2010), or the underlying dark ocean where the exported
13 organic matter is remineralized (Jenkins, 1982; Jenkins and Goldman, 1985; Stanley et al.,
14 2012).

15
16 DIC and O₂-based measurement approaches suggest that annual NCP is remarkably uniform
17 across the global ocean, being no lower in the subtropical gyres than other open ocean
18 environments (Emerson et al., 2008; Emerson and Stump, 2010; Emerson, 2014; Hamme and
19 Emerson, 2006; Hendricks et al., 2005; Jenkins and Doney, 2003; Munro et al., 2013; Quay et
20 al., 2009; 2012; Reuer et al., 2007; Spitzer and Jenkins, 1989; Stanley et al., 2010, 2012). This
21 similarity is surprising given the scarcity of major nutrients in the subtropical surface ocean and
22 the substantial density difference between the nutrient poor surface waters and underlying
23 nutrient-rich deep waters. Accordingly, most models do not predict it (see Emerson, 2014, for a

1 compilation). This unexpected result is bound up with two long-standing gaps in our
2 understanding of productivity in the subtropical ocean, which have been brought into focus by
3 studies at ocean time-series sites, particularly the Bermuda Atlantic Time-series Study (BATS)
4 in the northwestern North Atlantic subtropical gyre (the Sargasso Sea). Below, we describe these
5 problems and then propose a hypothesis to explain them as well as the overarching observation
6 of high subtropical NCP.

7

8 The Problems:

9 Problem 1: Missing carbon export

10 Integrating over the depth range of the euphotic zone at BATS, the high NCP rates estimated
11 from O₂ production and DIC consumption are not detected in the fluxes of organic matter from
12 the euphotic zone by sinking as measured with sediment traps, downward mixing of dissolved
13 and particulate organic carbon (POC) as measured by depth profiles of DOC and POC
14 concentration, or other processes (Michaels et al., 1994; Carlson et al., 1994). On average, NCP
15 at BATS based on the O₂ and DIC approaches described above is 2-5 mol C m⁻² yr⁻¹ (Table 1)
16 (Spitzer and Jenkins, 1989; Cianca et al., 2013; Stanley et al., 2012; see Emerson, 2014, for a
17 compilation). In contrast, the average annual rate of export production from 24 years of sinking
18 POC collected in surface-moored particle interceptor sediment traps (PITS) is 0.88 ± 0.14 mol C
19 m⁻² yr⁻¹ (Lomas et al., 2013), while the downward mixing of POC and DOC have been estimated
20 at 0-0.05 mol C m⁻² yr⁻¹ and 0.4-1.4 mol C m⁻² yr⁻¹, respectively (Michaels et al., 1994; Carlson
21 et al., 1994; Hansell and Carlson, 2001; Omand et al., 2015). Measured carbon export is thus
22 roughly 1.3-2.3 mol C m⁻² yr⁻¹, 43-77% of the NCP estimates.

23

1 The disagreement between measured euphotic zone NCP and the organic carbon measured
2 leaving the euphotic zone has been widely suspected to result from a tendency of sediment traps
3 to “undercollect” sinking POC. If so, then the BATS traps would need to miss as much as ~60%
4 of the organic matter export associated with NCP. Trap inaccuracies are thought to derive
5 primarily from three factors: hydrodynamic biases, contamination by zooplankton “swimmers”,
6 and in-trap solubilization of material after collection (Buesseler et al., 2007). With respect to trap
7 hydrodynamics, Buesseler et al. (2000) found no coherent difference between the export flux
8 captured by the PITS traps at BATS and that collected by neutrally buoyant sediment traps
9 (NBSTs) designed to minimize hydrodynamic biases (Valdes and Price, 2000; Valdez and
10 Buesseler, 2006). With respect to “swimmers,” the BATS traps are poisoned and covered with a
11 baffle to prevent direct feeding on the collected material, and swimmers are removed prior to
12 sample analysis. In any case, failure to remove swimmers would tend to yield an overestimate of
13 the sinking flux rather than an apparent under-collection of POC (Karl and Knauer, 1989). The
14 last possibility, solubilization, has proven difficult to assess, and data on this issue are scarce.
15 Buesseler et al. (2007) conclude based on the existing data that short-term trap deployments (1-3
16 days) and timely processing of samples after collection, both of which are standard practice for
17 the BATS trap program, minimize particle solubilization. However, this conclusion relies to
18 some degree on assumptions about the particles involved.

19

20 Problem 2: Missing nutrient supply

21 The second problem has been best characterized for the summertime surface mixed layer, the
22 base of which is typically <30 m deep during the summer at BATS (Lomas et al. 2013). The
23 summertime drawdown of DIC from the mixed layer is consistent with the high NCP measured

1 for the euphotic zone (Gruber et al., 1998). Yet the mixed layer is isolated from dissolved
2 nutrients, with more than 50 m of effectively nutrient-free euphotic zone water below it that
3 separates it from the nutrient-bearing dark subsurface waters. Thus, the mixed layer DIC
4 drawdown appears to occur without a circulation-based mechanism of major nutrient (nitrate and
5 phosphate) supply. These observations have led to a search for (1) unseen nutrient supply
6 mechanisms to fuel the needed export production (Hood et al., 2001; Houghton et al., 2018;
7 Johnson et al., 2010; Katija and Dabiri, 2009; McGillicuddy et al., 1998; Villareal and
8 Lipschultz, 1995) or alternatively (2) the production and export of organic matter with a much
9 lower N and P content than is typical for upper ocean biomass, such that no additional nutrient
10 supply is required (Martiny et al., 2013; Ono et al., 2001; Toggweiler, 1993).

11
12 With regard to a previously unrecognized nutrient supply mechanism, there are fewer options
13 than frequently assumed. The nutrients in the shallow subsurface of the subtropical ocean were
14 mostly replaced by the regeneration of organic matter exported from the surface, such that they
15 are paired with a DIC excess (and an O₂ deficit) determined by the stoichiometry of that organic
16 matter (as will be discussed in detail below in the context of “preformed” nutrient changes). As a
17 result, phytoplankton growth and carbon export driven by a greater-than-recognized supply of
18 dissolved nutrients from below would not drive a larger net deficit in DIC within the summer
19 mixed layer. Rather, the circulation- or mixing-based nutrient supply and resulting export
20 production would largely offset one another in their effects on DIC (Johnson et al., 2010; Lomas
21 et al., 2013). This is the case regardless of the specific physical mechanism, be it diapycnal
22 diffusion, salt fingering, eddies, or frontal effects. Thus, while mesoscale features such as eddies
23 have been shown to be an important source of nutrients to summertime Sargasso Sea surface

1 waters (McGillicuddy et al., 1998), they cannot explain the amplitude of the DIC drawdown in
2 the summer mixed layer at BATS. The only mechanisms of nutrient supply that have the
3 potential to explain the net DIC drawdown out of the summer mixed layer are biological
4 processes: (1) N₂ fixation (for N, possibly augmented by a contribution from atmospheric N
5 deposition) and (2) phytoplankton uptake of dissolved nutrients (N and P) from the subsurface
6 followed by migration into the mixed layer.

7

8 An argument has been made that, while a nutrient supply problem exists for the summer mixed
9 layer at BATS, it does not apply to the Sargasso Sea euphotic zone. The nitrate supply rate
10 estimated with the “³He flux gauge” is adequate (possibly more than adequate) to fuel the high
11 NCP estimated for the euphotic zone from DIC and O₂ budgets (Stanley et al., 2012; 2015).
12 However, several critical caveats must be recognized. First, as acknowledged by the originators
13 of the ³He flux gauge, an unknown fraction of the estimated nitrate supply may be consumed
14 along the obduction region of the northern margin of the North Atlantic subtropical gyre,
15 reducing the implied nitrate supply at BATS (Stanley et al., 2012 and references therein).
16 Second, as already described in the context of the mixed layer, nutrient supply from below would
17 occur with a stoichiometric burden of DIC excess and O₂ deficit, such that production
18 immediately fueled by it would not contribute to the net drawdown of DIC or the net production
19 of O₂. The only way around this problem is for the nutrients to be supplied during the winter
20 when O₂ can be taken up from the atmosphere and upwelled DIC can be mixed throughout the
21 euphotic zone, providing the baseline from which summertime DIC drawdown is calculated.
22 Even this possibility is limited by the nutrient stocks in the euphotic zone observed in the early
23 summer at BATS (Lomas et al., 2013), which are far too low to yield the rate of summer NCP

1 indicated by DIC and O₂. Thus, for the euphotic zone as for the mixed layer, circulation-driven
2 nutrient supply (even if sporadic) is not a viable driver of summer measurements of high NCP
3 (e.g., as observed by Luz and Barkan (2009) and Estapa et al. (2015)).

4

5 Previous studies have shown that the rate of N₂ fixation at BATS is far too low to provide the
6 “missing N” required to support the measured rate of NCP (Altabet 1988; Hansell and Carlson,
7 2001; Orcutt et al., 2001; Knapp et al., 2005). Moreover, in a biogeochemical model, if an
8 adequately high N₂ fixation rate is imposed to simulate the observed DIC drawdown, the model
9 also produces DON and DOC anomalies in late summer/early fall that are not observed in the
10 environment (Hood et al., 2001). In addition, even if adequate N were supplied by N₂ fixation, it
11 would not address the needed supply of P, which is also required for phytoplankton growth and
12 is present at extremely low concentrations in BATS surface waters (Wu et al., 2000; Ammerman
13 et al., 2003; Mather et al., 2008; Lomas et al., 2010). Thus, the available data indicate that N₂
14 fixation is not the answer.

15

16 Atmospheric N deposition represents an alternative source of N to surface waters that is not
17 stoichiometrically linked to carbon. However, the flux of atmospheric N to the BATS site is low
18 (Knap et al., 1986; Michaels et al., 1993; Altieri et al., 2016). More importantly, even if adequate
19 N were supplied to BATS surface waters via atmospheric deposition, this flux would supply N
20 but not P, which is typically present in low concentrations in atmospheric deposition (i.e., with a
21 N/P ratio of >30; see Kanikadou et al., 2012 and references therein).

22

1 Upward nutrient transport by phytoplankton migration has been identified as a significant
2 process in the open ocean (Villareal et al., 2014). Direct observations are limited to large diatoms
3 and the dinoflagellate *Pyrocystis*, and this process has not yet been shown to be significant in the
4 Sargasso Sea. Nonetheless, data from the subtropical North Pacific show that large migrating
5 diatom mats (*Rhizosolenia* spp.) can mediate a significant upward transport of nitrate from the
6 subsurface into the mixed layer, and high intracellular nitrate concentrations have been measured
7 for putative migrators in both the Pacific and Atlantic (Villareal and Carpenter 1994; Villareal
8 and Lipschultz, 1995; Villareal et al., 2014). While nitrate transport by *Pyrocystis* has not been
9 directly shown in the Sargasso Sea, its migration has been documented in these waters (Rivkin et
10 al., 1984), and, in a study focused on the North Pacific, this species has been estimated to
11 transport as much as $17 \mu\text{mol N m}^{-2} \text{d}^{-1}$ into the euphotic zone (Villareal et al., 2014). However,
12 even if this rate of nitrate supply were sustained throughout the year at ATS, it would account for
13 a flux of only $6.2 \text{ mmol N m}^{-2} \text{yr}^{-1}$. Approximately half of the NCP at BATS is unaccounted for
14 by the documented nutrient supply. This amounts to $\sim 1.5 \text{ mol C m}^{-2} \text{yr}^{-1}$, or $\sim 0.2 \text{ mol N m}^{-2} \text{yr}^{-1}$
15 assuming a C/N ratio of 7. Year-round migration by *Pyrocystis* would supply only 3-4% of the N
16 needed to fuel this outstanding portion of the NCP. In addition, *Rhizosolenia* mats are very rare
17 at BATS (Carpenter et al., 1977), and the abundance of the large migrating diatom, *Ethmodiscus*,
18 is also low ($0.03\text{-}4.7 \text{ cells m}^{-3}$; Swift, 1973; Villareal and Carpenter, 1994; Villareal et al., 1999).
19 The possibility that these migrating phytoplankton supply some quantity of subsurface nitrate to
20 BATS surface waters in summer cannot be ruled out, but the data in hand do not make a
21 convincing argument for the process as a central nutrient flux, especially in light of concerted
22 efforts to find it in the Sargasso Sea.

23

1 There is evidence from culture studies that phytoplankton smaller than *Pyrocystis*, *Rhizosolenia*,
2 and *Ethmodiscus* can display ascending behavior (Waite et al. 1997). However, this has not yet
3 been shown in the open ocean. This process operating alone at BATS would require small
4 phytoplankton to migrate 75-100 m to transport nitrate from below the euphotic zone into the
5 upper 20-40 m meters of the water column. Nitrogen isotopic analysis indicates that nitrate
6 assimilation by small eukaryotes occurs in the euphotic zone and on some occasions in the mixed
7 layer of the summertime Sargasso Sea (Fawcett et al., 2011). It is possible that the mixed layer
8 nitrate derives from migration of the small eukaryotes in question, although it would require
9 them to migrate very large distances in just a few days. The existing eukaryotic N content and
10 isotope data suggest that these small eukaryotes constitute ~25% of the euphotic zone biomass
11 and rely on nitrate for ~30% of their N in the summertime mixed layer and <10% by the fall
12 (Fawcett et al., 2011, 2014). Taking an upper bound for their growth rate of 0.5 d^{-1} (Goerike and
13 Welschmeyer, 1998; Cuvelier et al., 2010), a growing season of 210 days (7 months), and a
14 biomass C/N ratio of 7, this amounts to the removal of 7-11 μM DIC. Between April and
15 October, mixed layer DIC drawdown is ~26 μM (Gruber et al., 1998). Thus, even when small
16 eukaryotes are assumed to acquire all of their nitrate by migration into subsurface waters,
17 biological nitrate transport could fuel less than half of the DIC drawdown.

18

19 Lacking promising mechanisms of nutrient supply that could explain the summertime mixed
20 layer decline in DIC, the most straightforward explanation is that it results from the export of
21 organic matter with C/N and C/P ratios far higher than Redfield's values of ~7 and ~106 (e.g.,
22 Lomas et al., 2013; Martiny et al., 2013). Indeed, when ocean models are confronted with the

1 existing biogeochemical data, they predict exactly this sense of deviation from Redfield
2 stoichiometry in the subtropical regions (DeVries and Deutch, 2014; Teng et al., 2014).

3

4 However, the nature of this putative nutrient-poor organic carbon export is a mystery. Its
5 characteristics must be such that it is neither measured by sediment traps nor apparent in
6 calculations that consider the hydrographic transport of DOC and suspended POC from the
7 euphotic zone to the subsurface. Suspended particulate organic matter (POM) sampled by
8 filtration and (more importantly) sinking POM captured by sediment traps have a C/N ratio close
9 to 7 (Martiny et al., 2013; Schneider et al., 2003), suggesting no preferential export of C relative
10 to N. Moreover, their nutrient content aside, the magnitude of the measured downward flux of
11 these materials cannot account for either NCP from the euphotic zone or DIC drawdown in the
12 summer mixed layer (Michaels et al., 1994). Dissolved organic matter (DOM) produced in the
13 subtropical euphotic zone does have an appropriately low nutrient (e.g., N) content (Hansell and
14 Carlson, 2001). However, as with the sinking and downward mixing of POC, calculations of the
15 downward mixing of DOC indicate that it is too slow to explain the high rates of NCP (Michaels
16 et al., 1994; Carlson et al., 1994; Hansell and Carlson, 2001); it was included in the carbon
17 accounting above.

18

19 The hypothesis: Export of gel-like, nutrient-poor organic matter

20 DOM has strong chemical similarities with carbohydrate exuded by phytoplankton (Aluwihare
21 and Repeta, 1999), and this carbohydrate can develop a gel-like substance known as “transparent
22 exopolymer particles” (TEP) (Alldredge et al., 1993). Phytoplankton release DOC as roughly a
23 quarter of their organic carbon production (Teira et al., 2003), a significant fraction of which is

1 polysaccharide (Engel et al., 2004; Passow, 2002). This assembles to form TEP (Alldredge et al.,
2 1993; Chin et al., 1998; Engel et al., 2004), often a significant portion of POC (Beauvais et al.,
3 2003; Engel et al., 2004).

4
5 TEP often binds particles together, forming marine snow that may transport large quantities of
6 biomass-derived organic matter into the ocean interior (Passow, 2002). TEP itself is positively
7 buoyant (Azetsu-Scott and Passow, 2004; Mari, 1999; Mari et al., 2017), with a density inferred
8 from settling experiments of $0.70\text{-}0.84\text{ g cm}^{-3}$ (Azetsu-Scott and Passow, 2004). Nonetheless,
9 there is ample evidence for a role for TEP in sedimentation in the ocean (Kumar et al., 1998;
10 Passow, 2002; Passow et al., 2001; Riebesell et al., 1995). While adding denser organic matter
11 may or may not render TEP sufficiently negatively buoyant to sink, the addition of ballasting
12 material such as calcium carbonate, clays, and biogenic and lithogenic silica can produce
13 aggregates with densities that are adequate for slow sinking ($1.04\text{-}1.12\text{ g cm}^{-3}$; *SI 1.1*; Mari et al.,
14 2017). Because TEP-containing aggregates likely cover a broad compositional spectrum, with
15 TEP content varying widely, a continuum of sinking rates is to be expected, ranging from non-
16 sinking TEP-rich particles to rapidly sinking particles with a high proportion of ballast and only
17 a small proportion of TEP (Mari et al., 2017).

18
19 We propose that the unexpectedly high O_2 production and DIC uptake in subtropical surface
20 waters is largely due to the production and sinking of such nutrient-poor, gel-like organic matter
21 (Fig. 1). This gel-like organic matter (GLOM) may be TEP or a material exhibiting similar
22 physical characteristics and having a related composition. We further propose that GLOM
23 accumulates in the summertime mixed layer at BATS because of an increased proportion of

1 carbon-rich exudates from phytoplankton growing under nutrient limitation (Corzo et al., 2000;
2 Mari et al. 2017) and/or because heterotrophic bacteria lack the N and P to metabolize it. Upon
3 binding to an adequate number of denser particles, a fraction of it begins to sink, generating a
4 flux of carbon-rich organic matter out of the euphotic zone. Because this material is nutrient-
5 poor (Mari, 1999; Mari et al., 2017; Passow, 2002), it obviates the need for the as-yet-
6 unobserved nutrient supply to the BATS euphotic zone. Because particles with large proportions
7 of GLOM will sink very slowly and because GLOM is probably easily disaggregated and
8 dissolved, we propose that much of the exported GLOM is hydrodynamically excluded from
9 traps, is inadequately dense to settle into the brine-filled trap collection cups, and/or is not
10 preserved in them.

11
12 The POM collected in sediment traps at BATS has a C/N ~ 7 (Schneider et al., 2003), suggesting
13 that N-poor GLOM is not captured in the traps in significant quantities. Sediment traps have long
14 been suspected of excluding some sinking material for hydrodynamic reasons (Gardner, 2000;
15 Buesseler et al., 2007; see above), and low-density GLOM would be a prime candidate for such
16 under-collection (Buesseler et al., 2006). In addition, the brine solution added to the collection
17 cups of sediment traps prior to deployment (which typically has a density of $\sim 1.08\text{-}1.1\text{ g cm}^{-3}$)
18 acts as a physical barrier to particles and aggregates that are less dense than the brine. These
19 particles are then resuspended or broken up at the interface of the trap (Gardner, 2000). Indeed,
20 laboratory experiments have shown that the addition of brine (with a density $\leq 1.08\text{ g cm}^{-3}$) can
21 decrease the flux collected in the traps by 50% (Gardner and Zhang, 1997). GLOM, which we
22 expect to be less dense than the brine, would thus be preferentially excluded. We note that
23 because of its shallow remineralization, GLOM export should not affect the remineralization

1 ratios in the mid-depth and deep ocean, explaining the consistency of 400-4000 m data with the
2 remineralization of marine biomass with Redfield-like C-to-nutrient ratios (Anderson and
3 Sarmiento, 1994).

4

5 GLOM may also avoid detection as suspended material in the surface ocean. Much of the GLOM
6 that has an *in situ* size appropriate to be captured on filters with the typical pore sizes (e.g., 0.7 μm
7 pore size glass fiber filters) may be lost through disintegration on the filter. There is strong
8 evidence that this applies to TEP. A typical average concentration of TEP in the summertime
9 Sargasso Sea is 60-80 $\mu\text{g Xanthan equivalent L}^{-1}$ (Xeq L^{-1}) (Cisternas-Novoa et al., 2015; Estapa
10 et al., 2015), which translates to 3-4 $\mu\text{M C}$ using the conversion factor of 0.63 of Engel and
11 Passow (2001). This concentration is similar to or higher than typical measurements of total
12 suspended POC in the region (Lomas et al., 2013). Estapa et al. (2015) sampled simultaneously
13 for TEP and POC in the summertime Sargasso Sea, measuring average concentrations of TEP
14 and suspended POC of $\sim 4 \mu\text{M C}$ and $2.7 \pm 0.7 \mu\text{M C}$, respectively. Moreover, some TEP
15 concentrations were much higher, up to 200 $\mu\text{g Xeq L}^{-1}$ (or $\sim 11 \mu\text{M C}$), well above the POC
16 concentration in the corresponding samples. These data raise the possibility that a large fraction
17 of TEP is disaggregated by filtration, passes through the filter, and is binned into the much larger
18 DOC pool. While this is not required by our hypothesis, it would be consistent with it.

19

20 In a similar vein, NCP as measured with DIC and O_2 budgets at BATS is similar to ^{14}C
21 incubation-based measurements of net primary production (NPP) (Jenkins and Goldman, 1985),
22 whereas ecological and biogeochemical expectations are for NCP to be a small fraction (10-
23 25%) of NPP (Dugdale and Goering, 1967; Eppley and Peterson, 1979). GPP measurements rely

1 on filtration to separate ^{14}C -labeled POC from the ^{14}C -labeled DIC substrate. Thus, if the
2 production of TEP (or GLOM) is a major fate for C fixation during the summer, disaggregation
3 of this material upon filtration would lead to an underestimation of GPP (even disregarding the
4 possibility that many GLOM particles may be smaller than the typically used filter size of 0.7
5 μm ; Mari et al., 2017). This would then explain the long-troubling result of a very high
6 NCP/NPP ratio in the subtropical North Atlantic (Jenkins and Goldman, 1985; Luz and Barkan,
7 2009).

8

9 To summarize, given the ephemeral nature of GLOM proposed above, it could have gone
10 undetected at BATS. Nevertheless, there should be signs of its remineralization in the
11 subsurface. Below, we provide three forms of evidence for the remineralization of such low-
12 nutrient organic matter at 100-400 m depth near BATS. The first involves “preformed nitrate,”
13 the quantity of nitrate in excess of that expected from the respiration of typical marine biomass,
14 which is itself estimated from the apparent oxygen utilization. We report data from profiling
15 floats with nitrate and O_2 sensors, corroborated by the BATS program data, that indicate a
16 summertime decline in preformed nitrate at 80-400 m depth, consistent with the regeneration of
17 low-N organic matter. Second, the consolidation of multiple water column profiles of nitrate
18 oxygen isotopes ($^{18}\text{O}/^{16}\text{O}$) at BATS suggest summertime heterotrophic nitrate assimilation in
19 subsurface waters, an expected consequence of the respiration of N-poor organic matter. Third,
20 dark incubations of 140-200 m water samples from BATS yield heterotrophic nitrate
21 assimilation, consistent with N limitation of heterotrophic bacteria in these subsurface waters.
22 The first form of evidence (from the profiling floats) is the most compelling of non-Redfield
23 export, although it has an alternative interpretation in the form of biological nutrient transport;

1 these data do not prove the GLOM hypothesis, but they are fully consistent with it. The latter
2 two forms of evidence, both serendipitous, are more novel but also more speculative. While
3 intriguing to the authors, they are included not as definitive proof but rather to point to two
4 complementary avenues for pursuing this hypothesis and related concepts.

5

6 **Materials and methods**

7 *Profiling floats* – Observations were made with Teledyne/Webb Research APEX profiling floats
8 (Johnson et al., 2010; Riser and Johnson, 2008) fabricated at the University of Washington. The
9 floats were equipped with In Situ Ultraviolet Spectrophotometer (ISUS) optical nitrate sensors
10 (Johnson et al., 2013) produced at MBARI and Aanderaa 3830 and 4330 (float 7663) optical O₂
11 sensors (Tengberg et al., 2006). Nitrate concentrations were computed from the UV spectra
12 measured by the ISUS sensor with the TCSS algorithm (Sakamoto et al., 2009). These floats
13 were typically set to profile from 1000 m to the surface at 5-day intervals. An array of floats
14 (Fig. S1; Table S1) has been operating since late 2009 near BATS, giving five float years of data
15 for analysis that cover four full annual cycles. Data analysis was restricted to profiles within a
16 box bounded by 34°N to 28°N and 62°W to 73°W. Nitrate and O₂ data were quality controlled
17 before analysis (SI 1.2; Fig. S2).

18

19 Preformed nitrate was calculated from O₂ and nitrate concentrations and computed O₂ solubility
20 at *in situ* temperature and salinity, (O₂)_{Sol}, as Preformed Nitrate = [NO₃⁻] - 16/150 x [(O₂)_{Sol} -
21 (O₂)]. For comparison with the float data, annual preformed nitrate and phosphate (PO₄³⁻)
22 climatologies were also computed from 20 years of BATS nitrate, phosphate, and O₂
23 concentration data (with Preformed Phosphate = [PO₄³⁻] - 1/150 x [(O₂)_{Sol} - (O₂)]). The quality

1 controlled profiling float data used in this study are permanently archived within the SOCCOM
2 (Southern Ocean Carbon and Climate Observations and Modeling) float data archive
3 at doi:10.6075/JODR2SDD.

4
5 *The oxygen isotopic composition of water column nitrate* – Samples were collected onboard the
6 R/V *Atlantic Explorer* at the BATS site (31°40'N; 64°10'W) on the following cruises: B244 in
7 March 2009, B248 in July 2009, BV44 in October 2009, B253 in December 2009, B259 in June
8 2010, B260 in July 2010, B274 in October 2011, AE1203 in February 2011, B280 in April 2012,
9 B283 in July 2012, AE1220 in August 2012, B287 in November 2012, B292 in April 2013, B295
10 in July 2013, and B299 in November 2013. Samples were also collected onboard the R/V *Knorr*
11 from BATS during US GEOTRACES Intercalibration Cruise 1 in July 2008. Seawater was
12 collected unfiltered in 60 mL HDPE Nalgene bottles that were rinsed copiously with sample
13 water prior to filling and immediately frozen at -20°C.

14
15 Seawater nitrate+nitrite concentrations were determined by reduction to nitric oxide followed by
16 nitric oxide chemiluminescence detection (Braman and Hendrix, 1989) in a configuration with a
17 detection limit of ~0.01 µM. Samples from the surface to 500 m were analyzed for nitrite
18 concentration according to the colorimetric method of Strickland and Parsons (1968), with a
19 detection limit of ~0.005 µM. Nitrate concentration alone was calculated by difference.

20
21 The $\delta^{18}\text{O}$ and $\delta^{15}\text{N}$ of nitrate were determined by the 'denitrifier' method wherein denitrifying
22 bacteria lacking nitrous oxide (N_2O) reductase quantitatively convert sample nitrate and nitrite to
23 N_2O (Casciotti et al., 2002; Sigman et al., 2001). The isotopic composition of N_2O was measured

1 by GC-IRMS using a Thermo MAT 253 mass spectrometer and a purpose-built on-line N₂O
2 extraction and purification system (Weigand et al., 2016). The international reference materials,
3 IAEA-N3 and USGS-34, were used to determine the $\delta^{18}\text{O}$ of samples relative to Vienna Standard
4 Mean Ocean Water (VSMOW): $\delta^{18}\text{O}$, in ‰ vs. VSMOW, = $\{[(^{18}\text{O}/^{16}\text{O})_{\text{sample}}/(^{18}\text{O}/^{16}\text{O})_{\text{VSMOW}}] -$
5 $1\} \times 1000$, and the $\delta^{15}\text{N}$ of samples relative to N₂ in air: $\delta^{15}\text{N}$, in ‰ vs. air, =
6 $\{[(^{15}\text{N}/^{14}\text{N})_{\text{sample}}/(^{15}\text{N}/^{14}\text{N})_{\text{air}}] - 1\} \times 1000$.

7
8 Nitrate concentration and isotope data for individual BATS cruises are reported in Fawcett et al.
9 (2015), where analytical details can also be found. A total of 220 samples from BATS were
10 analyzed for $\delta^{18}\text{O}$ and $\delta^{15}\text{N}$, 3-7 times each. The pooled standard error for $\delta^{18}\text{O}$ and $\delta^{15}\text{N}$ was
11 0.12‰ and 0.04‰, respectively, for nitrate concentrations $\geq 0.5 \mu\text{M}$ (195 samples), and 0.22‰
12 and 0.14‰, respectively, for concentrations $< 0.5 \mu\text{M}$ (25 samples). Samples with nitrate
13 concentrations below $0.2 \mu\text{M}$ were not analyzed.

14
15 Here, we report $\delta^{18}\text{O}$ (and $\delta^{15}\text{N}$) data for nitrate only, after removal of nitrite (Granger and
16 Sigman, 2009). Nitrite typically declines to unquantifiable levels by ~ 300 m at BATS and is
17 always undetectable by 500 m (Fawcett et al., 2015); samples collected deeper than 500 m were
18 thus not treated with sulfamic acid to remove nitrite. This cut-off does not drive the measured
19 upward nitrate $\delta^{18}\text{O}$ increase observed to start at roughly 500 m (see below) as verified by
20 nitrate+nitrite $\delta^{18}\text{O}$ data from the relevant depth range (Fawcett et al., 2015). The isotopic impact
21 of nitrite removal is addressed at length in Fawcett et al. (2015).

22

1 Nitrate $\delta^{18}\text{O}$ was adjusted for the effect of increasing salinity from deep to shallow waters (by
2 ~ 1.4 psu between 800 m and 200 m; following Knapp et al., 2008; Fawcett et al., 2015)
3 according to: $\delta^{18}\text{O}_{\text{NO}_3(\text{salinity corr})} = \delta^{18}\text{O}_{\text{NO}_3} - (0.52 \times (\text{sal} - \text{sal}_m))$, where $\delta^{18}\text{O}_{\text{NO}_3}$ is the measured
4 $\delta^{18}\text{O}$ of sample nitrate, ‘sal’ is the measured salinity of that sample (<http://bats.bios.edu>), and
5 ‘sal_m’ is the mean salinity at 1000 m. The factor of 0.52 is the approximate slope of the
6 relationship between seawater $\delta^{18}\text{O}$ and salinity in the upper subtropical ocean (Bigg and
7 Rohling, 2000; LeGrande and Schmidt, 2006). Hereafter, “nitrate $\delta^{18}\text{O}$ ” refers to the salinity-
8 adjusted value. This adjustment tends to lower the $\delta^{18}\text{O}$ of thermocline water nitrate relative to
9 deeper water, which yields the most conservative interpretation of the data, as described below.

10

11 The mean seasonal nitrate concentration at each depth at BATS was calculated by averaging all
12 late winter/early spring data (4 vertical profiles) and all summer/fall data (12 vertical profiles).
13 The mean seasonal nitrate $\delta^{18}\text{O}$ profiles were generated by concentration-weighted averaging of
14 all late winter/early spring profiles and all summer/fall profiles. In all cases, error was calculated
15 according to standard statistical practices. Nitrate concentration and isotope data are archived at
16 <http://www.bco-dmo.org>

17

18 *Seawater incubation experiments* – Seawater was collected from 140 m, 160 m, and 200 m at
19 BATS in February 2012 (AE1203) and August 2012 (AE1220). Samples were collected
20 unfiltered in 1 L acid-washed HDPE Nalgene bottles and stored frozen until commencement of
21 the experiments in August 2014.

22

1 Seawater was thawed at room temperature (~22°C) in the dark, after which half the volume of
2 each 1 L sample was gently vacuum filtered, first through a combusted (450°C for 5 hours) 47
3 mm diameter glass fiber filter (nominal pore size of 0.7 µm), and then through a 47 mm diameter
4 0.2 µm pore size polycarbonate filter that had been soaked and copiously rinsed with ultra high
5 purity deionized water (DIW). All filtration glassware was acid-washed and combusted at 500°C
6 for 5 hours prior to use. Despite these precautions, filtration may have introduced some level of
7 DOC contamination to the filtered aliquots (Carlson and Ducklow, 1996); however, it was the
8 unfiltered aliquots that showed nitrate drawdown (see below), and these did not undergo any of
9 the filtration steps.

10

11 For each month and depth, ~225 mL of filtered and unfiltered seawater were decanted in
12 duplicate into 250 mL acid-washed HDPE Nalgene bottles. One of each of the treatments (i.e.,
13 February or August, 140 m, 160 m, or 200 m, and filtered or unfiltered) was placed in a bench-
14 top hood that receives ambient daylight and in which the overhead light was left on (“light
15 treatments”), and the others were placed in a drawer in the lab (“dark treatments”). On each day
16 of the four-week experiment, all of the Nalgene bottles were uncapped briefly and then shaken
17 vigorously to encourage exchange of CO₂ and O₂ between sample seawater and the headspace,
18 and ensure homogeneity of subsamples (see below).

19

20 Beginning on day 1, and continuing for 28 days at intervals of 12-24 hours (early in the
21 experiment) to ~4 days (later in the experiment), 10 mL aliquots of seawater from each 250 mL
22 bottle were subsampled into acid-washed 15 mL centrifuge tubes and immediately frozen for
23 later nitrate concentration and isotope analysis. In addition, 1 mL aliquots of each treatment were

1 pipetted into acid-washed 1.5 mL cryovials to which 25 μL of glutaraldehyde (Grade I, Sigma-
2 Aldrich) was added (1% v/v). Cryovials were gently agitated and incubated at 4°C for 1-2 hours
3 to allow the fixative to bind to cellular components, then frozen at -80°C for later flow
4 cytometric analysis.

5

6 Seawater nitrate concentration and $\delta^{18}\text{O}$ (and $\delta^{15}\text{N}$) for each time-point of the experiment were
7 analyzed as described above. We note that volume restrictions precluded the removal of nitrite
8 from samples, such that the reported nitrate concentration and isotope data are more accurately a
9 measure of the nitrate+nitrite pool.

10

11 Microbial community composition and cell abundance at each time-point was determined by
12 flow cytometric analysis of 250 μL of glutaraldehyde-preserved seawater. Heterotrophic bacteria
13 were identified by nucleic acid staining with SYBR Green I (1:7500) according to Marie et al.
14 (1997) and Gasol and Del Giorgio (2000). In brief, a blue laser (488 nm) was used for excitation
15 of SYBR Green I-stained and autofluorescent cells. Heterotrophic bacteria were discriminated
16 from the picoautotrophic cyanobacteria, *Prochlorococcus*, by gating the cell populations using
17 side scatter, green fluorescence (emission detection at 533 nm; indicative of relative nucleic acid
18 content), and red fluorescence (emission detection >670 nm; indicative of chlorophyll content).
19 Samples were analyzed using a BD Accuri C6 flow cytometer at a flow rate of 35 $\mu\text{L min}^{-1}$ and
20 with a core diameter of 16 μm . Polystyrene-based latex beads (0.91 μm) were used to assess
21 instrument performance and standardize scatter and fluorescence measurements.

22

23 **Results**

1 *Float data* – O₂ and nitrate concentrations for the upper 300 m of the water column measured in
2 2010-2013 by the profiling float array are shown in Fig. 2a-b. Deep, wind-driven mixing is
3 evident in the homogenization of the upper water column O₂ concentration and brief increase in
4 the surface nitrate concentration in late winter/early spring of 2010 and 2011. In 2012 and 2013,
5 the gradient in O₂ concentration and lack of an increase in surface nitrate suggest that spring
6 mixing did not penetrate as deeply as in the previous two years. Billheimer and Talley (2013)
7 note a near cessation of Eighteen Degree Water formation in 2012, which is reflected in these
8 float data. In the summer, thermal stratification of the upper water column sets in, the available
9 nitrate is rapidly consumed by phytoplankton, and O₂ is produced in the euphotic zone. The
10 decrease in O₂ below the euphotic zone is due to its consumption by heterotrophic bacteria
11 during the decomposition of organic matter sinking out of surface waters.

12

13 From the float-derived O₂ and nitrate concentration data, the concentration of preformed nitrate
14 was calculated (Fig. 2c). The preformed nitrate parameter was originally defined in order to
15 quantify the nitrate entering the deep ocean dissolved in the ventilating water, as opposed to
16 “regenerated” nitrate that is added to the subsurface by organic matter decomposition. However,
17 the partitioning of nitrate into these two origins is calculated based on the assumption of
18 decomposition of organic matter with the elemental stoichiometry of Redfieldian marine biomass
19 (i.e., C/N/P/-O₂ = 106/16/1/-150) (Anderson, 1995; Anderson and Sarmiento, 1994). Thus,
20 changes in preformed nitrate, despite its name, can result from decomposition that deviates from
21 Redfield stoichiometry. One could equivalently use the tracer “NO” as an indicator of non-
22 Redfieldian stoichiometric changes (Fig. S4; where “NO” (μM) = (150/16)*[NO₃⁻]_{measured} +

1 $[\text{O}_2]_{\text{measured}}$); Broecker 1974). However, the absolute value of this property in the surface mixed
2 layer interferes with the clarity of the seasonal thermocline changes.

3
4 The float-derived preformed nitrate concentrations are high in the spring and summer euphotic
5 zone (Fig. 2c). Since surface nitrate concentrations are below detection at this time (Fig. 2b), the
6 preformed nitrate maxima can be attributed to the production of O_2 above saturation by
7 phytoplankton (i.e., $[(\text{O}_2)] > [(\text{O}_2)_{\text{sol}}]$). Below the euphotic zone, a spring-to-fall decrease in
8 preformed nitrate is apparent, reaching values as low as $-2 \mu\text{M}$. This feature is biogeochemically
9 and physically separated from the surface layer, and at times penetrates deeper than 300 m.
10 Realistic variations in apparent oxygen utilization (i.e., $[(\text{O}_2)_{\text{sol}} - (\text{O}_2)]$) or the N/ $-\text{O}_2$
11 remineralization ratio for typical marine biomass are insufficient to explain a negative preformed
12 nitrate anomaly as low as $-2 \mu\text{M}$ (Emerson and Hayward, 1995; Abell et al., 2005).

13
14 The Aanderaa optode O_2 sensor used in this work is characterized by a relatively slow response
15 (Bittig et al., 2012) that will result in a bias in O_2 concentration in steep gradients. In some
16 conditions, this bias could contribute to the minimum in preformed nitrate. However, the
17 preformed nitrate minimum occurs in Eighteen Degree Water where O_2 gradients are low. For
18 this reason, we argue that biases resulting from slow sensor response time are not important (*SI*
19 *1.2*; Fig. S3).

20
21 The O_2 and nitrate concentration at 130 m and on the 26.4 isopycnal surface (which lies near 200
22 m) are shown in Fig. 3a-d. The mean rate of O_2 decline at 130 m after deep mixing is 35 ± 12
23 $\mu\text{M y}^{-1}$ (Fig. 3a). Assuming the decomposition of typical marine biomass, such a decrease calls

1 for $\sim 3.8 \mu\text{M}$ of nitrate production. The observed rate of nitrate increase is $<10\%$ of this (mean
2 nitrate production rate of $0.3 \pm 1.1 \mu\text{M y}^{-1}$; Fig. 3b; Table S2). At ~ 200 m, the mean rate of O_2
3 decline is $24 \pm 16 \mu\text{M y}^{-1}$ (Fig. 3c), and three quarters of the expected nitrate production is
4 observed ($2.2 \pm 1.1 \mu\text{M y}^{-1}$, instead of the $\sim 2.8 \mu\text{M y}^{-1}$ predicted by Redfield stoichiometry; Fig.
5 3d; Table S2).

6

7 *Oxygen isotopes of nitrate* – For both the winter/spring and summer/fall, the average $\delta^{18}\text{O}$ of
8 nitrate varies little between 1000 m and 600 m, and we observe no difference in absolute value
9 between the two seasonal profiles (Fig. 4a). In both seasons, nitrate $\delta^{18}\text{O}$ rises by more than 7‰
10 from ~ 150 m into the euphotic zone, due to isotopic fractionation during nitrate assimilation by
11 phytoplankton (Knapp et al., 2008; Fawcett et al., 2015). A weak upward nitrate $\delta^{18}\text{O}$ rise is also
12 observed in the subsurface, from 1.4‰ at 500 m to 1.7‰ at 200 m in the winter/spring, and
13 1.6‰ at 500 m to 2.1‰ at 200 m in the summer/fall. Most importantly for our study, student's t-
14 tests indicate that nitrate $\delta^{18}\text{O}$ at 200 m in the summer/fall is significantly higher than in the
15 winter/spring by an average of 0.4‰ ($p < 0.01$), 250 m ($p < 0.001$), 300 m ($p < 0.001$), and 400 m
16 ($p < 0.001$).

17

18 Like its $\delta^{18}\text{O}$, the $\delta^{15}\text{N}$ of nitrate also rises into the euphotic zone due to nitrate assimilation
19 (Fawcett et al., 2015; Fig. 4b). However, nitrate $\delta^{18}\text{O}$ and $\delta^{15}\text{N}$ have different depth gradients in
20 the subsurface. While nitrate $\delta^{18}\text{O}$ rises upward from ~ 500 m, nitrate $\delta^{15}\text{N}$ declines upward from
21 ~ 700 m to 200-150 m. The upward decline is due to the remineralization of newly fixed N,
22 which is low in $\delta^{15}\text{N}$ (Knapp et al., 2008; Fawcett et al., 2015). Also in contrast to nitrate $\delta^{18}\text{O}$,
23 neither nitrate $\delta^{15}\text{N}$ nor nitrate concentration show a seasonal change below the euphotic zone

1 (Fig. 4b,c).

2

3 *Seawater incubations* – In all but one treatment (February 200 m light), unfiltered sample nitrate
4 concentrations decreased in both the dark and the light during the experiments (Fig. 5a; Fig. S5a-
5 b), and the $\delta^{18}\text{O}$ of the remaining nitrate rose (Fig. 5b; Fig. S5c-d; the $\delta^{15}\text{N}$ of the remaining
6 nitrate also rose (not shown)). In contrast, no change was observed in the filtered samples.
7 Plotting the nitrate concentration and $\delta^{18}\text{O}$ data in “Rayleigh” space (i.e., nitrate $\delta^{18}\text{O}$ vs.
8 $\ln([\text{NO}_3^-]_{\text{measured}}/[\text{NO}_3^-]_{\text{initial}})$) for the dark experiments, which we take to be more representative
9 of the BATS subsurface than the light experiments, provides an estimate of the average oxygen
10 isotope effect for nitrate assimilation ($^{18}\epsilon$) of $3.5\text{‰} \pm 0.3\text{‰}$ (p-value < 0.001; 95% confidence
11 interval of 2.9‰ to 4.1‰; Fig. 5b; $^{18}\epsilon = (^{16}k/^{18}k - 1) \times 1000$, where ^{16}k and ^{18}k are the rate
12 coefficients of the reaction for ^{16}O - and ^{18}O -containing nitrate, respectively).

13

14 Flow cytometric analysis showed that heterotrophic bacterial abundance initially ranged from 3.5
15 $\times 10^4$ to 8.1×10^4 cells mL^{-1} , and was lowest in the deepest samples (Fig. 5c). After a ~4-day lag
16 period during which cell abundances decreased slightly, heterotrophic bacteria grew in all the
17 unfiltered incubation bottles along with the decline in nitrate concentration, increasing 3- to 5-
18 fold in abundance by day 20 to 28. *Prochlorococcus* initially comprised 5-8%, 2-7%, and 0.3-2%
19 of the total cell abundance in the unfiltered 140 m, 160 m, and 200 m seawater samples,
20 respectively, but declined to undetectable levels by day 5 in all cases. No other autotrophic (i.e.,
21 chlorophyll-containing) cells were detected in any of the experiments. For the filtered samples,
22 particle concentrations were below the quantification limit of the flow cytometric method, and no
23 increase in cell abundance was observed at any time during the experiments.

1

2 **Discussion**

3 *Evidence for subsurface remineralization of nutrient-poor organic matter*

4 A decrease in the concentration of O₂ without an increase in nitrate concentration has been
5 observed previously in the thermocline of the Sargasso Sea (Ono et al., 2001) as well as the
6 subtropical North Pacific (Abell et al., 2005). The high-resolution data from the profiling floats
7 deployed near BATS and the BATS hydrographic data from the upper 400 m indicates an even
8 greater spring-to-fall mismatch between O₂ consumption and nitrate (and phosphate) production
9 than observed by Ono et al. (Fig. 6a-c; Table S2).

10

11 This discrepancy causes a decrease in preformed nitrate (and phosphate) concentration in the
12 subsurface, indicating less than the expected amount of nitrate (and phosphate) production for
13 the amount of O₂ consumed. The decrease in preformed nitrate from early spring to fall spans the
14 water column from ~80 m to 300-400 m (Fig. 6a). This trend is mirrored in the BATS
15 hydrographic data for both nitrate and phosphate when many years of observations are combined
16 (Fig. 6b, c). Integrated from 80 and 300 m, preformed nitrate decreases by 0.88 mmol m⁻² d⁻¹
17 during the summer and early fall (Fig. S6). These data are consistent with the remineralization of
18 organic matter below the BATS euphotic zone that is rich in carbon and poor in N and P relative
19 to typical marine biomass; quantification is pursued below.

20

21 The DOC that accumulates in the summertime mixed layer at BATS is transported downward
22 upon wintertime mixed layer deepening (Carlson et al. 1994; Hansell and Carlson, 2001;
23 Goldberg et al., 2009). Its subsequent remineralization is expected to contribute to the O₂

1 consumption observed in the subsurface (Jenkins, 1982; Jenkins and Goldman, 1985; Stanley et
2 al., 2012) and possibly also to the negative preformed nitrate signal. However, seasonal data
3 from BATS show that elevated subsurface DOC concentrations resulting from wintertime mixed
4 layer deepening decline rapidly thereafter (typically within a month; Carlson et al. 1994; Hansell
5 and Carlson, 2001; Goldberg et al., 2009). The DOC decline does not overlap in time with the
6 summertime decline in O₂ or the accumulation of negative preformed nitrate (and phosphate) in
7 the subsurface (Fig. 3, Fig. 6), such that DOC remineralization cannot explain the O₂ or
8 preformed nutrient changes. Moreover, the rapidity of the subsurface DOC concentrations
9 decrease upon cessation of winter mixing begs the question of whether much of the DOC
10 concentration decline may be due to dilution by lateral and vertical exchange in the thermocline
11 and with underlying water as opposed to *in situ* remineralization. Finally, the existing data from
12 BATS suggest that the rate of surface DOC production is too low to explain the summertime
13 mixed layer DIC drawdown. Even if all of the summertime mixed layer DOC production (net
14 accumulation of 10-12 μM; Carlson et al., 1994; Hansell and Carlson, 2001) derived from excess
15 DIC fixation in the absence of nutrients, it could account for <50% of the observed DIC removal
16 (26 μM over the spring-summer-early fall; Gruber et al. 2008).

17

18 Letscher et al. (2016) made the observation that lateral transport of nutrients and carbon at non-
19 Redfieldian proportions helps to fuel NCP in the downwelling subtropical gyre regions,
20 including the Sargasso Sea surrounding BATS. The seasonal signals in nitrate, O₂ and preformed
21 nitrate that we report here should help to evaluate lateral transport on smaller scales than
22 addressed by the global ocean model used by Letscher et al. (2016). Preliminarily, we note that
23 our new nitrate data are consistent with prior data in indicating that nitrate is exceedingly scarce

1 in the surface mixed layer not only at BATS but in surrounding waters, including the waters to
2 the west and the north (Hansell and Follows, 2008; Moore et al., 2013; Jenkins et al., 2015). This
3 suggests that the calculations of Letscher et al. (2016) for the upper ~110-130 m, in isolation,
4 may not help to explain NCP and DIC drawdown in the ~30 m deep summer mixed layer.
5 Alternatively, high lateral N input to the BATS site may be occurring not as nitrate but rather as
6 DON. While we cannot attest to viability of this possibility, we are not familiar with data arguing
7 for substantial DON convergence in the surface mixed layer or euphotic zone of subtropical
8 regions such as the BATS site (Knapp et al., 2005; Hansell and Follows, 2008; Letscher et al.,
9 2013). Finally, consistent with our interpretation, Letscher et al. (2016) require non-Redfieldian
10 organic matter export to approach the observed NCP at BATS. Accordingly, while lateral
11 transport is likely critical to the resupply of nutrients to the upper waters of the subtropical gyres
12 on the large scale, we expect that the non-Redfieldian composition of exported organic matter
13 reconstructed here is fundamental to the high summertime NCP observed in the Sargasso Sea
14 and similar environments (Martiny et al., 2013; Ono et al., 2001).

15

16 *Evidence for nitrate assimilation by heterotrophic bacteria*

17 *The $\delta^{18}\text{O}$ of water column nitrate* – Laboratory and field data indicate that nitrification produces
18 nitrate with a $\delta^{18}\text{O}$ that is ~0-1‰ lower than that of the nitrate in the inflowing Antarctic
19 Intermediate Water and Subantarctic Mode Water (at 600-1200 m) (Buchwald et al., 2012;
20 Sigman et al., 2009). *In situ* nitrification accounts for a greater fraction of the nitrate at shallow
21 depths in the Sargasso Sea (Palter et al., 2005), where it would work to lower the $\delta^{18}\text{O}$ of the
22 nitrate pool (Rafter et al., 2013; Sigman et al., 2009), which should lead to an upward decrease in
23 nitrate $\delta^{18}\text{O}$ into the shallow subsurface as more regenerated, low- $\delta^{18}\text{O}$ nitrate is added. Instead,

1 nitrate $\delta^{18}\text{O}$ is observed to increase upward beginning at 400-500 m (Fig. 4a), to values $\sim 0.5\text{‰}$
2 higher than global deep nitrate, which is inconsistent with *in situ* nitrification acting alone.

3

4 A nitrate-consuming process such as nitrate assimilation is required to explain this $\delta^{18}\text{O}$ rise.
5 Nitrate assimilation by phytoplankton at the base of the euphotic zone at BATS is one
6 possibility. However, nitrate assimilation is likely limited to the upper ~ 120 m of the BATS
7 water column, and late winter deep mixing at BATS is not well-suited to propagate the signal to
8 depths greater than ~ 200 - 250 m, as any assimilation signal is held in a very low concentration of
9 nitrate near the base of the euphotic zone (Fig. 2b; Fig. 4c). As a related alternative, it is possible
10 that the subsurface $\delta^{18}\text{O}$ rise is a remnant geochemical signal of nitrate assimilation by
11 phytoplankton in higher-latitude surface waters that has been subducted into the BATS
12 thermocline. There are shallow subsurface waters in the subpolar North Atlantic with a
13 significantly elevated nitrate $\delta^{18}\text{O}$ (Marconi, 2017); however, it is not yet clear if this high
14 latitude signature can propagate and persist into the thermocline of the subtropical gyre.

15

16 While the year-round subsurface $\delta^{18}\text{O}$ elevation is potentially due to phytoplankton nitrate
17 assimilation, near BATS or further afield, the summertime increase in nitrate $\delta^{18}\text{O}$ observed at
18 200-400 m does not fit with the same explanation. Within the 200-400 m depth interval, nitrate
19 $\delta^{18}\text{O}$ increases from an average of 1.6‰ in the late spring to 2.0‰ in the summer and fall (Fig.
20 4a). The seasonal rise in $\delta^{18}\text{O}$ has the wrong sense to be explained by downward mixing or
21 wintertime subduction at higher latitudes of the signal of phytoplankton nitrate assimilation, as
22 this would drive a higher subsurface nitrate $\delta^{18}\text{O}$ in the winter and spring. Moreover, ventilation
23 of the thermocline from higher latitudes occurs over multiple years (Jenkins, 1998), masking

1 seasonal ventilation changes. Similarly, mode water formation and thermocline ventilation peak
2 in the winter (Kelly and Dong, 2013), such that any injection of nitrate with a high $\delta^{18}\text{O}$ occurs
3 in the wrong season to explain the observed seasonality.

4

5 As with phytoplankton, heterotrophic bacteria are known to fractionate the O (and N) isotopes of
6 nitrate during nitrate assimilation (Granger et al., 2010). Indeed, our dark seawater incubations
7 show this effect (Fig. 5b; Fig. S5c-d). Nitrate assimilation by heterotrophic bacteria degrading N-
8 poor organic matter, occurring *in situ*, may thus explain the spring-to-summer increase in nitrate
9 $\delta^{18}\text{O}$ between 400 m and 200 m depth at BATS. If the spring-to-fall decrease in preformed
10 nitrate and phosphate observed at BATS signals the remineralization of nutrient-poor organic
11 matter, it is within this time window that sporadic events of heterotrophic bacterial nitrate
12 assimilation should occur, and these events would work to raise the $\delta^{18}\text{O}$ of the nitrate, as we
13 observe in the BATS subsurface.

14

15 Importantly, the $\delta^{18}\text{O}$ elevation in nitrate can occur without net nitrate drawdown. This can
16 happen if the events of assimilation are alternated with the subsequent remineralization of the
17 heterotrophic biomass and the occurrence of nitrification through the summer, whenever organic
18 matter flux to the subsurface has an adequately low C/N to lead to net metabolic ammonium
19 production. In this context, the lack of a summertime rise in nitrate $\delta^{15}\text{N}$ at 200-400 m (Fig. 4b)
20 is consistent with *in situ* nitrate assimilation, as the re-nitrification of the assimilated nitrate
21 would yield no net $\delta^{15}\text{N}$ change. Moreover, the N isotope effect of heterotrophic nitrate
22 assimilation may be lower than its O isotope effect (Granger et al., 2010), such that the N
23 isotopic imprint of this process may be more difficult to detect. Finally, it is possible that any

1 weak $\delta^{15}\text{N}$ rise due to heterotrophic nitrate assimilation is overprinted by the export and
2 remineralization of particularly low- $\delta^{15}\text{N}$ N from N recycling (as well as N fixation and possibly
3 atmospheric N deposition) in the euphotic zone during the summer and fall (Fawcett et al., 2014;
4 Gobel et al., 2013; Knapp et al., 2010; Orcutt et al., 2001).

5

6 The float data suggest that approximately a quarter of the expected nitrate production is
7 “missing” at 200 m ($\sim 0.6 \mu\text{M}$; Table S2; Fig. 3c-d). Using the mean isotope effect for bacterial
8 nitrate assimilation estimated from the dark incubation experiments ($^{18}\epsilon = 3.5\text{‰} \pm 0.3\text{‰}$; Fig.
9 5b), we calculate that the 0.4‰ spring-to-fall nitrate $\delta^{18}\text{O}$ rise at 200 m requires the consumption
10 of $10 \pm 2\%$ of the ambient nitrate pool, or $0.26 \pm 0.05 \mu\text{M}$ (*SI 1.3*). This amounts to $\sim 40\%$ of the
11 “missing” nitrate, the remainder presumably being explained by the O_2 consumption associated
12 with the respiration of N-poor organic matter. One implication is that the net C/N supply ratio to
13 the heterotrophic bacterial community, calculated by averaging the float data from 130 m and
14 200 m, could be as high as 39 ± 17 (*SI 1.3*). While highly uncertain, a C/N ratio of this order is
15 consistent with N-limitation of the bacteria remineralizing carbon-rich organic matter
16 (Kirchman, 1994; Fagerbakke et al., 1996; Del Giorgio and Cole, 1998, and references therein;
17 Church, 2008), which then mechanistically justifies their assimilation of nitrate.

18

19 Shallower than 175 m, the sense of the seasonal $\delta^{18}\text{O}$ change appears to reverse, although the
20 seasonal distinction is statistically much weaker than between 200 and 400 m depth. We explain
21 these observations as the result of phytoplankton nitrate assimilation within the euphotic zone,
22 which yields substantial $\delta^{18}\text{O}$ elevation ($>5\text{‰}$ above ~ 120 m; Fig. 4a), and the downward
23 transport of this nitrate $\delta^{18}\text{O}$ elevation by mixing, which occurs dominantly in the wintertime

1 when the mixed layer deepens. We reason that without this large wintertime signal from vertical
2 mixing, a spring-to-fall $\delta^{18}\text{O}$ rise would also develop in the 100-200 m depth interval, coincident
3 with the decrease in preformed nitrate.

4
5 *Incubation of subsurface seawater* – The incubation experiments were not originally designed to
6 detect subsurface remineralization of low-N organic matter. Nevertheless, they provide an
7 unexpected complement to the nitrate $\delta^{18}\text{O}$ evidence for *in situ* heterotrophic bacterial nitrate
8 assimilation in the BATS thermocline. It is well-known that some marine heterotrophic bacteria
9 can assimilate nitrate (Allen et al., 2001). Given the observed nitrate drawdown and nitrate $\delta^{18}\text{O}$
10 rise during the experiments (Fig. 5a, b), the low background of particulate organic N in the
11 Sargasso Sea subsurface seawater, the recalcitrant nature of dissolved organic N in the
12 subsurface at BATS (Hansell and Carlson, 2001; Knapp et al., 2005), and the extremely low
13 ambient ammonium concentrations at the depths from which incubation seawater was collected
14 (Fawcett et al., 2014; Lipschultz, 2001; Treibergs et al., 2014), the evidence for nitrate
15 consumption and the increase in heterotrophic bacteria in the incubations is remarkably
16 consistent with an innate capacity for heterotrophic nitrate assimilation in the shallow subsurface
17 at BATS. Initial bacterial abundances were slightly lower than is typically observed in the BATS
18 subsurface ($4\text{-}8 \times 10^4$ cells mL^{-1} vs. $1\text{-}4 \times 10^5$ cells mL^{-1} ; Fig. 5c; Carlson and Ducklow, 1996),
19 likely due to the seawater samples having been frozen for ~ 2 years. We are nonetheless confident
20 that the heterotrophic bacteria that grew in the unfiltered seawater were present at the time of its
21 collection and are not the result of experimental contamination. The simplest pieces of support
22 for this assertion are that (1) no bacterial growth was observed in the filtered samples and (2)
23 growth was observed in every unfiltered sample, not only a subset of them. In addition, the flow

1 cytometry cytograms show the growth of bacterial populations in the unfiltered samples that
2 resemble native populations from the Sargasso Sea and other oligotrophic regions (Cavender-
3 Bares et al., 2001; Zubkov et al., 2004; 2007).

4

5 The heterotrophic nitrate assimilation rates observed in the incubations (averaging $0.04 \mu\text{M d}^{-1}$)
6 are too rapid to apply within the BATS thermocline; the entire $3.5 \mu\text{M}$ of “missing” nitrate
7 between 100-200 m (Fig. 3) could be accounted for by nitrate assimilation alone in a matter of
8 months, without even considering the effect of O_2 consumption without nitrate production (i.e.,
9 the subsurface decrease in preformed nitrate; Fig. 2c; Fig. 6a,b). Seawater incubations are
10 vulnerable to “bottle effects” that can lead to unrepresentative rates, but such incubations are
11 nevertheless useful for identifying and characterizing processes (e.g., Kirchman et al., 1991;
12 Carlson and Ducklow, 1996; Quay et al., 2010). In this vein, the key lessons from heterotrophic
13 nitrate assimilation in the incubations are (1) that the N content of the organic matter available
14 for remineralization in these subsurface waters is so low that heterotrophic bacteria are limited
15 by N, a situation that would not arise if the remineralization were dominantly of Redfieldian
16 marine biomass or its degradation products, and (2) that the resident bacteria readily turn to
17 nitrate assimilation to grow on this N-poor organic matter.

18

19 POM cannot have been the main C source to the heterotrophic bacteria in the incubations given
20 that its concentration in the Sargasso Sea thermocline is $<1 \mu\text{M C}$ and its C/N ratio is typically
21 <8 (Martiny et al., 2013). Rather, the concentration, C/N, and apparent bioavailability of DOM at
22 140-200 m depth at BATS are appropriate to explain the heterotrophic bacterial growth and
23 nitrate assimilation in the incubations, contingent on the bacterial growth efficiency (BGE; *SI*

1 1.4).

2

3 Assuming a mean bacterial C/N ratio of 5 (Gundersen et al., 2002) and a BGE of 14-49%
4 (Carlson and Ducklow, 1996; Pedler et al., 2014), we calculate that the observed bacterial nitrate
5 drawdown requires an average of 11-38 μM DOC. If we disregard the August 200 m dark
6 treatment in which the quantity of nitrate drawdown was more than double that of any other
7 treatment, this demand decreases to an average of 9-30 μM DOC. In the 100-250 m depth
8 interval at BATS, the DOC concentration varies seasonally between 50 μM (during summer and
9 autumn) and 65 μM (during late winter deep mixing) (Carlson et al., 1994). Of this,
10 approximately 10-20 μM is considered biologically available on the time-scale of hours to
11 months (Hansell, 2013; Zweifel et al., 1993). This range is roughly sufficient but on the low end
12 to explain the bulk of the nitrate consumption in our incubations unless BGE was consistently
13 high. It is possible that the mean bacterial C/N requirement over the course of our experiments
14 may have been less than 5. In a series of batch culture experiments, Vrede et al. (2002) found
15 that under conditions of C limitation and inorganic N availability, bacterial C/N declined (to as
16 low as 3.6). As with a higher BGE, a lower C/N would decrease the DOC requirement suggested
17 by the extent of bacterial nitrate consumption. We cannot rule out, however, that the bacterial
18 C/N requirement could have been >5 , as under conditions of N limitation, bacteria have been
19 shown to alter their C/N ratios (4.9 – 11; Vrede et al., 2002); if this were the case in our
20 incubations, it would imply a higher DOC requirement. Another possibility is that the labile
21 fraction of the DOC pool in the subsurface at BATS may be greater than 10-20 μM ; below we
22 describe several arguments for why this may be the case.

23

1 First, work on the role of DOC in the BATS carbon budget has focused on the ability of deep
2 mixing and thermocline ventilation to carry DOC produced in the euphotic zone into the shallow
3 subsurface, followed by net consumption of this pool. If this is the only process at work,
4 however, the stability of the deep DOC concentration through summer and fall (Carlson et al.
5 1994; Hansell and Carlson 2001) implies that there is little net respiration over this period, with
6 all net respiration occurring in the spring months immediately after deep mixing. Yet the
7 incubations suggest that the DOC pool is reactive, and no less reactive in August than in
8 February immediately following a deep mixing event. The implication is that DOC is being
9 respired continuously in the shallow subsurface, such that DOC must also be continuously
10 supplied. There is not an adequately vigorous circulation to carry N-poor DOC to these depths
11 from the euphotic zone in summer (Carlson et al., 1994). Rather, the solubilization of sinking
12 carbon-rich organic matter in the summertime subsurface could both fuel the observed O₂
13 drawdown and explain the greater inferred lability of the DOC in our incubations than is
14 suggested by the decrease in DOC concentration over the seasons at BATS or with increasing
15 ventilation age.

16

17 Second, in the presence of adequate nitrate and given sufficient time, heterotrophic bacteria may
18 consume some of the “semi-refractory” DOC pool once the labile DOC has been exhausted. A
19 number of studies suggest that surplus inorganic nutrients stimulate bacterial DOC consumption
20 (e.g., Kroer, 1993; Zweifel et al., 1993; Letscher et al. 2015), although others conclude that
21 inorganic nutrients do not enhance DOC utilization (e.g., Carlson and Ducklow, 1996;
22 Kirchman, 1990). Such experiments are typically conducted on the time-scale of days, whereas
23 our incubations lasted for four weeks, which perhaps allowed the consumption of less labile

1 DOC to begin while nitrate concentrations were still high (Del Giorgio and Cole, 1998). This is
2 supported by the findings of the year-long study by Pedler et al. (2014), who observed a diverse
3 bacterial assemblage continue to use semi-refractory DOC in seawater for the entirety of the
4 experiment once the more labile pool had been consumed. In addition, Letscher et al. (2015)
5 recently showed that surface ocean DOC is significantly more recalcitrant to remineralization by
6 the surface layer microbial community than it is to degradation by heterotrophic bacteria
7 occupying the upper mesopelagic (i.e., the depth range from which our samples were collected).
8 They also observed the concomitant consumption of nitrate by the subsurface microbial
9 community during its remineralization of surface DOC (Letscher et al. 2015).

10

11 Above, we have argued that the bioavailability of the *in situ* subsurface DOM pool at BATS
12 could explain the results of our incubations. However, we cannot completely rule out the
13 possibility that some DOC leached out of the HDPE bottles into the incubation seawater, that
14 there was some bacterial degradation of the HDPE bottles that resulted in DOC production
15 (Restrepo-Florez et al. 2014), or that freezing and thawing of sample seawater rendered the *in*
16 *situ* DOC pool more available to the bacteria. Samples for bulk DOC concentration analysis are
17 typically collected in glass bottles to ensure that potential contamination is avoided (Sharp et al.,
18 1995). However, it has been shown that polyethylene bottles do not introduce significant DOC
19 contamination to seawater left at room temperature for three months, provided the bottles are
20 soaked in 10% HCl and rinsed with sample prior to filling (Kepkay and Wells, 1992), as our
21 sample bottles were (see Materials and Methods). Moreover, to explain our observations, not
22 only would a significant quantity of DOC need to leach out of the HDPE bottles, but it would
23 need to be labile DOC, which is highly unlikely if sourced from the HDPE itself. Bacterial

1 degradation of the HDPE bottles themselves is also highly unlikely. The only available data
2 show losses of 0-1.6% of HDPE exposed to marine bacteria for months (Lobelle and Cunliffe,
3 2011; Artham et al., 2009). Moreover, it appears that very few strains of bacteria can actually
4 degrade HDPE, and there is no evidence that those bacteria that do degrade HDPE can actually
5 use it as a carbon source (Restrepo-Florez et al., 2014). Finally, there is no consensus as to the
6 effect of freezing on the lability of DOC, with studies reporting both an increase and a decrease
7 in DOM aromaticity after freezing (Chen et al., 2016; Peacock et al., 2015). Thus, while we
8 cannot rule out alteration of some fraction of the DOC pool due to sample freezing, this could
9 just as easily have decreased its lability as increased it. We conclude that our interpretation of the
10 data is far more plausible than that of marine bacteria consuming DOC sourced from
11 contamination, HDPE degradation or cryo-alteration, especially on the time scale of days and
12 weeks and at the level of the growth response observed in the bottles.

13

14 Regardless of the source of the DOC supporting nitrate drawdown in the incubations, sustained
15 heterotrophic nitrate assimilation at depth in the BATS water column is unlikely because
16 heterotrophic organisms respire most of the carbon in their diet, leaving N in excess of
17 requirements for growth even when the diet has a higher C/N ratio than their own biomass.
18 However, at times of low sinking particle flux and minimal zooplankton migration (e.g., during
19 the mid-summer), heterotrophic bacteria in the water column relying largely on an extremely
20 nutrient-poor carbon source may turn to nitrate assimilation for brief periods and/or at low levels,
21 as suggested by our data.

22

23 *Implications for the upper ocean carbon budget at BATS* – Above, we present three lines of

1 evidence for the remineralization of nutrient-poor (carbon-rich) organic matter in the BATS
2 subsurface. Below, we show that the rate of remineralization derived from the float data is
3 adequately high that, if this organic matter originates in the mixed layer, it can account for the
4 high NCP at BATS.

5

6 The profiling float data indicate that nitrate is regenerated at <10% of the expected rate at 130 m
7 and ~75% of the expected rate at 200 m (Fig. 3a-d). We interpret these changes with depth to
8 derive from a changing ratio of remineralization of GLOM and “normal” (Redfieldian) sinking
9 organic matter, with the lower rate of nitrate regeneration at 130 m indicating the greater
10 importance of GLOM in the shallower remineralization depths. Making an end-member
11 assumption that the organic matter being remineralized in the BATS subsurface (i.e., nutrient-
12 free GLOM) is essentially pure carbohydrate (as is true of TEP) with a remineralization ratio of
13 C to O₂ of 1 to 1 ($\text{CH}_2\text{O} + \text{O}_2 \leftrightarrow \text{CO}_2 + \text{H}_2\text{O}$), the decline in preformed nitrate integrated from 80
14 m to 300 m equates to a subsurface remineralization rate of $3.0 \text{ mol C m}^{-2} \text{ yr}^{-1}$ (i.e., $-0.88 \times -$
15 $150/16 \times 1/1 \times 365 \times 1/1000$). If, instead, the remineralization ratio of C to O₂ of Redfieldian
16 marine biomass is used, the decline in preformed nitrate equates to a subsurface remineralization
17 rate of $2.1 \text{ mol C m}^{-2} \text{ yr}^{-1}$ (i.e., $-0.88 \times -150/16 \times 106/150 \times 365 \times 1/1000$). Either case could
18 account for essentially all of the NCP at BATS inferred from euphotic zone O₂ production and of
19 the carbon export required to explain the DIC drawdown from the surface mixed layer (Table 1).
20 A smaller amount of subsurface GLOM remineralization would be implied by the data if it were
21 associated with heterotrophic nitrate assimilation (which our nitrate $\delta^{18}\text{O}$ data suggest may
22 explain $0.26 \mu\text{M}$ (~40%) of the seasonal preformed nitrate decline). However, even if the rate of
23 N-poor GLOM export is 60% of that calculated above ($1.3\text{-}1.8 \text{ mol C m}^{-2} \text{ yr}^{-1}$, depending on the

1 stoichiometry used), when added to previously measured POC and DOC export fluxes, it would
2 approximately match O₂ measurements of NCP and explain most of the summer mixed layer
3 DIC drawdown in the Sargasso Sea.

4

5 *Implications for subtropical ocean NCP* – GLOM export out of the euphotic zone of the
6 subtropical gyres would account for the high rates of NCP measured in these regions. If GLOM
7 is proportionally more important to export production in these regions than in high latitude and
8 upwelling systems, its export weakens the NCP gradients measured across the global ocean,
9 helping to explain the observed uniformity of NCP (Emerson, 2014 and references therein).
10 However, the shallow remineralization of GLOM has an important consequence with regard to
11 its role in the biological pump (Koeve, 2005). A large fraction of the excess CO₂ resulting from
12 subsurface remineralization of GLOM would have the opportunity to escape back to the
13 atmosphere upon wintertime deep mixing (to ~250-300 m depth near BATS; Lomas et al., 2013),
14 making it less important in ocean CO₂ storage (Fig. 1). In addition, because nutrient-poor GLOM
15 is likely metabolized dominantly by bacteria, its production and sinking is poorly suited to fuel
16 the growth of zooplankton or the upper trophic levels that rely on them, helping to explain the
17 infertility of the subtropical gyres with regard to fisheries. Our hypothesis of a form of carbon
18 export ill-suited to reach the deep ocean and its sediments can also explain why deep sediment
19 trap and sediment respiration-based reconstructions suggest much stronger spatial gradients (e.g.,
20 between the subtropical gyres and upwelling regions) than are suggested by surface ocean NCP
21 measurements (Honjo et al., 2008). At the same time, the hypothesis requires that shallow
22 sediment traps are obfuscating our large-scale view of the ocean by failing to capture significant
23 fluxes of ephemeral, less physically robust forms of organic matter.

1

2 *Conclusions* – After decades of study, the export of organic carbon from the upper subtropical
3 North Atlantic near Bermuda remains enigmatic, with regard to the nutrients that fuel it, its
4 biological and geochemical identity, and its detection in the interior. Having summarized these
5 unknowns, we propose here that there is an as-yet underappreciated contribution to export by a
6 form of low-nutrient, sinking organic carbon. In our proposal, gel-like organic matter (GLOM)
7 rich in carbon but poor in N and P, akin to TEP, is produced by phytoplankton under nutrient
8 limitation, and a portion sinks into the shallow subsurface, where it is respired by heterotrophic
9 bacteria (Fig. 1). As a source of preliminary support, we have presented evidence for the
10 subsurface remineralization of carbon-rich organic matter exported from the euphotic zone at
11 BATS, and have shown that the calculated rate of this carbon-rich export is adequate to explain
12 the observed drawdown of DIC in the summertime mixed layer.

13

14 Our proposal helps to explain a range of otherwise challenging observations from the Sargasso
15 Sea near Bermuda. In particular, the production of GLOM in surface waters, which requires a
16 minimal supply of nutrients, can explain the drawdown of mixed layer DIC and production of
17 euphotic zone O₂ at BATS that have long been recognized to be unsupported by an observed
18 source of nutrients. Moreover, because of the proposed physical characteristics of GLOM, its
19 export can explain the longstanding failure to account for the measured NCP of the mixed layer
20 and euphotic zone with sediment trap-based measurements of carbon export.

21

22 There is much to be done to test the GLOM hypothesis and, if it is correct, understand its
23 implications. As one important avenue, our incubation experiments should be repeated to include

1 measurements of DOC, POC, their chemical compositions and other properties. Another
2 approach could be the measurement of ^{14}C -labeled DOC excreted over the course of ^{14}C
3 incubation-based measurements of NPP, although we note that adequately interrogating the
4 GLOM hypothesis requires that such data be collected from many experiments conducted over
5 different seasons. It is likely (and implied by our incubations) that the remineralization of GLOM
6 in the subsurface at least partly involves its breakdown to DOC prior to oxidation to CO_2 . If so,
7 higher DOC remineralization rates are required than if the only source of DOC to the subsurface
8 is by circulation transport as dissolved (not sinking) material (Carlson et al., 1994). Accordingly,
9 studies of subsurface DOM remineralization will help in assessing the importance of GLOM.
10 Furthermore, if GLOM (such as TEP) is important in NCP, investigations will be required to
11 understand the mechanisms and sensitivities of production, survival, and export of this material
12 into the ocean interior. Such studies would have the benefit of the tools and findings of prior
13 work on TEP and similar materials (e.g., Azetsu-Scott and Passow, 2004; Beauvais et al., 2003;
14 Chin et al., 1998; Cisternas-Novoa et al., 2015; Corzo et al., 2000; Engel and Passow, 2001;
15 Engel et al., 2004; Kumar et al., 1998; Mari et al., 2017; Passow, 2002, Passow et al., 2001), but
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17

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19

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21

1 **Table and figure captions**

2

3 **Table 1.** Compilation of geochemical estimates of NCP at BATS.

4

5 **Fig. 1.** The production and fate of typical marine organic matter compared to that proposed for
6 nutrient-poor GLOM. a) In the sunlit upper ocean (the euphotic zone), phytoplankton growth
7 (“phyto”) is fueled by the upward supply of subsurface nitrate (NO_3^-). This growth fixes
8 atmospheric CO_2 dissolved in surface waters into biomass with a C:N ratio of 106:16. This
9 organic matter is transported into the subsurface through its consumption and repacking as fecal
10 pellets by zooplankton and the higher trophic levels that rely on them (e.g., fish), and by
11 aggregation and passive sinking. Here, the shaded grey area represents the idealized
12 remineralization-driven decline in the flux of sinking organic matter with depth in the water
13 column (i.e., the “Martin curve”). The decrease in organic matter flux below the euphotic zone is
14 due largely to its remineralization by heterotrophic bacteria (shown as dark grey cylinders),
15 which consumes oxygen (O_2) and produces CO_2 and nitrate in approximate ratios of O_2 :C:N of -
16 150:106:16. Much of the CO_2 produced above the base of the winter mixed layer will escape
17 back to the atmosphere during deep winter mixing, whereas excess CO_2 deriving from
18 remineralization below the winter mixed layer will be retained in the deep ocean on 100-1000
19 year timescales. A small fraction of the organic matter produced in the surface escapes
20 remineralization in the water column and is buried in deep ocean sediments, resulting in the
21 geologic sequestration of carbon, contributing to the atmospheric reservoir of O_2 and providing
22 an indicator in the sedimentary record. b) Under conditions of nutrient limitation that are
23 characteristic of the summer and fall at BATS, phytoplankton produce carbohydrates (“ CH_2O ”)
24 in surface waters that assemble to form nutrient-poor GLOM (i.e., with a C:N \gg 106:16 and C:P

1 >> 106:1). A portion of this GLOM sinks slowly into the shallow subsurface where it is respired
2 by heterotrophic bacteria, resulting in the consumption of O₂ without the production of the
3 quantity of nitrate expected from the decomposition of typical marine biomass. During times
4 when GLOM export dominates the flux of sinking organic matter, heterotrophic bacteria
5 degrading GLOM may consume nitrate to satisfy their N requirements. Because GLOM is
6 remineralized in the shallow subsurface, much of the excess CO₂ produced during its
7 decomposition will have the opportunity to escape back to the atmosphere upon wintertime deep
8 mixing rather than being stored in the deep ocean. Similarly, GLOM will not contribute to the
9 flux of organic carbon to the seabed. Finally, we expect GLOM to be dominantly metabolized by
10 heterotrophic bacteria such that it will contribute little to fueling higher trophic levels. Panels a
11 and b should be taken as end-members, with biomass-derived sinking organic matter and GLOM
12 often associated with one another in the sinking flux.

13

14 **Fig. 2.** Profiling float-derived concentrations (0-300 m) from 2010-2013 of a) oxygen; b) nitrate;
15 and c) preformed nitrate, where preformed nitrate = $[\text{NO}_3^-] - 16/150 \times [(\text{O}_2)_{\text{sol}} - (\text{O}_2)]$.

16

17 **Fig. 3.** Profiling float-derived concentrations of a) oxygen at 130 m; b) nitrate at 130 m; c)
18 oxygen at 200 m; and d) nitrate at 200 m.

19

20 **Fig. 4.** Depth profiles of a) nitrate $\delta^{18}\text{O}$ and b) nitrate concentration at BATS, averaged for the
21 late winter/early spring (February to April, 4 profiles) and summer/fall (June to December, 12
22 profiles). Note that in panel a, the 1‰ to 6‰ range of the x-axis has been expanded (left of the
23 vertical dashed line). Error bars indicate ± 1 S.D. about the mean at each depth. Nitrate
24 concentration and isotope data for individual BATS cruises averaged here are reported in

1 Fawcett et al. (2015).

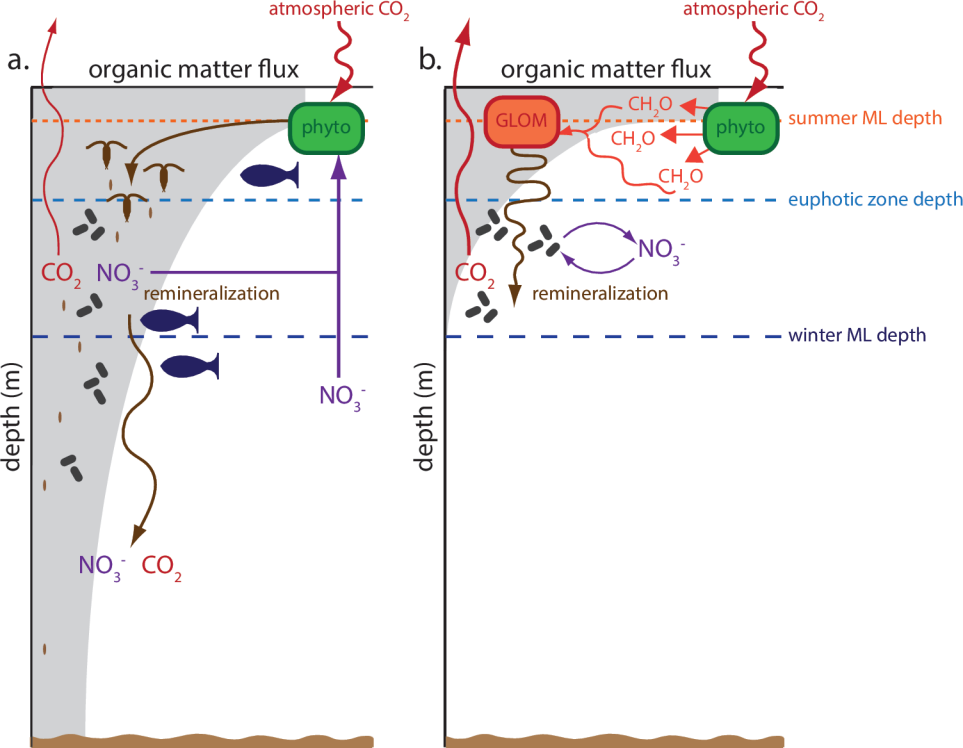
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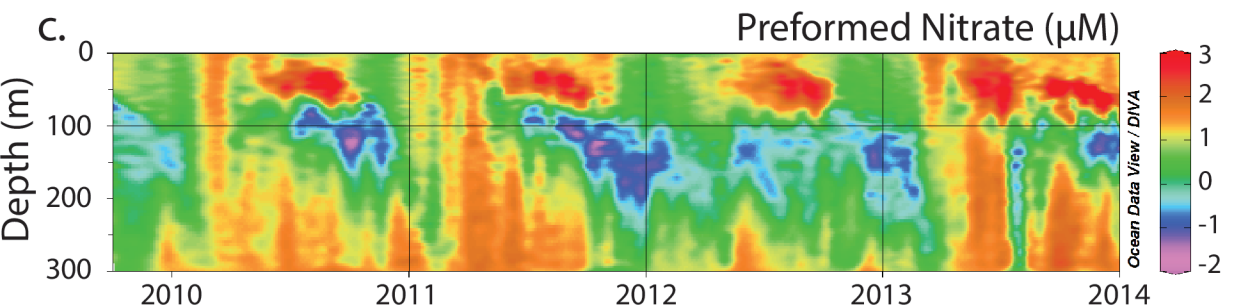
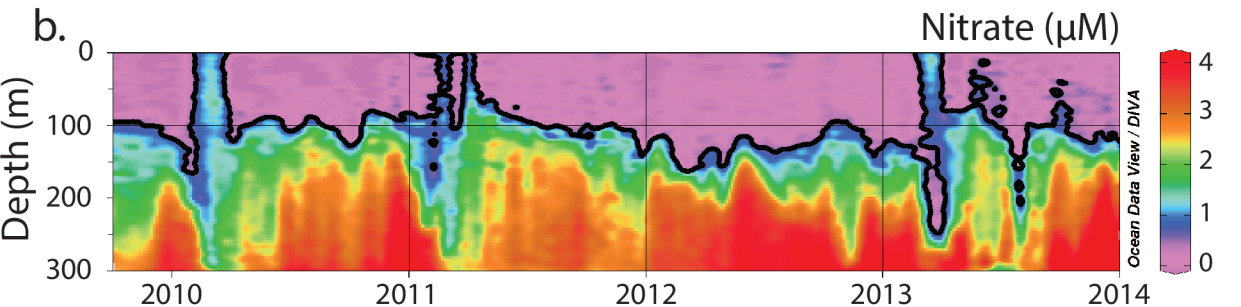
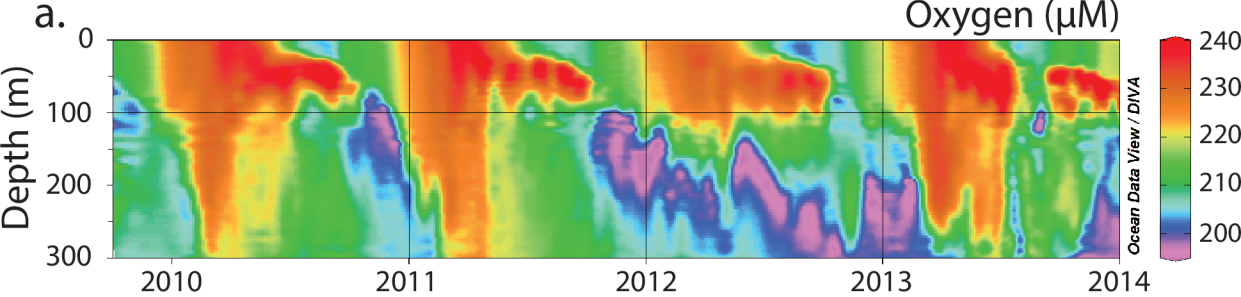
3 **Fig. 5.** Results of incubations of shallow subsurface water from the Sargasso Sea. a) Nitrate
4 concentration over the four-week dark incubation experiments in the unfiltered treatments (filled
5 symbols), with the corresponding filtered treatments indicated by the crosses and dashed lines. b)
6 Nitrate $\delta^{18}\text{O}$ in the dark experiments plotted in Rayleigh space, with the slope of the linear
7 regression providing an estimate of the average oxygen isotope effect ($^{18}\epsilon$) of heterotrophic
8 bacterial nitrate assimilation. Error bars indicate ± 1 S.D. of replicate ($n = 2-3$) measurements. c)
9 Flow cytometry counts showing the abundance of heterotrophic bacteria in the unfiltered
10 incubation bottles during the experiment. No growth was detected in the filtered samples. For
11 nitrate concentration and oxygen isotope data from the light experiments, see Fig. S5.

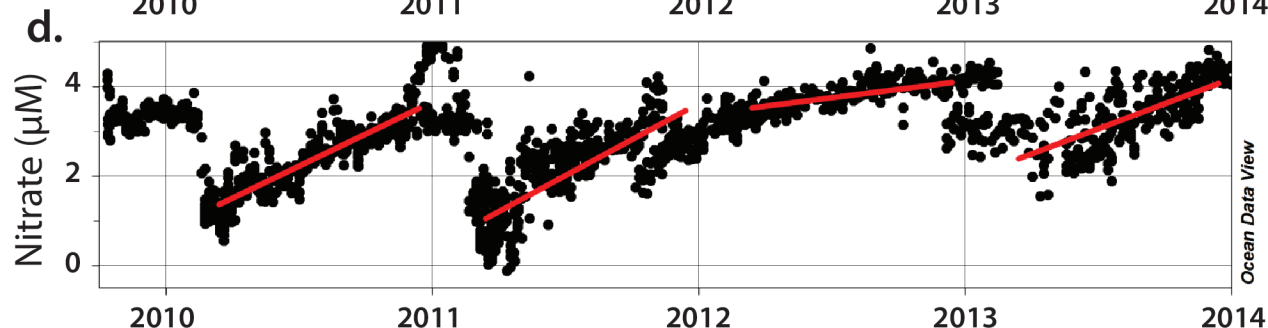
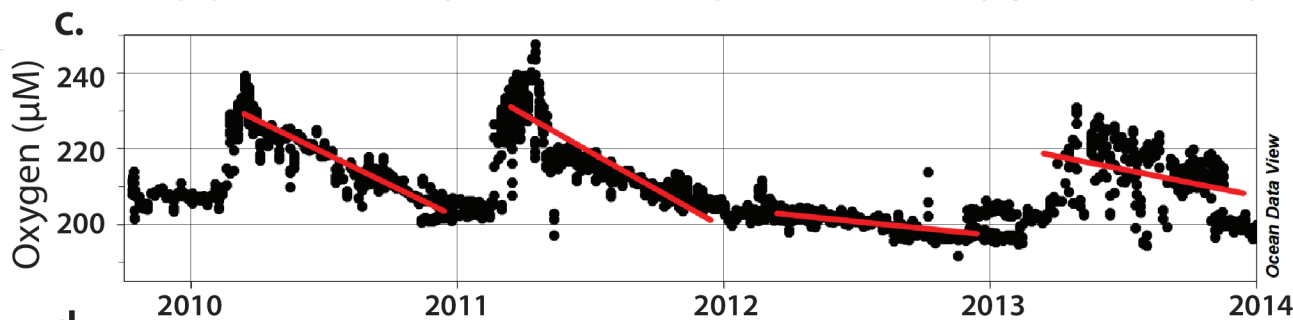
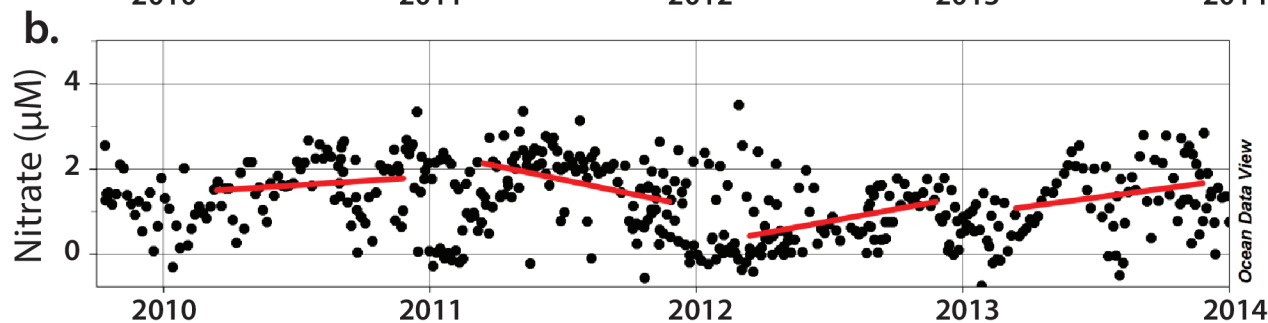
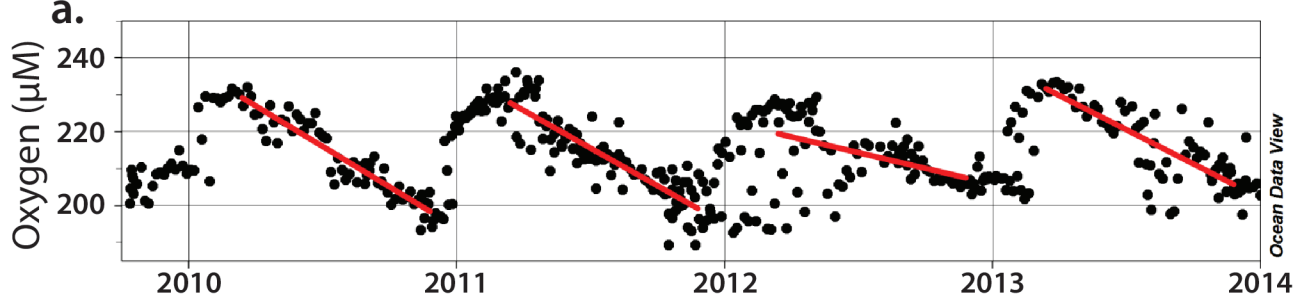
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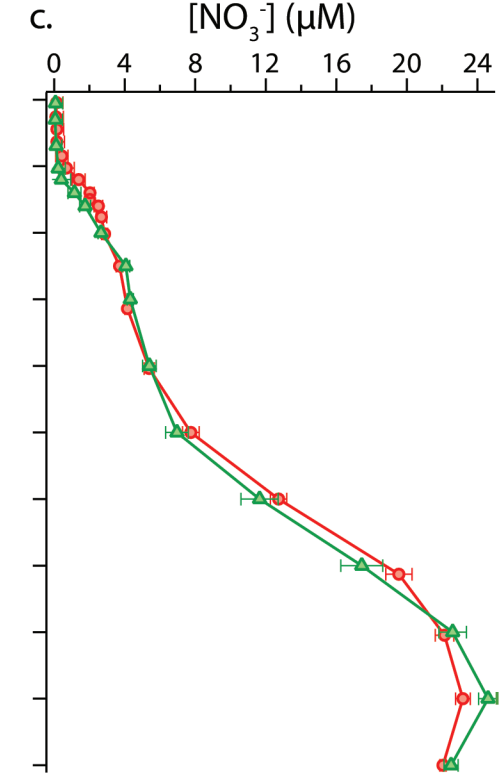
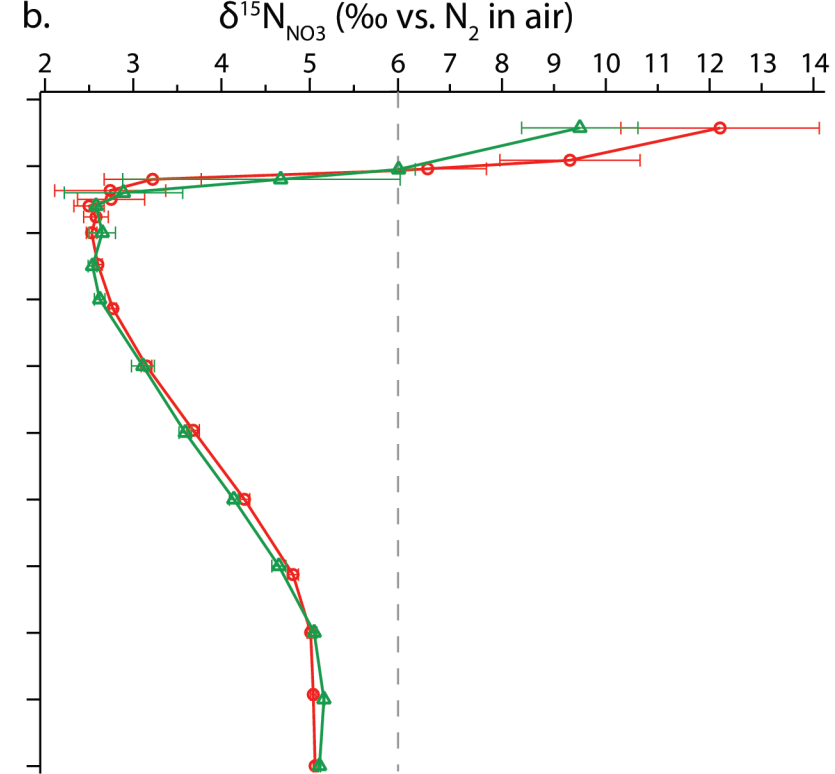
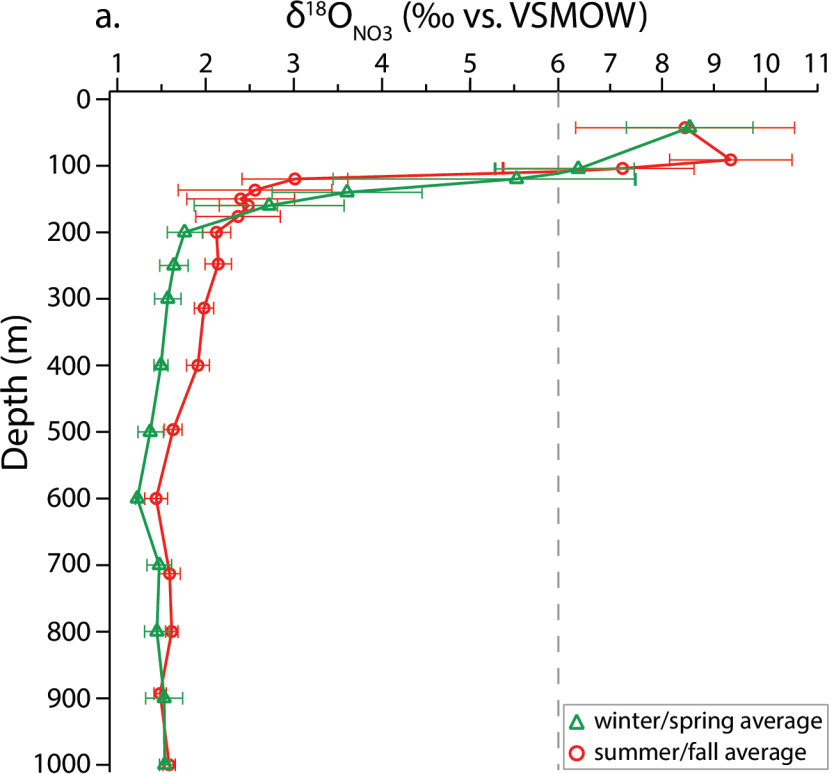
13 **Fig. 6.** a) Average preformed nitrate concentration in the upper 400 m measured by the profiling
14 float array from 2010-2013; annual average preformed b) nitrate and c) phosphate concentrations
15 in the upper 400 m derived from measurements at BATS from 1988-2008 (<http://bats.bios.edu>).
16 Contours are included to emphasize the similarity between the float data and BATS data.

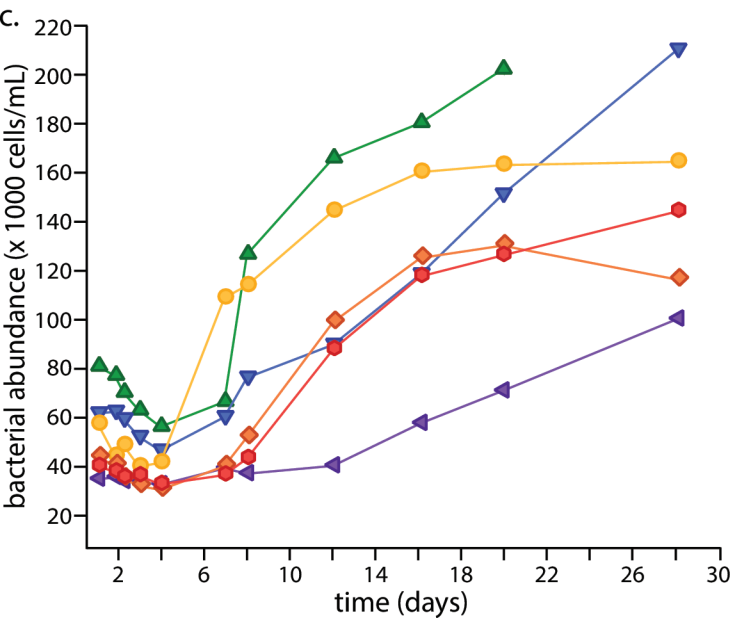
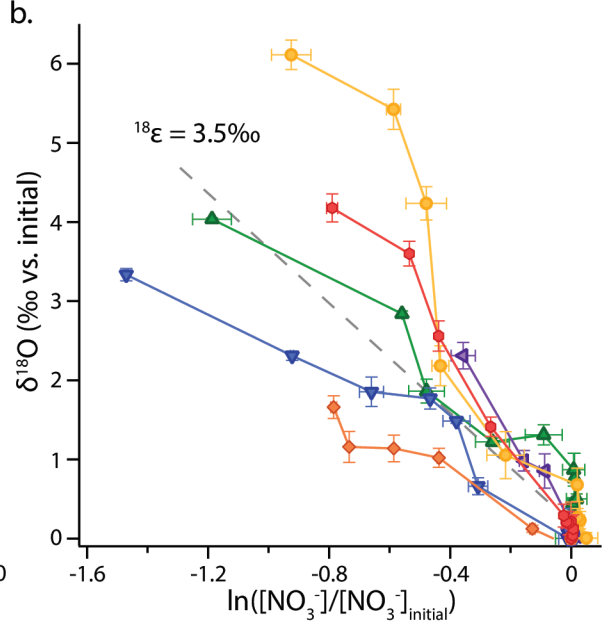
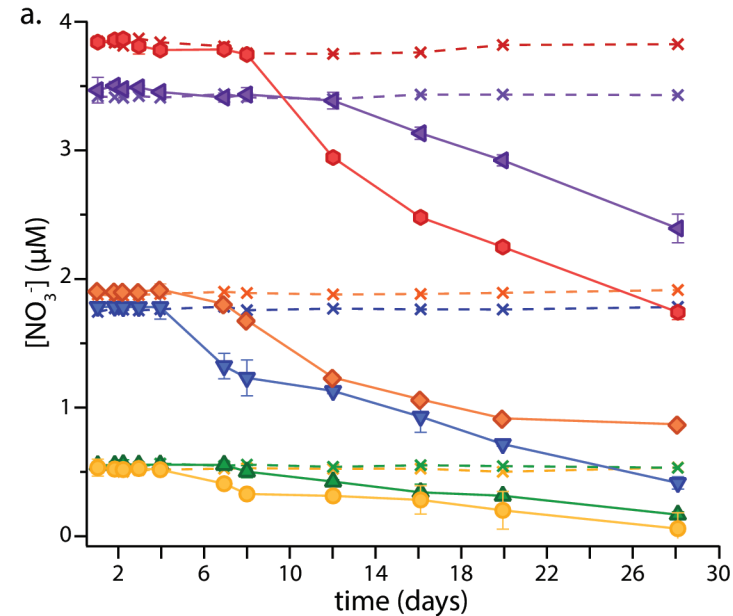
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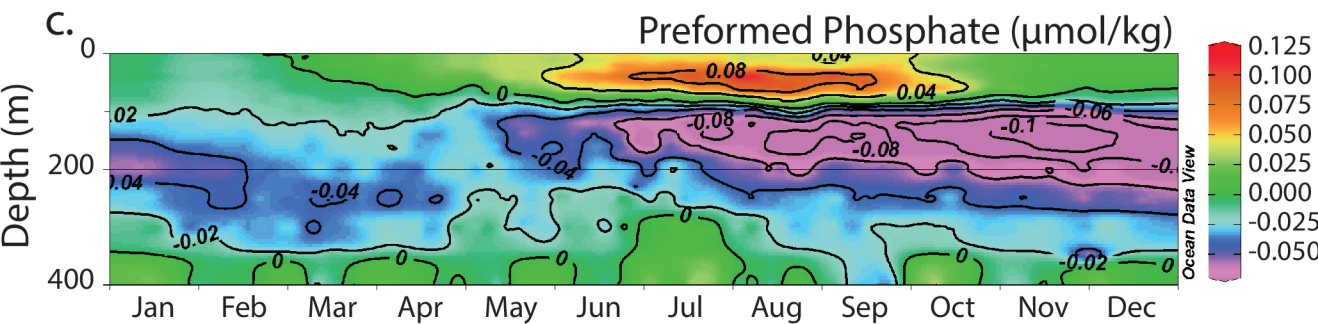
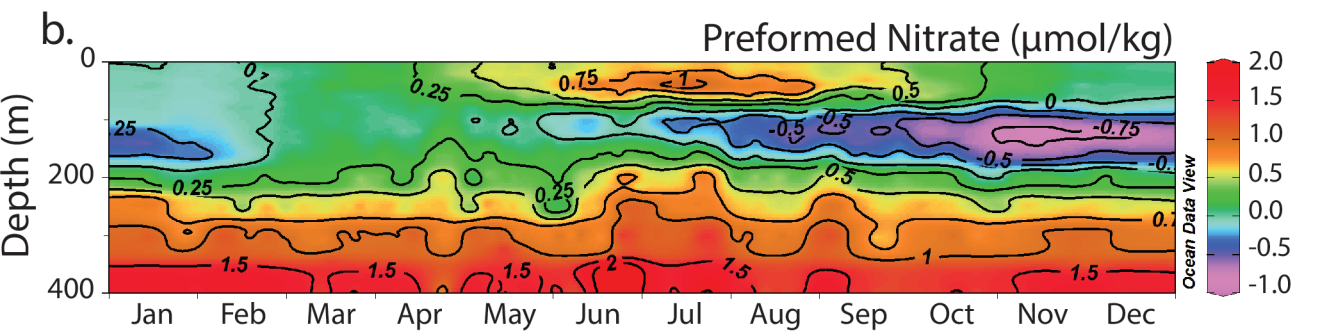
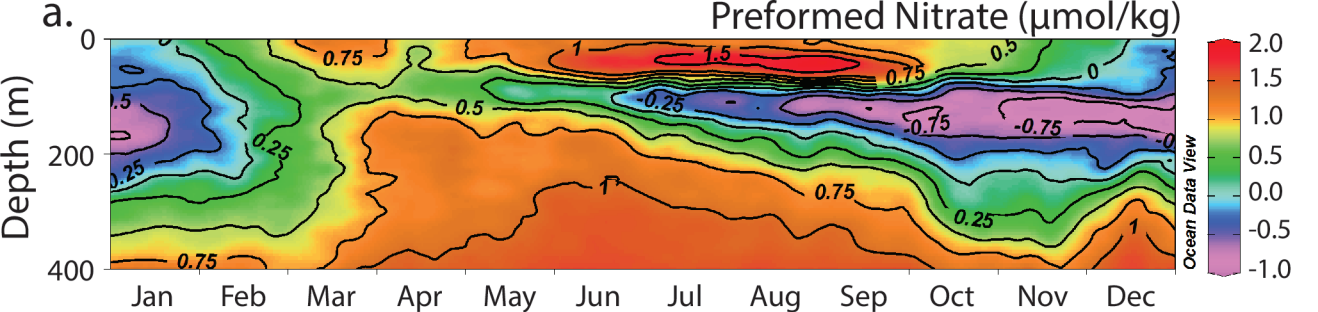


Table 1: Geochemical estimates of NCP from BATS

NCP (mol C m⁻² yr⁻¹)	Geochemical technique	Reference
3.6 ± 0.7	Oxygen utilization rate (OUR) using ³ He ventilation	Jenkins, 1980
3.4	O ₂ mass balance	Jenkins, 1982
3.3-4.8	Oxygen utilization rate (OUR) using seasonal drawdown in O ₂ stock	Jenkins and Goldman, 1985
3.6 ± 0.6	Euphotic zone O ₂ production	Spitzer and Jenkins, 1989
3.2 ^a	Seasonal (April to December) drawdown of DIC and DOC (integrated from 0-150 m)	Michaels et al, 1994
2.3 ± 0.9 ^b	Carbon isotope mass balance (mixed layer DIC)	Gruber et al., 1998
2.5 ± 0.5	Oxygen utilization rate (OUR) using ³ He ventilation	Jenkins, 1998
2.1-2.5 ^c	Oxygen utilization rate (OUR) using seasonal drawdown in O ₂ stock (100-250 m)	Hansell and Carlson, 2001
2.6-3.5	Summertime mixed layer DIC drawdown estimated from global DIC, pCO ₂ , and alkalinity	Lee, 2001
1.5 ± 0.4	Modeled remineralization rate (100-250 m) using mean annual DIC cycle	Ono et al., 2001
5.0 ± 1.0 ^d	Oxygen utilization rate (OUR) using drawdown in O ₂ stock (100-2800 m)	Jenkins and Doney, 2003
1.1-2.8	Mixed layer net O ₂ production (O ₂ /Ar)	Luz and Barkan, 2009
2.1 ± 0.5	Oxygen utilization rate (OUR) using ³ He ventilation	Stanley et al., 2012
2.8	Euphotic zone O ₂ production	Cianca et al., 2013
4.3 ± 0.9 ^e	Upward physical nitrate flux computed from ³ He flux	Stanley et al., 2015
3.0 ± 1.0	Average of estimates compiled in Table 1	
3.8 ± 1.2	<i>Average of available geochemical measurements</i>	<i>Emerson, 2014</i>

^aDecreases to 2.6 mol C m⁻² yr⁻¹ if only DIC is considered

^bMixed layer estimate. Increases to 3.8 mol C m⁻² yr⁻¹ if extrapolated to the euphotic zone assuming 40% of NCP occurs below the mixed layer

^cIncreases to 3.3-3.9 mol C m⁻² yr⁻¹ if the 100-400 m depth interval is considered

^dBased on numerical integration of aphotic zone oxygen consumption rates between 100 m and 2800 m

^eAssumes C:NO₃⁻ of 106:16