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**Modeling Atlantic Fe  
and PO<sub>4</sub> limitations  
on N<sub>2</sub>-fixation**

V. J. Coles and  
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# Modeling the impact of iron and phosphorus limitations on nitrogen fixation in the Atlantic Ocean

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## Abstract

The overarching goal of this study is to simulate subsurface  $N^*$  (sensu, Gruber and Sarmiento, 1997) anomaly patterns in the North Atlantic Ocean and determine the basin wide rates of  $N_2$  fixation that are required to do so. We present results from an Atlantic implementation of a coupled physical-biogeochemical model that includes an explicit, dynamic representation of  $N_2$  fixation with light, nitrogen, phosphorus and iron limitations, and variable stoichiometric ratios. The model is able to reproduce nitrogen, phosphorus and iron concentration variability to first order. The latter is achieved by incorporating iron deposition directly into the model's detritus compartment which allows the model to reproduce sharp near surface gradients in dissolved iron concentration off the west coast of Africa and deep dissolved iron concentrations that have been observed in recent observational studies. The model can reproduce the large scale  $N^*$  anomaly patterns but requires relatively high rates of surface nitrogen fixation to do so ( $1.8 \times 10^{12}$  moles  $N yr^{-1}$  from  $10^\circ N$ – $30^\circ N$ ,  $3.4 \times 10^{12}$  moles  $N yr^{-1}$  from  $25^\circ S$ – $65^\circ N$ ). In the model the surface nitrogen fixation rate patterns are not co-located with subsurface gradients in  $N^*$ . Rather, the fixed nitrogen is advected away from its source prior to generating a subsurface  $N^*$  anomaly. Changes in the phosphorus remineralization rate (relative to nitrogen) linearly determine the surface nitrogen fixation rate because they change the degree of phosphorus limitation, which is the dominant limitation in the Atlantic. Phosphorus remineralization rate must be increased by about a factor of 2 (relative to nitrogen) in order to generate subsurface  $N^*$  anomalies that are comparable to the observations. We conclude that  $N_2$  fixation rate estimates for the Atlantic (and globally) may need to be revised upward, which will help resolve imbalances in the global nitrogen budget suggested by Codispoti et al. (2001) and Codispoti (2006)<sup>1</sup>.

<sup>1</sup>Codispoti, L. A.: Implications of an oceanic fixed nitrogen sink that exceeds 400 Tg N/A for homeostasis in the fixed-nitrogen inventory, Biogeosci. Discuss., in review, 2006.

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## 1 Introduction

Oceanic uptake of anthropogenic CO<sub>2</sub> is driven by a combination of physical processes, i.e., the “solubility pump” (Doney et al., 2004), and biologically mediated uptake, i.e., the “biological pump” (Sarmiento and Bender, 1994). In a steady-state ocean where biogeochemical cycling occurs in Redfield proportions (Redfield et al., 1963), biologically-driven export does not result in a significant net carbon export to the deep ocean (Broecker, 1991). However, there are two very important processes in the ocean that can result in non-Redfield elemental cycling and net carbon export. These are atmospheric dinitrogen (N<sub>2</sub>) fixation and denitrification. In regions where carbon fixation and export are limited by the availability of nitrogen (N), the addition of fixed N from the atmosphere due to N<sub>2</sub>-fixation can allow carbon fixation to proceed and a net carbon export can result. Estimates suggest that this process can drive a significant fraction of the total, global carbon export (Hood et al., 2000). The removal of N from the ocean due to denitrification has the opposite effect. Thus, it may be very important to represent the impact of these processes in prognostic regional and global ocean carbon cycle models. However, the magnitudes of these sources and sinks are still poorly constrained by observations (Codispoti et al., 2001; Gruber and Sarmiento, 2002; Capone et al., 2005; Codispoti, 2006<sup>1</sup>; Gruber, 2006). Because nitrogen fixation and denitrification determine the total fixed nitrogen inventory in the oceans both locally (on seasonal and annual timescales) and globally (on geological timescales) they also determine the degree to which the oceans are either nitrogen or phosphorus limited (Tyrrell, 1999).

The factors that control pelagic nitrogen fixation are still not fully understood. Rueter (1982) and Rueter et al. (1992) argued that iron (Fe) is a primary factor limiting N<sub>2</sub>-fixation due to the high Fe requirement of the nitrogenase enzyme which is required for nitrogen fixation. This idea is supported by the early experimental work of Paerl et al. (1994) and the general global correspondence between regions of high iron/dust deposition and high N<sub>2</sub>-fixation, e.g., N<sub>2</sub>-fixation rates are generally higher in the Atlantic than in the Pacific (Capone et al., 1997). Similarly, Walsh and Steidinger (2001)

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argue that Fe deposition events stimulate *Trichodesmium* blooms and N<sub>2</sub>-fixation in the Gulf of Mexico. In contrast, the in situ stoichiometric studies of Sañudo-Wilhelmy et al. (2001) suggest that phosphorus (P) limits *Trichodesmium* growth and N<sub>2</sub>-fixation in the central north Atlantic. The potential for P limitation is supported by the fact that inputs of N to the surface ocean due to N<sub>2</sub>-fixation are not accompanied by a stoichiometric equivalent input of P. Thus, N<sub>2</sub>-fixation tends to increase the N:P ratio in the water which leads to phosphorus limitation. It appears that these conflicting results have been reconciled at some level by Mills et al. (2004) who showed that Fe and P co-limit nitrogen fixation in the eastern tropical Atlantic, implying that both P and Fe limitations can be important, and that switching from one to another will likely occur depending upon local availability and recycling of Fe and P. Finally, it should also be noted that light availability and temperature are clearly important factors controlling N<sub>2</sub>-fixation (Capone et al., 1997; Coles et al., 2004; Hood et al., 2004a), i.e., significant rates are observed only in warm tropical and subtropical waters and generally under stratified conditions where mean irradiance levels in the mixed layer are very high.

Only a handful of modeling studies have been undertaken that include explicit representations of N<sub>2</sub>-fixation along with N, P and/or Fe limitations. Tyrrell (1999) developed a simplified box model with both nitrogen and phosphorus cycling where the N<sub>2</sub>-fixation rate is a function of the supply ratio of inorganic nitrogen and phosphorus. In a three-dimensional model developed by Neumann (2000) N<sub>2</sub>-fixation rate depends upon phosphorus supply, temperature and irradiance. The models of Tyrrell (1999) and Neumann (2000) can distinguish between nitrogen and phosphorus control of phytoplankton and diazotroph growth. However, both assume a fixed Redfield stoichiometry. In contrast, Fennel, et al. (2002) developed a 1-dimensional model with N<sub>2</sub>-fixation and both N and P limitations that allows for different stoichiometric ratios in different organic matter compartments (i.e., high N:P ratio for diazotrophs) and differential export of N and P from the upper ocean. However, they did not provide any mechanism for enhancing phosphorus supply to the surface. As a result, the diazotrophs in this model form a subsurface maximum associated with the phosphocline rather than growing

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5 near the surface. In two related global modeling efforts Moore et al. (2002a; 2002b) and Moore et al. (2004) included diazotrophs in a multi-species, multi-element biogeochemical model with N, P and Fe limitations. The large-scale spatial patterns of diazotroph biomass and N<sub>2</sub>-fixation in these studies suggest that they are determined primarily by mixed-layer depth and light, and/or temperature (as in Hood et al., 2004a). However, Moore et al. (2002b) and Moore et al. (2004) report widespread Fe and P limitation, with the latter particularly pronounced in the Atlantic in Moore et al. (2004) where the N/P ratio is fixed.

10 Clearly, there are some important questions that have not been fully addressed with these models. The fact that N<sub>2</sub>-fixation naturally leads to P limitation presents a serious dilemma in modeling studies, i.e., some means of supplying a stoichiometric equivalent amount of P to near-surface waters must be invoked in models in order to allow N<sub>2</sub>-fixation to proceed at significant rates. This problem becomes particularly significant if one attempts to reproduce the high rates of pelagic N<sub>2</sub>-fixation that have been reported in recent studies (Gruber and Sarmiento, 1997, 2002; Capone et al., 2005; Gruber, 2006). One potential mechanism for supplying additional phosphorus to the surface is “phosphorus mining”, i.e., the idea that large diazotrophs like *Trichodesmium* can migrate down to the phosphocline periodically to satisfy their phosphorus requirement (Karl et al., 1992). Another, perhaps more likely, mechanism is that phosphorus availability can be maintained at the surface to support diazotrophic growth by simply recycling it faster than nitrogen (Wu et al., 2000; Hood et al., 2006<sup>2</sup>). In addition, Dyhrman (2006) has suggested that diazotrophs have access to dissolved organic phosphorus (DOP) pools that are not available to other phytoplankton species. However, none of these P supply mechanisms have been quantified (Hood et al., 2000).

25 Another important challenge is Fe-cycling. Many biogeochemical models that carry Fe do not produce realistic Fe profiles at depth (e.g., Dutkiewicz et al., 2005; but

<sup>2</sup>Hood, R. R., Laws, E. A., Armstrong, R. A., et al.: Pelagic functional group modeling: Progress, challenges and prospects, Deep-Sea Res., Part II, 3rd Special Issue on the U.S. JGOFS Synthesis and Modeling Project, accepted, 2006.

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see also Parekh et al., 2005). Finally, a major challenge that has not yet been fully addressed in prognostic modeling studies is whether or not it is possible to simultaneously reproduce surface rates of pelagic  $N_2$ -fixation (Gruber and Sarmiento, 1997, 2002; Capone et al., 2005; Gruber, 2006) and observed subsurface N:P ratio anomaly patterns (i.e.,  $N^*$ , Michaels et al., 1996; Gruber and Sarmiento, 1997; Hansell et al., 2004). This is a critical constraint for modeling because  $N^*$  may provide a spatially and temporally integrating constraint on the relatively rare observations of  $N_2$ -fixation rate, export and relative recycling rates of N and P.

In this paper we present results from a 3-dimensional, coupled, physical-biogeochemical model with a dynamic representation of  $N_2$ -fixation that also includes N, P and Fe mineral cycling and limitations. Our goal is to determine whether the  $N^*$  patterns observed in the Atlantic (e.g., Gruber and Sarmiento, 1997) can be reproduced in a numerical model that includes an explicit, dynamic representation of nitrogen fixation, and whether the distribution and biomass of diazotrophs that gives rise to a reasonable  $N^*$  signal is consistent with observations (e.g., Carpenter and Romans, 1991). We also seek to assess the sensitivity of the modeled  $N^*$  distributions to the relative rates of detrital nitrogen and phosphorus ( $D_N$  and  $D_P$ ) remineralization, and determine the degree to which the patterns of diazotroph biomass and nitrogen fixation rate are dependent on phosphorus and iron supply and distribution.

## 2 The model

### 2.1 Biogeochemical model

The biogeochemical model used in this study is a modified version of the Hood et al. (2001) model (see also Coles et al., 2004; Hood et al., 2004a) that includes iron and phosphorus cycling and limitations (Fig. 1). Here we provide a general description of the model. The equations are presented in detail along with the parameter values and sources in Appendices A and B, respectively.

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The biogeochemical model is N-based, and includes six essential components: Inorganic nutrients (DIN, DIP, DIFe), organic nutrients (DON, DOP, DOFe), detritus ( $D_N$ ,  $D_P$ ,  $D_{Fe}$ ), phytoplankton, a diazotroph patterned after *Trichodesmium*, and heterotrophs. The latter is intended to represent all heterotrophic processes, which are dominated by microbes (i.e., bacteria and protozoa). The phytoplankton, diazotroph, and heterotroph compartments have fixed stoichiometric ratios, but these ratios can be different from one another. In the simulations presented in this paper, phytoplankton and heterotrophs have Redfield P:N stoichiometry, whereas the diazotrophs are substantially more N-rich as observed (D. G. Capone, personal communication, Table 1). The Fe:N ratios for phytoplankton and heterotrophs were converted from published Fe:C ratios for these groups (Tortell et al., 1999; Fung et al., 2000, respectively) using appropriate conversion factors. Diazotrophs were parameterized to have a substantially higher Fe:N ratio (and therefore Fe requirement) than phytoplankton or heterotrophs using conversion factors from Sañudo-Wilhelmy et al. (2001).

In contrast, the inorganic nutrient, organic nutrient and detritus compartments have variable stoichiometric ratios, which are determined by excretion and egestion by heterotrophs, exudation and mortality by and of phytoplankton, diazotrophs and heterotrophs, and remineralization of dissolved and particulate organic matter (DOM and POM). Remineralization of the DOM pools is driven by heterotroph-mediated microbial consumption and degradation and also by direct remineralization to the DIM pools. Heterotroph consumption rate is determined largely by biomass, which varies in space and time and it will lead to variable-ratio excretion and egestion if the heterotrophs consume organic matter that has a different stoichiometric ratio than its own mass. In contrast, remineralization of the POM pools is driven only by direct remineralization, which allows us to fully and independently control the remineralization length scales of sinking organic  $D_N$ ,  $D_P$ ,  $D_{Fe}$ . It is assumed that the remineralization rate of  $P > Fe > N$  for dissolved organic matter and detritus (Table 1). The specific remineralization rate values used for our main run solution (Table 1) were determined a posteriori, i.e., they were tuned to reproduce observed levels and patterns of P and Fe limitation as observed

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by Mills et al. (2004) and also to give reasonable concentrations at the surface and approximately correct  $N^*$  patterns. The degree of enhancement of P and Fe recycling relative to N that was required to achieve these solutions is discussed in Sect. 3.1.4 below.

5 Iron deposition at the surface mixed layer is specified using the model-generated climatological dust fluxes from Mahowald et al. (2003); Luo et al. (2003), which take into account both wet and dry deposition. It is assumed that a fraction of this dust is iron (Table 1) and that some fraction of this iron is bioavailable following previous modeling studies (Christian et al., 2002a; Moore et al., 2002a, Table 1). This bioavailable fraction  
10 of the Fe is added directly to the  $D_{Fe}$  compartment of the biogeochemical model. Fe is added to  $D_{Fe}$  because it is logical to do so (i.e., dust is a form of detritus), and also because it results in much more realistic vertical DIFe profiles (subsurface maximum) compared to those that are obtained when Fe is by added directly to the DIFe pool (surface maximum). Following Christian et al. (2002b), Fe is “scavenged” from the  
15 DIFe pool at a rate that is proportional to the total detritus concentration (Eq. A22) using a scavenging rate specified in Table 1 and a non-dimensional multiplier specified in Eq. (A22). Our formulation differs from Christian et al. (2002b) in that we do not specify a maximum upper rate and we use a multiplier set to give  $D_{tot}$  ( $=D_N+D_P+D_{Fe}$ ) concentrations that vary approximately between 0 and 1.

## 20 2.2 Physical model

The 3-D general circulation model is the Hybrid Coordinate Ocean Model (HYCOM) (Bleck, 2002; Chassignet et al., 2003; Halliwell, 2004). This model is formulated with an arbitrary vertical coordinate that transitions from depth based coordinates at the surface to isopycnal layers in the ocean interior. (The model can also transition to  
25 sigma coordinate layers in coastal areas, however we have not used this feature in the simulations described in this paper.) This flexibility allows for enhanced vertical resolution in the mixed layer and euphotic zone while retaining the properties of an isopycnal model for advection of tracers in the ocean interior. Surface forcing occurs through mo-

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mentum transfer from atmospheric winds, and thermodynamic fluxes of freshwater and heat with the uppermost model layer. Below the surface layer, advection and diffusion occur along isopycnal surfaces, with diapycnal mixing leading to an explicit exchange of mass and tracers between layers.

Our Atlantic basin implementation stretches from 35° S to 65° N on a Mercator grid, with 24 vertical layers, and 2° spatial resolution. This version is based on a similar Miami Isopycnal Coordinate Ocean Model implementation that has been shown to appropriately simulate Atlantic ecosystem and mixed layer dynamics (Coles et al., 2004; Hood et al., 2004a). River runoff is incorporated in the simulations as a precipitation freshwater flux to the surface for the four largest Atlantic Basin rivers with modest P and Fe loads of 0.4 mmol/m<sup>3</sup> and 0.007 mmol/m<sup>3</sup> respectively. Atmospheric forcing comes from the COADS dataset (da Silva et al., 1994), and surface salinity and temperature are relaxed back to the World Ocean Atlas 1994 (WOA94) monthly values to minimize model drift (Levitus and Boyer, 1994; Levitus et al., 1994a).

The model is initialized and relaxed at the boundaries to the WOA94 temperature, salinity, nitrate, and phosphate (Conkright et al., 1994; Levitus and Boyer, 1994; Levitus et al., 1994b). Initial DFe is set proportionally to DIN using an Fe:DIN ratio of 25:1 (μmol:mol) (c.f., Fung et al., 2000). The phytoplankton, diazotroph, heterotroph, DOM and Detrital pools are initialized at low and spatially uniform values using Redfield ratios for the DOM and Detrital pools. At the southern, northern and Mediterranean Sea boundaries the inorganic nutrient concentrations are relaxed below 25 m to seasonal averages. It is particularly important to initialize and relax to the observed nutrient fields at both the northern and southern open ocean boundaries in the Atlantic because these are significant sources of low N\* water, and failing to include the low N\* Arctic inflow will tend to underestimate the amount of nitrogen fixation required to maintain the observed N\* conditions.

The physical model is spun up for 25 years before the biogeochemical model is initialized and run. The modeled N\* and nutrient distributions have a rapid initial adjustment, stabilizing after 20 years, though they continue to adjust gradually as the model physics

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and biogeochemical cycles equilibrate. The simulations presented here are shown after 30 years for the remineralization sensitivity runs (Sect. 3.5), and 60 years for all other simulations. For the latter, the nutrient concentrations in the  $N^*$  maximum are changing by less than .003% in the last five years.

### 5 2.3 Model runs and tuning procedures

The main run solution, which we refer to in the following discussion as the NSTAR run, was tuned to reproduce the observed  $N^*$  anomaly distribution and magnitude in the Atlantic as well as possible. The in situ data used for comparison with the model are described below.  $N^*$  was calculated following Gruber and Sarmiento (1997) (hereafter GS97):

$$N^* = \left( \text{NO}_3^- - 16\text{PO}_4^{-3} + 2.9 \right) \times 0.87 \quad (1)$$

with the model-generated  $N^*$  calculated by setting  $\text{NO}_3 = \text{DIN}$  and  $\text{PO}_4 = \text{DIP}$ . The model was also tuned to reproduce surface chlorophyll, and both surface and subsurface absolute DIN, DIP and DIFe concentrations (see Sect. 3.1). The primary tuning parameters were: the natural mortality rate of the diazotrophs ( $s_T$ ), which is used to modulate the nitrogen fixation rate; the sinking rate of detritus ( $w_s$ ), which modulates all nutrient export length scales simultaneously and is used to balance the inputs of nitrogen from nitrogen fixation (following the tuning procedure described in Hood et al., 2004); and the relative remineralization rates of the  $D_N$ ,  $D_P$  and  $D_{\text{Fe}}$  ( $e_{\text{DN}}$ ,  $e_{\text{DP}}$  and  $e_{\text{DFe}}$ , respectively), which separately determine the export length scales for N, P and Fe and therefore the relative availability of DIN, DIP and DIFe in the upper ocean and also the distribution of  $N^*$  at depth (see Sect. 3.5). It should be noted that these parameters are not independent because diazotroph biomass and nitrogen fixation rate (and also phytoplankton biomass and primary production rate) are also strongly influenced by  $w_s$ ,  $e_{\text{DN}}$ ,  $e_{\text{DP}}$  and  $e_{\text{DFe}}$  which determine the surface nutrient concentrations.

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### 3 Results and discussion

#### 3.1 Surface fields

The primary goal in this study is to reproduce the  $N^*$  patterns in the Atlantic (e.g., Gruber and Sarmiento 1997) with a prognostic numerical model that includes an explicit, dynamic representation of nitrogen fixation. In addition we seek to assess the sensitivity of the modeled  $N^*$  distributions to the relative rates of  $D_N$  and  $D_P$  remineralization, and also the influence of phosphorus and iron limitations on nitrogen fixation rate. However it is important to capture the gross patterns of the seasonal evolution of surface chlorophyll, as well as the surface and subsurface nutrient distributions in order to determine whether or not the biogeochemical cycles in the model are reasonable. Here we evaluate the model performance relative to climatological surface data.

Figures 2a and b show the NSTAR run surface chlorophyll concentration in spring and late summer, to be compared with Figs. 2c and d which are climatologies for the same time periods constructed from SeaWiFs satellite ocean color data from 1997–2005. The model generally reproduces the seasonal cycles of upwelling, the North Brazil Current retroflexion and the spring bloom to the far north of the domain. The model physics are consistent with other coarse resolution models, with the Gulf Stream hugging the North American shelf too far north of Cape Hatteras. This tends to advect biological quantities to the north rather than to the east in the model domain, and leads to the tongue like spring bloom, rather than the more zonal orientation in the observations. This is not an error that greatly affects the Atlantic tropics however, where the bulk of the nitrogen fixation occurs well south of the northern subtropical gyre boundary.

The model also tends to overestimate equatorial upwelling and underestimate coastal upwelling, as is generally the case for coarse ocean models, leading to excess phytoplankton biomass along the equator and low phytoplankton biomass along the coastlines (see also Hood et al., 2004a). The NSTAR run is less oligotrophic in the subtropical gyre than the observations, which may be due to insufficient export

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in this region. The model also doesn't reproduce the very high chlorophyll concentrations observed in the Amazon plume and North Brazil Current retroreflection. This latter discrepancy is, at least to some degree, due to the fact that the satellite climatology overestimates chlorophyll in this area due to the high values of colored dissolved organic matter associated with the river plume (Del Vecchio and Subramaniam 2004).

Modeled  $N_2$ -fixation rate for the spring and fall seasons in the NSTAR run is shown in Fig. 3. For comparison with our previous model results, see Fig. 7 in Hood et al. (2004a). The NSTAR run reproduces many of the same temporal and spatial patterns generated in previous runs that did not include phosphorus or iron limitations, but there are also some substantial differences. Both solutions generate elevated rates off northwest Africa at  $\sim 10^\circ N$  and in the Gulf of Guinea with markedly higher rates in spring in the latter. Both solutions also produce elevated rates in two zonal bands situated to the north and south of the equator in spring, and both have much higher rates overall in the North Atlantic in summer. In general, the comparison between the NSTAR run and our previous model results show strong similarities in the  $N_2$  fixation rate patterns in tropical waters with, perhaps, somewhat more realistic seasonality in the NSTAR run because the rates do not decrease as much in the winter (Hood et al., 2004a) and the rates are also lower in the Gulf of Guinea (Fig. 3).

However, there are some dramatic differences between the NSTAR run and our previous model results in the subtropics. For example, in late summer there is a minimum in the  $N_2$ -fixation rate in the southern Sargasso and in the Gulf of Mexico in the NSTAR run (Fig. 3b), whereas there are maxima in the rates in these regions in Hood et al. (2004a). The rates are reduced in the NSTAR run in these regions due to phosphorus limitation, which suppresses  $N_2$ -fixation over a broad area of the subtropics that extends from the Gulf of Mexico to  $\sim 30^\circ W$  (see discussion below). Another aspect of the NSTAR run that differs significantly from Hood et al. (2004a) are the high  $N_2$ -fixation rates north of  $30^\circ N$  in the late summer and Fall (Fig. 3b). Observations show *Trichodesmium* extending further north off of Africa during summer and fall (Herandez-León et al., 1999; Tyrrell et al., 2003) as in the NSTAR run. However, *Trichodesmium*

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colony concentrations and rates of  $N_2$ -fixation decline dramatically from south to north approaching Bermuda in the western Atlantic, and it is generally believed that concentrations and rates remain low further north (Carpenter and Romans, 1991; Orcutt et al., 2001). The model however, allows  $N_2$ -fixation to occur north of Bermuda in the summer and fall (Fig. 3) because this region becomes highly stratified during this time of year and depleted in DIN, i.e., ideal conditions for diazotrophic growth. Because there is a strong meridional temperature gradient in this region, the model predicted  $N_2$ -fixation rate can be reduced in this region by the addition of a simple temperature constraint on the diazotroph growth rate (e.g., following Moore et al., 2002a). We did not impose this constraint because there is evidence for elevated rates of  $N_2$ -fixation in these waters, i.e., spring time whole water incubations show significant rates of nitrogen fixation north of  $35^\circ$  N that are comparable to those generated in this region by the model in spring, suggesting that microbial diazotrophs are active in the region (J. Montoya, unpublished observations and see also Langlois et al., 2005). For this reason, we have chosen to retain the signal in the NSTAR solution. This area does not contribute significantly to basin wide  $N_2$ -fixation because of its transient nature and relatively modest rates, i.e., the region from  $30^\circ$  N northward, and east of  $30^\circ$  W represents only 4% of the basin total nitrogen fixation rate.

Because the physical model differed between this study and the solutions presented in Hood et al. (2004a) and Coles et al. (2004) we repeated the NSTAR run without phosphorus and iron limitation (NOLIM run) in order to determine specifically their influence (Fig. 4). The NOLIM run is identical to the NSTAR simulation except that N, P, and Fe are all cycled identically, with the same initial conditions, source functions, and half saturation coefficients. The nutrient ratios in the compartments are also set to 1:1:1. The NOLIM simulation has a greater basin averaged nitrogen fixation rate ( $6.5 \times 10^{12}$  mol  $N$  yr $^{-1}$ ) than the NSTAR run ( $3.4 \times 10^{12}$  mol  $N$  yr $^{-1}$ , Table 1) indicating that phosphorus and/or iron limitation does play a significant role in limiting the basin averaged rate. The primary differences between these runs is that nitrogen fixation in the western North Atlantic extends farther north in fall without P and Fe limitations (com-

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pare Figs. 3b and 4b), i.e., generating high rates throughout the subtropical gyre and in the Gulf of Mexico. This also increases chlorophyll concentrations in response to the additional input of new nitrogen without phosphorus limitation of either diazotrophy or primary production (not shown).

It is not possible to determine which of the  $N_2$ -fixation patterns in the NSTAR and NOLIM simulations is more correct based on the sparse observations (see Fig. 5 in Hood et al., 2004a), though the addition of the P and Fe limitations does appear to depress  $N_2$ -fixation in the Sargasso Sea and Gulf of Mexico more than it should. On the other hand, some aspects of the NSTAR simulation appear to be more realistic, i.e., less seasonality in the tropics, lowered rates in the Gulf of Guinea and high rates extending further north in the eastern subtropical Atlantic as discussed above. Regardless, adding phosphorus and iron cycling provides several important benefits. The addition of these nutrients allow us to estimate the basin-wide  $N_2$ -fixation rate using  $N^*$  as the constraint, rather than tuning to observed diazotroph biomass as in Coles et al. (2004) and Hood et al. (2004a). The latter approach is flawed because of the limited availability of biomass observations and the general lack of data on diazotrophs other than *Trichodesmium*. We find that tuning our model to  $N^*$  gives rates of  $N_2$ -fixation that are substantially higher than tuning to *Trichodesmium* biomass and these rates are more consistent with earlier geochemical rate estimates derived from observed  $N^*$  using inverse methods (e.g., Gruber and Sarmiento, 1997; see Sect. 3.6 and Table 1 below). As shown in Coles et al. (2004), these high geochemically-derived rates can be reproduced without phosphorus and iron limitations. However, adding these limitations has the additional advantage of allowing prognostic calculations that take into account potential changes in the availability of these nutrients that may arise in the future in response to changing atmospheric dust fluxes (e.g., Mahowald and Luo, 2003), or enhanced river runoff, for example, that might be associated with longer time-scale climate variability and global warming.

Figures 5a and b show the NSTAR model run surface iron concentration for spring and summer, and Figs. 5c and d show surface concentrations from a compilation of

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Parekh et al. (2005), with additional observations from Bergquist and Boyle (2006) and Bowie et al. (2002). The observations show all available data in small triangles, with larger circles showing data collected over three month periods in spring (Fig. 5c) and late summer (Fig. 5d). These data indicate that near-surface Fe concentrations off of the West African coast vary widely, between  $\sim 0.2$ – $2.5$  nM depending upon the cruise and time of year, with the highest concentrations occurring at about  $20^\circ$  N. The model surface fields show maxima in the same general area with surface concentrations varying from  $\sim 0.10$  nM at the northern and southern limits of the domain to  $>1.0$  nM off the coast of West Africa. These concentrations are consistent with the patterns in the modeled dust deposition fields (Luo et al., 2003; Mahowald et al., 2003) and the higher reported in situ surface concentrations (Bowie et al., 2002; C. Measures and B. Landing, unpublished data; and see also Anderson and Henderson, 2005).

To the west, the model surface concentrations are lower and broadly consistent with the observed values, near 0.2 nM with higher values along the equator, and diminishing to north and south. There is a suggestion in the data from  $35^\circ$  N,  $75^\circ$  W of decreasing surface iron concentrations from spring to summer, which is consistent with the patterns in the model, however the data density is insufficient to constrain the model beyond giving a reasonable range of concentrations. The model has lower surface iron concentrations in the northern Gulf of Mexico and in the northwestern Atlantic in general in the fall (Fig. 5b). This arises because the dominant supply of iron to these regions comes from enhanced vertical mixing in wintertime, coupled with maximum springtime dust deposition. Deposition is lowest north of  $30^\circ$  N during fall and winter, so the low model concentrations there are consistent with the deposition fields.

Figure 6 shows surface phosphate concentrations in spring and fall for the NSTAR model runs compared to the Louanchi and Najjar climatology (Louanchi and Najja, 2000; Louanchi and Najjar, 2001). This comparison suggests that the model slightly underestimates spring surface phosphorus concentrations (by  $0.1 \text{ mmol m}^{-3}$ ). However, the climatology shows higher surface phosphorus concentrations than vertical sections through the area (shown below), suggesting that the historical database may

be somewhat high, perhaps due to averaging and smoothing, relative to more recent measurements. Similar discrepancies are observed in Fall (Figs. 6b and d). In general, the model suggests that the Gulf of Mexico and central Sargasso Sea have vanishingly low concentrations, which extend over to the southwestern African coast and into the northern Gulf of Guinea, which is consistent with sections from individual cruises through these regions (see below and also Wu et al., 2000 for phosphate observations from the Sargasso Sea). Equatorial and southern hemisphere concentrations are higher as a result of upwelling, and wintertime mixing.

### 3.2 Nutrient limitation patterns

Recent observational work (Mills et al., 2004; Moore et al., 2006) has clarified the role of various macro and micronutrient limitations in modulating phytoplankton, diazotroph, and bacterial growth rates in the Atlantic Ocean. The Mills et al. (2004) study shows the general dominance of nitrogen in limiting phytoplankton growth in the tropical Atlantic (4° N–11° N, 35° W–18° W), while iron and phosphorus are observed to co-limit diazotrophy in the same region. In contrast, in a complimentary study Moore et al. (2006) demonstrate iron limitation of primary productivity during the spring bloom in the central (temperate) North Atlantic. These studies allow us to identify whether the model is reproducing the appropriate elemental (nitrogen vs. phosphorus vs. iron) limitations on growth rates. Figures 7 and 8 show the greatest limitation on phytoplankton and diazotroph growth respectively in the spring and fall seasons. Darker shades indicate more severe nutrient limitation, while the color indicates the limiting nutrient. Where the 2 (or 3) limitations are within 15% of each other, it is assumed that they are co-limiting, and they are indicated with a different shade.

Figures 7a and b show phytoplankton growth limited by nitrogen in the Atlantic in the model in both spring and late summer. Nitrogen is less limiting in upwelling zones, such as the equator and coastal North Africa. In the west, off South America, where nitrogen fixation rates are high, nitrogen is also slightly less limiting. Phytoplankton growth rate is substantially reduced (to .2 of the maximum) in the northern part of the

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domain in the subtropical gyre.

Figure 8 shows the corresponding limitation maps for spring and summer diazotroph growth, and hence  $N_2$ -fixation rate. By definition, nitrogen never limits diazotroph growth. The model predicts modest co-limitation of  $N_2$ -fixation by both iron and phosphorus over most of the tropics ( $20^\circ N$  to  $10^\circ S$ ), with stronger limitations in the western Atlantic. Phosphorus becomes a stronger limiting factor in the southern Sargasso Sea and Gulf of Mexico, particularly in fall. This is consistent with measurements from the Bermuda Atlantic Time Series (BATS) station and vicinity that reveal extremely low phosphorus concentrations in surface waters in the Sargasso Sea (Wu et al., 2000; Ammerman et al., 2003), suggesting strong potential for phosphorus limitation. Similarly low surface phosphorus concentrations are observed in the central Gulf of Mexico (J. O'Neil, personal communication). Iron limitation becomes increasingly severe to the northeast, where dust deposition rates diminish, and there is no advective supply. Thus, the model reproduces a co-limitation of diazotroph growth rate by phosphorus and iron in the tropics as observed by Mills et al. (2004), though the magnitude of the limitation may be somewhat lower than the observations suggest. This may in part be due to other factors that limit growth, such as light which are included in the model and reduce the uptake of surface nutrients but are not considered in the experimental design of Mills et al. (2004). The model also reproduces iron limitation of primary productivity during the spring bloom in the central (temperate) North Atlantic as demonstrated by Moore et al. (2006) (not shown). It is perhaps unsurprising that the limitations are reasonable because the model generates surface concentrations of nitrogen, iron and phosphorus (Figs. 5 and 6) that are close to observations, and the model limitations are based on the ambient concentrations relative to half-saturation constants specified from literature values (see Appendix B).

### 3.3 Subsurface fields

The surface ocean adjusts rapidly in the model to the seasonal and interannual variability in ecosystem dynamics and atmospheric flux fields. However, the conditions

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at depth integrate over longer time and space scales. Here we use selected vertical sections to show the model nutrient fields at depth and to validate the remineralization length scales chosen for nitrogen, phosphorus, and iron. It should be noted, however, that Bates and Hansell (2004) have shown that there is significant temporal variability in  $N^*$  at BATS, with excess nitrogen values varying by as much as a factor of 3 during different time periods, perhaps associated with the phase of the North Atlantic Oscillation (see also Hood et al., 2001). The WOCE A20 line (July, 1997) shown here (Fig. 9c) was evaluated by Bates and Hansell (2004), and shown to have lower excess nitrogen than a more recent (May 2001) section along the same meridian. Thus, the comparison of these observations with the climatological model is necessarily qualitative.

Figure 9 shows a comparison of phosphate concentrations for the NSTAR run and a WOCE meridional section along  $52^\circ W$  (A20, July 1997). The section (Fig. 9) cuts through the Sargasso Sea, and regions with high observed and simulated nitrogen fixation rates. Surface phosphate is below  $0.025 \text{ mmol m}^{-3}$  everywhere in the observations which is significantly lower than the Louanchi and Najjar (2000) climatology. The model has similarly low surface concentrations, though somewhat higher than observed in the northern and southern extremes of the section. A subsurface maximum ( $>2.2 \text{ mmol m}^{-3}$ ) with southern hemisphere origins is centered at 800 m in the observations, and also in the model, though the model maximum is weaker ( $>1.8 \text{ mmol m}^{-3}$ ). This maximum also is maintained farther to the north in the model, which will tend to cause the model to underestimate N:P ratios and  $N^*$ . However, the concentrations match to within  $0.2 \text{ mmol m}^{-3}$  throughout most of the water column. The phosphocline is shallow (100 m) at the southern edge of the section in both model and observations, it deepens to nearly 500 m at  $30^\circ N$  in the observations with a secondary seasonal nutricline higher in the water column. The model phosphocline is generally shallower, which may reflect an absence of deep mixing in the northern gyre in winter. The model also captures aspects of the impact of the mode water formed in the Gulf Stream extension on the phosphocline in the upper 400 m of the water column in the northern part of the section, and it generates a secondary near-surface seasonal phosphocline,

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though shallower than the observations.

In general, the good comparison between the modeled and observed subsurface phosphorus distributions suggests that the remineralization length scale for phosphorus in the model (122 m with a nominal sinking rate of 55 m/d, see Appendix B) is approximately correct. Further sensitivity tests of the importance of this length scale are discussed below in Sect. 3.5.

A vertical section of dissolved inorganic iron concentration along 28° W for the NSTAR run is shown in Fig. 10a. This section is chosen for comparison with a vertical section of dissolved Fe along WOCE line A16N sampled from May to August, 2003 (not shown, Measures and Landing, unpublished data; see Fig. 1 in Anderson and Henderson, 2005) it is also compared with the deep profiles from Bergquist and Boyle (2006) (Fig. 10b). The Measures and Landing section shows a very shallow ferrocline at 150–200 m between the equator and 20° N marking the transition from surface concentrations of ~1 nM to >2 nM below. This feature is also observed in the profile at 10° N, 45° W from Bergquist and Boyle (2006) (Fig. 10b), although their subsurface maximum is only half the Measures and Landing observations. Bergquist and Boyle attribute the feature to remineralization of organic matter in the low oxygen zone of the eastern Atlantic slowly ventilated shadow zone. The model reproduces this feature as long as iron is input from the atmosphere as detrital iron ( $D_{Fe}$ ) that must remineralize prior to becoming bioavailable (Fig. 10a). If iron flux at the surface enters the model dissolved iron pool directly, the surface concentrations cannot be taken up quickly enough to show a surface minimum and subsurface maximum. Nonetheless, the model ferrocline between 0–20° N is still more diffuse than the observations. The subsurface Fe concentration maximum in the model is comparable to the Measures and Landing data (~1.5 to >2 nM Fe), but higher than the maximum in the Bergquist and Boyle (2006) 10° N profile (~1–1.3 nM, Fig. 10b). The latter are from a profile located farther to the west, where presumably iron deposition rates are lower. The subsurface maximum between 0–20° N in the model is also somewhat deeper (centered at ~1000 m) than observed. In the model, this high Fe feature is primarily a result of surface dust depo-

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sition, which is co-located with the highly productive coastal and open ocean (Guinea Dome) upwelling region off western Africa. The high production rate results in greater export and higher sinking rates.

At 30° N the model profile compares well with the Landing and Measures section, showing maximum Fe concentrations below 600 m, and roughly comparable concentrations (~1.2–1.8 nM). However, these concentrations are substantially higher at depth than reported by Bergquist and Boyle (2006) at 30° N (~0.6 nM, Fig. 10b). Spatial heterogeneity in the deep iron distributions (below 1000 m) is suggested in the observations, and in the model to a lesser extent; more data are needed to resolve the robustness of these features.

### 3.4 N\* comparisons

Given the limited information on the spatial and temporal distribution of nitrogen fixation rate and biomass of diazotrophic organisms, the anomaly in the nitrogen to phosphorus ratio at depth provides, arguably, the most robust means of evaluating the correctness of the model's nitrogen fixation rate. The major caveat is that in order to make a meaningful comparison to the observed N\*, the model must form the anomaly in the right place and deposit it over a similar depth range as observed. Achieving this is a significant challenge because it requires 1) modeling the temporal and spatial variability and the magnitude of the surface N<sub>2</sub> -fixation rate, 2) exporting the N:P ratio anomaly that is created by N<sub>2</sub>-fixation with the correct remineralization length scale which varies in space and time and, 3) advecting and diffusing this anomaly correctly with the physical model so it end up in the right place. Here, we calculate N\* (according to Eq. 1 above) on the 26.5  $\sigma_\theta$  density surface in the NSTAR run (Fig. 11a), and compare it to N\* computed from the GLODAP bottle database (Key et al., 2004) using Ocean DataView (Fig. 11b). For comparison, the annually averaged nitrogen fixation rate is shown in Fig. 11c. We choose this isopycnal surface in order to compare the model fields and the updated observations with the fields presented in Gruber and Sarmiento (1997).

The model shows a similar pattern to the observations on this density surface, with

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maximum  $N^*$  concentrated in the subtropical gyre, and extending northward into the eastern subpolar gyre. However, the model  $N^*$  maximum is somewhat low, and extends too far up the coast of North America following the misplaced path of the Gulf Stream. The  $N^*$  pattern also spreads eastward along  $15^\circ N$  in both the observations and the model, though more weakly in the latter. A tongue of low  $N^*$  water entering the model from the northeast stems from the model boundary conditions which specify low N:P ratios at depth in this region, but this feature is not represented in the GLODAP database. Nonetheless, it is clear from comparisons with Fig. 11c that the surface nitrogen fixation rate pattern in the model bears little resemblance to the pattern of  $N^*$  at depth. The highest  $N_2$ -fixation rates should be spatially related to the  $N^*$  distribution if particulate export is remineralized at depth in the same region where the nitrogen fixation occurred. Instead, the highest rates of  $N_2$  fixation occur in the model, on average, just north of the equator off the coast of Africa, centered on  $20^\circ W$ , and also in the Caribbean and off the north coast of South America centered at  $15^\circ N$ ,  $55^\circ W$  (Fig. 11c). These are regions of relatively low and very homogeneous  $N^*$  on the  $26.5 \sigma_\theta$  density surface (Fig. 11a), suggesting that advection of both the fixed nitrogen at the surface, and the exported particulate nitrogen and phosphorus have a strong influence on the  $N^*$  distribution pattern. The export from the euphotic zone is concentrated in regions of tropical and equatorial upwelling, except for a local maximum collocated with the Caribbean nitrogen fixation maximum (not shown). The particles that give rise to the  $N^*$  signal must then advect horizontally over large distances and ultimately accumulate in the western tropical and subtropical north Atlantic. This conclusion is consistent with the results of Siegel and Deuser (1997) who showed that export trajectories of individual sinking particles have a very strong horizontal component, and also with our unpublished analyses of previous model results which show that export flux at depth usually bears little resemblance to surface productivity patterns.

One of the features of the model-generated  $N_2$ -fixation pattern that has not been verified by observations is the maximum in the Gulf of Guinea, and off coastal Africa, as discussed in Hood et al. (2004a). While *Trichodesmium* has been observed anecdotally

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in these regions, the magnitude of the signal in the model is comparable to the more well-known western tropical North Atlantic signal. One argument against significant  $N_2$ -fixation in these regions is the subsurface  $N^*$  signal reported in Gruber and Sarmiento (1997) (see their Figs. 3a, b). Their maps, as well as their mixing analysis comparing  $N^*$  to salinity, suggests that any modest increases in nitrogen fixation south of  $15^\circ N$  were associated with mixing of high and low  $N^*$  watermasses, and not high rates of  $N_2$  fixation. The updated observations (Fig. 11b) show local maxima in  $N^*$  associated with the Congo Plume, as well as in the eastern Gulf of Guinea, and well south of  $15^\circ N$ . The model is consistent with these patterns, and it shows that high nitrogen fixation rates in the eastern Atlantic are not inconsistent with the  $N^*$  pattern at depth.

$N^*$  signals on isopycnal and depth surfaces are not straightforward to interpret. A significant fraction of the  $N^*$  signal can be explained by passive advection and mixing on isopycnal surfaces, as the regions of sources and sinks for  $N^*$  are relatively localized. However, one must also consider that in the tropical Atlantic the  $26.5 \sigma_\theta$  isopycnal surface varies in depth from 400 m in the central subtropical gyre to nearly 100 m at the eastern boundary and in the equatorial zone. The source of  $N^*$  comes from the export of nitrogen rich organic matter from the upper ocean and this remineralization process is a function of depth primarily (Martin et al. 1987), not density. This can be seen in zonal sections of  $N^*$  (not shown) which exhibit  $N^*$  maxima at about 500 m across the basin, and if anything deepen to the east. As a result, the model and data both show organic matter remineralization occurring on different density surfaces in both zonal and meridional directions, with high  $N^*$  in the western Atlantic preferentially accumulating on shallower surfaces (e.g.,  $26.75 \sigma_\theta$ ), and the eastern Atlantic signal preferentially accumulating on deeper density surfaces (e.g.,  $27.25 \sigma_\theta$ ). Because of these differences, analysis of  $N^*$  maps drawn on isopycnal surfaces tend to further exaggerate the false impression that most of the  $N_2$ -fixation in the North Atlantic occurs in the west. This is consistent with the analysis of Hansell et al. (2004), who also noted the difference in density at which  $N^*$  patterns emerge.

According to this argument the eastern Atlantic nitrogen fixation signal should appear

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to be more important on deeper isopycnal surfaces due to the zonal uplift of the isopycnals in the east. This is confirmed in Fig. 12, which shows a comparison between the NSTAR run (Fig. 12a) and the observational data (Fig. 12b) on the  $27.03 \sigma_\theta$  surface. Indeed, neither the model nor the observations show a western intensified  $N^*$  pattern on this density surface. Note also that an additional discrepancy between the modeled and observed  $N^*$  patterns is revealed on this density surface, i.e., in NSTAR simulation shows high  $N^*$  values spreading into the Gulf of Guinea, rather than a front at  $5\text{--}10^\circ \text{N}$  as in the observations. This occurs because of errors in the circulation pattern of this coarse resolution model. The salinity field on this surface also shows a very diffusive pattern, rather than a well-developed salinity front across  $5\text{--}10^\circ \text{N}$  as observed, and the same effect occurs in the  $N^*$  distribution. Regardless, it is clearly incorrect to interpret  $N^*$  maps as being indicative of patterns of  $N_2$ -fixation sources because the signals accumulate far from their origins and in regions dictated by the circulation. This problem is compounded by the fact that  $N^*$  maps drawn on the  $26.5 \sigma_\theta$  isopycnal surface tend miss the anomaly on the eastern side of the basin, further enhancing the incorrect impression that most of the  $N_2$ -fixation in the North Atlantic occurs in the west.

The  $N^*$  anomaly plotted on vertical sections better illustrates the spatial and depth signatures of the remineralization signal. Figures 13a and b show vertical sections of  $N^*$  along  $52^\circ \text{W}$  in July from the NSTAR run compared with observations (WOCE line A20), completed in summertime. The magnitude of the  $N^*$  anomaly in the model is approximately correct ( $\sim 3.5\text{--}4.0$ ) but the maximum is somewhat deep compared to the observations. This suggests that either the sinking rate of detritus is a bit too fast, or that the remineralization of detritus is too slow in this region of the north Atlantic. The model also fails to represent the low  $N^*$  signal at 800 m in the observations, however this is probably due to shelf/slope denitrification, which is not represented in the model. The model does not have high  $N^*$  values in the upper ocean at the southern boundary of the section, along the coast of South America in the North Brazil Current retroflexion region. This suggests that the inflow from the southern hemisphere is underestimating  $N^*$  or net nitrogen fixation.

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To the east, along 28°W, we compare the modeled N\* in January (Fig. 13c) with the WOA initial conditions (Fig. 13d). Both sections show a near surface N\* minimum centered at 200 m, and initiating from the south. The model and observations both show an N\* maximum between 200 and 1500 m, however the model maximum extends farther south than the observations, as a result of the coarse resolution thermocline tropical – subtropical interactions which manifest themselves in the salinity field as discussed above. The most intense N\* signal is centered at 20°N in both the model and observations. The WOA nutrient climatology is noisier than the model, as might be expected. However, to first order the model does represent the basin wide N\* patterns.

One important caveat in these simulations is the possibility that differential remineralization of nitrogen and phosphorus can itself give rise to the N\* signal observed in the model. That is, differential remineralization could generate local positive and negative N\* anomalies in the absence of inputs of new nitrogen from N<sub>2</sub>-fixation or removal of nitrogen by denitrification simply by remineralizing N and P at different places in the water column. However, differential remineralization obviously cannot generate a net positive or negative anomaly over the entire basin all by itself. To explore these effects we carried out a simulation where N<sub>2</sub>-fixation is eliminated in the model (referred to hereafter as the NOFIX run), though the relaxation to nutrient climatologies at the northern and southern boundaries, and Straits of Gibraltar are maintained. Figure 14 shows the NOFIX N\* pattern on the 52°W section. The differential remineralization generates a strong negative N\* anomaly in the upper ~200 m of the water column due to the more rapid remineralization of phosphorus. This negative anomaly is not apparent in the observations (Fig. 13b). The differential remineralization also generates a weak positive anomaly that is distributed over the entire water column below 200 m with a maximum centered at ~500 m. Thus, the differential remineralization alone can create positive and negative anomalies, but without N<sub>2</sub>-fixation the N\* distribution is unrealistic. This does however indicate that remineralization parameterization will affect the magnitude of the basinwide nitrogen fixation signal, so the processes are not independent. In the absence of boundary condition effects, the N\* anomalies should

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sum to zero over the entire model domain. However, because we have maintained the relaxation to nutrient climatologies in the NOFIX run we expect some influences from, for example, the Mediterranean Sea outflow, which is represented in the model through nutrient relaxation to a positive  $N^*$  anomaly at the Straits of Gibraltar. This outflow probably contributes to the positive  $N^*$  anomaly at depth in Fig. 14, and it clearly influences the  $N^*$  distribution further east (not shown). We note here as an aside that representing this outflow is important to the basinwide  $N^*$  pattern, and that not including the outflow would result in an overestimation of the Atlantic Basin nitrogen fixation signal, i.e., much higher rates of  $N_2$  fixation would have to be invoked to reproduce the observed  $N^*$  anomalies without the positive  $N^*$  source from the Straits of Gibraltar.

### 3.5 Sensitivity of $N^*$ to remineralization length scale

A number of simulations are performed to examine the sensitivity of the modeled  $N^*$  and  $N_2$ -fixation rate to the remineralization length scale, which is set by the sinking rate ( $w_s$ ) and the remineralization rate ( $e_{DX}$ ) of detritus. (As we discussed in Sect. 2.1, in our model the sinking rate is assumed to be same for  $D_N$ ,  $D_P$  and  $D_{Fe}$ , whereas the remineralization rates are different with  $e_{DP} > e_{DFe} > e_{DN}$ . Thus, the differences in the remineralization length scales are dictated by differences in the detritus remineralization rates.) Specifically, in these sensitivity runs we varied the phosphorus remineralization rate so that  $D_P$  remineralizes between 1 and 3.5 times faster than nitrogen (In the NSTAR run the remineralization rate of  $D_P$  is set so that phosphorus remineralizes 2.5 times faster than nitrogen). One consequence of changing the phosphorus recycling rate is that it changes the availability of phosphorus in the surface water, which alters the nitrogen fixation rate between the runs. Figure 15a shows the results of the sensitivity experiments; as the phosphorus remineralization scale is lengthened, nitrogen fixation rate decreases, and the subsurface maximum  $N^*$  decreases. To remove the effect of the variability in the source of nitrogen, we also plot the  $N^*$  maxima normalized to the basin averaged  $N_2$ -fixation rate. The normalized  $N^*$  maximum is relatively constant across the range of different remineralization length scales, except for

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the simulation where nitrogen and phosphorus remineralization length scales are set equal to 400 m. In this simulation, nitrogen fixation rates are quite low, and are below a level that can maintain a subsurface  $N^*$  maximum. These sensitivity runs reveal the tight coupling between the phosphorus remineralization length scale, and the nitrogen fixation rate. Because nitrogen fixation is phosphorus limited over much of the subtropical gyre, the phosphorus remineralization length scale sets the nitrogen fixation rate and ultimately determines the intensity of the  $N^*$  signal. In addition, Fig. 15b shows that the maximum  $N^*$  value at  $52^\circ$  W is tightly correlated with annual nitrogen fixation rate averaged over the entire basin ( $r^2=0.99$ ). This tight correlation suggests that the relationship between maximum  $N^*$  in vertical sections and basin averaged nitrogen fixation rate can be exploited to directly estimate the nitrogen fixation rate based on observed  $N^*$ .

### 3.6 Nitrogen fixation and export rates

$N_2$ -fixation is important because of its role as an N source in the oceanic fixed nitrogen budget (Gruber and Sarmiento, 1997; Codispoti et al., 2001; Gruber and Sarmiento, 2002; Codispoti, 2006<sup>1</sup>; Gruber 2006), and also because of its potential to drive a net carbon export (Hood et al., 2000). Here, we compare basin wide estimates of nitrogen fixation derived from our NSTAR run with previous observational and model based estimates for a domain that extends from  $10^\circ$ – $30^\circ$  N (following Gruber and Sarmiento 1997) and also over our entire model domain. In addition we estimate the potential carbon sink associated with this source of new nitrogen the tropical North Atlantic.

Table 1 shows that the NSTAR run  $N_2$  fixation rate estimate for the GS97 domain in the North Atlantic is consistent with previous geochemical estimates ( $1.8 \times 10^{12}$  mol N yr<sup>-1</sup>). It should be noted, however, that a substantially lower N:P ratio (45) was assumed in the NSTAR run compared to the high value (125) used in the GS97 calculation. When the GS97 rate estimate is recalculated using an N:P of 45 a much higher basin wide rate is obtained ( $3.2 \times 10^{12}$  mol N yr<sup>-1</sup>) that is not consistent with our NSTAR run estimate for the GS97 domain (Capone et al., 2005). As we dis-

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cussed in Sect. 3.1.4, the  $N^*$  anomaly that accumulates at depth on isopycnal surfaces in the North Atlantic bears little relation to the pattern of  $N_2$  fixation at the surface. It is more appropriate then to integrate the surface nitrogen fixation over a larger domain that contributes to the subsurface signal. If we assume that the equator acts as a barrier to the surface advection of diazotrophically fixed nitrogen, then the  $N_2$  fixation rate that gives rise to the North Atlantic  $N^*$  signal in the model is substantially larger ( $2.8 \times 10^{12}$  mol N/yr) and is closer though still low relative to the recalculated GS97 estimate that is derived when an N:P ratio of 45 is used (Table 1). Thus, it appears that our model-generated rate estimate is consistent, after all, with GS97 as long as we account for the fact that contributions to the  $N^*$  anomaly can be derived from  $N_2$ -fixation that happens over a much larger domain than  $10^\circ N$ – $30^\circ N$ . In general, these results support the idea that the GS97 rate estimate significantly underestimates the basin wide  $N_2$ -fixation rate in the north Atlantic (and also globally) because the N:P ratio used in the calculation is too high (Capone et al. 2005).

Our model-estimated basin wide  $N_2$ -fixation rate derived by tuning to the observed  $N^*$  is  $\sim 2.5X$  higher than our previous rate estimates derived by tuning to *Trichodesmium* biomass (Coles et al., 2004). We now attribute this difference to significant underestimation of diazotroph biomass, i.e., the simulated *Trichodesmium* colony concentrations were, in many regions, much too low and the biomass variability was too strongly seasonal with  $N_2$ -fixation dropping to nearly zero in winter even at low latitudes (Hood et al., 2004). In addition, contributions from other diazotrophic species were ignored. The major advantage of tuning to  $N^*$  is that, presumably,  $N_2$ -fixation inputs from all potential sources are included, even if they are not occurring at exactly the right times and places. Note that the Capone et al. (2005) rate estimate for the  $10^\circ N$ – $30^\circ N$  domain is comparable to our new model-derived estimate. These higher rates are now emerging from direct measurements of  $N_2$ -fixation because recent surveys have, indeed, revealed higher and more seasonally persistent rates of  $N_2$ -fixation and because inputs from other diazotrophic species are now being accounted for. Our rate estimate is also substantially higher than that of Hansell et al. (2004) largely because they

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assumed that significant  $N_2$ -fixation occurs over a much smaller area of the Atlantic than is simulated by our model. As we have shown, contributions to the  $N^*$  anomaly in the western north Atlantic are derived from a much larger area that includes substantial contributions from the eastern side of the basin and the tropics. Finally, we note that the  $N_2$ -fixation rate estimates derived from our  $N^*$ -tuned model for the entire Atlantic domain are nearly double the GS97 estimate (Table 1), but still lower than rate estimates derived from nitrogen isotope ratios (Capone et al., 2005). Again, these results suggest that current geochemical estimates of Atlantic (and therefore global)  $N_2$ -fixation (Gruber and Sarmiento, 1997, 2002; Gruber, 2006) may be too low.

In terms of carbon, we can estimate export production for the NSTAR run by integrating the total nitrogen export out of the euphotic zone and converting to carbon units using a C:N ratio of 106:16 (i.e., assuming that all export is derived from phytoplankton-based carbon fixation). This gives  $1.16 \times 10^{15} \text{ gC yr}^{-1}$  for our entire Atlantic model domain ( $62^\circ \text{ N} - 20^\circ \text{ S}$ ). This falls between estimates based on nutrient uptake ( $0.58 \times 10^{15} \text{ gC yr}^{-1}$  from Louanchi and Najjar, 2000, for the domain  $66^\circ \text{ N} - 10^\circ \text{ S}$ , adjusted following Lee, 2001) and estimates based on carbon inventories ( $1.5 \times 10^{15} \text{ gC yr}^{-1}$  from Lee (2001), adjusted by using 2/3 of the  $40^\circ \text{ N} - 40^\circ \text{ S}$  production to bring the range closer to that of our model domain). All of these are significantly less than the export flux estimate of Laws et al. (2000) ( $7.4 \times 10^{15} \text{ gC yr}^{-1}$ , but note that this is for the entire Atlantic).

If we convert the NSTAR run total  $N_2$ -fixation rate to carbon units using a C:N ratio of 106:16 (i.e., assuming that the new nitrogen input ultimately fuels phytoplankton-based carbon fixation) it gives  $0.27 \times 10^{15} \text{ gC yr}^{-1}$ , which is 23% of the total export production integrated annually for the NSTAR run (Table 1). Thus, our model results indicate that  $N_2$ -fixation fuels a significant fraction of the export production in the Atlantic ( $62^\circ \text{ N} - 20^\circ \text{ S}$ ). This result is consistent with similar estimates reported in Karl et al. (1997), Capone et al. (1997) and Coles et al. (2004).

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## 4 Summary and conclusions

The focus of this study has been on incorporating phosphorus and iron limitations into a model with an explicit dynamic representation of N<sub>2</sub>-fixation. Although this study focuses more on the process of nitrogen fixation and its role in basin wide biogeochemistry, it is important to note that the model reproduces reasonable surface chlorophyll patterns as compared with satellite-derived distributions, as well as reasonably good agreement with surface and subsurface nitrogen, phosphorus and iron distributions. While surface concentrations of phosphorus remain somewhat low in the model relative to a nutrient climatology (Louanchi and Najjar, 2000), they are in close agreement with synoptic WOCE sections. In terms of nutrient limitation, the model reproduces a co-limitation of diazotroph growth rate by phosphorus and iron in the tropics as observed by Mills et al. (2004) and iron limitation of primary productivity during the spring bloom in the central (temperate) North Atlantic as demonstrated by Moore et al. (2006).

Iron is of particular interest given the recent emphasis on modeling iron biogeochemistry in marine systems. The representation employed here is very simple, and similar to prior efforts (e.g., Christian et al., 2002a; Moore et al., 2002a, 2002b, 2004; Christian et al., 2002b; Dutkiewicz et al., 2005; Parekh et al., 2005; Moore and Doney, 2006<sup>3</sup>). One modest change involves the addition of surface atmospheric fluxes of iron to the detrital iron pool rather than to the dissolved iron pool. This is necessary to maintain low surface iron concentrations with a sharp subsurface ferrocline that has been observed in direct measurements off of the west coast of Africa in high Fe/dust deposition regions (Measures and Landing, unpublished observations, see Anderson and Henderson, 2005; Bergquist and Boyle, 2006). This model also maintains reasonable iron concentrations at depth without setting an arbitrary concentration for ligand bound dissolved iron that is not scavenged.

<sup>3</sup>Moore, J. K. and Doney, S. C.: Iron availability limits ocean nitrogen inventory stabilizing feedbacks between marine denitrification and nitrogen fixation, *Global Biogeochem. Cycles*, submitted, 2006.

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We find that our model can reproduce subsurface N\* distributions with reasonable, though relatively high rates of surface N<sub>2</sub>-fixation. One important conclusion of this study is that surface nitrogen fixation is not co-located with subsurface gradients in N\*. Rather, the fixed nitrogen is advected far from its source prior to generating a subsurface N\* anomaly. We conclude that subsurface N\* distributions cannot be used to infer where surface nitrogen fixation occurs, except on very broad (i.e., basin wide) scales. It should be noted, however, that the model misses certain aspects of the subsurface N\* anomaly. In general, the maximum N\* anomaly in the model is deposited somewhat too deeply. There is also some evidence in the observed N\* distributions of sinks for N\* perhaps associated with African or South American continental shelf denitrification, which are not included in the model. The model also has a rather poor representation of the boundary between thermocline water with southern and northern sources, and as a result, tends to advect the northern hemisphere N\* signal too far to the south and east into the Gulf of Guinea.

Modeling the subsurface N\* distribution is particularly challenging because it involves simulating not only the magnitude and time-space pattern of the surface N<sub>2</sub>-fixation rate, but also the relative remineralization length scales of nitrogen and phosphorus in the context of advection and diffusion in 3 dimensions. Our sensitivity studies reveal that changes in the phosphorus remineralization rate (relative to nitrogen) linearly determine the surface nitrogen fixation rate. Without enhanced phosphorus remineralization (relative to nitrogen) the basin averaged nitrogen fixation rate is very modest, and cannot sustain the observed subsurface N\* maximum. Phosphorus remineralization rate has to be increased by about a factor of 2 (relative to nitrogen) to allow high enough rates of N<sub>2</sub>-fixation to sustain subsurface N\* anomalies that are comparable to the observations. We also show that the magnitude of the subsurface N\* anomaly is tightly related to the basin averaged N<sub>2</sub>-fixation rate, which provides a potential means for estimating the basin averaged rate using observed N\* anomalies.

The 10°N–30°N N<sub>2</sub>-fixation rate generated by the model when it is tuned to N\* ( $\sim 1.8 \times 10^{12}$  moles N yr<sup>-1</sup>) is comparable to geochemical estimates derived from ob-

served N\* anomalies (Gruber and Sarmiento, 1997; Gruber and Sarmiento, 2002; Gruber, 2006) even though these geochemical estimates assume a much higher N:P ratio for diazotrophs. When a more reasonable N:P ratio is adopted the geochemical estimates give much higher rates ( $3.2 \times 10^{12}$  moles N yr<sup>-1</sup>, Capone et al., 2005). Our model generates comparably high rates ( $2.8 \times 10^{12}$  moles N yr<sup>-1</sup>) when we integrate over the entire North Atlantic domain. We therefore suggest that current geochemical estimates of N<sub>2</sub>-fixation rate in the North Atlantic include contributions from diazotrophs that occur well outside of the area of integration (i.e., 10° N–30° N), which is consistent with our finding that the fixed nitrogen is advected far from its source prior to generating a subsurface N\* anomaly. Furthermore, these findings indicate that current geochemical estimates substantially underestimate N<sub>2</sub>-fixation rates in the Atlantic. We calculate a total of  $3.4 \times 10^{12}$  moles N yr<sup>-1</sup> of N<sub>2</sub>-fixation from 25° S–65° N, which supports 23% of the new production in the Atlantic.

If we scale our northern hemisphere Atlantic estimate of nitrogen fixation globally following GS97 (i.e., multiply  $2.8 \times 10^{12}$  by a factor of 4), then we obtain a global rate on the order of  $11 \times 10^{12}$  moles N yr<sup>-1</sup>. These nitrogen fixation rate estimates are based on our efforts to reproduce observed N\* anomalies in the Atlantic and should therefore include a representation of all N<sub>2</sub> fixation sources. This is less than half of the recently revised global denitrification rate ( $28 \times 10^{12}$  moles N yr<sup>-1</sup>) estimated by Codispoti (2006)<sup>1</sup>. However, if we scale global estimates of shelf denitrification ( $21 \times 10^{12}$  moles N yr<sup>-1</sup>, Codispoti, 2006<sup>1</sup>), by the fraction of global shelf area in the North Atlantic (~10%), then shelf denitrification in the North Atlantic could be as high as  $2.1 \times 10^{12}$  moles N yr<sup>-1</sup>, or nearly as high as our open ocean nitrogen fixation estimate. Because our results suggest that the N:P anomalies associated with nitrogen fixation can be advected far from source regions before they appear at depth, it is possible that our “net” estimate of Atlantic (and thus global) nitrogen fixation underestimates the total rate by 50%.

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## Appendix A

### Biogeochemical model equations

In this appendix we provide the specific model equations along with brief definitions of the terms. The reader is referred to Sect. 2 for a general description of the model.

The phytoplankton equation includes terms for growth, mortality, and grazing;

$$\frac{\partial P}{\partial t} = \alpha U_P P - s_P P - h_P C_M H \cdot P \quad (\text{A1})$$

where growth is modeled with co-limitation of light and nutrient availability;

$$U_P = \mu_P e^{-I/I_{\beta P}} \left(1 - e^{-I/I_P}\right) \left[ \min \left( \frac{\text{DIN}}{\text{DIN} + \text{PKS}_N}, \frac{\text{DIP}}{\text{DIP} + \text{PKS}_P}, \frac{\text{DIFe}}{\text{DIFe} + \text{PKS}_{\text{Fe}}} \right) \right] \quad (\text{A2})$$

The *Trichodesmium* equation includes growth, mortality, and grazing;

$$\frac{\partial T}{\partial t} = \alpha G_T T - s_T T - h_T C_M H \cdot T \quad (\text{A3})$$

here growth is a function of light limitation and phosphorus or iron limitations;

$$G_T = \mu_T \left(1 - e^{-I/I_T}\right) \left[ \min \left( \frac{\text{Ph}}{\text{Ph} + \text{TKS}_P}, \frac{\text{DIFe}}{\text{DIFe} + \text{TKS}_{\text{Fe}}} \right) \right] \quad (\text{A4})$$

Ph = max (DIP, DOP) under the assumption that diazotrophs can take up DOP as well as DIP.

The heterotroph equation includes grazing on other compartments, as well as itself;

$$\frac{\partial H}{\partial t} = R_H F (g_{e_{\text{DON}}} h_{\text{DON}} C_M H \cdot \text{DON} + g_{e_D} h_D C_M H \cdot D_N + g_{e_P} h_P C_M H \cdot P + g_{e_T} h_T C_M H \cdot T + g_{e_H} h_H C_M H^2) - h_H C_M H^2 \quad (\text{A5})$$

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where  $R_{HF}$  expresses the most limiting nutrient in available prey for heterotrophs;

$$R_{HF} = \min \left( \frac{R_{NF}}{R_{NH}}, \frac{R_{PF}}{R_{PH}}, \frac{R_{FeNF}}{R_{FeNH}} \right) \quad (A6)$$

$$R_{NNF} = \frac{DON + D_N + P + H + T}{DON + D_N + P + H + T} = 1 \quad (A7)$$

$$R_{PNF} = \frac{DOP + D_P + R_P NPP + R_P NHH + R_P NTT}{DON + D_N + P + H + T} \quad (A8)$$

$$5 \quad R_{FeNF} = \frac{DOFe + D_{Fe} + R_{Fe} NPP + R_{Fe} NHH + R_{Fe} NTT}{DON + D_N + P + H + T} \quad (A9)$$

and prey preference is expressed as;

$$h_{DON} = \Phi_{DON} / \Theta \quad (A10)$$

$$h_D = \Phi_D / \Theta \quad (A11)$$

$$h_P = \Phi_P / \Theta \quad (A12)$$

$$10 \quad h_T = \Phi_T / \Theta \quad (A13)$$

$$h_H = \Phi_H / \Theta \quad (A14)$$

with;

$$\Theta = \Phi_{DON} DON + \Phi_D P + \Phi_P P + \Phi_T P + \Phi_H P + HKS \quad (A15)$$

15 The dissolved inorganic nitrogen equation includes terms for phytoplankton and diazotroph uptake, excretion from food uptake as well as from food that was unusable due to stoichiometry, and inputs from remineralization of dissolved and particulate material;

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$$\begin{aligned}
\frac{\partial \text{DIN}}{\partial t} = & -U_P P - G_T T + R_H F_Y ((a e_{\text{DON}} - g e_{\text{DON}}) h_{\text{DON}} C_M H \cdot \text{DON} \\
& + (a e_D - g e_D) h_D C_M H \cdot D_N + (a e_P - g e_P) h_P C_M H \cdot P \\
& + (a e_T - g e_T) h_T C_M H \cdot T + (a e_H - g e_H) h_H C_M H^2) \\
& + X_{\text{HFNY}} (a e_{\text{DON}} h_{\text{DON}} C_M H \cdot \text{DON} + a e_D h_D C_M H \cdot D_N + a e_P h_P C_M H \cdot P \\
& + a e_T h_T C_M H \cdot T + a e_H h_H C_M H^2) + e_{\text{DON}} \text{DON} + e_{D_N} D_N
\end{aligned} \tag{A16}$$

where;

$$X_{\text{HF}} = \max(0, 1. - R_{\text{HF}}) \tag{A16.1}$$

and uptake by diazotrophs is limited by DIN;

$$U_T = \mu_T \left(1 - e^{-I/I_T}\right) \left[ \min \left( \frac{\text{DIN}}{\text{DIN} + \text{TKS}_N}, \frac{\text{DIPh}}{\text{DIPh} + \text{TKS}_P}, \frac{\text{DIFe}}{\text{DIFe} + \text{TKS}_{\text{Fe}}} \right) \right] \tag{A17}$$

The dissolved inorganic phosphorus equation follows the DIN equation except for the diazotroph uptake, which may be from the DOP pool if DOP concentrations are higher than DIP concentrations;

$$\begin{aligned}
\frac{\partial \text{DIP}}{\partial t} = & R_P N H R_H F_Y ((a e_{\text{DON}} - g e_{\text{DON}}) h_{\text{DON}} C_M H \cdot \text{DON} + (a e_D - g e_D) h_D C_M H \cdot D_N \\
& + (a e_P - g e_P) h_P C_M H \cdot P + (a e_T - g e_T) h_T C_M H \cdot T + (a e_H - g e_H) h_H C_M H^2) \\
& + X_{\text{HFPY}} (a e_{\text{DON}} h_{\text{DON}} C_M H \cdot \text{DON} + a e_D h_D C_M H \cdot D_N + a e_P h_P C_M H \cdot P \\
& + a e_T h_T C_M H \cdot T + a e_H h_H C_M H^2) - R_P N P U_P P - \overbrace{R_P N T G_T T}^{\text{DIP} \geq \text{DOP}} + e_{\text{DOP}} \text{DOP} + e_{D_P} D_P
\end{aligned} \tag{A18}$$

where;

$$X_{\text{HFP}} = R_{\text{PNF}} - (R_{\text{HF}} \cdot R_{\text{PNH}}) \tag{A18.1}$$

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The dissolved inorganic iron equation again follows DIP, but with an additional scavenging term;

$$\begin{aligned} \frac{\partial \text{DIFe}}{\partial t} = & R_{\text{Fe}} e_{\text{NH}} R_{\text{H}} F \gamma ((ae_{\text{DON}} - ge_{\text{DON}}) h_{\text{DON}} C_{\text{M}} H \cdot \text{DON} + (ae_{\text{D}} - ge_{\text{D}}) h_{\text{D}} C_{\text{M}} H \cdot D_{\text{N}} \\ & + (ae_{\text{P}} - ge_{\text{P}}) h_{\text{P}} C_{\text{M}} H \cdot P + (ae_{\text{T}} - ge_{\text{T}}) h_{\text{T}} C_{\text{M}} H \cdot T + (ae_{\text{H}} - ge_{\text{H}}) h_{\text{H}} C_{\text{M}} H^2) \\ & + X_{\text{HFFe}} \gamma (ae_{\text{DON}} h_{\text{DON}} C_{\text{M}} H \cdot \text{DON} + ae_{\text{D}} h_{\text{D}} C_{\text{M}} H \cdot D_{\text{N}} + ae_{\text{P}} h_{\text{P}} C_{\text{M}} H \cdot P \\ & + ae_{\text{T}} h_{\text{T}} C_{\text{M}} H \cdot T + ae_{\text{H}} h_{\text{H}} C_{\text{M}} H^2) - R_{\text{Fe}} e_{\text{N}} U_{\text{P}} P - R_{\text{Fe}} e_{\text{N}} T G_{\text{T}} T - S_{\text{KVG}} + e_{\text{DOFe}} \text{DOFe} \\ & + e_{\text{DFe}} D_{\text{Fe}} \end{aligned} \quad (\text{A19})$$

where;

$$X_{\text{HFFe}} = R_{\text{Fe}} e_{\text{N}} F - (R_{\text{H}} F \cdot R_{\text{Fe}} e_{\text{NH}}) \quad (\text{A19.1})$$

and scavenging is represented as;

$$S_{\text{KVG}} = -K_{\text{Fe}} [1 + k (D_{\text{N}} + D_{\text{P}} + D_{\text{Fe}})] \text{DIFe} \quad (\text{A20})$$

where k is an arbitrary constant that doubles scavenging at high detrital concentrations.

The dissolved organic nitrogen equation includes terms for excretion following the DIN equation, as well as inputs from mortality and growth of phytoplankton and diazotrophs and losses from remineralization;

$$\begin{aligned} \frac{\partial \text{DON}}{\partial t} = & R_{\text{H}} F (1 - \gamma) ((ae_{\text{DON}} - ge_{\text{DON}}) h_{\text{DON}} C_{\text{M}} H \cdot \text{DON} + (ae_{\text{D}} - ge_{\text{D}}) h_{\text{D}} C_{\text{M}} H \cdot D_{\text{N}} \\ & + (ae_{\text{P}} - ge_{\text{P}}) h_{\text{P}} C_{\text{M}} H \cdot P + (ae_{\text{T}} - ge_{\text{T}}) h_{\text{T}} C_{\text{M}} H \cdot T + (ae_{\text{H}} - ge_{\text{H}}) h_{\text{H}} C_{\text{M}} H^2) \\ & + X_{\text{HFN}} (1 - \gamma) (ae_{\text{DON}} h_{\text{DON}} C_{\text{M}} H \cdot \text{DON} + ae_{\text{D}} h_{\text{D}} C_{\text{M}} H \cdot D_{\text{N}} + ae_{\text{P}} h_{\text{P}} C_{\text{M}} H \cdot P \\ & + ae_{\text{T}} h_{\text{T}} C_{\text{M}} H \cdot T + ae_{\text{H}} h_{\text{H}} C_{\text{M}} H^2) + (1 - \beta) s_{\text{P}} P + (1 - \beta) s_{\text{T}} T + (1 - \alpha) U_{\text{P}} P \\ & + (1 - \alpha) G_{\text{T}} T - h_{\text{DON}} C_{\text{M}} H \cdot \text{DON} - e_{\text{DON}} \text{DON} \end{aligned} \quad (\text{A21})$$

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The dissolved organic phosphorus equation follows the DON equation, but includes an uptake term for diazotrophy in cases where DOP concentrations exceed DIP;

$$\begin{aligned}
 \frac{\partial \text{DOP}}{\partial t} = & R_{\text{PNHR}_\text{HF}} (1 - \gamma) ((ae_{\text{DON}} - ge_{\text{DON}}) h_{\text{DON}} C_{\text{MH}} \cdot \text{DON} \\
 & + (ae_{\text{D}} - ge_{\text{D}}) h_{\text{D}} C_{\text{MH}} \cdot D_{\text{N}} + (ae_{\text{P}} - ge_{\text{P}}) h_{\text{P}} C_{\text{MH}} \cdot \text{P} + (ae_{\text{T}} - ge_{\text{T}}) h_{\text{T}} C_{\text{MH}} \cdot \text{T} \\
 & + (ae_{\text{H}} - ge_{\text{H}}) h_{\text{H}} C_{\text{MH}} H^2) + X_{\text{HFP}} (1 - \gamma) (ae_{\text{DON}} h_{\text{DON}} C_{\text{MH}} \cdot \text{DON} + ae_{\text{D}} h_{\text{D}} C_{\text{MH}} \cdot D_{\text{N}} \\
 & + ae_{\text{P}} h_{\text{P}} C_{\text{MH}} \cdot \text{P} + ae_{\text{T}} h_{\text{T}} C_{\text{MH}} \cdot \text{T} + ae_{\text{H}} h_{\text{H}} C_{\text{MH}} H^2) + R_{\text{PNP}} (1 - \beta) s_{\text{P}} \text{P} \\
 & + R_{\text{PNT}} (1 - \beta) s_{\text{T}} \text{T} + R_{\text{PNP}} (1 - \alpha) U_{\text{P}} \text{P} + R_{\text{PNT}} (1 - \alpha) G_{\text{T}} \text{T} - \overbrace{R_{\text{PNT}} G_{\text{T}} \text{T}}^{\text{DOP} > \text{DIP}} - h_{\text{DON}} C_{\text{MH}} \cdot \text{DOP} \\
 & - e_{\text{DOP}} \text{DOP}
 \end{aligned} \tag{A22}$$

The dissolved organic iron equation follows the DON equation;

$$\begin{aligned}
 \frac{\partial \text{DOFe}}{\partial t} = & R_{\text{FeNHR}_\text{HF}} (1 - \gamma) ((ae_{\text{DON}} - ge_{\text{DON}}) h_{\text{DON}} C_{\text{MH}} \cdot \text{DON} \\
 & + (ae_{\text{D}} - ge_{\text{D}}) h_{\text{D}} C_{\text{MH}} \cdot D_{\text{N}} + (ae_{\text{P}} - ge_{\text{P}}) h_{\text{P}} C_{\text{MH}} \cdot \text{P} + (ae_{\text{T}} - ge_{\text{T}}) h_{\text{T}} C_{\text{MH}} \cdot \text{T} \\
 & + (ae_{\text{H}} - ge_{\text{H}}) h_{\text{H}} C_{\text{MH}} H^2) + X_{\text{HFFe}} (1 - \gamma) (ae_{\text{DON}} h_{\text{DON}} C_{\text{MH}} \cdot \text{DON} + ae_{\text{D}} h_{\text{D}} C_{\text{MH}} \cdot D_{\text{N}} \\
 & + ae_{\text{P}} h_{\text{P}} C_{\text{MH}} \cdot \text{P} + ae_{\text{T}} h_{\text{T}} C_{\text{MH}} \cdot \text{T} + ae_{\text{H}} h_{\text{H}} C_{\text{MH}} H^2) + R_{\text{FeNP}} (1 - \beta) s_{\text{P}} \text{P} \\
 & + R_{\text{FeNT}} (1 - \beta) s_{\text{T}} \text{T} + R_{\text{FeNP}} (1 - \alpha) U_{\text{P}} \text{P} + R_{\text{FeNT}} (1 - \alpha) G_{\text{T}} \text{T} - h_{\text{DON}} C_{\text{MH}} \cdot \text{DOFe} \\
 & - e_{\text{DOFe}} \text{DOFe}
 \end{aligned} \tag{A23}$$

The N-detritus equation includes terms for egestion, as well as mortality, and losses from remineralization, grazing, and sinking;

$$\begin{aligned}
 \frac{\partial D_{\text{N}}}{\partial t} = & R_{\text{HF}} ((1 - ae_{\text{DON}}) h_{\text{DON}} C_{\text{MH}} \cdot \text{DON} + (1 - ae_{\text{D}}) h_{\text{D}} C_{\text{MH}} \cdot D_{\text{N}} \\
 & + (1 - ae_{\text{P}}) h_{\text{P}} C_{\text{MH}} \cdot \text{P} + (1 - ae_{\text{T}}) h_{\text{T}} C_{\text{MH}} \cdot \text{T} + (1 - ae_{\text{H}}) h_{\text{H}} C_{\text{MH}} H^2)
 \end{aligned}$$

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$$\begin{aligned}
& +X_{HF}((1 - ae_{DON}) h_{DON}C_M H \cdot DON + (1 - ae_D) h_D C_M H \cdot D_N \\
& + (1 - ae_P) h_P C_M H \cdot P + (1 - ae_T) h_T C_M H \cdot T + (1 - ae_H) h_H C_M H^2) + \beta s_P P \\
& + \beta s_T T - e_{DN} D_N - h_D C_M H \cdot D_N - w_S D_N
\end{aligned} \tag{A24}$$

where

$$w_S = \min(75., \max(w_{SS}, D_N * 4444 + 10.555)) \tag{A25}$$

which provides a linear ramp from 15m/day to 75 m/d, increasing as a function of detrital concentration;

The P-detritus equation follows the N-detritus equation;

$$\begin{aligned}
\frac{\partial D_P}{\partial t} &= R_P N R_H F ((1 - ae_{DON}) h_{DON} C_M H \cdot DON + (1 - ae_D) h_D C_M H \cdot D_N \\
& + (1 - ae_P) h_P C_M H \cdot P + (1 - ae_T) h_T C_M H \cdot T + (1 - ae_H) h_H C_M H^2) \\
& + X_{HFP} ((1 - ae_{DON}) h_{DON} C_M H \cdot DON + (1 - ae_D) h_D C_M H \cdot D_N \\
& + (1 - ae_P) h_P C_M H \cdot P + (1 - ae_T) h_T C_M H \cdot T + (1 - ae_H) h_H C_M H^2) + R_{PNP} \beta s_P P \\
& + R_P N T \beta s_T T - e_{DP} D_P - h_D C_M H \cdot D_P - w_S D_P
\end{aligned} \tag{A26}$$

The Fe-detritus equation also follows the N-detritus equation;

$$\begin{aligned}
\frac{\partial D_{Fe}}{\partial t} &= R_{Fe} N R_H F ((1 - ae_{DON}) h_{DON} C_M H \cdot DON + (1 - ae_D) h_D C_M H \cdot D_N \\
& + (1 - ae_P) h_P C_M H \cdot P + (1 - ae_T) h_T C_M H \cdot T + (1 - ae_H) h_H C_M H^2) \\
& + X_{HFFe} ((1 - ae_{DON}) h_{DON} C_M H \cdot DON + (1 - ae_D) h_D C_M H \cdot D_N \\
& + (1 - ae_P) h_P C_M H \cdot P + (1 - ae_T) h_T C_M H \cdot T + (1 - ae_H) h_H C_M H^2) + R_{FeNP} \beta s_P P \\
& + R_{FeNT} \beta s_T T - e_{DFe} D_{Fe} - h_D C_M H \cdot D_{Fe} - w_S D_{Fe} + Fe_{SOL} Fe_{\%} DST
\end{aligned} \tag{A27}$$

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**Table 1.** Schematic diagram of the biogeochemical model with N<sub>2</sub>-fixation and Fe and P limitations. Single boxes with multiple colors represent model constituents with fixed elemental ratios, whereas compartments with multiple boxes represent model constituents with variable element ratios. DIX = dissolved inorganic nutrients, DOX = dissolved organic nutrients, DX = organic detritus, P = phytoplankton, T = Trichodesmium, and H = heterotroph.

Source	10 <sup>12</sup> mol N/year	Notes
Tropical Atlantic Domain, 10°N–30°N		
NSTAR run (this paper)	1.8	This study: tuned to observed N* distribution
NSTAR run (this paper)	2.8	This study: surface nitrogen fixation integrated over the northern hemisphere only.
Coles et al. (2004)	0.55	Tuned to surface Trichodesmium biomass observations
Coles et al. (2004)	2.15	Tuned to GS97 N* rate estimates
Hansell et al. (2004)	0.31	Geochemically based, excludes Gulf of Mexico, and assumes smaller volumes for active N <sub>2</sub> fixation.
GS97	2.0	Based on observed N*, using an N:P ratio for diazotrophs of 125.
GS97	3.2	Based on observed N*, but using an N:P ratio for diazotrophs of 45 (from Capone et al., 2005).
Capone et al. (2005)	1.6–2.4	Extrapolated from surface N <sub>2</sub> fixation rate measurements
Whole Atlantic Domain		
NSTAR run	3.4	This study: Tuned to N* maximum
Capone et al. (2005)	5.5–8.7	Isotopic N calculation
Lee et al. (2002)	2.0	Carbon inventory

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**Table B1.** Model parameters.

Description	Symbol	Value	Units	Source
Growth efficiency for H on P	$g_{eP}$	0.2	Dimensionless	Hood et al. (2001)
Assimilation efficiency for H on P	$ae_P$	0.7	Dimensionless	Hood et al. (2001)
Growth efficiency for H on D	$g_{eD}$	0.2	Dimensionless	Hood et al. (2001)
Assimilation efficiency for H on D	$ae_D$	0.7	Dimensionless	Hood et al. (2001)
Growth efficiency for H on T	$g_{eT}$	0.2	Dimensionless	Hood et al. (2001)
Assimilation efficiency for H on T	$ae_T$	0.7	Dimensionless	Hood et al. (2001)
Growth efficiency for H on DON	$g_{eDON}$	0.2	Dimensionless	Hood et al. (2001)
Assimilation efficiency for H on DON	$ae_{DON}$	1.0	Dimensionless	Hood et al. (2001)
Heterotrophic max. consumption rate	$C_M$	6.4	$d^{-1}$	Hood et al. (2001)
Heterotrophic saturation constant	$HKS$	0.8	$mmol\ m^{-3}$	Hood et al. (2001)
Heterotrophic preference for P	$\Phi_P$	0.4286	Dimensionless	Hood et al. (2004)
Heterotrophic preference for D	$\Phi_D$	0.0	Dimensionless	Hood et al. (2004)
Heterotrophic preference for H	$\Phi_H$	0.2857	Dimensionless	Hood et al. (2004)
Heterotrophic preference for T	$\Phi_T$	0.05	Dimensionless	Hood et al. (2004)
Heterotrophic preference for DON	$\Phi_{DON}$	0.2175	Dimensionless	Hood et al. (2004)
Maximum phytoplankton growth rate	$\mu_P$	3.22	$d^{-1}$	Hood et al. (2001)
Maximum <i>Trichodesmium</i> growth rate	$\mu_T$	0.23	$d^{-1}$	See below <sup>1</sup>
Phytoplankton natural mortality rate	$s_P$	0.05	$d^{-1}$	Hood et al. (2001)
<i>Trichodesmium</i> natural mortality rate	$s_T$	0.01	$d^{-1}$	See below <sup>2</sup>
Partitioning of P and T senescence	$\beta$	0.25	Dimensionless	Hood et al. (2001)
Partitioning of P and T production	$\alpha$	0.7	Dimensionless	Hood et al. (2001)
Phytoplankton light saturation param.	$I_P$	30	Watts $m^{-2}$	See below <sup>3</sup>
T light saturation parameter	$I_T$	70	Watts $m^{-2}$	See below <sup>4</sup>
Partitioning of excretion to DIN	$\gamma$	0.75	Dimensionless	Hood et al. (2001)
P photoinhibition parameter	$I_{\beta P}$	400	Watts $m^{-2}$	Hood et al. (2001)
N:N ratio for H	$R_{NNH}$	1	Dimensionless	See below <sup>5</sup>
P:N ratio for H	$R_{PNH}$	0.0625	Dimensionless	Redfield et al. (1964)
Fe:N ratio for H	$R_{FeNH}$	$3.75 \times 10^{-5}$	Dimensionless	See below <sup>6</sup>
P:N ratio for T	$R_{PNT}$	0.02222	Dimensionless	See below <sup>7</sup>
Fe:N ratio for T	$R_{FeNT}$	$2.236 \times 10^{-4}$	Dimensionless	See below <sup>8</sup>
P:N ratio for P	$R_{PNP}$	0.0625	Dimensionless	Redfield et al. (1964)
Fe:N ratio for P	$R_{FeNP}$	$2.981 \times 10^{-5}$	Dimensionless	See below <sup>9</sup>
Sat. const. for DIN uptake by P	$PKS_N$	0.5	$mmol\ m^{-3}$	Hood et al. (2001)
Sat. const. for DIN uptake by T	$TKS_N$	0.5	$mmol\ m^{-3}$	Hood et al. 2001
Sat. const. for DIP uptake by P	$PKS_P$	0.0030	$mmol\ m^{-3}$	See below <sup>12</sup>
Sat. const. for DIP uptake by T	$TKS_P$	0.0077	$mmol\ m^{-3}$	See below <sup>10</sup>
Sat. const. for DiFe uptake by P	$PKS_{Fe}$	$1.0 \times 10^{-5}$	$mmol\ m^{-3}$	See below <sup>11</sup>
Sat. const. for DiFe uptake by T	$TKS_{Fe}$	$1.0 \times 10^{-4}$	$mmol\ m^{-3}$	See below <sup>13</sup>
Fe scavenging rate constant	$K_{Fe}$	$12.5 \times 10^{-5}$	$d^{-1}$	See below <sup>14</sup>
Enhanced $D_p$ recycling rate	$e_{DP}$	0.35	$d^{-1}$	See below <sup>15</sup>
Enhanced DOP recycling rate	$e_{DOP}$	0.14	$d^{-1}$	See below <sup>16</sup>
Enhanced $D_N$ recycling rate	$e_{DN}$	0.14	$d^{-1}$	See below <sup>17</sup>
Enhanced DON recycling rate	$e_{DON}$	0.14	$d^{-1}$	See below <sup>18</sup>
Enhanced $D_{Fe}$ recycling rate	$e_{DFe}$	0.35	$d^{-1}$	See below <sup>19</sup>
Enhanced DOFe recycling rate	$e_{DOFe}$	0.14	$d^{-1}$	See below <sup>20</sup>
Sinking rate of Detritus	$W_{SS}$	12.0	meters $d^{-1}$	See below <sup>21</sup>
Iron solubility	$Fe_{SOL}$	0.01	$d^{-1}$	See below <sup>22</sup>
Fe fraction in dust	$Fe_{\%}$	0.035	Dimensionless	See below <sup>23</sup>

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**Table B1.** Continued.

- <sup>1</sup> Equivalent to a doubling time of  $\sim 3$  days, from Capone et al. (1997).
- <sup>2</sup> Increased slightly from  $.025 \text{ d}^{-1}$  in Hood et al. (2004) to maintain *Trichodesmium* colony concentrations in the western tropical and subtropical Atlantic that are consistent with those observed there by Carpenter and Romans (1991).
- <sup>3,4</sup> Decreased from 40 and 80  $\text{Watts m}^{-2}$ , respectively, in Hood et al. (2001) to compensate for deeper mixed layers generated by HYCOM.
- <sup>5</sup> Equals 1 by definition. Included for clarity in Eq. (A6).
- <sup>6</sup> Derived from  $\text{Fe:C} = 7.5$  ( $\mu\text{mole:mole}$ ) for heterotrophic bacteria from Tortell et al. (1999) using a C:N ratio of 32:6.4 ( $\text{mole:mole}$ ) from Vrede et al. (2002).
- <sup>7</sup> D. G. Capone, personal communication (2004).
- <sup>8</sup> Derived using *Trichodesmium* Fe colony $^{-1}$  (median =  $8 \text{ pmol Fe colony}^{-1}$ ) and N colony $^{-1}$  (median =  $32 \mu\text{mol N colony}^{-1}$ ) from Sañudo-Wilhelmy et al. (2001).
- <sup>9</sup> Using  $\text{Fe:C} = 4.5$  ( $\mu\text{mol:mol}$ ) for phytoplankton from Fung et al. (2004) and assuming a Redfield C:N = 106:16.
- <sup>10</sup> Mean from J. Sohm personal communication (2004), unpublished kinetic studies on *Trichodesmium* in Atlantic waters.
- <sup>11</sup> From Christian et al. (2002b), assuming phytoplankton in the model are dominated by small forms.
- <sup>12</sup> Assuming half saturation constants for P uptake are similar for phytoplankton and *Trichodesmium*.
- <sup>13</sup> Assuming half saturation constants for Fe uptake are similar for *Trichodesmium* and phytoplankton, following Moore et al. (2002a).
- <sup>14</sup> Set a posteriori to give best fit to published Fe profiles in the Wu et al. (2000) and Landing et al. (2004).
- <sup>15,16</sup> Set a posteriori to relieve P limitation and provide reasonable near-surface DIP concentrations.
- <sup>17,18</sup> Applied to ensure  $D_N$  and DON remineralization at depth.
- <sup>19,20</sup> Applied to ensure  $D_{Fe}$  and DOf<sub>e</sub> remineralization at depth and adjusted to reproduce observed deep Fe profiles from Wu et al. (2000) and Landing et al. (2004).
- <sup>21</sup> Set a posteriori to give reasonable near-surface DIN, DIP and DIFe concentrations.
- <sup>22</sup> Values typically assumed to be between 1 and 10% of the Fe in dust soluble/bioavailable, cf., Moore et al. (2002a); Christian et al. (2002a); Duce and Tindale (1991).
- <sup>23</sup> Following Christian et al. (2002a).

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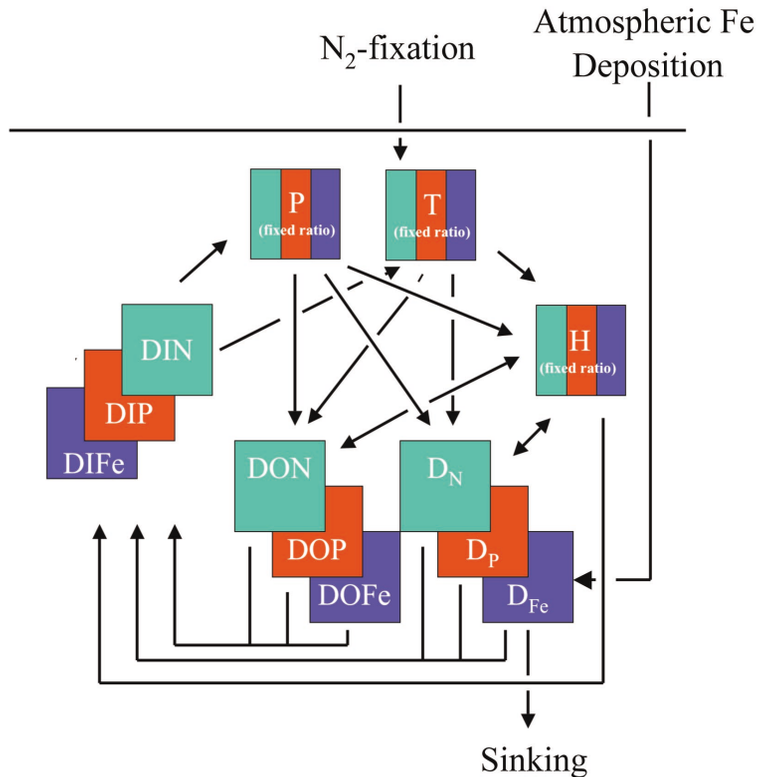
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**Fig. 1.** Basin wide nitrogen fixation rates from the NSTAR run. These are computed over the domain of GS97 ( $10^\circ N$ – $30^\circ N$ ) and also over the entire North Atlantic for the NSTAR run. For comparison we include previous estimates of the basin wide  $N_2$ -fixation rate in the Atlantic from: 1) prognostic model runs tuned to observed *Trichodesmium* biomass and GS97 rates (Coles et al., 2004); 2) geochemical estimates based on observed  $N^*$  anomalies (GS97, Hansell et al., 2004) and carbon drawdown (Lee et al., 2002); and 3) observational estimates based on direct rate measurements (Capone et al., 2005).

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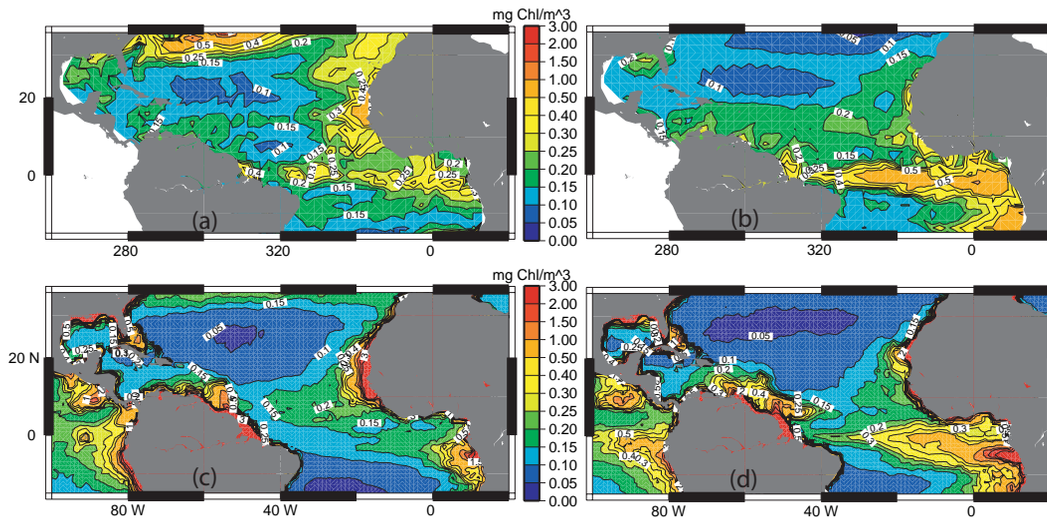
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**Fig. 2.** Surface phytoplankton chlorophyll ( $\text{mg Chl/m}^3$ ) for (a) spring (April) and (b) fall (August) for the NSTAR model run and (c) spring (April) and (d) fall (August) for a climatology based on the SeaWiFS satellite data from 1998 through 2005.

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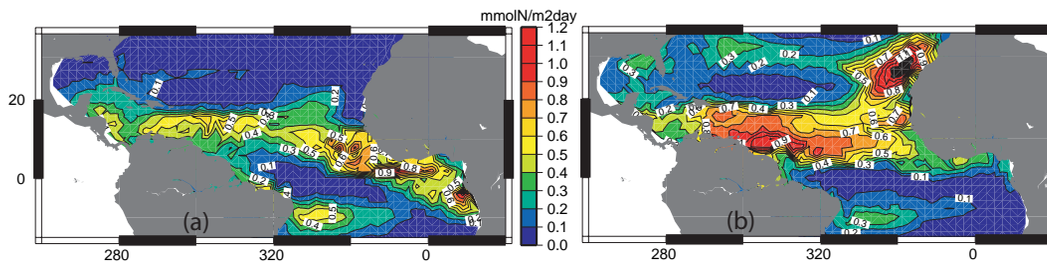


Fig. 3. Surface nitrogen fixation rates for (a) spring (April) and (b) fall (August) are shown for the NSTAR run.

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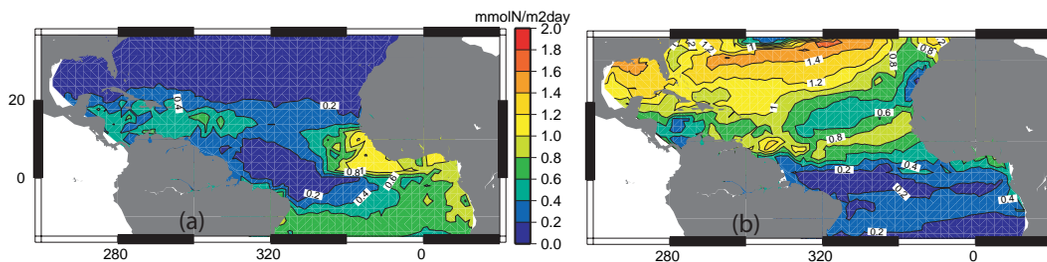
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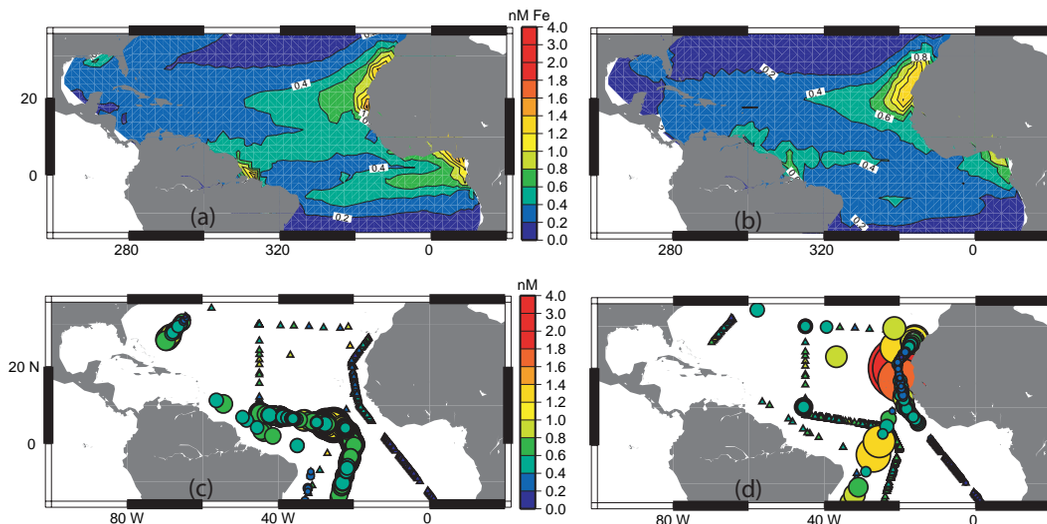


**Fig. 4.** Surface nitrogen fixation rates for **(a)** spring (April) and **(b)** fall (August) are shown for the NOLIM run.

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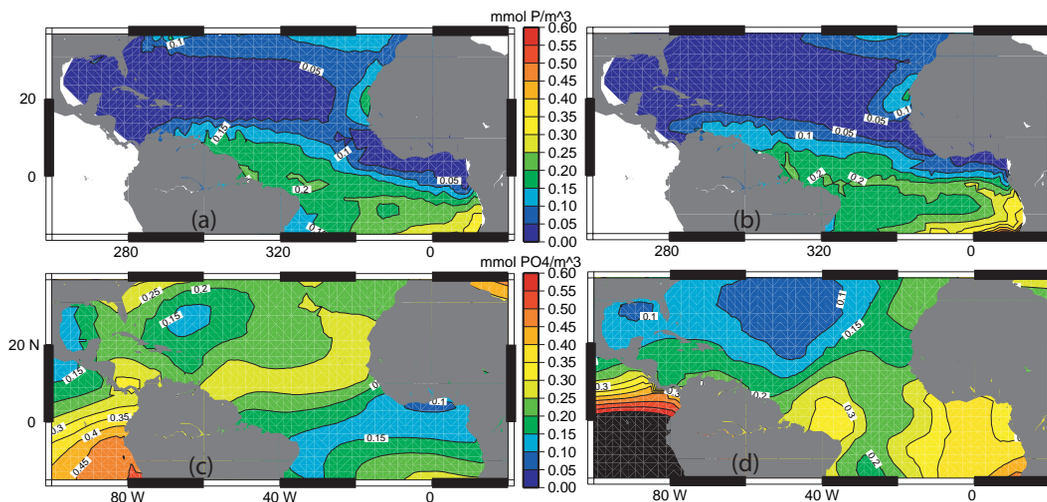
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**Fig. 5.** Surface iron concentrations in nM for the model NSTAR run for (a) spring (April) and (b) fall (August), and the available observations (Parekh et al., 2005; Bergquist and Boyle, 2006; Bowie et al., 2002) for (c) spring (March, April, May) and (d) fall (July, August, September). Because of the paucity of observational data, the surface iron observations are shown in entirety as small triangles, and the seasonal data are indicated as larger circles proportional to concentration.

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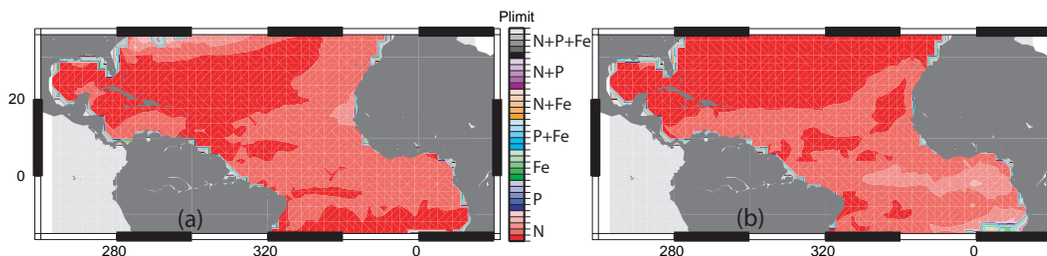
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**Fig. 6.** Surface phosphorus concentrations in  $\text{mmol/m}^3$  for the model NSTAR run for **(a)** spring (April) and **(b)** fall (August), and a nutrient climatology (Louanchi and Najjar, 2000) for **(c)** spring (April) and **(d)** fall (August).

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**Fig. 7.** Phytoplankton growth limitations for the NSTAR model run for **(a)** spring (April) and **(b)** fall (August). Each color indicates a different limiting nutrient, and the intensity of the shade reflects the intensity of the limitation, from low limitations (nearly white) to severe limitation (intense color). Co-limitation of the nutrients is indicated where the degree of limitation is within 20% for more than one nutrient, and is indicated by a shade of color.

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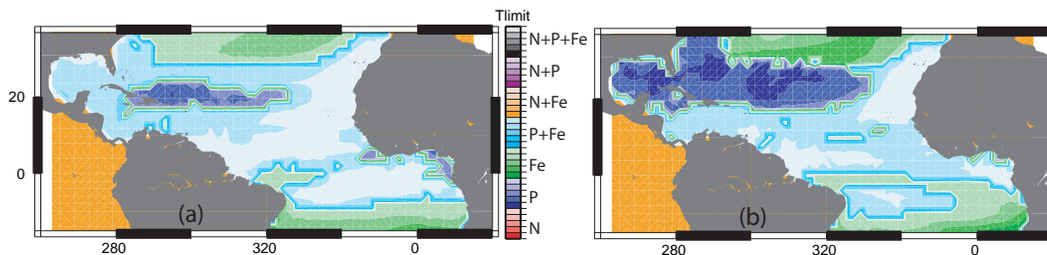
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**Fig. 8.** Diazotroph growth limitation for the NSTAR model run for (a) spring (April) and (b) fall (August). Each color indicates a different limiting nutrient, and the intensity of the shade reflects the intensity of the limitation, from low limitations (nearly white) to severe limitation (intense color). Co-limitation of the nutrients is indicated where the degree of limitation is within 20% for more than one nutrient, and is indicated by a shade of color.

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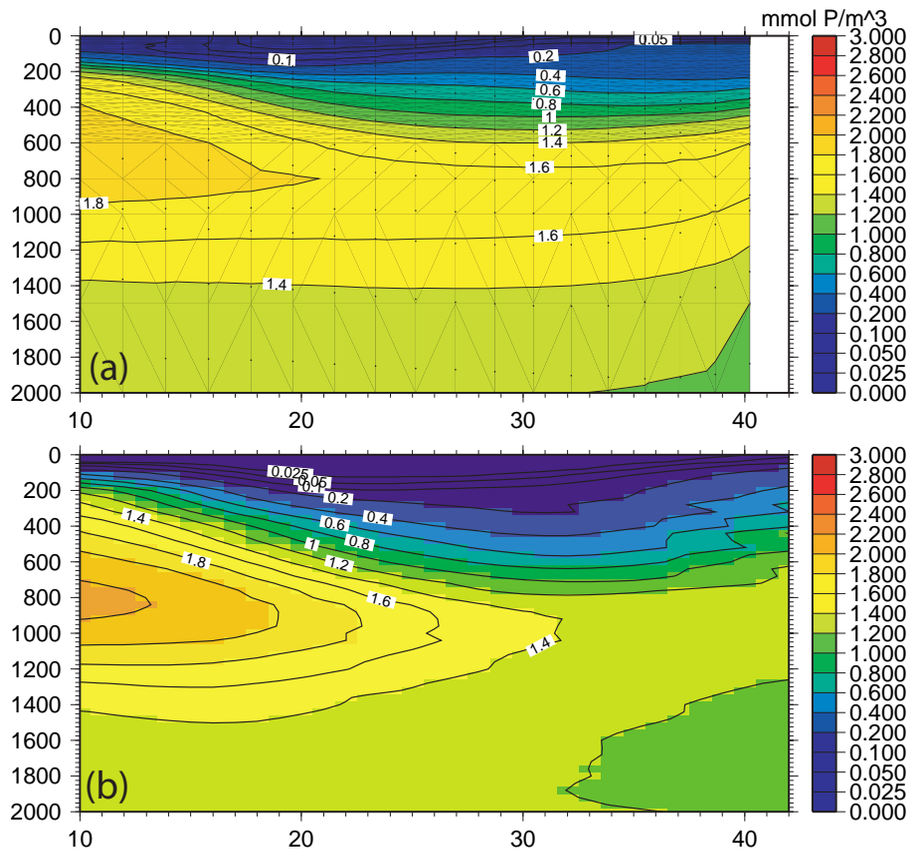
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**Fig. 9.** Vertical sections of phosphate in July for (a) the NSTAR model at 52° W, and (b) for the WOCE A20 section at 52° W.

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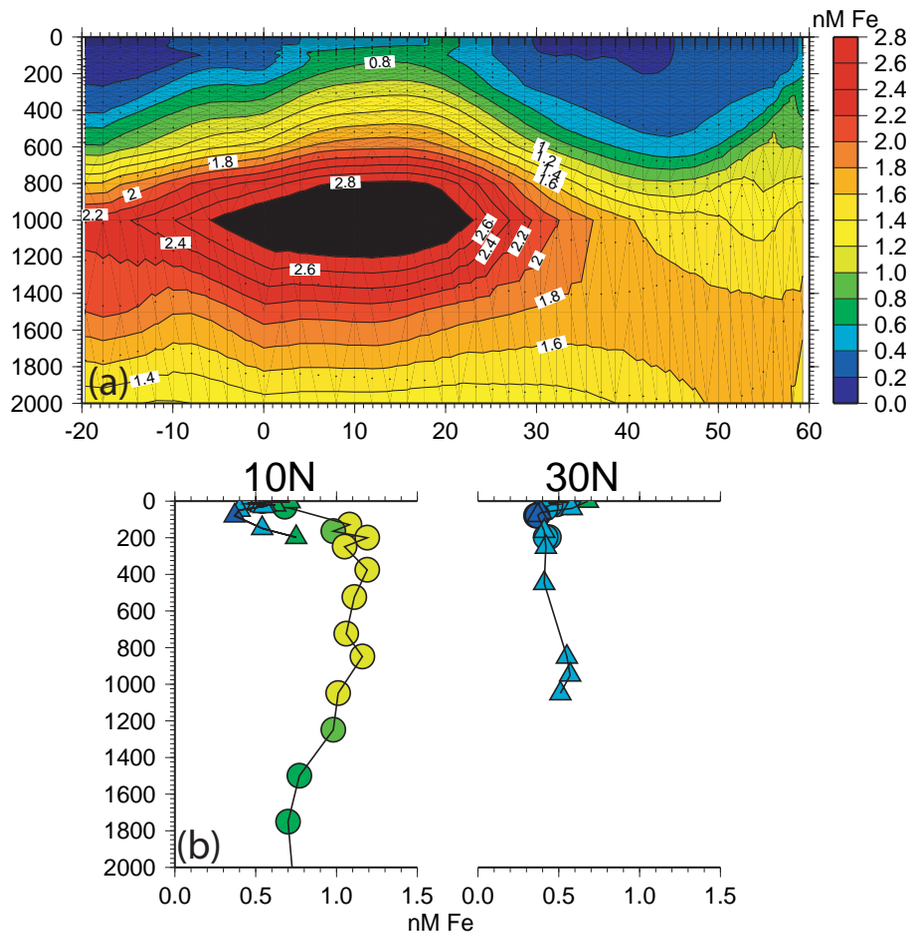
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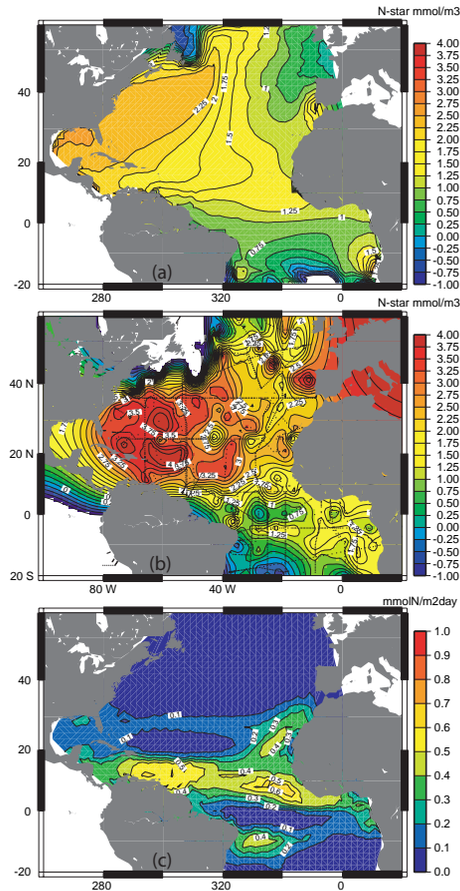
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**Fig. 10.** (a) Vertical section of dissolved inorganic iron nM for the NSTAR model along 28° W. (b) two deep profiles from Berquist and Boyle 2005 at 10° N, 45° W, and 30° N, 45° W.

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**Fig. 11.** Isopycnal maps of  $N^*$  from (a) the annually averaged NSTAR run at  $26.8 \sigma_\theta$ , (b) the GLODAP bottle database on  $26.5 \sigma_\theta$ , (c) annual average nitrogen fixation rate integrated over the euphotic zone in the NSTAR run.

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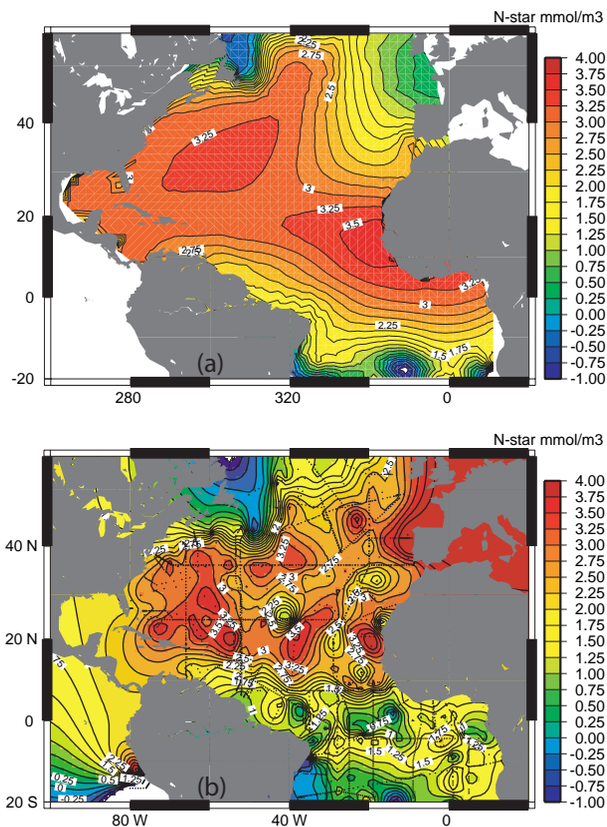


Fig. 12. Isopycnal maps of N\* from (a) the annually averaged NSTAR run at 27.03  $\sigma_\theta$ , (b) the GLODAP bottle database on 27.03  $\sigma_\theta$ .

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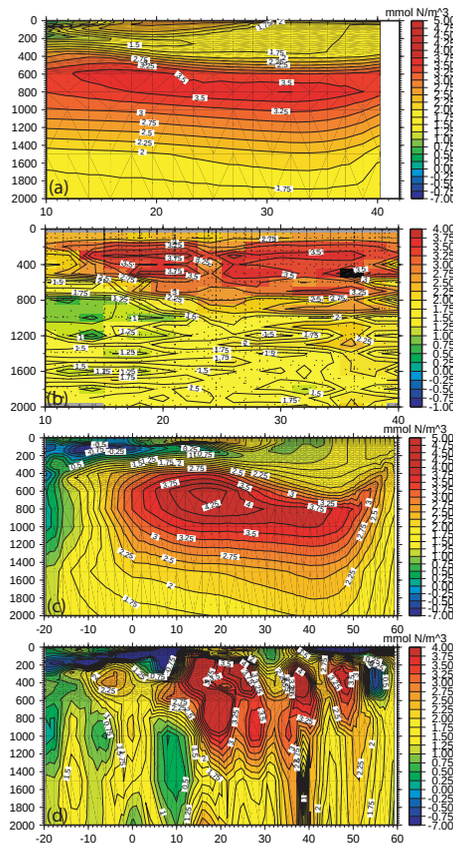
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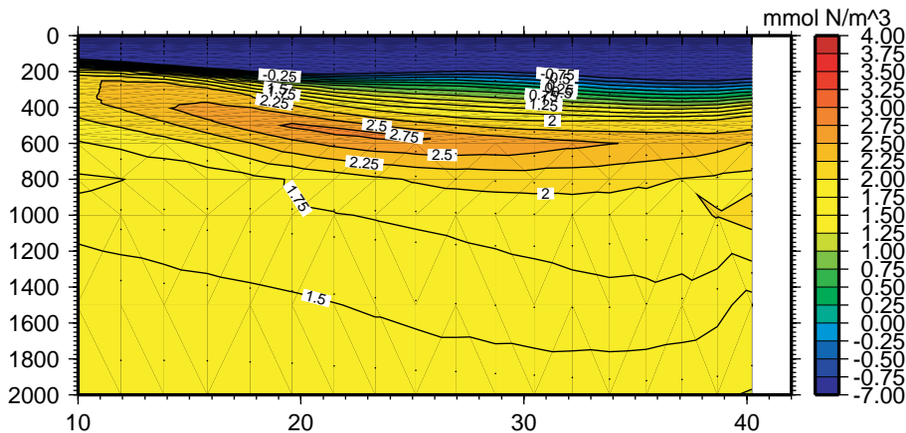
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**Fig. 13.** (a) Vertical section of N\* for the NSTAR model along 52° W, (b) N\* from WOCE A20 at 52° W, (c) NSTAR model run along 28° W, (d) N\* from WOA initial conditions at 28° W.

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**Fig. 14.** Vertical section of N\* for the NOFIX model along 52° W.

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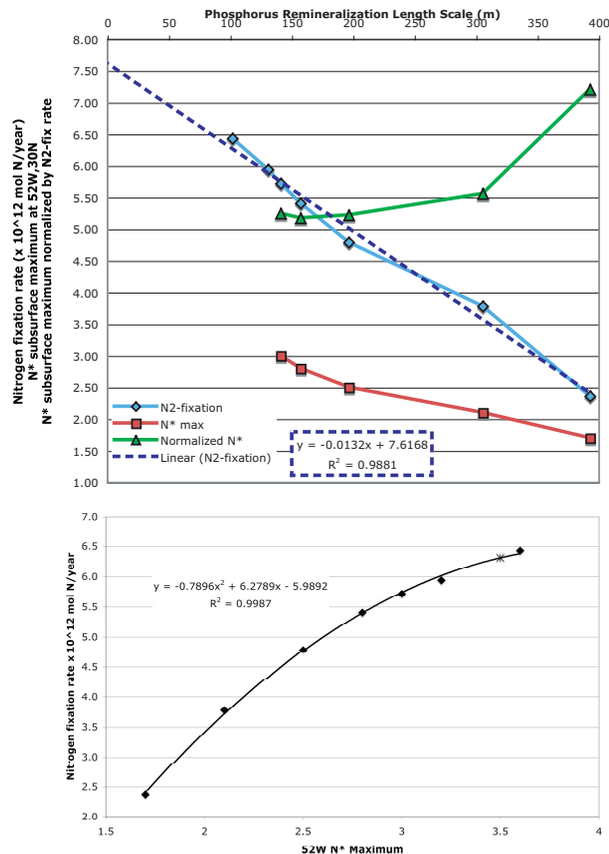
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**Fig. 15. (a)** Sensitivity experiments to phosphorus remineralization length scale. Blue symbols show basin averaged nitrogen fixation rate, red symbols show maximum subsurface N\* values at 20° N, 52° W, green symbols show N\* normalized by the basin averaged nitrogen fixation rate. **(b)** Simulated relationship between N\* maximum at 20° N, 52° W, and basin averaged nitrogen fixation rate.

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