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Seasonal dynamics of SAR11 populations in the euphotic and mesopelagic zones of the northwestern Sargasso Sea

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Bacterioplankton belonging to the SAR11 clade of α -proteobacteria were counted by fluorescence in situ hybridization (FISH) over eight depths in the surface 300 m at the Bermuda Atlantic Timeseries Study (BATS) site from 2003 to 2005. SAR11 are dominant heterotrophs in oligotrophic systems; thus, resolving their temporal dynamics can provide important insights to the cycling of organic and inorganic nutrients. This quantitative time-series data revealed distinct annual distribution patterns of SAR11 abundance in the euphotic (0–120) and upper mesopelagic (160– 300 m) zones that were reproducibly correlated with seasonal mixing and stratification of the water column. Terminal restriction fragment length polymorphism (T-RFLP) data generated from a decade of samples collected at BATS were combined with the FISH data to model the annual dynamics of SAR11 subclade populations. 16S rRNA gene clone libraries were constructed to verify the correlation of the T-RFLP data with SAR11 clade structure. Clear vertical and temporal transitions were observed in the dominance of three SAR11 ecotypes. The mechanisms that lead to shifts between the different SAR11 populations are not well understood, but are probably a consequence of finely tuned physiological adaptations that partition the populations along physical and chemical gradients in the ecosystem. The correlation between evolutionary descent and temporal/spatial patterns we describe, confirmed that a minimum of three SAR11 ecotypes occupy the Sargasso Sea surface layer, and revealed new details of their population dynamics.

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Introduction

Over two decades ago, small subunit rRNA gene clones recovered from the surface waters of the Sargasso Sea revealed the presence of SAR11, a diverse clade of heterotrophic a-proteobacteria [\(Giovannoni](#page-11-0) et al., 1990). Since then SAR11 genes have been found ubiquitously distributed throughout the World's oceans. They represent approximately one quarter of all rRNA genes identified in clone libraries from marine environments (Rappé [and Giovannoni, 2003](#page-12-0)). The prevalence of SAR11

cells in oceanic systems has been confirmed quantitatively by fluorescence in situ hybridization (FISH) cell counts and account for approximately one-third of the prokaryotic cells in the surface waters of the northwestern Sargasso Sea (Morris et al[., 2002\)](#page-12-0).

Field et al[. \(1997\)](#page-11-0) were the first to identify ecotypes of the SAR11 clade using 16S rRNA gene sequencing and RNA hybridization probes. The ecotypes had highly depth-specific distributions. One SAR11 ecotype, later identified as subclade II, was more abundant in the upper mesopelagic, whereas the original SAR11 ecotype (subclade Ia) was more prevalent in the euphotic zone. They concluded that the differing depth profiles were indicative of niche partitioning. Later work with additional sequences described the structure of the SAR11 clade in greater detail (Suzuki et al[., 2001;](#page-12-0) Rappe´ [and Giovannoni, 2003\)](#page-12-0). Morris et al[. \(2005\)](#page-12-0) Received 19 May 2008; revised 20 October 2008; accepted Rappé and Giovannoni, 2003). Morris *et al.* (2005)
1 November 2008; published online 4 December 2008 then used terminal restriction fragment length

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polymorphisms (T-RFLP) to confirm earlier findings and to show that SAR11 subclade Ib was also found in surface waters.

The analogous discovery of ecotypes of Prochlorococcus (Moore et al[., 1998\)](#page-12-0) was supported by genome sequences and physiological characteristics that helped to explain their differing vertical distributions. In particular, adaptations to high-light and low-nutrients were observed in the shallow ecotypes, whereas deeper ecotypes were suited to low-light, high-nutrient environments. Studies of ecotype distributions help provide oceanographers with an understanding of how phylogenetic microdiversity impacts estimates of productivity and nutrient cycling, and are being used to develop new models to explore relationships between climate change, community structure and biogeochem-ical cycles [\(Follows](#page-11-0) et aI ., 2007; Hood et aI [., 2006](#page-12-0)).

Resolving the dynamics of SAR11 ecotypes in the ocean surface is particularly important because the flux of organic carbon through bacterioplankton is comparable to daily primary production in this environment [\(Ducklow, 2000](#page-11-0)), where SAR11 are the dominant heterotrophs. Insights into the interactions between SAR11 populations and nutrient fields have emerged from interpretations of genomic data, proteomic data and the study of cultured isolates ([Giovannoni](#page-11-0) et al., 2005a). These studies have shown that the SAR11 cells have lightdependent proton pumps and unusual nutritional requirements for reduced sulfur compounds and glycine ([Giovannoni](#page-11-0) et al., 2005b; [Tripp](#page-12-0) et al., [2008a, b](#page-12-0)). They appear to have sacrificed nutritional versatility for a streamlined metabolism ([Giovan](#page-11-0)noni et al[., 2005a](#page-11-0)). These findings were largely developed through study of cultured SAR11 isolates. However, little is known about how natural populations of SAR11 and its ecotypes vary through time and over depth in response to physical and chemical variability. Temporal variability of hydrography and biogeochemical processes is highly complex. Consequently, progress in understanding the correspondence between biogeochemical processes and microbial diversity is constrained by the practical problem of implementing experimental designs that sample with sufficiently fine scale to resolve processes (for example, Karl et al[., 2003\)](#page-12-0).

The goal of this study was to quantitatively resolve temporal patterns in SAR11 cell densities throughout the euphotic and mesopelagic zone. Here, we present a 3-year time-series of SAR11 cell distributions, as determined by FISH, collected from the upper 300 m at the Bermuda Atlantic Timeseries (BATS) site. We frame these dynamics in the context of physical and biogeochemical data from other long-term studies at the site. To resolve SAR11 ecotype population dynamics in greater detail, we used 16S rRNA gene sequence and T-RFLP data to measure the relative abundance of SAR11 subclades Ia, Ib and II, and used these data to model the ecotype population's dynamics. The data reveal

in the ocean surface layer and show that they occur reproducibly each year.

Materials and methods

Study site and sample collection

Samples were collected aboard the R. V. Weatherbird II at the BATS site $(31^\circ 40' N, 64^\circ 10' W)$. All cruises were conducted as part of the larger BATS program and sampled at least monthly with biweekly sampling each year between January and March. This sampling strategy has been successful in revealing the major temporal microbial and biogeochemical patterns [\(Carlson](#page-11-0) et al., 1996; [Steinberg](#page-12-0) et al[., 2001](#page-12-0); Morris et al[., 2005](#page-12-0)). A broader assessment of the BATS biogeochemical data is presented elsewhere (see Deep Sea Research II volumes 43 (2–3) and 48 (8–9) for relevant studies).

many new features of SAR11 population dynamics

Samples for FISH were collected from eight depths over the surface 300 m from January 2003 to December 2005. Seawater was fixed with $0.2 \mu m$ filtered formalin (10% concentration) and stored in liquid nitrogen until processing (within 2 weeks of collection). No appreciable decay of probe signal was detected when stored in liquid nitrogen for periods up to 6 months.

High molecular weight DNA was collected regularly during BATS cruises (97 separate casts) from 1 and 200 m from August 1991 to January 1994, and September 1997 to January 2004. An additional 36 profiles over 6–8 depths were sampled periodically throughout this time-series ([Table 1\)](#page-2-0). All 341 DNA samples were collected through CTD rosette or in situ pumps (McLane Research Laboratories Inc., East Falmouth, MA, USA). Sixteen to seventy liters of seawater were filtered onto $0.2 \mu m$ Supor filters (Pall Life Sciences, East Hills, NY, USA) and extracted as described earlier ([Giovannoni](#page-11-0) et al., [1996\)](#page-11-0).

Ancillary data used to provide the context for the annual patterns of mixed layer depth (MLD) and primary production are available from the public data set [\(http://www.bios.edu\)](http://www.bios.edu). The MLD during experimental periods was determined as the depth where potential density (sigma-t) of the water was equal to sea surface sigma-t plus an increment in sigma-t equivalent to a $0.2 \degree C$ temperature decrease ([Sprintall and Tomczak, 1992\)](#page-12-0). Contour plots were created in Ocean Data View (Schlitzer, R., [http://www.awi-bremerhaven.de/GEO/ODV,](http://www.awi-bremerhaven.de/GEO/ODV) 2004). Tukey–Kramer HSD statistical analyses were performed using the statistical package JMP (SAS Institute Inc., Cary, NC, USA).

FISH

Samples $(10 \,\mathrm{ml})$ were filtered onto $0.2 \,\mathrm{\mu m}$ 25 mm polycarbonate filters and stored desiccated at $-$ 20 $^{\circ}\mathrm{C}$ until hybridization. Filters were cut into quarters

Depth (m)	Month relative to timing of deep mixing													
	-1	Deep mixing	$+1$	$+2$	$+3$	$+4$	$+5$	$+6$	$+7$	$+8$	$+9$	$+10$		
			հ	6			6	8						
40														
80														
120														
160														
200	10													
250				ົ						o				
300				റ	∩		ົ		ົ	Ω				

Table 1 Annual composite of all 341 DNA samples collected between 1991 and 2004 from which T-RFLP data were generated

The values represent the number of samples collected for each month relative to the time of deep mixing at each depth throughout the DNA timeseries. 1 m and 200 m samples were sampled at the highest frequency and other depths were sampled periodically. Blanks mean no samples exist for that month or depth.

and hybridization reactions were performed on onequarter membrane sections as described in [Morris](#page-12-0) et al[. \(2002\)](#page-12-0). Briefly, four Cy3-labelled oligonucleotide probes targeting the SAR11 clade including SAR11 152R-Cy3 (5' ATTAGCACAAGTTTCCY CGTGT), SAR11 441R-Cy3, (5'-TACAGTCATTTTCT TCCCCGAC); SAR11-542R (5'-TCCGAACTACGCTA GGTC) and SAR11-732R (GTCAGTAATGATCCAGA AAGYTG). A nonsense probe 338F- (TGAGGATGC CCTCCGTCG) served as a negative control. Forty microliters of hybridization solution containing $2 \text{ ng } \mu l^{-1}$ of probe in a buffer of 900 mmol l^{-1} NaCl, 15% formamide, 20 mmol l^{-1} Tris-HCl (pH 7.4) and 0.01% sodium dodecyl sulfate (SDS) was spotted onto each quarter filter and incubated at $37 \,^{\circ}\text{C}$ in a Techne hybridizer (HB-1D, NJ) for 16 h. Optimal hybridization stringency was achieved by washing the membranes in hybridization wash $(20 \text{ mmol})^{-1}$ Tris-HCl (pH 7.4), 150 mmol l^{-1} NaCl, 5 mmol l^{-1} EDTA and 0.01% SDS) for two 10-min intervals at experimentally determined temperatures of dissociation (SAR11, 55 °C; 338F, 50 °C) in Coplin jars. The hybridized filters were then air-dried and mounted on dry slides using a mounting media containing $1.67 \,\mathrm{\mu g\,ml^{-1}}$ 0.6-diamidino-2-phenylindole dihydrochloride (DAPI) in Citiflour solution (Ted Pella Inc., Redding, CA, USA).

Microscopy and image analysis

Total prokaryotic cell abundance was determined through DAPI direct counting through epifluorescence microscopy [\(Porter and Feig, 1980\)](#page-12-0). Ten to twelve fields of view encompassing $\geqslant500$ cells per sample were counted from one of the filter quarters. Detection of CY3-positive cells and their ratio to DAPI-positive cells was aided by image analysis using an Olympus AX70 microscope $(x 1000)$ (Olympus, Japan) equipped with a Toshiba 3CCD video camera (IK-TU40A Toshiba, Japan), a computer assisted frame grabber and appropriate dichroic filters [\(Morris](#page-12-0) et al., 2002). Briefly, exposure times of 1 and 5 seconds were used for DAPI and CY3 images, respectively. CY3 images were segmented

with Image Pro Plus software (Media Cybernetics, Bethesda, MD, USA) and overlaid onto corresponding segmented DAPI images. Objects with overlapping signals in both Cy3 and DAPI images were counted as probe positive. The negative control was determined similarly and subtracted from the positive probe counts to correct for autofluorescence and non-specific binding. The percent probe positive cells were determined for 300 to >1000 DAPIpositive cells. The ratio of probe-positive to DAPIpositive cells from the hybridized filter membrane was then multiplied by total DAPI count to derive SAR11 abundances (Morris et al[., 2002](#page-12-0)). The coefficient of variability for positive probe enumeration was on the order of 15% in the euphotic zone where SAR11 comprised $>30\%$ of total DAPI counts and increased as SAR11 densities decreased over depth.

T-RFLP analysis of 16S rDNA

T-RFLP analysis was used for comparison of bacterial community composition between samples (Liu et al[., 1997\)](#page-12-0). Ribosomal RNA genes from mixed communities were amplified by PCR with Taq polymerase (Fermentas Inc., Glen Burnie, MD, USA) and the commonly used bacterial primers, 8F-FAM (5'-AGRGTTYGATYMTGGCTCAG-3'), and 519R (5'-GWATTACCGCGGCKGCTG- 3') [\(Morris](#page-12-0) $et al., 2005$. The 8F-FAM primer was $5'$ end-labeled with the phosphoramidite fluorochrome 5-carboxyfluorescein (6-FAM). Amplifications were performed with $5 \text{ ng of template DNA per } 50 \mu l$ reaction using the following conditions: 30 cycles, denaturation at 94 °C for 15 s, annealing at 55 °C for 30 s and elongation at 72 °C for 1 min. Acetamide was added to a final concentration of 5% to reduce $G + C$ content-induced bias.

PCR products were restricted for 6 hours at 37° C in reactions containing 10 mmol $\rm l^{-1}$ MgCl $_{\rm 2}$, and 10 U BsuRI restriction enzyme (Fermentas, MD, USA). PCR and restriction digest were cleaned up according to QIAquick PCR purification kit instructions (Qiagen, Valencia, CA, USA). Terminal restriction fragments (T-RFs) were resolved on an ABI 3100 Genetic Analyzer (Applied Biosystems, Forest City, CA, USA), and ABI Genescan software was used to size fragments based on MapMarker 1000 size standards (Bioventures, Murfeesboro, TN, USA). According to [Kitts \(2001\)](#page-12-0) large and small T-RFLP fragments are more comparable and accurate when integrated peak areas are used compared with peak height. To be consistent with our earlier study (Morris et al[., 2005\)](#page-12-0) we used the peak area for the determination of percent contribution of the different fragments in all downstream analyses. T-RFLP data was standardized according to [Osborn](#page-12-0) et al. [\(2006\)](#page-12-0).

T-RFLP fragments were putatively identified using the sequences from our 16S rDNA clone libraries and publicly available sequences from marine samples. The T-RFs were predicted from the available sequence data using ARB ([Ludwig](#page-12-0) et al[., 2004](#page-12-0)) and TRF-CUT (Ricke et al[., 2005](#page-12-0)). T-RF matches were verified by applying T-RFLP to identified clones (Morris et al[., 2005\)](#page-12-0). We focused specifically on T-RFLP fragments that putatively identified members of the SAR11 clade. One or more T-RFs within each electropherogram represented the predominant SAR11 subclades, Ia, Ib and II and were putatively designated as follows: SAR11 subclade Ia, 113 bp, SAR11 subclade Ib, 227, 228 and 293 bp and SAR11 subclade II 292 bp. The sum of all T-RF areas for each subclade was normalized by total SAR11 peak area to determine the relative contribution of each subclade. The phylogenetic affiliations assigned to the T-RFLP fragments in this study were the same as described in [Morris](#page-12-0) et al. [\(2005\)](#page-12-0) except that fragments 292 and 293, which coincided with subclade II and Ib, respectively, were 1 bp longer than reported by Morris et al[. \(2005\).](#page-12-0) This slight discrepancy is likely due to subtle differences in base pair calling and standardization between the two studies.

Cloning and sequencing

Clone libraries were constructed from 40 m samples collected in the vicinity of the BATS site in March 2004 and July 2004. Ribosomal RNA genes were amplified from environmental DNA for cloning by PCR as described above with the exception that triplicate PCRs were pooled following amplification. A single band of the predicted length was excised and purified from a 1% agarose gel using a QIAquick PCR purification kit (Qiagen). Clone libraries were constructed using the pGEM-T-Easy (Promega Corporation, Madison, WI, USA) vector following the manufacturer's instructions. Transformations were submitted for plating and single-pass sequencing at the High-Throughput Genomics Unit (University of Washington, Seattle, WA, USA).

Partial 16S rRNA gene clone sequences were obtained and added to an aligned database containing greater than 350 SAR11 sequences maintained

with the ARB software package ([Ludwig](#page-12-0) *et al.*, [2004\)](#page-12-0). Evolutionary distance analysis methods identified phylogenetic relationships within the SAR11 dataset, using the program PAUP* 4.0 beta 10 [\(Swofford, 2002\)](#page-12-0). The sequence from 'Candidatus Pelagibacter ubique' ([Giovannoni](#page-11-0) et al., 2005a) was used as a type of strain for alignment and fragment designation.

Nucleotide sequence accession numbers

Gene sequences were deposited in Genbank and given accession numbers EU984307 through EU984496.

Results

SAR11 abundance measurements revealed regular annual dynamics of cell density increase and redistribution in the upper 300 m that coincided with water column mixing and stratification (Figure 1a). SAR11 densities were significantly greater within the euphotic zone than in the upper mesopelagic zone. The general temporal trend for all years demonstrated a significant decrease in surface SAR11 densities upon onset of deep mixing in winter (Figure 1a). As the MLD began to shoal in

Figure 1 Contours of SAR11 cell densities (E8 cells L⁻¹) determined through FISH (a) and percentage of DAPI-positive cells identified as SAR11 (b) in the surface 300 m at BATS from January 2003 to December 2005. Mixed layer depth (MLD) estimate (dashed white line) was based on a variable sigma-t (potential density) criterion ([Sprintall and Tomczak, 1992\)](#page-12-0). MLD was determined as the depth where sigma-t is equal to sea surface sigma-t plus an increment in sigma-t equivalent to a $0.2 \text{ }^{\circ}\text{C}$ temperature decrease.

spring of each year, the SAR11 densities increased significantly (Student's *t*-test, *P* value $\langle 0.0001 \rangle$ from winter lows to mean cell densities of 2.2×10^8 cells l⁻¹. During periods of summer stratification $(MLD<50 \text{ m})$ the SAR11 maximum was most pronounced just below the MLD between 40 and 80 m and reached cell densities as high as $2.5-3.25 \times 10^8$ cells l^{-1} each year. Within the upper mesopelagic (160–300 m) SAR11 cell densities increased significantly during and following the maximum MLD, and persisted at elevated densities for months after restratification ([Figure 1a](#page-3-0)).

The contribution of SAR11 to total DAPI cell counts also varied through time and over depth. The percentage of DAPI-positive cells identified as SAR11 ranged from a mean of $33 \pm 8\%$ in the surface 120 m $(n=176)$ to \sim 13 ± 8% within the upper mesopelagic zone $(n=176)$ ([Figure 1b](#page-3-0)). In the mesopelagic zone the percent contribution of SAR11 cells to total DAPI counts increased to as high as 30% after deep mixing and then decreased to 10% during summer-stratified periods.

Annual variability of depth-integrated SAR11 stocks

An annual linear harmonic model was fit to the temporal trend of the depth-integrated SAR11 data—it explained 44% of the variability with significance $>99\%$ confidence interval (CI) throughout this time-series. Annual winter deep mixing extended the maximal MLD below the euphotic zone (0.1% light level \sim 140 m) to depths between 190 m to 240 m [\(Figure 1a\)](#page-3-0). Deep mixing entrains nutrients into the surface waters stimulating spring phytoplankton blooms that reached a productivity maximum a month after maximal MLD (Figure 2a). The annual deep mixing and subsequent phytoplankton blooms initiate important annual organic and inorganic nutrient dynamics at this site (see [Steinberg](#page-12-0) et al., 2001; [Hansell and Carlson,](#page-12-0) [2001\)](#page-12-0). The timing of deep mixing varied between early January and late February during this study. Therefore, to compare SAR11dynamics in the context of the mixing event, we collapsed the 3-year time-series into a single 12-month composite. We aligned each year's data to the timing of the deep mixing event and determined the integrated SAR11 means for the corresponding months prior to and following deep mixing (Figure 2). Integrated SAR11 composite (0-300 m) revealed a significant two-fold increase in SAR11 stocks following deep mixing (Figure 2b).

Significant regular annual dynamics of SAR11 were observed within the euphotic zone ([Figures 1](#page-3-0) [and 2c\)](#page-3-0). The integrated SAR11 stocks (0–120 m) revealed significant accumulation rates of 0.78 E6 cells $l^{-1}d^{-1}$ ($r^2 = 0.72$; $n = 5$), 0.99 E6 cells $l^{-1}d^{-1}$ $(r^2 = 0.81; n = 6)$ and 1.45 E6 cells $l^{-1}d^{-1}$ $(r^2 = 0.68;$ $n = 6$), in the springs of 2003, 2004 and 2005, respectively. The euphotic zone SAR11 maximum lagged the deep mixing event by 2 months

Figure 2 Annual composite of the depth-integrated means of (a) primary production (0–140 m) and SAR11 stocks for each month plotted relative to the deep convective mixing event. The various integration depth horizons for SAR11 include (b) the surface 300 m (c) the euphotic zone (0-120 m) (d) and the upper mesopelagic zone (160–300 m). The negative and positive values on the x axis refer to the month (s) prior to or following the deep convective mixing event. The gray bar is the reference and it represents the period of convective mixing for each year. The error bars represent s.e. of the 3-year mean for each corresponding month.

(Figure 2c). After the spring maximum, the euphotic zone SAR11 stocks decreased in the third and fourth months following deep mixing (Figure 2c). Each year after stratification, SAR11 stocks increased again, suggestive of a secondary summer maximum. However, the interannual variability of the summer maxima rendered this composite trend statistically insignificant (Figure 2c).

 $-$ 0.005 substitutions/site

Figure 3 Neighbor-joining 16S rRNA phylogenetic tree showing terminal restriction fragment sizes and relationships among representative members of the SAR11 clade. Clones obtained from this study are shown as squares and triangles for 2004, 40 m spring and summer clone libraries, respectively. Representative sequences derived from public databases are labeled with clone names and accession numbers. Bootstrap proportions over 60% from the neighbor-joining analysis are shown. SAR11 Ia, SAR11 Ib and SAR11 II subclade designations were determined by terminal restriction sites and by tree topology. Rickettsia bellii (U11014) was used as the outgroup. Terminal restriction fragment sizes of the 16S rRNA gene from clones are shown to the right of each subclade. Scale bar, 0.05 substitutions per nucleotide position. SAR11 subclade Ia, 113 bp, SAR11 subclade Ib, 227, 228 and 293 bp and SAR11 subclade II 292 bp.

Regular annual SAR11 dynamics were also observed within the upper mesopelagic zone. Mesopelagic SAR11 stocks remained \leqslant 30 \times 10¹¹ cells m⁻² from late summer $(>6$ months post mixing) through the mixing event of each year contributing $\langle 10\%$ of the SAR11 stocks in the surface 300 m. However, following deep mixing, mesopelagic SAR11 stocks increased >4 fold from the annual minimum accounting for \sim 25% of the total 0–300 m integrated SAR11 stock ([Figure 2d](#page-4-0)).

16S rDNA terminal restriction fragment designation Our 2004 spring and summer 40 m clone libraries added to existing 16S rDNA sequence data for 1 m

and 200 m (Field et al[., 1997](#page-11-0); [Suzuki](#page-12-0) et al., 2001; Morris et al[., 2005](#page-12-0)). From each library, 94 clones were screened, of which 28 and 38 were identified as members of the SAR11 clade from March 2004 and July 2004, respectively. Subclade designations for fragments predicted from sequence data and for those observed by T-RFLP analyses were determined as described by Morris et al[. \(2005\)](#page-12-0). T-RFLP assignments are shown in Figure 3 (see Materials and methods). We designated fragment 292 bp as a proxy for SAR11 subclade II; however, a few of the clones belonged to the phylum Cyanobacteria. The contribution of SAR11 subclade II was most pronounced below the euphotic zone $(\geq 160 \,\mathrm{m})$ ([Table 2](#page-6-0)), where the cyanobacteria are expected to

288

Depth	Observed T-RFLP	Putative SAR11	Month relative to the timing of deep mixing											
M	Fragment length (bp)	Subclade	-1	Mixing event	$+1$	$+2$	$+3$	$+4$	$+5$	$+6$	$+7$	$+8$	$+9$	$+10$
$\bf{0}$ 40 80 120 160 200 250 300	113	Ia	37 (6) 39 (4) 47 (9) 40(6) 21(9) 10(6) 9(2) 8(1)	39(5) 33 (7) 38(12) 29 (9) 28 (11) 17(8) 15(14) 11(7)	45(9) 37(10) 33(4) 29(13) 24 (9) 13(9) 11(5) 11(6)	51(9) 42(21) 45(14) 44 (2) 24(10) 15(8) 13(3) 12(7)	58 (9) 47(10) 43(10) 38 (11) 24(10) 13(10) 17(6) 20(5)	54 (11) 49(4) 43(11) 41(10) 25(17) 17(10) 17(6) 18(6)	61(10) 51(11) 40 (12) 39 (9) 22(10) 14(6) 17(6) 13(5)	63 (14) 53 (10) 37(4) 39 (10) 19(9) 16(3) 18(8) 16(5)	53 (12) 48(5) 38(3) 37(9) 24(6) 14(5) 12(4) 14(11)	42(9) 43 (7) 35(4) 33(14) 26(14) 15(11) 12(1) 14(6)	41(8) 43(5) 37(4) 22(2) 7(10) 12(6) 10(1) 11(4)	40(7) 43(5) 38(4) 30(14) 12(1) 11(4) 4(6) 8(1)
$\bf{0}$ 40 80 120 160 200 250 300	227, 228 and 293	Ib	58 (6) 60(1) 52 (9) 56 (3) 43 (7) 25(9) 21(12) 14(5)	54 (6) 56 (11) 51(5) 49 (7) 47(4) 37(13) 28(8) 25(10)	48 (7) 53(5) 51(2) 45(2) 50(5) 28 (14) 24 (6) 28 (9)	43(5) 47(16) 49 (14) 48(4) 36(2) 35(11) 29(14) 28 (10)	34(6) 44(8) 49(6) 47(8) 34(7) 44 (15) 29(7) 32(14)	38(8) 43 (4) 50(10) 47 (12) 26(2) 33 (10) 31(1) 30(11)	36 (10) 41 (14) 51(10) 50(8) 39 (14) 31(11) 26(8) 29 (18)	28(10) 39 (13) 52(2) 52(3) 46(15) 32(12) 27(8) 25(16)	40(12) 47(12) 53 (6) 51(8) 40(11) 25(3) 20(2) 18(3)	47 (11) 44 (8) 62(8) 55(8) 35(18) 27(12) 25(6) 30(10)	51(8) 46(11) 55(10) 47(7) 29(21) 19(7) 20(1) 22 (9)	53(7) 46(11) 53(11) 47(8) 30(2) 21(5) 19(2) 16(5)
0 40 80 120 160 200 250 300	292	\mathbf{I}	5(8) 2(3) 1(2) 4(7) 37(16) 65 (12) 70 (10) 78 (6)	7 (6) 11(6) 10(9) 22(8) 24(7) 46 (19) 57 (22) 64 (16)	7(5) 11(5) 16(3) 26(15) 27(4) 59 (22) 65(11) 61 (14)	6(6) 10(4) 7(1) 8(1) 40(11) 50(10) 59 (17) 60 (17)	8(5) 9(3) 8(2) 15(10) 42(11) 43 (14) 55(10) 48 (19)	8(7) 8(2) 6(8) 12(2) 49(15) 51 (14) 52(5) 52(16)	2(5) 8 (8) 8(8) 11(6) 39 (18) 55 (7) 58 (14) 58 (23)	9(6) 8(8) 11(2) 9(8) 35(23) 52 (12) 55(15) 60(11)	8(8) 6(8) 10(1) 12(10) 36(17) 61 (5) 68 (2) 68 (8)	11(8) 16(7) 3(5) 12(6) 38 (31) 59 (22) 63 (5) 56(16)	8 (7) 11(10) 8(7) 32(10) 64 (30) 70 (12) 70 (1) 67(13)	7(7) 9(10) 9(5) 23 (22) 58(2) 68 (7) 77 (4) 76 (6)

Table 2 Mean relative contribution (%) matrix determined from SAR11 T-RFLP fragments representative of each subclade

The annual mean composite represents means percen^t contribution for each subclade over each depth for each month relative to the time of annual mixing. The mean composite was generated from 341 T-RFLP samples collected over a 10-year period at BATS. Each year's data was aligned to the timing of the deep mixing event and are presented as the month prior to mixing through the tenth month following mixing. Values in parenthesis represen^t one s.d. of the mean. Values in bold did not have any representative samples and values were interpolated from previous and following month following. Values in italic were samples that only contained one sample and were averaged with the months prior and following to smooth data. The values for each ecotype in the same depth—month field add up to 100%.

be inconsequential ([DuRand](#page-11-0) *et al.*, 2001). However, we cannot rule out the possibility that SAR11 subclade II was overestimated in the euphotic zone due to a small contribution of cyanobacteria T-RF 292 bp. The clone libraries demonstrated a spring-tosummer transition in the relative contribution of the subclades: the spring library contained 32% subclade Ia, 50% subclade Ib and 17% subclade II; the summer library was comprised of 66% subclade Ia, 21% subclade Ib and 13% subclade II ([Figure 3](#page-5-0)).

Vertical and temporal variation in SAR11 ecotype distributions

The SAR11 oligonucleotide FISH probes hybridize broadly to the SAR11 clade. To model the contributions of the SAR11 subclades to the quantitative SAR11 cell counts, we incorporated the T-RFLP data from a decade of BATS samples. The percent contribution of each SAR11 subclade was determined from T-RFLP patterns from 1 and 200 m samples (90 and 88 samples, respectively). The decade of SAR11 T-RFLP data was collapsed to a single annual composite, again with the 12 months of each year aligned to the timing of the deep mixing event. The data yielded clear trends over depth and through time (Figure 4). The percent contribution of SAR11 subclades Ia and Ib to SAR11 surface populations averaged $48 \pm 12\%$ and $45 \pm 12\%$, respectively, whereas subclade II averaged only 7±7% of SAR11 surface populations (Figure 4a). Maxima for subclades Ia and Ib at 1 m alternated between periods of mixing (late winter early spring) and stratification (summer), resulting in a significant inverse relationship between the two subclades (Figure 4a). The seasonal transition from SAR11subclade Ib to subclade Ia in the surface waters corresponded to water column stratification and MLD shoaling to $< 50 \text{ m}$ ([Figures 1a and 4a](#page-3-0)). In contrast, subclades Ia and Ib contributed less to SAR11 populations at 200 m, where they averaged $14 \pm 7\%$ and $29 \pm 12\%$, respectively. SAR11 subclade II, dominated at 200 m, averaging $57 \pm 15\%$ of total SAR11 populations. The periodicity in relative contribution of the various subclades was less pronounced at 200 m compared with surface samples (Figure 4b).

Quantitative modeling of temporal variation in SAR11 ecotype distributions

T-RFLP patterns at 1 and 200 m indicated that combining the qualitative T-RFLP data with the quantitative FISH data could resolve the dynamics of the three major SAR11 subclades. However, the FISH data were collected from 2003 to 2005 and T-RFLP data was collected between 1991 to 2004 with an additional 36 depth profiles sampled at depths other than 1 and 200 m at a lower temporal resolution. We generated a T-RFLP percent contribution matrix from the 341 samples by determining

Figure 4 Annual composite of the percent contribution (%) of SAR11 Subclade Ia (open circles) Ib (closed circles) and II (closed triangles) to total SAR11 T-RFLP peak area (see text for details) for the 1 m (a) and 200 m (b) samples at BATS. The annual composite was generated by aligning each of the 10 years of data to the timing of the deep mixing event and taking the mean of the month prior and 10 months following the deep mixing event. The error bars represent s.e.m.

the mean subclade contribution for each depth and month, aligned to the timing of each year's deep mixing event ([Table 2](#page-6-0)). For the fifth month (post mixing) there was no data for the 40 m and 80 m depths; thus, we interpolated data between the fourth and sixth months (bold in [Table 2\)](#page-6-0).

The T-RFLP percent composition matrix ([Table 2](#page-6-0)) was used to factor the quantitative FISH data, producing models of the population dynamics of the subclades in units of cell density [\(Figure 5](#page-8-0)). [Figure 6](#page-8-0) shows the three SAR11 subclade densities integrated over various depth horizons presented as an annual composite relative to the timing of deep mixing. The asterisks in [Figure 6](#page-8-0) indicate months in which all three subclades were significantly different from each other (Tukey–Kramer HSD at 95% CI).

The integration of all SAR11 subclades over the surface 300 m predicted that subclade II was

Figure 5 Contours of modeled subclade cell densities in the surface 300 m from 2003 through 2005 for subclade Ia (a), Ib (b) and II (c). The data reported in this figure were determined by multiplying the quantitative SAR11 densities in [Figure 1a](#page-3-0) by the percent contribution matrix presented in [Table 2](#page-6-0). White dashed line represents MLD as described in [Figure 1](#page-3-0) and is used to examine distribution patterns in the context of mixing and stratification.

significantly less abundant than the other two subclades (Tukey–Kramer HSD at 95% CI), except for the first month post mixing. Subclade Ib stocks were greater than subclade Ia in the month prior through to the month following deep mixing and then became similar in magnitude within the upper 300 m (Figure 6a).

To further examine how the subclade dynamics varied over depth we partitioned the modeled data into three depth horizons—the euphotic zone (that is, 0–120 m and 0–40 m) and the mesopelagic zone (160–300 m). During periods of mixing the SAR11 densities and stocks were dominated by subclade Ib (Figures 5b and 6b), but, as the MLD shoaled to $<$ 50 m, the euphotic zone stocks of SAR11 subclades Ia and Ib became comparable (0–120 m) (Figure 6b). However, focus-

Figure 6 Annual composite of 3 years of modeled SAR11 data aligned relative to the timing of the deep convective mixing event. The data were integrated over various depth horizons (a) 0–300 m, (b) the euphotic zone $0-120$ m, (c) the surface 40 m of the euphotic zone and (d) the upper mesopelagic zone 160–300 m. Open circles are SAR11 subclade Ia, filled circles are SAR11 subclade Ib and triangles are SAR11 subclade II. Gray bar is a reference that represents the timing of the deep mixing in the composite plot. Error bars represent s.e. Asterisk represents months when all subclade stocks were significantly different from each other by Tukey–Kramer HSD (95% CI).

ing on the surface 40 m revealed a striking transition from subclade Ib dominance during the mixing period to subclade Ia dominance during the summer-stratified period (5–6 months post mixing) (Figure 6c).

Subclade II dynamics were relatively minor within the euphotic zone but became a significant contributor in the upper mesopelagic zone (160–300 m). Members of subclade II were significantly

291

greater than subclade Ia, but not subclade Ib, in the mesopelagic during the 2 months following deep mixing (Tukey–Kramer HSD at 90% CI) ([Figures 5c and 6d](#page-8-0)). Cell densities of subclade II were greatest and its relative contribution most significant between 200 and 300 m [\(Figure 5c](#page-8-0)). Integrated stocks of subclade Ib, and to a lesser extent Ia, were dynamic in the mesopelagic zone ([Figure 6d\)](#page-8-0); however, the majority of their contribution was localized between 160 and 200 m [\(Figure 5](#page-8-0)).

Discussion

Abundance

The high relative abundance of SAR11 cells we report for the BATS site is consistent with previous studies in the Sargasso Sea (Morris et al[., 2002;](#page-12-0) [Malmstrom](#page-12-0) et al., 2004), the eastern North Atlantic subtropical gyre (Mary et al[., 2006\)](#page-12-0) and the coastal oligotrophic Mediterranean Sea [\(Alonso-Saez](#page-11-0) et al., [2007\)](#page-11-0). The ubiquity and numerical dominance of SAR11, supported by genomic and culture evidence revealing their metabolism (Rappé et al[., 2002;](#page-12-0) [Giovannoni](#page-11-0) et al., 2005a) and cell-specific radioactive substrate uptake experiments ([Malmstrom](#page-12-0) et al[., 2004;](#page-12-0) Mou et al[., 2007](#page-12-0)) is compelling evidence that these cells play a major role in the oxidation of organic carbon in the oceans.

Evidence for ecotypes: vertical stratification and seasonality

Ecotypes are similar phylogenetic subgroups of bacteria that differ in physiological details that determine niche specificity ([Cohan, 2006\)](#page-11-0). Patterns of variation in time and space are often used to identify ecotypes. Subclades Ia and Ib are 3–4% different in 16S rRNA sequence, making them different species by commonly employed standards. SAR11 subclade II differs in 16S rRNA gene sequence by 10–11% from the SAR11 Ia and Ib, which is more than enough sequence divergence to justify placing them in a different genus or family. It is assumed that divergence of ecotypes is caused by selection for specific traits, although in most cases the traits are unknown [\(Koeppel](#page-12-0) et al., 2008; Hunt *et al.*, 2008). The best example is *Prochlor*ococcus, where the distributions of ecotypes and some traits that contribute to niche specificity are both known (Moore et al[., 1998](#page-12-0)).

The pronounced vertical and seasonal patterns in the SAR11 subclade distributions we observed provide evidence of at least three SAR11 ecotypes. All three of these ecotypes were reported previously (Suzuki et al[., 2001;](#page-12-0) Morris et al[., 2005\)](#page-12-0), but this analysis revealed many previously unseen details of their ecology. Our new findings show that SAR11 subclade Ia is not just a surface ecotype, but an ecotype that blooms in the extreme environment of the summer mixed layer, where sunlight (UV) is

intense [\(Zafiriou](#page-12-0) et al., 2008) and macronutrients are scarce [\(Steinberg](#page-12-0) et al., 2001). Recent genome sequence data from a Sargasso Sea summer isolate of SAR11 subclade Ia (strain HTCC7211; unpublished data) indicated that they have nutrient acquisition systems that are not present in coastal strains. Likewise, subclade II is not only a mesopelagic ecotype; it specifically blooms in the upper mesopelagic zone following deep mixing. SAR11 Ib is prominent throughout the surface 300 m year round, but becomes dominant in the euphotic zone during periods of water column instability and blooms in the mesopelagic after mixing. We observed a rhythmic succession each year of subclade Ia replacing Ib in the upper 40 m with the progression of stratification, with the maximum in Ib populations moving downward to 80m at the height of summer stratification ([Figures 5 and 6](#page-8-0)). There could be finer patterns of evolutionary divergence within the SAR11 clade at BATS, for example the three subclades we delineate could each be a composite of related ecotypes that further subdivide the habitat, but the data could not resolve this.

The partitioning of microbial populations with depth is a dominant theme in microbial plankton ecology. For example, partitioning between the euphotic and aphotic regions has been amply documented for bacterioplankton processes ([Carlson](#page-11-0) et al[., 2004](#page-11-0); [Reinthaler](#page-12-0) et al., 2006) and for many major microbial plankton clades [\(Gordon and](#page-11-0) [Giovannoni, 1996](#page-11-0); [Giovannoni](#page-11-0) et al., 1996; [Wright](#page-12-0) et al[., 1997](#page-12-0); DeLong et al[., 2006\)](#page-11-0). Evolutionary adaptive radiation into ecotypes that are specialized to specific depth horizons has also been previously described as niche partitioning for SAR11 and Prochlorococcus (Field et al[., 1997](#page-11-0); [Moore](#page-12-0) et al., [1998\)](#page-12-0). Many factors that impact microbial metabolism, including light, temperature and organic and inorganic nutrient availability, vary in consistent patterns with depth. Below, we discuss the potential significance of one of the least obvious of these factors—the export and vertical stratification of dissolved organic matter (DOM).

We note however, that all of our evaluations were made in two dimensions—depth and time—because we had inadequate spatial data to assess the impact of horizontal mixing and advective processes to the results. We, therefore, cannot rule out the possibility that horizontal advection interfered with some of the temporal variability we observed. Nonetheless, we demonstrated that the dynamics of SAR11 stocks fit an annual harmonic model with a high level of significance, indicative of regular seasonal variability.

Geochemical patterns at BATS and SAR11 metabolism The factors that determine the distributions of SAR11 populations are unknown, but information is accumulating that may eventually lead to a mechanistic understanding of how physical, chemical and biological processes combine to cause the reproducible population dynamics we observed. So far the genomic and physiological properties that distinguish the ecotypes of SAR11 have not been reported; most of what is known was determined from the study of coastal isolates from subclade Ia [\(Giovannoni](#page-11-0) et al., 2005a; Tripp et al[., 2008a, b](#page-12-0)) and metagenomic and metaproteomic data ([Venter](#page-12-0) *et al.*, [2004; Wilhelm](#page-12-0) et al., 2007; Sowell et al[., 2008\)](#page-12-0). These organisms have the smallest genomes known for free-living heterotrophic cells, an apparent consequence of genome streamlining and reduction driven by the selection for efficient growth in oligotrophic ocean habitats [\(Giovannoni](#page-11-0) et al., [2005a\)](#page-11-0). Mary et al[. \(2006\)](#page-12-0) found that SAR11 dominated the low nucleic acid fraction of bacterioplankton in the North Atlantic subtropical gyre, consistent with the prediction that small genomes are common to this clade.

The unusual requirements of SAR11 for reduced sulfur compounds (Tripp et al[., 2008a\)](#page-12-0) and glycine or serine (Tripp et al[., 2008b\)](#page-12-0) may play a role in spatial and temporal control of SAR11 populations, although too little chemical data exists at present to test this hypothesis. The proteome composition of SAR11 appears to be another facet of a survival strategy related to small cell dimensions [\(Sowell](#page-12-0) et al[., 2008](#page-12-0)). Recent environmental proteomic data from a summer surface sample at BATS demonstrate that SAR11 ecotypes highly express genes for highaffinity transport systems, involved with phosphate, amino acid and carbohydrate uptake, relative to other genes ([Wilhelm](#page-12-0) et al., 2007; [Sowell](#page-12-0) et al., [2008\)](#page-12-0). The unusual regulation of the SAR11 glyoxylate cycle reported by Tripp et al[. \(2008b\)](#page-12-0) supports the interpretation that osmolytes, amino acids and other precursors are important sources of DOM fueling SAR11 metabolism.

Interactions between bacterioplankton cells and the natural DOM pool are important, but difficult to directly measure, although DOM quality has been shown to affect the community structure of heterotrophic bacterioplankton [\(Cottrell and Kirchman,](#page-11-0) [2000\)](#page-11-0). Although it has not yet been possible to directly measure heterotrophic interactions between microbial populations and DOM at a comprehensive molecular level, described patterns of DOM concentration, composition and reactivity in the surface layer suggest they are factors that influence distributions of SAR11 populations.

Photoautotrophy and subsequent food web interactions provide a source of new DOC to the surface waters. A significant portion of daily DOM production is labile, cycled rapidly (that is, minutes to hours) ([Nagata, 2000](#page-12-0); [Carlson, 2002\)](#page-11-0) and does not accumulate appreciably in stratified surface waters (that is, labile monomers $\langle 5 \text{ nM} \rangle$ ([Keil and Kirch](#page-12-0)[man, 1999\)](#page-12-0). The tight coupling between production and contemporaneous uptake of labile compounds largely fuels heterotrophic bacterial production within the euphotic zone [\(Ducklow, 2000\)](#page-11-0) leaving behind a recalcitrant or 'semi-labile' pool of DOM resistant to rapid microbial degradation by surface populations [\(Hansell and Carlson, 2001;](#page-12-0) [Carlson](#page-11-0) et al[., 2004](#page-11-0)). Members of SAR11 ecotypes Ia and Ib appear to be competitive at utilizing labile organic compounds under low inorganic nutrient conditions. Previous work has demonstrated that surface SAR11 out-compete other bacterioplankton for labile compounds such as amino acids, sugars, aromatic monomers and DMSP ([Malmstrom](#page-12-0) et al., [2004;](#page-12-0) Mary et al[., 2006](#page-12-0); Mou et al[., 2007](#page-12-0)).

Our findings also suggest that SAR11 ecotypes II and Ib are implicated in the continued diagenetic alteration and remineralization of euphotic zone produced semi-labile DOM after it is exported from the surface to the upper mesopelagic ([Figure 5c\)](#page-8-0). Although resistant to rapid microbial remineralization within the euphotic zone, the seasonally produced semi-labile DOM is more 'bioreactive' compared to the resident mesopelagic DOM [\(Gold](#page-11-0)berg et al[., 2008\)](#page-11-0). Once exported to the upper mesopelagic, a fraction of the surface semi-labile DOM is oxidized and diagenetically altered ([Carlson](#page-11-0) et al[., 1994](#page-11-0); [Hansell and Carlson, 2001](#page-12-0); [Carlson](#page-11-0) et al[., 2004](#page-11-0); [Goldberg](#page-11-0) et al., 2008). A bloom of SAR11, marine Actinobacteria, SAR202 and OCS116 occurs in the mesopelagic ([Morris](#page-12-0) et al., [2005\)](#page-12-0) coincident with this spring episode of DOM oxidation ([Hansell and Carlson, 2001\)](#page-12-0), suggesting that the exported DOM provides substrates that drive the spring increases in SAR11 subclade we report here, as well as blooms of other bacterioplankton ([Carlson](#page-11-0) et al., 1996; [Steinberg](#page-12-0) et al., 2001; [Carlson](#page-11-0) et al., 2004).

It seems very likely that the supply, quality and quantity of DOM is different for each of the SAR11 subpopulations we observed, but further experimentation will be needed to measure patterns in the use of specific organic compounds by SAR11 populations.

Further considerations

The time-series data presented here demonstrate the power of combining quantitative FISH abundance data with relative abundance T-RFLP data to assess transitions of microbial ecotypes in time and space. Several recent time-series studies have demonstrated recurring interannual patterns in bacterioplankton community structure linked to regular ecosystem variability (Morris et al[., 2005](#page-12-0); [Fuhrman](#page-11-0) et al[., 2006](#page-11-0)), but the relative numbers provided by ARISA and T-RFLP methods do not provide accurate estimates of cell abundance or biomass. FISH provides a quantitative measure of cell density (Morris et al[., 2002\)](#page-12-0), but it is a targeted approach. The combination of the two forms of data, as we do here, provides estimates of cell numbers that are often critical for geochemical models, but also yield insights into population dynamics.

Significant insight can be gained from such a combination of these approaches. However, there are limitations that must be considered. First,

The ISME Journal

despite 3 years of relatively high frequency sampling, the temporal coverage is still considered short to predict long-term trends by traditional time-series analyses. Second, primer specificity, high 'species' richness and biases associated with PCR amplification and the restriction digest can lead to over or underrepresentation of specific ribotypes or the generation of 'pseudo-T-RFs' (Suzuki et al[., 1998;](#page-12-0) Egert and Friedrich, 2003; Castle and Kirchman, 2004). Third, phylogenetic resolution by T-RFLP may not be sufficient to resolve genetically distinct populations, thus the distributions of any one of the three SAR11 ecotypes we present may be the sum of additional distinct subclades of SAR11.

Comparative studies of the physiological properties and genomes of cultured isolates and metagenomics have been effective tools for identifying specific mechanisms that link microbial ecotypes with environmental variability. These tools, combined with controlled experiments using natural assemblages and isolates, will test if factors such as macronutrient availability, organic matter quality or light spectrum are responsible for controlling the response of SAR11 ecotypes. Further insights provided by the continued combination of field and experimental approaches will lead to a better understanding of the relationship between gene and genome sequence diversity and the biogeochemical properties of microbial plankton cells.

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