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ORIGINAL ARTICLE

Seasonal dynamics of active SAR11 ecotypes in the oligotrophic Northwest Mediterranean Sea

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A seven-year oceanographic time series in NW Mediterranean surface waters was combined with pyrosequencing of ribosomal RNA (16S rRNA) and ribosomal RNA gene copies (16S rDNA) to examine the environmental controls on SAR11 ecotype dynamics and potential activity. SAR11 diversity exhibited pronounced seasonal cycles remarkably similar to total bacterial diversity. The timing of diversity maxima was similar across narrow and broad phylogenetic clades and strongly associated with deep winter mixing. Diversity minima were associated with periods of stratification that were low in nutrients and phytoplankton biomass and characterised by intense phosphate limitation (turnover time < 5 h). We propose a conceptual framework in which physical mixing of the water column periodically resets SAR11 communities to a high diversity state and the seasonal evolution of phosphate limitation competitively excludes deeper-dwelling ecotypes to promote low diversity states dominated (> 80%) by SAR11 *la*. A partial least squares (PLS) regression model was developed that could reliably predict sequence abundances of SAR11 ecotypes ($Q^2 = 0.70$) from measured environmental variables, of which mixed layer depth was quantitatively the most important. Comparison of clade-level SAR11 rRNA:rDNA signals with leucine incorporation enabled us to partially validate the use of these ratios as an *in-situ* activity measure. However, temporal trends in the activity of SAR11 ecotypes and their relationship to environmental variables were unclear. The strong and predictable temporal patterns observed in SAR11 sequence abundance was not linked to metabolic activity of different ecotypes at the phylogenetic and temporal resolution of our study.

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Introduction

An important goal in microbial oceanography is to link the extensive diversity of marine bacteria (Venter *et al.*, 2004; Sogin *et al.*, 2006) to environmental conditions (Morris *et al.*, 2005; Fuhrman *et al.*, 2006; Gilbert *et al.*, 2012), ecosystem function (Fuhrman, 2009; Galdos *et al.*, 2011) and climate sensitivity (Eiler *et al.*, 2009; Giovannoni and Vergin, 2012). Combined time-series of microbial community diversity and oceanographic parameters have emerged as a useful approach to understand how prokaryotic communities are distributed across ecological niches (Carlson *et al.*, 2009; Hugoni *et al.*, 2013). Temporal dynamics of marine bacteria over

short timescales are reasonably well studied and strong seasonal transitions in community structure are evident across different environments (Alonso-Sáez *et al.*, 2007; Campbell *et al.*, 2009; Lami *et al.*, 2009). However, there are few extended microbial time-series (> 4 years), which are necessary to establish predictable and statistically robust patterns in microbial community dynamics.

The long-term microbial studies conducted to date have revealed annually recurring patterns of surface bacterial communities. In Californian coastal waters these patterns can be reliably predicted from seasonality in environmental parameters (Fuhrman *et al.*, 2006; Chow *et al.*, 2013). In the Sargasso Sea bacterial dynamics exhibit a strong seasonal component that appears to be driven by convective overturning (Morris *et al.*, 2005; Carlson *et al.*, 2009; Treusch *et al.*, 2009). Repeatable seasonality in bacterial diversity is also evident from the Western English Channel (Gilbert *et al.*, 2012) and appears to be a general feature common of mid- to high-latitude environments (Ladau *et al.*, 2013). Although seasonal

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cycles of bacterial diversity are an emerging paradigm in microbial oceanography, the factors controlling these cycles and their significance for ecosystem function are poorly understood. Furthermore, it is unknown whether seasonality in the metabolic activity of different bacterioplankton clades follow similar cycles.

Estimating the metabolic activity of phylogenetically defined bacteria from mixed communities is challenging. Ribosomal RNA (rRNA) is commonly used to characterise active microbial populations (Troussellier *et al.*, 2002; Gentile *et al.*, 2006) and the frequently observed discrepancies in profiles of rRNA and ribosomal RNA genes (rDNA; Moessender *et al.*, 2001; Lami *et al.*, 2009) have been interpreted to represent different metabolic states (Treusch *et al.*, 2009; Brettar *et al.*, 2011). The basis of this approach is the observation that cell ribosome content increases with growth rate (DeLong *et al.*, 1989; Fegatella *et al.*, 1998; Kerkhoff and Kemp, 1999). More recent application of this principle has used rRNA/rDNA ratios as an *in-situ* proxy for growth/activity rates (Campbell *et al.*, 2011; Campbell and Kirchman, 2013; Hunt *et al.*, 2013).

Owing to the association of particular metabolic activities to certain clades, changes in the activity of specific, and in particular dominant, bacterial communities have the potential to affect ocean carbon and nutrient cycling. In this context the SAR11 clade of the Alphaproteobacteria is of particular interest because it is a dominant component of marine bacteria (Morris *et al.*, 2002). Ubiquitously distributed throughout the ocean (Giovannoni and Rappé, 2000) SAR11 typically accounts for 15–35% of all prokaryotic plankton (Malmstrom *et al.*, 2005, 2007; Lefort and Gasol, 2013). Furthermore, SAR11 are known to take up a variety of monomeric substrates including amino acids, glucose, ATP and dimethylsulfonylpropionate (Malmstrom *et al.*, 2004; Alonso-Sáez and Gasol, 2007; Laghdass *et al.*, 2012), and appears to be an important organism for a variety of biogeochemical functions. In oligotrophic environments such as the NW Mediterranean Sea, SAR11 can dominate bacterial production. Contributions towards total leucine assimilation may reach 60–70% in nutrient-limited summer months (Laghdass *et al.*, 2010, 2012). The apparent importance of SAR11 to ecosystem function in oligotrophic environments may stem from adaptations to low substrate conditions: a streamlined genome (Giovannoni *et al.*, 2005a; Grote *et al.*, 2012) and the use of light-harvesting proteorhodopsins (Giovannoni *et al.*, 2005b) may convey significant ecological advantages.

Despite these important generalisations concerning the abundance and activity of SAR11 as a clade, it is a diverse group (Field *et al.*, 1997; Garcia-Martinez and Rodriguez-Valera, 2000; Suzuki *et al.*, 2001; Brown and Fuhrman, 2005) comprised of multiple sub-clades (Morris *et al.*, 2005; Rusch *et al.*, 2007; Brown *et al.*, 2012). 16S rRNA

gene sequencing and molecular fingerprinting approaches have shown SAR11 sub-clades to be partitioned by season, depth and biome (Field *et al.*, 1997; Garcia-Martinez and Rodriguez-Valera, 2000; Stingl *et al.*, 2007; Carlson *et al.*, 2009; Beman *et al.*, 2011; Brown *et al.*, 2012). This ecological niche separation of SAR11 sub-groups *Ia*, *Ib* and *II*, led to their classification as ecotype-like clades, with potential for fine-scale phylogenetic structure within these groups (Vergin *et al.*, 2013a). The clear temporal and spatial patterns observed among SAR11 communities are thought to result from the variable response of specific ecotypes to different environmental conditions (Beman *et al.*, 2011; Carlson *et al.*, 2009; Brown *et al.*, 2012). However, there are no direct attempts to link seasonal trends in SAR11 ecotype abundance and activity over an extended microbial time-series. To understand the role of these clades in microbial community assembly and ecosystem function (Koeppel *et al.*, 2008) it is necessary to differentiate patterns of abundance and activity in SAR11 ecotypes.

In the present study we sequenced rRNA genes (named rDNA hereafter) and rRNA of surface bacterial communities from a 7-year time-series at monthly resolution in the NW Mediterranean. Short-term (<2 years) studies in coastal waters of the NW Mediterranean have shown important seasonal transitions among bacterial communities and general significance of the entire SAR11 clade (Schauer *et al.*, 2003; Alonso-Sáez *et al.*, 2007; Lami *et al.*, 2009). The aim of the current study was to characterise the seasonal dynamics of SAR11 diversity and ecotype relative abundance and activity at an offshore site. Our main objectives were to determine (i) if total bacterial diversity and that within the SAR11 clade varies seasonally (ii) if dominant ecotypes show specific seasonal patterns in relative abundance and (iii) whether certain biogeochemical parameters constrain ecotype abundance and activity.

Materials and methods

Sampling

Surface seawater (3–5 m) was collected at the Microbial Observatory of the Laboratoire Arago located at 47°27'136 N, 03°32'360 E in the North-West Mediterranean. Samples were collected approximately every month from July 2004 to April 2011. Occasionally samples were not collected due to poor weather conditions. Full sampling details are given in Supplementary Information and exact sampling dates are summarised in Supplementary Table 1. Depth profiles of conductivity, temperature and pressure were obtained with a Sea Bird SBE 19 profiler.

Chemical and physical data

Nutrients. Concentrations of nitrate, nitrite, phosphate and silicate were measured with a Skalar

auto-analyser following a previously described protocol (Treguer and Le Corre, 1975).

Ammonium. Dissolved ammonium was measured by spectrophotometry at 630 nm following conversion to indophenol via a monochloroamine intermediate (Solorzano, 1969).

Chlorophyll. One litre of seawater was collected on a GF/F filter at low pressure (<0.2 bar). Measurements were conducted according to previously published protocols (Yentsch and Menzel, 1963; Strickland and Parsons, 1997). Extended protocol details can be found in Supplementary Information.

Mixed layer depth (MLD). Raw seabird data were processed to bin temperature and salinity data at 0.25 m intervals. MLD is based on a 0.1 °C and/or 0.02-sigma difference from a reference threshold (typically an average of the upper 3 m).

Microbial abundances

Unfiltered seawater samples were fixed at a final concentration of 1% glutaraldehyde, incubated for 15 min at ambient temperature in the dark, frozen in liquid nitrogen and stored at -80 °C. Cytometry analyses were performed on a Becton Dickinson FacsCalibur. Full details of preparation and analysis of cytometry samples can be found in Supplementary Information.

Microbial rate measurements

Bacterial heterotrophic production. The incorporation of ³H-leucine was measured using the centrifugation method (Smith and Azam, 1992; Kirchman, 1993). Extended protocol details can be found in Supplementary Information.

SAR11 bacterial production. To determine bacterial production by SAR11 cells, fluorescence *in-situ* hybridisation was coupled with microautoradiography (MICRO-CARD-FISH). Identical to measurements for bulk bacteria, tritiated leucine was used as a precursor for SAR11 bacterial production (for example, Laghdass *et al.*, 2012). Extended protocol details can be found in Supplementary Information.

Phosphate turnover. These measurements were performed during 2012 at a location in the NW Mediterranean in close proximity (42°31'N, 03°11'E) to station Microbial Observatory of the Laboratoire Arago (47°27'N, 03°32'E). The turnover of phosphate by microbial populations was determined using a concentration series bioassay (Wright and Hobbie,

1966; Zubkov *et al.*, 2007). Extended protocol details can be found in Supplementary Information.

Nucleic acid extraction, reverse transcription-PCR and pyrosequencing

The extraction of nucleic acids (DNA and RNA) was performed according to a previously published method (West *et al.*, 2008). A combination of mechanic (freeze-thaw) and enzymatic cell lysis techniques were applied directly to Sterivex cartridges, followed by extraction using an AllPrep DNA/RNA kit (Qiagen, Venlo, Netherlands). The RNA samples were tested for the presence of contaminating genomic DNA by PCR and then reverse-transcribed with random primers using the SuperScript III Reverse Transcriptase kit (Invitrogen, Carlsbad, CA, USA). We are unable to account for potential biases associated with reverse transcription efficiency and RNA secondary structure within the present data set. The amplification of the V1-V3 region of the 16S rRNA gene was performed with universal bacterial primers 28F (5'-TTTGATCNTGG CTCAG-3') and 519R (5'-GTNTTACNGCGGCKG CTG-3'), followed in pyrosequencing in a commercial laboratory (Research and Testing Laboratory, Lubbock, TX, USA) using a Roche 454 GS-FLX system (Roche, Basel, Switzerland) with titanium chemistry. All sequences have been submitted to the sequence read archive under the Bioproject accession number: PRJNA235253.

Sequence data analyses

Sequences were analysed as previously described (Blanquer *et al.*, 2013). Briefly, the reads were first filtered by removing low quality reads and subsequently trimmed to remove those with ≥3% of bases with Phred values <27 (0.2% per-base error probability). This is recommended to ensure that when clustering at 97% the influence of erroneous reads is minimised (Huse *et al.*, 2010; Kunitz *et al.*, 2010). Sequences were then clustered at a 97% threshold using Uclust algorithm (Edgar, 2010) and each operational taxonomic unit (OTU) classified by comparison to the Greengenes database (DeSantis *et al.*, 2006). Sequence analyses were conducted with Pyrotagger (Kunitz and Hugenholtz, 2010).

A phylogenetic tree was constructed with the sequence representatives of the abundant SAR11 OTUs (<0.2%). Sequences included in the tree were deposited in GenBank under accession numbers KJ094527-KJ094552. Representative sequences for each OTU together were aligned using MUSCLE (Edgar, 2004) and phylogenetic analyses were completed with the programme PHYLIP (Felsenstein, 2008). DNADIST was used to calculate genetic distances using a Kimura-2 model. The distance tree was estimated with the phylogenetic algorithm FITCH in the PHYLIP programme.

To compare bacterial communities for diversity analysis, all samples were randomly re-sampled to the size of the sample containing fewest sequences ($n = 446$) using Daisy Chopper (Gilbert *et al.*, 2009). The Shannon diversity index (H') was calculated using the software PAST (Hammer *et al.*, 2001).

Statistical analyses and partial least squares regression (PLSR)

Unpaired *t*-tests. Samples were grouped into high or low diversity regimes using a constant statistical threshold determined empirically across the 7-year data set. The two categories contained sample populations of 13 and 12, respectively. Average values and standard deviations for each measured environmental parameter were determined and unpaired *t*-tests were used to calculate statistical differences between individual parameters within the two groups. The null hypothesis (H_0) was rejected if the *P*-value was < 0.05 (GraphPad, GraphPad Software Inc, San Diego, CA, USA). A relatively large number of independent comparisons were made and we therefore applied Benjamini–Hochberg False Discovery Rate to calculate an adjusted *P*-value (Benjamini and Hochberg, 1995), which we denote as P^* in Table 1. Results are statistically significant if $P^* < 0.05$.

Co-variance (COV). To determine the nature of COV between the rRNA/rDNA ratios of different ecotypes, the following formula was used:

$$\text{COV}(x,y) = \left[\sum (x_i - \bar{x})(y_i - \bar{y}) / (n - 1) \right]$$

Where x and y are dependent and independent variables and n is the number of observations.

Values > 0 and < 0 indicate positive and negative COV, respectively.

PLSR. Models were developed based on projection to latent structures using PLSR. PLSR was selected as a predictive tool over multiple linear regression because the latter is only valid when factors are not significantly redundant (co-linear). PLSR linearly extracts few latent variables that are most useful in modelling the response. We used partial least squares (PLS) to predict multiple Y values (SAR11 ecotype sequence abundances) from a set of X variables (environmental parameters). The model was constructed using a 4-year sub-set (2007–2011) of the 7-year data, for which all environmental parameters were available (Supplementary Table 1). All analyses were performed with UnscramblerX (Camo, Oslo, Norway). Input data were auto-scaled by dividing each x and y value by its standard deviation. We used the Non-Linear Iterative Partial Least Squares algorithm. Cross-validated R^2 values (Q^2) were taken as a measure of the models Goodness of Fit (Wold *et al.*, 2001). The PLSR model we developed was optimised with a Q^2 value of 0.70 at three latent factors (Supplementary Figure 1). Weighted regression coefficients were calculated for each x -variable and SAR11 sequence abundance data set (that is, $Y_{1,4}$) to ascertain the most important measured environmental variables describing SAR11 ecotype abundance.

Inter-quartile range. Using the inter-quartile range principle [$Q_{75} + c(Q_{75} - Q_{25})$], where Q_{25} and Q_{75} are lower and upper quartiles and c is a positive constant, one of the $\mu\text{CARD-FISH}$ data points was identified as a potential outlier. This single data point was removed when calculating regression coefficients.

Results

Sequence distributions

We pyrosequenced 57 16S rDNA and 50 16S rRNA

Table 1 Comparison of biogeochemical variables between low and high diversity regimes

Parameter	Low diversity	High diversity	P	P^*
Day length	12.8 ± 0.5	9.1 ± 1.2	< 0.05	< 0.05
Temperature (°C)	19.5 ± 3.1	14.2 ± 2.0	< 0.05	< 0.05
Salinity (p.s.u.)	37.9 ± 0.3	38.2 ± 0.2	< 0.05	< 0.05
Mixed layer	12.9 ± 9.1	82.9 ± 31.2	< 0.05	< 0.05
Chlorophyll ($\mu\text{g l}^{-1}$)	0.18 ± 0.19	0.38 ± 0.23	< 0.05	NS
Nitrate ($\mu\text{mol l}^{-1}$)	0.15 ± 0.23	1.80 ± 1.47	< 0.05	< 0.05
Nitrite ($\mu\text{mol l}^{-1}$)	0.02 ± 0.01	0.12 ± 0.08	< 0.05	< 0.05
Ammonium ($\mu\text{mol l}^{-1}$)	0.17 ± 0.23	0.10 ± 0.08	NS	NS
Phosphate ($\mu\text{mol l}^{-1}$)	0.04 ± 0.01	0.06 ± 0.06	NS	NS
Silicate ($\mu\text{mol l}^{-1}$)	0.64 ± 0.41	1.8 ± 0.9	NS	NS
Bacterial abundance ($10^8 \text{ cells l}^{-1}$)	6.5 ± 2.7	4.8 ± 0.9	NS	NS
Bacterial production ($\text{pmol l}^{-1} \text{ h}^{-1}$)	25.9 ± 21.4	5.5 ± 3.6	NS	NS
<i>Synechococcus</i> abundance ($10^7 \text{ cells l}^{-1}$)	3.2 ± 4.0	1.1 ± 1.5	NS	NS
<i>Prochlorococcus</i> abundance ($10^6 \text{ cells l}^{-1}$)	3.7 ± 6.4	5.9 ± 4.4	NS	NS
Picoeukaryote abundance ($10^6 \text{ cells l}^{-1}$)	1.8 ± 2.1	7.3 ± 8.8	NS	NS
Nano-eukaryote abundance ($10^5 \text{ cells l}^{-1}$)	7.2 ± 3.3	1.3 ± 1.2	NS	NS

All averages are arithmetic means and errors are ± 1 standard deviation. The two-tailed *P*-values are computed from Welch's *t*-test. The null hypothesis was rejected if *P*-value was < 0.05 . A Benjamini–Hochberg False Discovery Rate correction was applied to calculate an adjusted P^* value. Non-significant results are denoted as NS.

samples collected as part of an offshore monthly time-series in the North-West Mediterranean. A total 66 705 16S rDNA and 57 524 rRNA sequences were obtained after performing quality filtering. Sequences of rRNA and rDNA from all sampling dates were pooled and clustered to yield a total of 2222 OTUs. The mean number of sequences obtained per sample was 1170 ± 470 for rDNA and 1150 ± 411 for rRNA. Descriptive statistics summarising the central tendencies and dispersion are provided in Supplementary Table 2. The vast majority of sequences were assigned to either members of the SAR11 clade or classified as members of the Cyanobacteria phylum. Classification of Cyanobacteria into lower taxonomic ranks was not possible using the sequenced region. The contribution of SAR11 to total sequences had medians of 78 and 45% in the rDNA and rRNA fractions compared with 13 and 46% for Cyanobacteria.

Total diversity and empirical identification of diversity regimes

Diversity in each individual sample (Shannon index) was estimated using the rDNA data set after random re-sampling. The Shannon diversity index ranged from 1 to 3.5 and a distinct seasonal cycle in diversity was observed with maxima occurring in the winter and minima in the summer (Figure 1). Diversity within the SAR11 group was calculated in a similar way without re-sampling. The median number of SAR11 sequences in diversity calculations was 344 ($-0.68\sigma = 209$; $+0.68\sigma = 388$). The SAR11 Shannon index ranged from 0.03 to 2.3. The number of SAR11 OTUs ranged from 1 to 65.

The observed seasonality in bacterial diversity was used to define an empirical framework of high and

low diversity regimes. Individual sampling dates were classified as high or low diversity by comparison to a constant statistical threshold of 2.08 ± 0.55 (arithmetic mean $\pm 1\sigma$). The exception to this is the high diversity peak of 2005/2006, which was slightly below the threshold criterion, nevertheless it was included to ensure at least one representative from the different diversity regimes were available for each sampling year. Partitioning the data in this way identified 25 sampling dates consisting of the 13 highest and 12 lowest estimates of diversity (Figure 1).

Biogeochemical conditions characterising the diversity regimes

Pairwise *t*-tests were used to determine whether the operationally defined diversity regimes were also segregated based on biogeochemical conditions (Table 1). The most significant differences in biogeochemical parameters among these regimes ($P < 0.0001$) were deeper mixed layers and lower temperatures at times of high diversity. In these samples, nitrate ($P = 0.001$), chlorophyll ($P = 0.03$) and nitrite ($P = 0.0003$) concentrations were significantly higher. The magnitude of the differences and significance levels for environmental comparisons are summarised in Table 1.

SAR11 ecotype distributions

SAR11 *Ia* ecotypes were the dominant SAR11 clade in all samples and accounted for $>60\%$ of total SAR11 rDNA sequences in high and low diversity regimes (Figure 2). Percentages of SAR11 *Ib* and SAR11 *II* rDNA increased from 3.3 ± 2.8 and 2.4 ± 2.8 at times of low diversity to 13.4 ± 4.2 and 6.6 ± 3.2 at

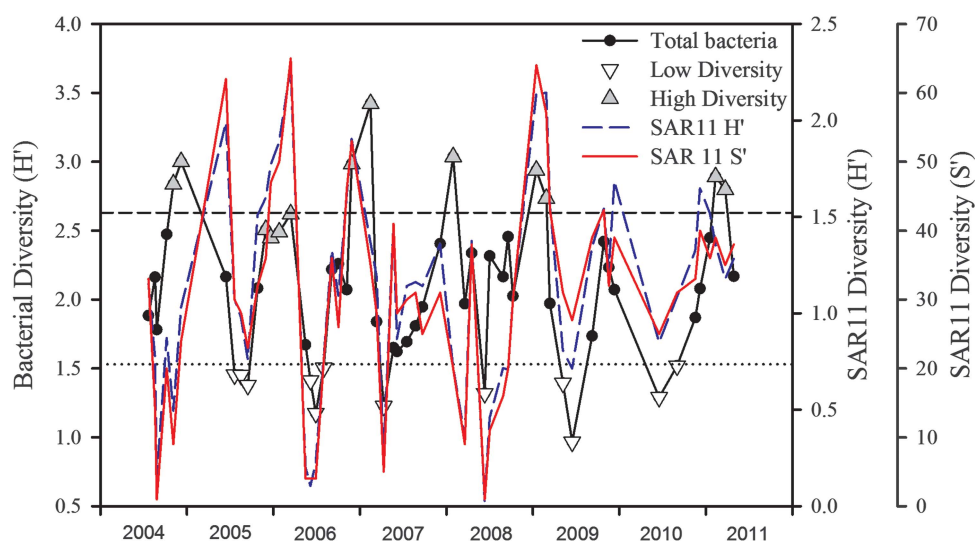


Figure 1 Seasonal cycles in total and SAR11 diversity. Shannon diversity index is calculated from tag sequence data of ribosomal RNA gene copies (rDNA). The horizontal dashed and dotted lines are empirical thresholds for high and low diversity regimes, which are marked by grey and white triangles, respectively. The dashed blue and solid red lines are SAR11 diversity expressed as Shannon index (H') and OTU richness (S'), respectively.

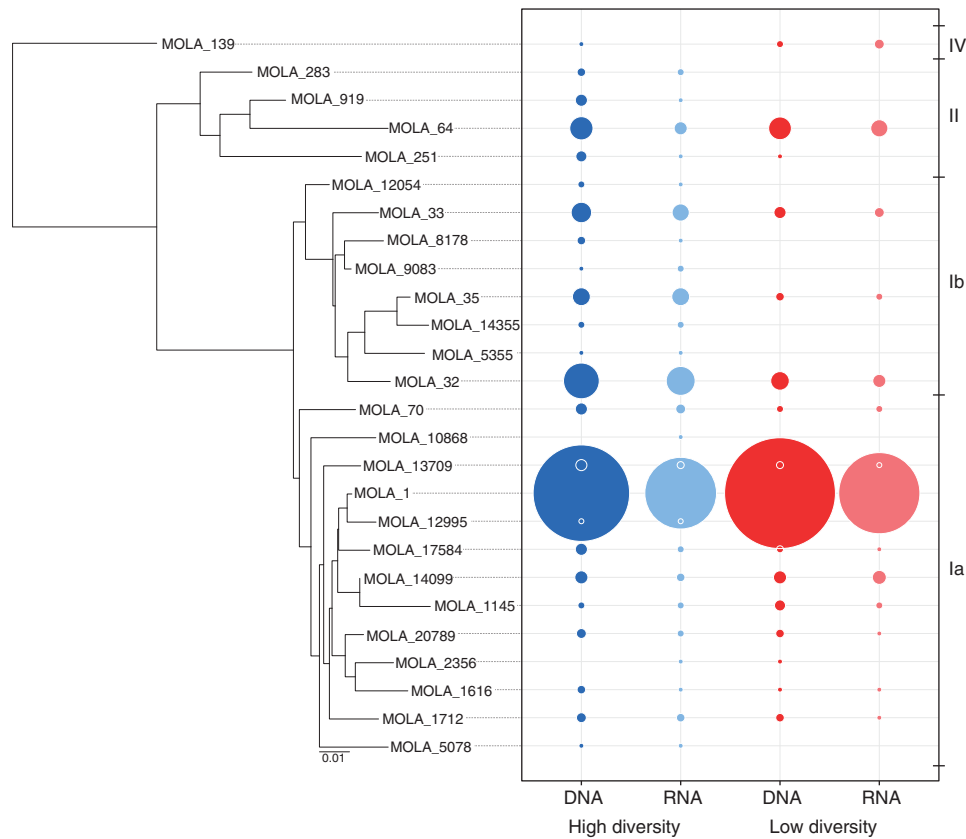


Figure 2 Phylogenetic relationship and relative abundance of SAR11 OTUs. Only SAR11 OTUs representing >0.2% of total SAR11 OTU sequences are included. Bubble size is scaled to sequence abundance of OTUs relative to total SAR11 communities. DNA and RNA is 16S rRNA gene copies and 16S rRNA, respectively. High and low diversity sub-groups were empirically defined; see Figure 1 and text for further details.

times of high diversity, but these differences were not statistically significant. When combining relative abundances of clades *Ib* and *II*, the differences between high and low diversity regimes were significant and indicated that the temporal dynamics of *Ib* and *II* are independent of *Ia*.

Month to month differences in ecotype distributions were also evident (Figure 3). Both the rDNA and rRNA sequences of ecotype *Ia* showed maximum abundances of 81–87% from April to October, a period corresponding to higher leucine incorporation ($16\text{--}42\text{ pmol l}^{-1}\text{ h}^{-1}$) and extremely rapid phosphate turnover times (<5 h, Figure 3a). Ecotypes *Ib* and *II* displayed consistently inverse patterns to *Ia* but were broadly consistent to each other with peaks from November to March. During this period leucine incorporation was lower ($5.0\text{--}7.4\text{ pmol l}^{-1}\text{ Hr}^{-1}$) and phosphate turnover was longer ranging from 30 to 130 h with highest values found in December and January (Figure 3).

PLSR model

The relative abundance of SAR11 ecotypes from both rDNA and rRNA sequences were combined with biogeochemical observations to develop a model that described SAR11 ecotype dynamics

based on PLS regression analysis (Figure 4). The biogeochemical variables included in the model are listed in Table 1. Using a three-factor component model with non-transformed input variables, the model was capable of describing 70% of observed variance in the temporal dynamics of SAR11 ecotype abundance. The goodness of fit for individual ecotype groupings based on rRNA abundances was higher (0.76–0.78) than those based on rDNA abundances (0.63–0.66; Figure 4a). All data were centred and auto-scaled to compute weighted regression coefficients for the biogeochemical input variables (Figure 4b). Regression coefficients were uniformly larger for rRNA abundance data, although similar to rDNA. The physical parameters of MLD, temperature and salinity were quantitatively the most important variables for describing temporal trends of SAR11 ecotype abundance.

rRNA/rDNA activity ratios

Sequences were grouped to calculate rRNA/rDNA ratios of the SAR11 clade and compared with estimates of SAR11 leucine incorporation (Figure 5). The $\mu\text{CARD-FISH}$ samples were collected as part of a separate project and were only available for the year 2008 (Laghdass *et al.*, 2012). SAR11 leucine

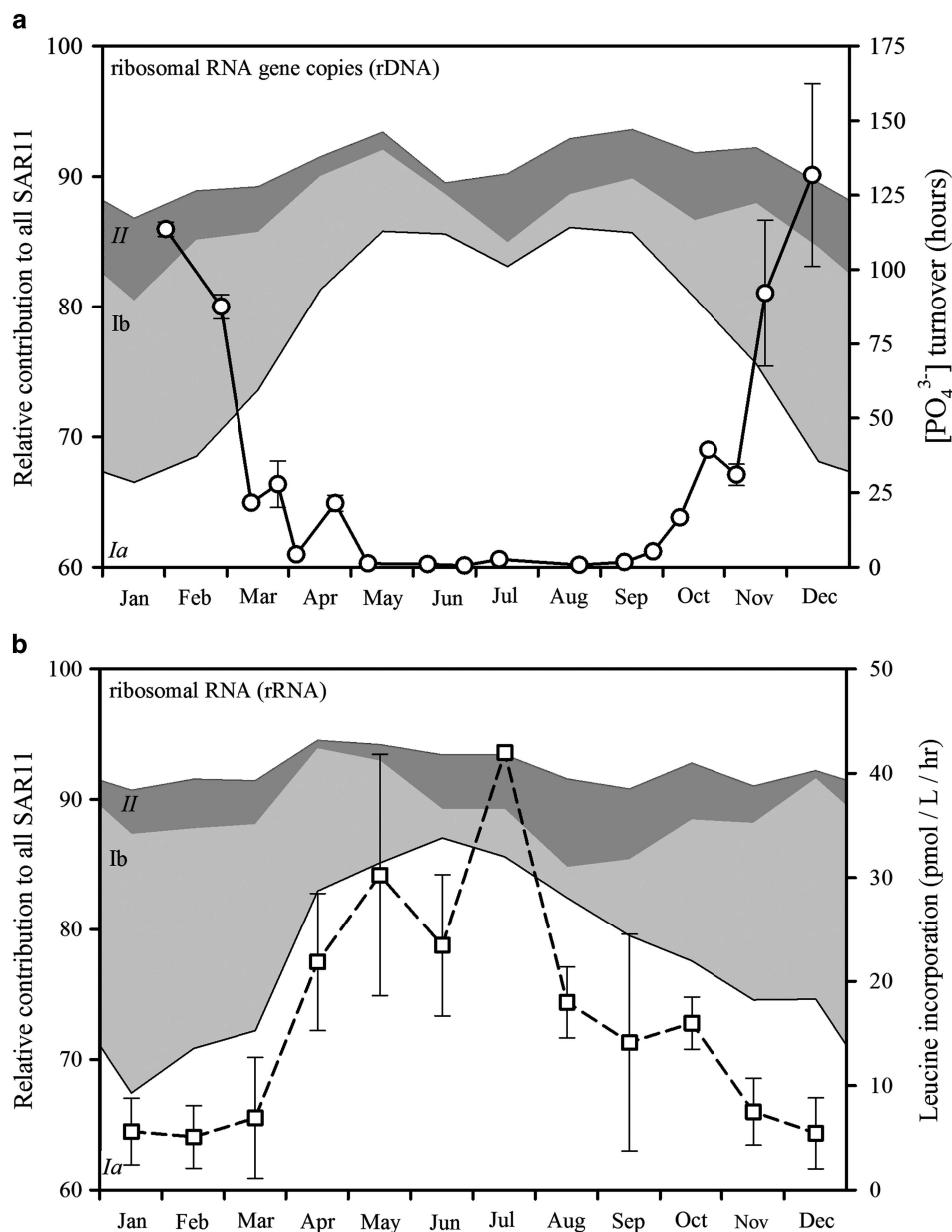


Figure 3 Monthly averages of SAR11 ecotype abundance and microbial rate processes. The monthly averages of sequence abundance are calculated from (a) ribosomal RNA gene copies and (b) ribosomal RNA tag-sequence data. The abundance of sequences in SAR11 ecotype classifications Ia, Ib and II are cumulative sequences within the phylogenetically defined sub-clades (Figure 2) expressed relative to total SAR11 sequences. The difference of SAR11 sequences not included in these sub-clade classifications is represented by the blank area at the top of panels. White, light grey and dark grey areas represent SAR11 Ia, Ib and II, respectively. White circles and solid line in panel a are phosphate turnover times determined in the NW Mediterranean during 2012, error bars are analytical errors on triplicate measurements. White squares and dashed lines in panel b are leucine incorporation rates temporally binned with 1-month resolution. Error bars are standard errors on monthly measurements integrated over the 7-year time-series.

incorporation and rRNA/rDNA ratios were positively correlated ($R=0.86$; $P<0.05$). The rRNA/rDNA ratios of SAR11 ecotypes were not correlated with respective abundances (Figure 6a). The PLSR modelling approach was unable to capture temporal dynamics of activity ratios with a satisfactory goodness of fit (Supplementary Figure 2). However, rRNA/rDNA ratios of all individual ecotype pairings (Ia vs Ib, Ib vs II, and Ia vs II) exhibited positive covariance (methods) indicating that they were

generally active at the same times and their activity ratios were not inversely related (Figure 6). SAR11 ecotype rRNA/rDNA ratios were compared over the range 0–1.5. Data that exceeded 1.5 were atypical (<4%) and not considered to be representative of the general trends within the data set. The largest positive correlation was found between Ia and Ib ($R=0.56$; $P<0.001$), followed by Ib and II ($R=0.47$; $P<0.01$) and Ia and II ($R=0.38$; $P<0.05$).

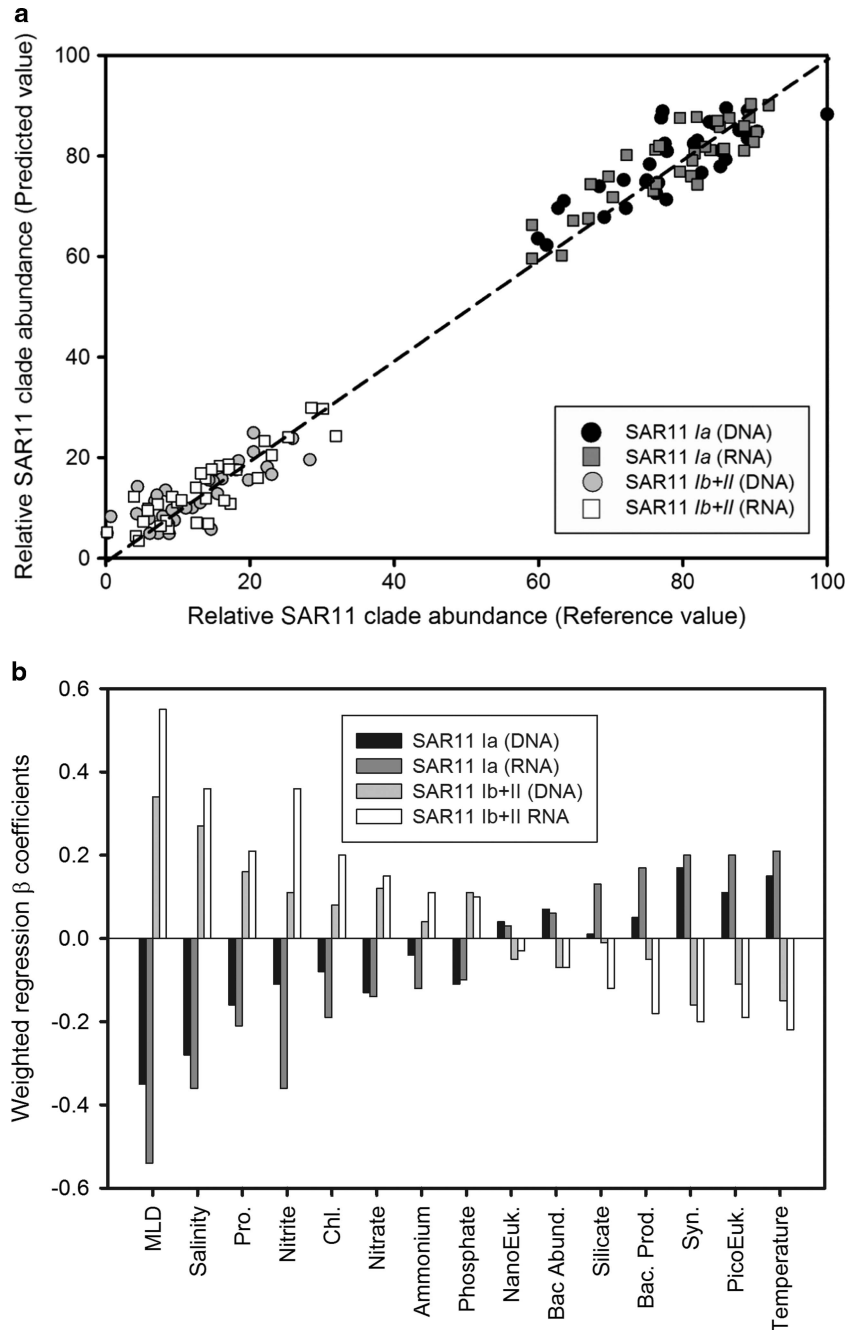


Figure 4 Partial least squares regression model of SAR11 ecotype abundance. (a) The relationship between modelled and observed SAR11 ecotype distributions from ribosomal RNA gene copies and ribosomal RNA tag-sequence data. (b) The weighted regression coefficient of each biogeochemical parameter used to model SAR11 ecotype patterns. Positive values indicate that the input variable is positively correlated to specific output variables. MLD, mixed layer depth; Pro., *Prochlorococcus*; Chl., chlorophyll; NanoEuk., nanoeukaryotes; Bac. Abund., bacterial abundance; Bac. Prod., bacterial productivity; Syn., *Synechococcus*; PicoEuk., picoeukaryotes, Temp., temperature.

Discussion

Mixing controls surface bacterioplankton diversity maxima at broad phylogenetic levels

A striking result of our time-series analysis was the observation that levels of diversity measured by the Shannon Index, a metric that considers equitability as well as richness, showed repeatable minima occurring in the summer and

maxima during winter months. These trends are remarkably similar when all OTUs were compared against only those within the SAR11 clade. Our findings are in agreement with the emerging paradigm of strong and predictable seasonality in total prokaryotic diversity (Gilbert *et al.*, 2012; Ladau *et al.*, 2013). However, the concordance between the dynamics of the entire community that contains a physiologically diverse assemblage of

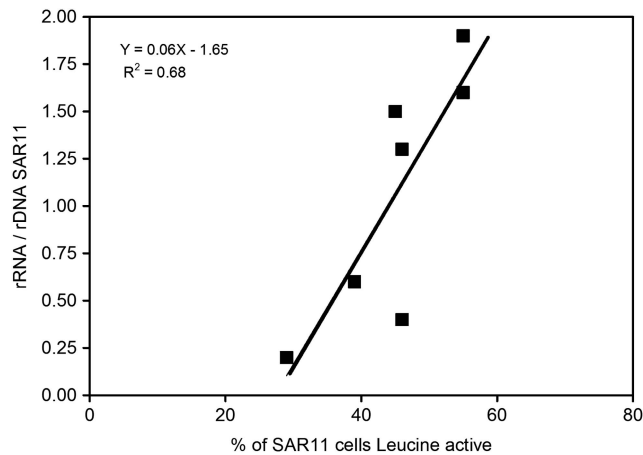


Figure 5 Leucine incorporation and RNA/DNA ratios in the SAR11 clade. The ratio of ribosomal RNA (rRNA) to ribosomal DNA gene copies (rDNA) were calculated by summing the sequences for all SAR11 clusters. SAR11-active cells were calculated as the percentage of total SAR11 cells assimilating tritiated leucine.

OTUs, to that of a narrower clade such as SAR11, suggests significant forcing by physical mechanisms which act similarly across broad phylogenetic levels.

We propose water column mixing to be the primary mechanism regulating seasonal augmentation of diversity in the NW Mediterranean. Periods of high and low bacterial and SAR11 ecotype diversity were repeatedly associated with deep and shallow mixed layers, respectively (Table 1). Furthermore, MLD was quantitatively the most important environmental parameter for predictive modelling of measured SAR11 ecotype sequence abundance. Presumably, stratification breakdown directly enhances diversity simply by mixing vertically segregated populations in a ‘community-merging’ event. Microbial populations frequently exhibit environmental niche partitioning as a function of depth (Field *et al.*, 1997; West and Scanlan, 1999; DeLong *et al.*, 2006; Carlson *et al.*, 2009; Treusch *et al.*, 2009) and convective overturning has been shown to be important for the temporal and vertical distributions of certain bacterial lineages (Morris *et al.*, 2005; Treusch *et al.*, 2009; Carlson *et al.*, 2009; Eiler *et al.*, 2009; Chow *et al.*, 2013; Vergin *et al.*, 2013b). In addition, we find seasonal maxima of SAR11 diversity is unrelated to the activity of these ecotypes estimated from their rRNA/rDNA ratios, further supporting the idea of translocation from biogeochemically distinct niches throughout the water column. Similar to findings in the English Channel (Gilbert *et al.*, 2012) we observe a strong correlation between diversity and day-length (data not shown). Although a good-predictor of bacterial diversity, we suggest that day-length is a non-mechanistic correlate of seasonality. Changes in day-length integrate strongly seasonal features, including mixing.

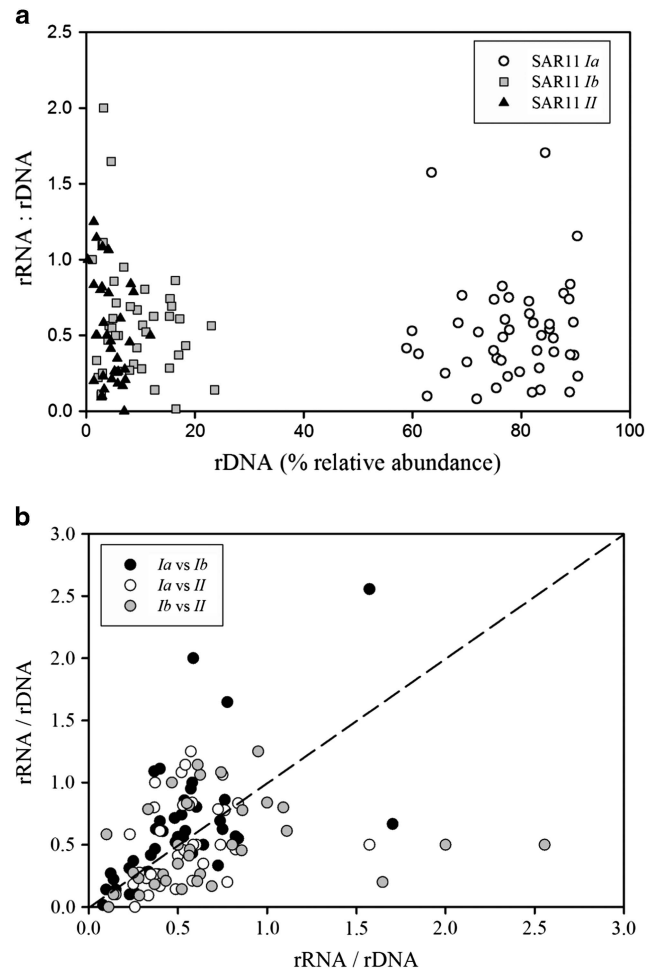


Figure 6 rRNA/rDNA activity ratios and abundance of SAR11 ecotypes. (a) The relationship between ecotype activity (rRNA/rDNA) and relative abundance (rDNA%) for Ia (white circles), Ib (grey squares) and II (black triangles) and (b) relationship of rRNA to rDNA ratios between SAR11 ecotypes. Dashed line is 1:1. Correlation statistics are Ia and Ib ($R = 0.56$; $P < 0.001$); Ib and II ($R = 0.47$; $P < 0.01$); and Ia and II ($R = 0.38$; $P < 0.05$).

Subsequent to the process of *community merging*, mixing the water column may lead to two other mechanistic processes; ‘*ex-novo* substrate merging’ and ‘*de-novo* substrate induction’, which act to consolidate high diversity systems. Substrate merging relies on the fact that the wide spectrum of substrates available for bacterial growth exhibits vertical structure (Carlson *et al.*, 2004; Reinthaler *et al.*, 2006). Mixing increases the range of substrates, and thus bacterial diversity, by rapidly expanding the number of available niches in the surface layer. Substrate induction may result from the production of new organic substrates by eukaryotic blooms stimulated by macronutrient resupply upon mixing. At our study site there is a strong association of enhanced chlorophyll features with deeper mixed layers. The potential range of *de novo* substrates produced from variable eukaryotic sources (Nelson *et al.*, 2013; Sarmiento *et al.*, 2013) could also expand the number of bacterial niches

(Landa *et al.*, 2013) in the surface ocean. However, in the NW Mediterranean peaks in chlorophyll tend to succeed those of bacterial and SAR11 diversity, suggesting it has a second order effect. Water column mixing (and the combination of populations that inhabited different strata during summer months) is therefore the most parsimonious explanation to explain the origin of winter diversity maxima in the NW Mediterranean, and perhaps those observed in other temperate environments (for example, see Gilbert *et al.*, 2012; Ladau *et al.*, 2013; Vergin *et al.*, 2013b). *Ex-novo* substrate merging and *de novo* substrate induction are suggested as secondary mechanisms related to physical mixing that act to consolidate high diversity states for some interval subsequent to the initial community-merging event.

Although it is intuitive that physical mixing of depth separated niche space enhances SAR11 ecotype diversity, it is more difficult to envisage how shoaling of the mixed layer could directly select against certain ecotypes. Stratification must influence SAR11 ecotype dynamics indirectly through competitive selection (for example, growth, substrate limitation, grazing, viral lysis). We thus propose a framework in which physical mixing is a common mechanism responsible for periodically 'resetting' bacterial ecosystems to a high diversity state. The subsequent progression towards low diversity communities commonly observed (Treusch *et al.*, 2009; Gilbert *et al.*, 2012; Chow *et al.*, 2013) is driven by seasonal evolution of local environmental conditions, leading to the dominance of more oligotrophic and high light-adapted SAR11 ecotypes. These environmental conditions may vary between regions and affect bacterial sub-groups in different ways. However, the widely reported winter maxima in bacterial diversity (Gilbert *et al.*, 2012; Ladau *et al.*, 2013) suggest increases in diversity due to vertical mixing are a widespread phenomenon in temperate environments. The seasonal diversity patterns observed in the NW Mediterranean and other temperate (Gilbert *et al.*, 2012; Ladau *et al.*, 2013) and high latitude environments (Ghiglione *et al.*, 2012) highlight the importance of adequate temporal resolution when attempting to synthesise global patterns of bacterial diversity.

Environmental selection of low diversity regimes

In the NW Mediterranean we propose inorganic nutrient limitation, specifically that of phosphate, as a plausible factor driving SAR11 towards a low diversity state. SAR11 *Ia* and *Ib/II* relative abundances were inversely correlated with nutrient concentrations. In the NW Mediterranean phosphate concentrations are very low (38 ± 17 nM) and frequently close to analytical detection limits. In such extreme oligotrophic environments turnover is a more reliable metric of phosphate limitation. Low diversity SAR11 communities occur congruently with periods of very rapid (<5 h) phosphate turnover. In addition,

SAR11 ecotypes were differentially correlated with *Synechococcus* abundance. *Synechococcus* are known to have high cell-specific uptake rates for PO_4^{3-} (Casey *et al.*, 2009; Michelou *et al.*, 2011) and may enhance phosphate limitation, promoting the development of low diversity SAR11 communities dominated by group *Ia*. In support of our environmental data, genomic sequences point to the adaptation of SAR11 *Ia* to rapid [PO_4^{3-}] turnover through enhanced expression of high-affinity phosphate transporters (Wilhelm *et al.*, 2007; Sowell *et al.*, 2008) or the ability to hydrolyse organic phosphorous substrates (Grote *et al.*, 2012). Unfortunately sequence information from groups *Ib* and *II* is currently lacking so the hypothesis of phosphorus as a segregating factor for SAR11 ecotypes remains circumstantial.

The dominance of SAR11 *Ia* also corresponds to seasonal maxima in leucine incorporation rates (Figure 3b). Dissolved organic matter accumulates in highly stratified surface layers (Williams, 1995; Hansell and Carlson, 1998) including our study site (Laghdass *et al.*, 2012). Low diversity communities dominated by SAR11 *Ia* thus appear capable of significant dissolved organic carbon utilisation at low nutrient concentrations, resulting in enhanced bacterial production relative to other times of year. Temperature effects and leucine uptake by co-occurring bacterial groups, including cyanobacteria, may also contribute to enhanced bacterial metabolism during summer.

Predictive model of SAR11 ecotype sequences

The available biogeochemical observations were used to calibrate a PLSR model that was capable of describing 70% of cross-validated variance (Q^2) in the temporal dynamics of SAR11 ecotype groupings. Mixed layer depth emerged as the most important environmental control. Overall, regression coefficients between the measured biogeochemical parameters and tag sequence data were larger for the rRNA fraction, perhaps indicating tighter bottom-up control on active SAR11 ecotypes. SAR11 *Ia* dominates in shallow mixed layers of mid-late summer that are depleted in inorganic nutrients, have low chlorophyll biomass and picoplankton populations dominated by *Synechococcus*. Groups *Ib* and *II* are associated with deeper mixed layers enhanced in nutrients and phytoplankton biomass. It is presently unclear however if ecotypes *Ib* and *II* are specifically adapted to enhanced phytoplankton biomass and nutrients in the same way as other bacterial clades; for example, *Roseobacter* (Le Fort and Gasol, 2013). The persistence of these clades for several months may be a result of repeated 'physical-seeding' to the surface throughout the mixing season.

Activity of SAR11 ecotypes from rRNA/rDNA ratios

The ratio of rRNA and ribosomal RNA gene copies (rRNA/rDNA) is frequently used to estimate *in-situ*

activity, or even growth rates, of specific bacterial OTUs (Campbell *et al.*, 2011; Campbell and Kirchman, 2013; Hunt *et al.*, 2013). However, a recent report has questioned the reliability of rRNA as an indicator of metabolic state (Blazewics *et al.*, 2013). To validate the utility of rRNA/rDNA in describing *in-situ* variation of SAR11 ecotype activity we compared tag sequence data to direct measurements of leucine incorporation (Laghdass *et al.*, 2012), which is a commonly used measure of heterotrophic production by bacteria (Kirchman *et al.*, 1985). The fraction of SAR11 incorporating leucine was positively correlated with SAR11 rRNA/rDNA ratio. Although limited in number, our data set covers the full range of rRNA/rDNA ratios (Campbell *et al.*, 2009, 2011) and leucine incorporation rates (for example, Malmstrom *et al.*, 2007) typically reported for SAR11. To the best of our knowledge we have made the first attempt to correlate rRNA/rDNA as an *in-situ* activity metric against a direct measure of bacterial metabolism at an identical, albeit broad, phylogenetic resolution. It is difficult to speculate if the relationship observed for SAR11 would hold for other bacterial lineages or at increasing phylogenetic resolution. From the limited observations we make a first-order assumption that rRNA/rDNA and leucine incorporation exhibit similar covariance at ecotype level and therefore apply this ratio to investigate patterns in SAR11 ecotype activity.

The rRNA/rDNA ratios of SAR11 ecotypes did not display any coherent seasonal patterns or relationships with the measured biogeochemical variables (Supplementary Figure 2; Table 1). Furthermore, ecotype abundance and activity were uncorrelated (Figure 6a). It might be expected that correlations between ecotype activity and abundance are characterised by an undefined time lag. However, applying a one-month time lag [+1 sampling event] (for example, Fuhrman *et al.*, 2006) to our data set failed to improve activity-abundance correlations. Decoupling of activity and abundance has been observed elsewhere (Campbell and Kirchman, 2013). The temporal resolution afforded by a monthly time-series may be too coarse to capture the transient nature of abundance–activity relationships in SAR11 ecotypes.

Alternatively, it is possible that despite differences in 16S RNA gene phylogeny (Vergin *et al.*, 2013a) and gene content (Grote *et al.*, 2012), SAR11 ecotypes share enough metabolic potential and overlapping niche space to maintain simultaneously active populations. In support of this SAR11 ecotype activities are statistically indistinguishable from each other in low and high diversity settings, regardless of the strong biogeochemical gradients that exist. Positive covariance in SAR11 ecotype activities across the 7-year data set (Figure 6b) also suggest that activity of different ecotypes may respond similarly to changes in environmental conditions. However, the clear seasonal partitioning

of SAR11 ecotype abundances are not systematically related to specific ecotype activity, as presently determined from rRNA/rDNA ratios. Although we provide preliminary evidence that *potential* metabolic activity (rRNA abundance) is linked to *realised* protein synthesis (leucine incorporation) in SAR11, the use of rRNA/rDNA ratios as a reliable activity measure remains ambiguous (Blazewics *et al.*, 2013).

Through our analysis we demonstrate how physical oceanographic processes and biogeochemical forcing factors combine to explain the well-documented seasonal cycles in bacterial diversity. Temporal instability in environmental conditions is the main factor driving patterns of bacterial diversity in the NW Mediterranean. We suggest that adequate spatial and temporal global coverage would reveal bacterial diversity peaks in winter at temperate to high latitudes, rather than seasonally stable low latitude environments. Although our analysis specifically focuses on SAR11 dynamics we expect the same mechanisms to be relevant for other bacterial clades. In the NW Mediterranean phosphate limitation appears to govern seasonal transitions to low-diversity SAR11 communities. Phosphate may be of similar importance in other oligotrophic environments that are characterised by seasonal dominance of SAR11 ecotype *Ia*. PLSR modelling demonstrated strong and predictable temporal patterns in SAR11 ecotype abundance that were not reflected in seasonal dynamics of metabolic activity. Future studies targeting the fine-scale phylogenetic structure of SAR11 communities, at increased temporal resolution, may help to further define the complex patterns observed between SAR11 ecotype abundance and activity.

Conflict of Interest

The authors declare no conflict of interest.

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