

ORIGINAL ARTICLE

Quantitative analysis of a deeply sequenced marine microbial metatranscriptome

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The potential of metatranscriptomic sequencing to provide insights into the environmental factors that regulate microbial activities depends on how fully the sequence libraries capture community expression (that is, sample-sequencing depth and coverage depth), and the sensitivity with which expression differences between communities can be detected (that is, statistical power for hypothesis testing). In this study, we use an internal standard approach to make absolute (per liter) estimates of transcript numbers, a significant advantage over proportional estimates that can be biased by expression changes in unrelated genes. Coastal waters of the southeastern United States contain 1×10^{12} bacterioplankton mRNA molecules per liter of seawater (~ 200 mRNA molecules per bacterial cell). Even for the large bacterioplankton libraries obtained in this study ($\sim 500\,000$ possible protein-encoding sequences in each of two libraries after discarding rRNAs and small RNAs from >1 million 454 FLX pyrosequencing reads), sample-sequencing depth was only 0.00001%. Expression levels of 82 genes diagnostic for transformations in the marine nitrogen, phosphorus and sulfur cycles ranged from below detection ($<1 \times 10^6$ transcripts per liter) for 36 genes (for example, phosphonate metabolism gene *phnH*, dissimilatory nitrate reductase subunit *napA*) to $>2.7 \times 10^9$ transcripts per liter (ammonia transporter *amt* and ammonia monooxygenase subunit *amoC*). Half of the categories for which expression was detected, however, had too few copy numbers for robust statistical resolution, as would be required for comparative (experimental or time-series) expression studies. By representing whole community gene abundance and expression in absolute units (per volume or mass of environment), ‘omics’ data can be better leveraged to improve understanding of microbially mediated processes in the ocean.

The ISME Journal (2011) 5, 461–472; doi:10.1038/ismej.2010.141; published online 16 September 2010

Subject Category: integrated genomics and post-genomics approaches in microbial ecology

Keywords: metatranscriptomics; marine; bacterioplankton; gene expression; biogeochemistry

Introduction

Metatranscriptomics is a powerful tool for capturing gene expression patterns in natural microbial communities without previous assumptions as to the ongoing activities or dominant taxa (Poretsky *et al.*, 2005; Frias-Lopez *et al.*, 2008). In contrast to metagenomics, which provides an inventory of the community gene pool, metatranscriptomics identifies which of those genes are being transcribed in a given ecological context, including under experimentally manipulated conditions (Gilbert *et al.*, 2008; Poretsky *et al.*, 2010).

The advent of second-generation sequencing has increased metatranscriptome library sizes by orders of magnitude (Poretsky *et al.*, 2005, 2009b; Frias-Lopez *et al.*, 2008; Urich *et al.*, 2008; Hewson *et al.*, 2009a), yet how deeply a community transcriptome is ‘covered’ by the sequence library

remains a critical issue. If too shallow, libraries will be dominated by transcripts from metabolic pathways shared by most cells and poor in those representing specialized biogeochemical pathways (Hewson *et al.*, 2009b; Poretsky *et al.*, 2009b). As a consequence, unique expression patterns within a community may be missed, and comparative analyses between communities can be insensitive.

Variability in sample-sequencing depth between community metatranscriptomes, regardless of coverage level, further limits the power of comparative analyses by restricting assessments to relative data (that is, as a proportion of the transcriptome; Figure 1). This is problematic because changes occurring in the abundance of some mRNAs in response to shifting conditions (van de Peppel *et al.*, 2003; Hannah *et al.*, 2008; Robinson and Oshlack, 2010) leads to changes in the percent representation of other mRNAs whose absolute abundance has not changed (Figure 1). Comparative analyses based on ratios of mRNA copies to DNA copies (that is, relative abundance in metatranscriptomes vs relative abundance in metagenomes; Frias-Lopez *et al.*, 2008) does not solve this problem, as both are similarly affected by unknown and possibly

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Received 10 May 2010; revised 1 July 2010; accepted 12 July 2010; published online 16 September 2010

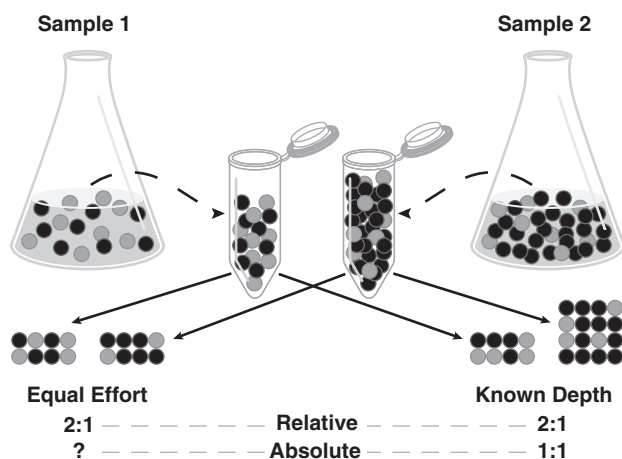


Figure 1 Effect of sample-sequencing depth on quantification of transcripts (or genes) in environmental samples. ‘Equal-effort’ sequences the same number of reads per sample volume, regardless of the size of the mRNA pool, and therefore conveys only relative abundance. ‘Known-depth’ sequences a known proportion of the transcript pool (50% for both, in this example), and therefore also conveys absolute copy numbers per sample volume. The latter is more relevant to biogeochemical rate measurements, as mRNAs of biogeochemical interest (gray dots) can make up different proportions in community transcriptomes yet have identical numbers in the environment.

different sample-sequencing depths. Thus as currently obtained, metatranscriptomic data provide information on enrichment or depletion of a transcript category in the community transcriptome, but not on the absolute abundance of these transcript categories per volume or mass of environment (Figure 1), which is the most relevant comparison for biogeochemical studies and ecosystem modeling. Metagenomic-, metaproteomic- and environmental microarray-based studies suffer these same proportional data constraints.

In this study, we report the deep sequencing of two replicate metatranscriptomes from southeastern United States coastal seawater to characterize microbial gene expression and address three critical questions about sequencing effort: (1) What was the sample-sequencing depth for the bacterioplankton community transcriptome? We used an internal mRNA standard to estimate the number of transcripts in the natural sample compared with the number of transcripts sequenced. (2) What was the abundance of transcripts representing key bacterial transformations in the nitrogen (N), phosphorus (P) and sulfur (S) cycles in coastal seawater? We used BLAST analysis normalized to internal standard recovery to estimate absolute transcript numbers for over 80 diagnostic steps in marine elemental cycles. (3) Was the sequencing strategy sufficient to detect differences in gene transcription between samples? We examined the effect of coverage depth on statistical comparisons of biogeochemically diagnostic transcripts in the metatranscriptomes.

Materials and methods

Sample collection

Two replicate seawater samples (FN56 and FN57) were collected at Marsh Landing, Sapelo Island, Georgia, USA (31°25′4.08N, 81°17′43.26W; <http://www.simo.marsci.uga.edu>) on 6 August 2008 at 2330 hours local time, 1 h before high tide and 3 h after sunset. These samples are part of a multiyear time series of the Sapelo Island Microbial Observatory (<http://www.simo.marsci.uga.edu>), in which collections are made every 3 months and each collection set consists of duplicate samples from four consecutive high tides (2 days and 2 nights). Surface water (5.75 l from a depth of 0.5 m) was pumped directly through 3 and 0.22 µm filters. The 0.22 µm filter was placed in a Whirl-Pak bag (Nasco, Fort Atkinson, WI, USA) and immediately flash frozen in liquid N₂. Total time from the start of filtration to freezing was 10 min. Although the samples are considered biological replicates within the larger time series, we note that there was 8 min between the end of the first collection (sample FN56) and the start of the second (sample FN57). Nutrient data are collected monthly at station GCE6 (~3 km from Marsh Landing) as part of the Georgia Coastal Ecosystems Long Term Ecological Research program (<http://gce-lter.marsci.uga.edu>).

RNA processing and sequencing

RNA processing in preparation for pyrosequencing was carried out as previously described (Poretsky *et al.*, 2009a, b), with the exception of the addition of an *in vitro* transcribed standard to the extraction tube before beginning the extraction. The standard was constructed by linearizing a pGem-3Z plasmid (Promega, Madison, WI, USA) with *ScaI* restriction enzyme (Roche, Penzberg, Germany) and cleaned with a phenol/chloroform/isoamyl alcohol extraction. Complete digestion of the plasmid was confirmed with a 1% agarose gel. The DNA fragment was then *in vitro* transcribed using the Riboprobe *in vitro* Transcription System (Promega) according to the manufacturer’s protocol; an SP6 RNA polymerase was used to create a 994 nt long RNA fragment. The pGem plasmid had another internal T7 promoter region, but it was present in the reverse complement sequence during *in vitro* transcription and aRNA amplification (see below), and did not interfere. Residual DNA was removed with RQ1 RNase-Free DNase, and the RNA was cleaned with a phenol/chloroform/isoamyl alcohol extraction. The RNA standard was quantified with a Nanodrop Spectrophotometer (Thermo Scientific, Wilmington, DE, USA), and correct fragment size was confirmed with an Experion automated electrophoresis system (Bio-Rad, Hercules, CA, USA).

RNA standard (25 ng; 4.7×10^{10} copies) was added to a 50 ml conical tube containing 8 ml RLT lysis buffer (Qiagen, Valencia, CA, USA) and 3 g of RNA

PowerSoil beads (Mo-Bio, Carlsbad, CA, USA). The sample filters were removed from -80°C storage, shattered and added to the extraction tubes. RNA was then extracted using an RNEasy kit (Qiagen), and any residual DNA was removed using the Turbo DNA-free kit (Applied Biosystems, Austin, TX, USA). To reduce the number of rRNAs in the pyrosequencing reads, total RNA was treated in two ways to enrich for mRNA. Epicentre's mRNA-Only isolation kit (Madison, WI, USA) was first used to decrease rRNA contamination enzymatically. The samples were then treated with MICROBExpress and MICROBEnrich kits (both from Applied Biosystems), which couple an oligonucleotide rRNA probe with magnetic separation to enrich for mRNA. Successful reduction of rRNA was confirmed by running both pre- and post-treated samples on an Experion automated electrophoresis system (Bio-Rad). To obtain enough mRNA for pyrosequencing, the samples were linearly amplified using the MessageAmp II-Bacteria kit (Applied Biosystems). The amplified RNA was then converted to cDNA using the Universal RioboClone cDNA synthesis system with random primers (Promega), which produced cDNAs primarily in the size range of 200–600 bp. Residual reactants and nucleotides from cDNA synthesis were removed from the sample using the QIAquick PCR purification kit (Qiagen), and gel-based size selection was used to select fragments in the 250–500 bp range. cDNAs from each replicate sample were loaded into 1/2 of each of four GS-FLX plates for 454 pyrosequencing. Sequences are deposited in the CAMERA database (<http://camera.calit2.net/about-camera/full-datasets>) under accession name 'CAM_PROJ_Sapelo2008'.

Read annotation

Duplicate clusters were identified using an online program (Gomez-Alvarez *et al.*, 2009). Ribosomal RNA sequences were identified with a BLASTn search against the small and large subunit SILVA database (<http://www.arb-silva.de>) with a bit score cutoff ≥ 50 ; sequences identified as rRNA were then removed from further consideration. To identify small, nonprotein encoding RNAs (Shi *et al.*, 2009), all nonribosomal reads were compared with the RFam database (<http://rfam.janelia.org>) using BLASTn with a bitscore ≥ 40 , and hits were considered putative small RNAs if the best hit in the RefSeq database was a hypothetical protein or if the RFam alignment was ≥ 95 nt (Supplementary Figure S1).

Remaining reads were annotated using BLASTx searches against the NCBI RefSeq and Clusters of Orthologous Genes (COG) databases (Tatusov *et al.*, 2003) with a bit score cutoff ≥ 40 . Taxonomic binning was based on RefSeq hit. Collector's curves were produced from a custom script in the R environment (R Development Core Team, 2009). Read coverage of proteorhodopsin PU1002_03206

gene bin and the internal standard was assessed by assembling reads against the reference sequence using Geneious version 4.8 (Biomatters, Auckland, New Zealand) with gaps and default scoring (word length = 18, maximum gap size = 1, maximum gaps per read = 20, maximum mismatches = 20 and maximum ambiguities = 4) and the consensus sequence representing the majority nucleotide at each position. Sequence variation at each nucleotide position was determined using a custom script in R with the BioStrings package (Pages *et al.*, 2009).

Elemental cycle transcripts

Reference diagnostic genes representing transformations in the N, P and S cycles were selected from marine alphaproteobacteria, gammaproteobacteria and bacterioidetes genomes (the three most common taxa in marine metagenomic libraries), or from other taxa if these three groups did not contain an ortholog to the gene of interest. These reference sequences were used as query sequences in BLASTX analysis against the metatranscriptomic data (bitscore ≥ 40 , E-value $< 10^{-3}$) and redundant hits were removed. Remaining hits were manually checked with BLASTX against the RefSeq database and discarded if the top three hits were not to a similar annotation as the original reference gene.

Statistics

Pairwise statistical comparisons were carried out with Xipe, a bootstrapped difference of means calculation developed by Rodriguez-Brito *et al.* (2006), using 20 000 bootstrap iterations and 95% confidence intervals, or with 2×2 contingency tables and the Fisher's exact test (White *et al.*, 2009) using $P < 0.05$. Subsampled libraries for Xipe analyses were created by sampling without replacement using R (R Development Core Team, 2009). The Benjamini–Hochberg correction was used to adjust the Fisher's exact test P -values as a control for the false discovery rate using the R package 'multtest' (Strimmer, 2008), and only those genes with an adjusted P -value < 0.05 were considered significant. A simulation analysis of Fisher's exact test significance threshold as a function of count number was carried out using an R script that ran 2×2 contingency tables at incrementing count values for library sizes of 125 000 reads.

Results and discussion

Sequence libraries

cDNAs derived from two replicate coastal bacterioplankton samples (samples FN56 and FN57 in the Sapelo Island Microbial Observatory series; <http://simo.marsci.uga.edu>) were sequenced in four GS-FLX 454 runs (Margulies *et al.*, 2005), with four technical sequencing replicates per biological replicate. Over a million reads averaging 210 nt

Table 1 Summary statistics for coastal ocean metatranscriptome datasets

	FN56	FN57	Combined
Total reads	1 067 363	1 114 536	2 181 899
rRNA	466 834 (44%)	623 804 (56%)	1 090 638 (50%)
psRNA	100 437 (9%)	25 213 (2%)	125 650 (6%)
Possible proteins	500 092 (47%)	465 519 (42%)	965 611 (44%)
RefSeq Hits	255 280	260 739	516 019
RefSeq Genes	96 573	109 395	168 669
RefSeq Taxa	1707	1761	1909
COG hits	162 925	170 593	333 518
Unassigned	244 812	204 780	449 592

Abbreviation: COG, Clusters of Orthologous Groups.

Percentages are of total reads.

RefSeq Hits, number of reads with significant homology to the RefSeq database; RefSeq Genes, number of unique accession numbers within those hits; Unassigned, = number of possible proteins that did not have a significant hit to either the RefSeq or COG databases (47% of possible proteins).

in length were obtained per sample (Table 1). After removal of rRNAs and putative small RNAs (Shi *et al.*, 2009), there were ~500 000 possible protein encoding reads in each library (Table 1).

Sample-sequencing depth

Sample-sequencing depth is defined here as the percent of mRNA molecules present in a sample that is represented in the sequence library. The greater the sequencing depth of an mRNA pool, the more thorough the representation of microbial gene transcription. Further, if the volume or weight of the sample is also known, information on the sample-sequencing depth allows absolute transcript abundance to be calculated for a given quantity of the environment, not just proportional abundance in the community transcriptome. To estimate sample-sequencing depth, a known number of artificial RNA sequences serving as an internal standard was added immediately before cell lysis at the initiation of nucleic acid extraction. This approach may have some biases, for example if the internal standard is more susceptible to degradation than natural mRNA or if the efficiency of release of natural mRNA from cells is <100%, but it provides a consistent accounting across samples through extraction, processing and sequencing steps. Similar approaches have been successfully applied to qPCR (Coyne *et al.*, 2005) and microarray studies (Hannah *et al.*, 2008).

A total of 4014 internal standards were identified in the FN56 sequence library out of 4.7×10^{10} copies added before cell lysis, leading to an estimate of 1.0×10^{12} bacterioplankton mRNA molecules per liter of coastal seawater (Table 2). Two other estimates of the size of the community transcriptome were derived using literature values for mRNA content of marine bacterioplankton (Table 2), and these were in reasonable agreement with the

internal standard method. The sample-sequencing depth was therefore ~0.00001%, or 1 in 10^7 transcripts, with FN57 sequenced slightly deeper than FN56 (Table 2).

Direct cell counts indicated 4.2×10^9 bacterioplankton cells per liter in the seawater samples, and therefore an average of 190 mRNA transcripts per cell (Table 2). Laboratory cultures of *Escherichia coli* in exponential growth phase have ~1400 transcripts per cell (Neidhardt and Umberger, 1996). The sevenfold lower estimate for coastal bacterioplankton was not unexpected, however, because the cells are considerably smaller in size (Azam and Hodson, 1977) and have much lower growth rates (Ducklow, 2000) than laboratory-grown *E. coli*. On the basis of this per cell abundance estimate, it can be deduced that transcript copy number was lower than gene copy number for most of the bacterial and archaeal genes present in this coastal ocean.

Coverage depth

Coverage depth is defined here as the percent of the unique mRNAs present in a sample that is represented in the sequence library. Sample-sequencing depth and coverage depth are not strictly coupled, as a low richness/high evenness community transcriptome will be well covered even with shallow sample sequencing.

We evaluated coverage depth for the coastal metatranscriptomes in terms of taxa, functional gene categories and genes. Taxonomic coverage, as assessed by a collector's curve of NCBI taxonomy bins at the species or strain level, was approaching saturation for the library (Figure 2, inset); indeed, 75% of the total taxonomic richness emerging from this analysis would have been discovered with <15% of the sequencing effort. Saturating coverage was also found for functional gene assignments based on best hits to the COGs database (Table 1); 75% of the total richness would have been found with <10% of the sequencing effort (Figure 2, inset). However, these coverage assessments are constrained by the composition of the reference database, as apparent richness can be no higher than the number of reference bins available for transcript assignment. We found 1909 taxon bins represented in the metatranscriptomic libraries out of 8054 entries in the NCBI taxonomy database, and 3298 COGs out of 5666 entries in the COG database.

When coverage was assessed based on gene assignments in the RefSeq database (>6 million accession numbers), transcripts binned to over 168 000 genes (Table 1), and the collector's curve indicated that the metatranscriptome library was far from saturating (Figure 2). Singletons made up 59% of the sequences (Supplementary Figure S2), and abundant transcripts (>10 hits accession number⁻¹) and highly abundant transcripts (>100 hits accession per number) composed only 3% and 0.5% of the library, respectively. Although RefSeq binning

Table 2 Estimation of the number of bacterioplankton mRNA molecules in coastal seawater and sequencing depth of the metatranscriptomic libraries

Calculation method	Sample	mRNA molecules per liter	mRNA molecules per cell ^b	Sequencing depth (%)
Internal standard ^a	FN56	1.0×10^{12}	238	0.000009
	FN57	0.6×10^{12}	142	0.000015
Extracted RNA mass ^c	FN56	0.2×10^{12}	48	0.000043
	FN57	0.4×10^{12}	95	0.000020
Per cell RNA content ^d	FN56	2.6×10^{12}	619	0.000003
	FN57	2.6×10^{12}	619	0.000003

^aThe libraries contained 4014 (FN56) and 6865 (FN57) copies of the internal standard out of a total of 500 092 (FN56) and 465 519 (FN57) potential protein encoding sequences. The standard was added at 4.7×10^{10} copies per 5.75 liter of seawater just before cell lysis for total RNA extraction (see Materials and methods for details).

^bCell numbers in the $3 \mu\text{M}$ filtrate averaged 4.2×10^9 per liter based on epifluorescence microscopy.

^cExtraction yields were 14.4 (FN56) and 32.9 (FN57) μg total RNA from 5.75 liter of seawater. Total RNA is assumed to contain 4% mRNA by mass (Neidhardt and Umberger, 1996) and bacterial mRNAs are assumed to average 924 nt (Xu *et al.* (2006)).

^dMarine bacterial cells are assumed to contain 5.7 fg total RNA per cell (mid point of 1.9–9.5 fg range reported by Simon and Azam (1989)). See footnote c for estimate of percent mRNA by mass and footnote b for cell counts per litre.

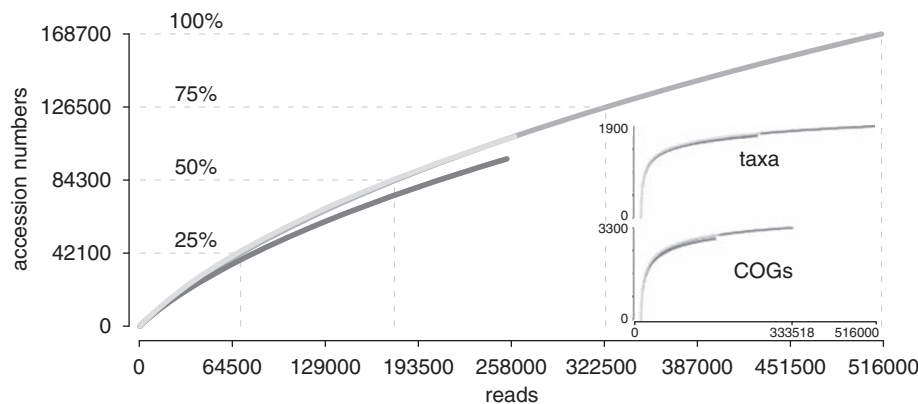


Figure 2 Collector's curve of gene richness as a function of reads analyzed. Light gray: FN56; dark gray: FN57; medium gray: combined libraries. Dashed lines indicate the number of reads needed to reach quarter percentiles of the total richness of the combined library. Inset: collector's curves for taxonomic and functional gene category (COG) richness, with the y axis corresponding to the number of unique reference organisms or COG numbers.

could overestimate transcript richness (if identical transcripts bin to different reference genes because of differences in the region sequenced or because of sequencing errors), it also underestimates richness (if a variety of sequence variants bin to the same reference gene; Figure 3). In any event, despite efforts to sequence more deeply than typical, our libraries exhibited the same low coverage that has been reported in previous metatranscriptomic analyses of marine bacterioplankton communities (Frias-Lopez *et al.*, 2008; Poretsky *et al.*, 2010; Stewart *et al.*, 2010).

Metatranscriptomes might be expected to have lower richness compared with metagenomes if expression is limited to a small fraction of the bacterial genome at any one time. In this case, they would also have higher coverage than metagenomes for the same size sequence library (Gilbert *et al.*, 2008). However, for this coastal metatranscriptome, the distribution of hits per gene (Supplementary Figure S2) did not indicate dominance by a limited number of highly transcribed genes (Figure 2).

Similarly, a synchronized clonal population of *Bacillus anthracis* expressed 40–80% of genes under all growth conditions tested (Passalacqua *et al.*, 2009), suggesting that the population's transcriptome was only slightly less rich than its genome. Even allowing for significant advances in sequencing technology, the extremely low sample-sequencing depth found in this study suggests that most natural community transcriptomes will continue to be undersampled.

Microdiversity

Assembly of transcripts from the most highly expressed genes (>1000 reads for some) revealed significant variation within reference bins (Hollibaugh *et al.*, in press). For example, the 2259 reads that binned to the *Pelagibacter ubique* HTCC1002 proteorhodopsin gene (PU1002_03206; 1 of 28 proteorhodopsin bins in the libraries) had high sequence diversity (Figure 3). That this observed diversity was in fact real biological variation was substantiated by

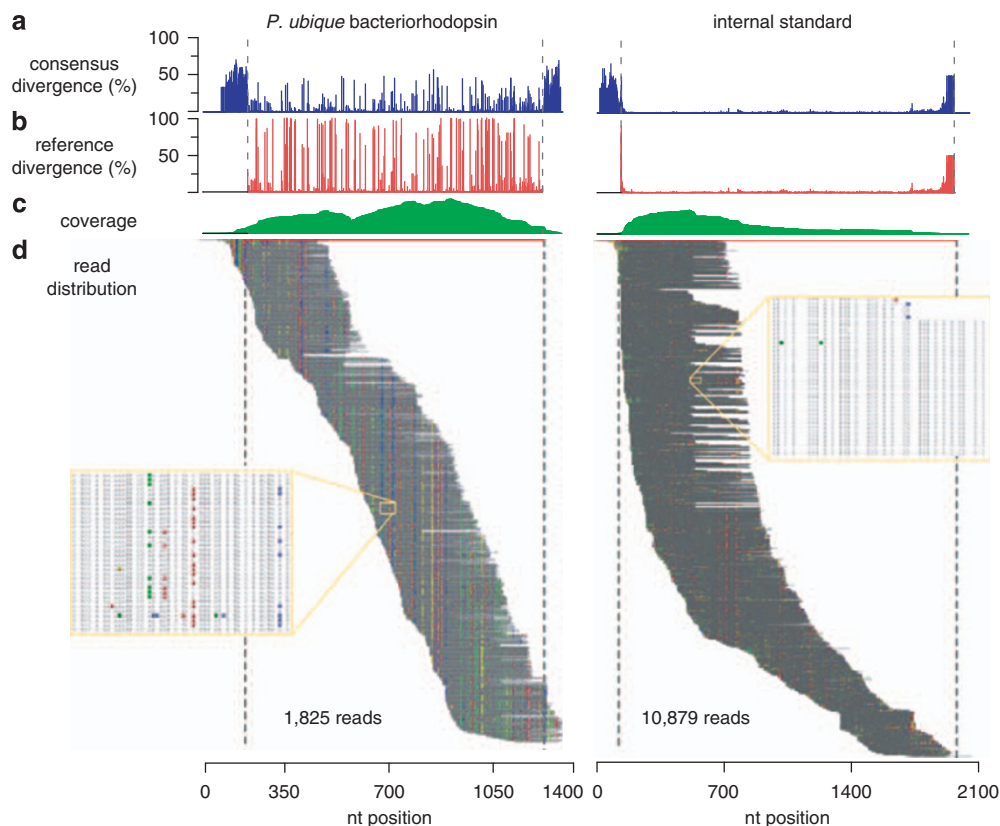


Figure 3 Assembly of 1825 reads (out of 2259 total) binning to the *P. ubiquus* HTCC1002 proteorhodopsin gene PU1002_03206 (left), and of 10 879 reads (out of 10 879 total) binning to the internal transcript standard (right). (a) Percent nucleotide divergence from the consensus sequence. (b) Percent nucleotide divergence from the reference sequence. (c) Coverage by nucleotide position. (d) Read assembly to the reference gene (shown in red), with dashed lines indicating start and end positions of the reference. Note that the reference gene lengths are extended by assembly gaps. Divergence from the consensus sequence (that is, the majority nucleotide at a given position) is indicated as follows: A = red, T = green, C = blue and G = yellow. Insets show close-up regions of assemblies.

an assembly of the internal standard reads (Figure 3), which indicated a mean sequencing error rate in this study of 3.7 ± 7.4 per 1000 bp compared with a mean sequence variation rate of $97.6 (\pm 28.0)$ per 1000 bp for transcripts binning to PU1002_03206. Although high diversity in proteorhodopsin genes has been found previously in the ocean (Rusch *et al.*, 2007; Campbell *et al.*, 2008), the metatranscriptomic data revealed simultaneous expression of scores of microdiverse sequence variants. Transcriptome coverage estimates based on gene binning to the RefSeq database are considerable underestimates of the true sequence richness.

Detection of biogeochemically informative transcripts

We determined the absolute abundance of transcripts for key genes representing the P cycle (25 diagnostic genes), N cycle (50 genes) and S cycle (7 genes) (Figure 4). Transcripts were found for 56% of genes surveyed. Most P and sulfur cycle transformations were represented by at least one transcript. N cycle expression was dominated by ammonia transporter and ammonia monooxygenase transcripts, which had the highest copy numbers of

any gene category (2.7×10^9 transcripts per liter; Figure 4); many other N cycle genes were not detected at all.

To examine detection of biogeochemically diagnostic mRNAs in theoretically smaller libraries, the full metagenomic libraries were randomly subsampled *in silico* to generate subsets. The majority of the elemental cycle transcripts detected in the full libraries were still evident in smaller libraries. For example, > 80% of the P-cycle related genes would have had at least one hit in a library one-fourth the size (Supplementary Figure S3).

Statistical resolution

Comparative metatranscriptomics seeks to differentiate transcript abundance between samples, for example, across natural environmental gradients or in response to experimental manipulations. We examined the statistical power of comparative analyses as a function of library size, starting first with broad categories of gene function as represented by COG assignments. Subsets of each of the replicate libraries were generated *in silico* and the

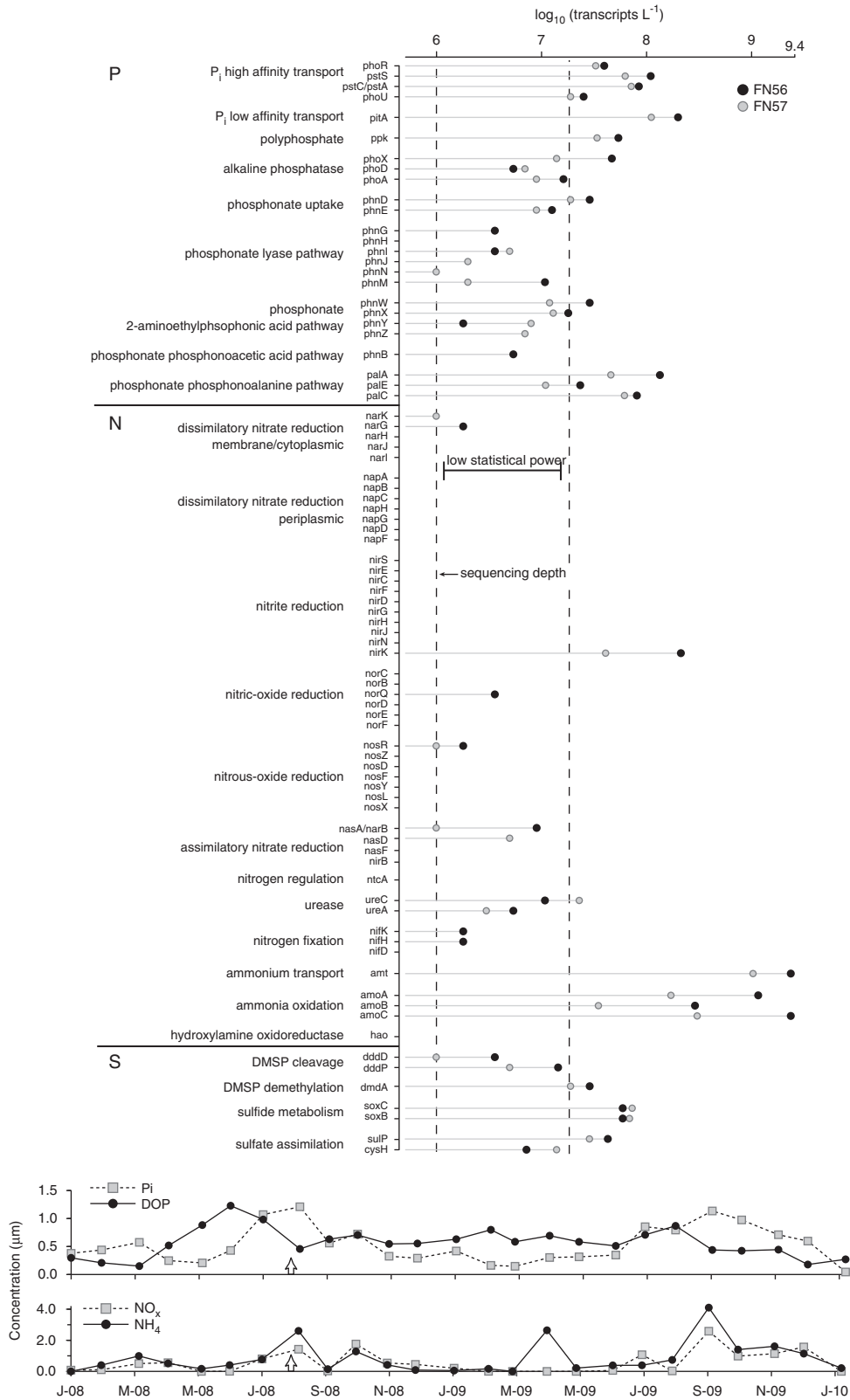


Figure 4 Copy numbers of phosphorus, nitrogen and sulfur cycle transcripts in a coastal ocean microbial community. The left line represents the limit of detection for this study, and together with the right line defines the region where copy numbers are too low for robust statistical analysis (that is, where the fold-difference requirement is > 2). Symbols indicate copy numbers in biological duplicates. Bottom graphs show monthly nutrient concentrations for GCE-LTER station six. The arrows mark the date of sample collection.

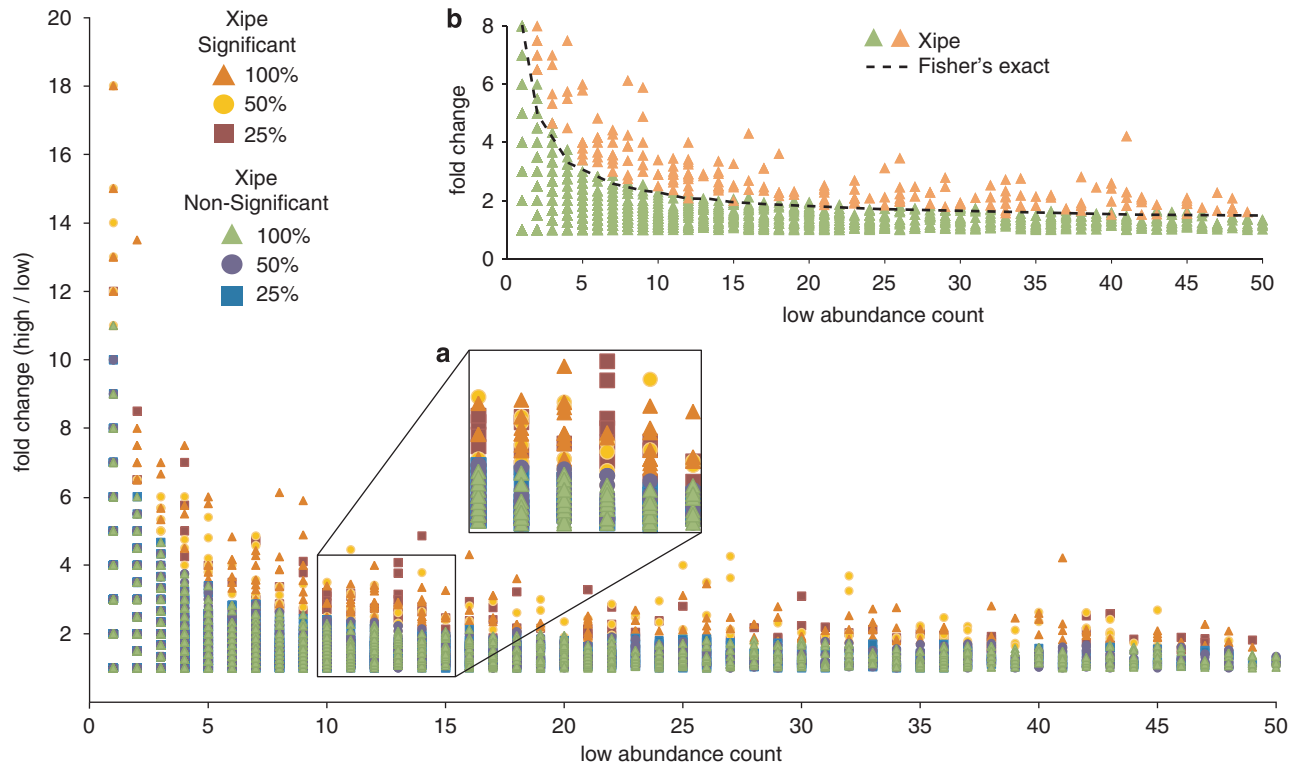


Figure 5 Minimum fold difference required for statistical significance (Xipe, $P < 0.05$) as a function of both the count in the lower abundance sample and the library size. Samples and subsamples were from the combined libraries (FN56 and FN57). Marker color is based on the statistical outcome (significant or nonsignificant) and library size (percent of full library). (a) Zoom of region in the main figure. Note that the minimum fold-difference for significance is independent of the three library sizes analyzed. (b) An alternative analysis of the significance threshold using contingency tables and Fisher's exact test. The minimum fold-difference threshold at which a low abundance count is significant by the Fisher's exact test is plotted as a dotted black line. The results from the Xipe analysis (main figure) at the 100% library size are also shown in inset B for direct comparison with the Fisher's exact test.

fold-difference criteria (high abundance count/low abundance count) needed for statistically significant differences were compared using a resampling method based on difference of medians (Xipe; Rodriguez-Brito *et al.*, 2006). Even for libraries one-fourth of the original size, there was little effect on the fold-difference threshold required for a COG category to be considered significantly different between samples (Figure 5). This was true as well for an alternate statistical approach using contingency tables and Fisher's exact test (White *et al.*, 2009) (Figure 5, inset b), and also when analyzing libraries much smaller than the original (for example, the average fold-difference threshold for significance in a 10 000 read library was $< 1\%$ greater than in a 500 000 read library).

Library size, however, had a direct impact on the number of counts in a transcript category, thereby affecting the power of statistical comparisons. Transcript categories with low copy numbers (defined here as ≤ 15 hits in the lower abundance sample) required from 2- to 8-fold difference between the two samples for statistical significance (Figure 5). For smaller *in silico* subsets of the libraries or for more specific transcript annotation categories (for example, RefSeq gene

bins), both of which result in lower counts per category, the power to detect statistical differences between two samples decreased. For example, 17 out of the 25 genes that mediate key steps in the marine P cycle fell into a low-count category even with the full-size library (Figure 4), and nearly all would do so if the library was one-fourth of the original size. For metatranscriptomic libraries of the magnitude obtained in this study ($> 1\,000\,000\,454$ FLX reads), only those transcripts present at concentrations $> 1 \times 10^6$ per liter had a good probability of being detected, and only those present at concentrations $> 1.5 \times 10^7$ per liter (which would exclude all singletons and other low-count transcript categories) could be compared across samples with good statistical power.

Replication

The need to improve sample-sequencing depth competes with the need for replication in comparative metatranscriptomic analyses. Two important sources of variability that can be quantified through replication include technical variation during sample processing/sequencing, and natural biological variation within the environment sampled.

For the first type, 454 pyrosequencing is prone to artifacts in which single DNA fragments are sequenced more than once ('duplicate sequences'). Although artifactual duplicates are recognized in metagenomes as sequences with identical 5'-sequence and high identity throughout (Dinsdale *et al.*, 2008; Gomez-Alvarez *et al.*, 2009), true duplicate sequences can arise in metatranscriptomes from discrete mRNAs from highly expressed genes. In this study, 24% of RefSeq reads were 'duplicates' (same start site and $\geq 90\%$ identity). As each replicate biological sample was sequenced as four technical replicates (independent emulsion PCRs and sequencing runs), and assuming that artifactual duplicates arise during the emulsion PCR step (Gomez-Alvarez *et al.*, 2009; Stewart *et al.*, 2010), artifactual duplicates should have uneven distributions across the four 454 runs, whereas natural duplicates should be evenly distributed. We found that most duplicate clusters averaged $\sim 25\%$ per technical sequencing replicate (Supplementary Figure S4), and a statistical comparison of COG assignments for all six within-sample pairwise combinations of the technical sequencing replicates indicated that only 0.2% fit the pattern for artifactual duplicates (significantly higher in one technical replicate compared to the other three). For the transcript with the highest copy number in the combined library (Rac prophage; ZP_03400590), removal of duplicate reads would have decreased the count by 98% (from 6235 to 111 hits) despite evidence from technical replicates that many of these are natural (Supplementary Figure S5). Duplicate removal from metatranscriptomic libraries based on sequence start position and percent identity (Gomez-Alvarez *et al.*, 2009; Stewart *et al.*, 2010) may therefore produce systematic underestimates of abundance for the most highly transcribed genes in the community, and statistical analysis of technical replicates is a recommended alternative.

For the second type of variation, within-treatment biological variability sets the false-positive rate against which differences in gene expression patterns across treatments or environments can be evaluated (Poretsky *et al.*, 2010). In this study, patchiness in community gene transcription patterns was detectable in paired coastal seawater samples separated by ~ 300 m (based on tidal flushing rates past a fixed collection point). At the level of functional gene categories, pairwise comparisons indicated significant differences between the samples for 461 of 3298 COGs (14%) (Xipe, $P < 0.05$). Only nine significant COGs contained sequences from a putative artifactual duplicate cluster (see above), highlighting the benefit of technical replicate averaging for reducing spurious differences from sequencing artifacts. In accordance with other studies of environmental sequence libraries (Rodriguez-Brito *et al.*, 2006), as well as our observations above, decreasing the library size had a major influence on the number

of significant differences that were detectable (Supplementary Table S1).

Differences between replicate samples at the individual gene level (that is, transcripts binned by RefSeq hits) were also examined, using Fisher's exact test coupled with a correction for the false discovery rate (Strimmer, 2008) to control for type I errors arising when simultaneously conducting large numbers of statistical tests (in this case, for $> 186\,000$ different RefSeq bins). Eighty-three (0.05%) of the gene bins were statistically different between the two samples ($P < 0.05$ with Benjamini-Hochberg correction), including those representing phage genes, ammonia oxidation genes and various genes for light-driven energy acquisition (Supplementary Figure S6). The replicate samples therefore established within-treatment variability (Figure 4) for future between-treatment comparisons.

Microbial gene expression in a coastal ocean

Transcripts from the combined library binned to genes from 1909 reference organisms. Thirty-three percent of the sequences had best hits to alphaproteobacteria genes (with roseobacters accounting for 11% and *P. ubique* for 7%) and 27% had best hits to gammaproteobacteria genes (Figure 6). Unexpectedly, 4% of the transcripts binned to the two archaeal genomes of *Nitrosopumilus maritimus* (3.3%) and *Cenarchaeum symbiosum* (0.1%). Although Archaea are often abundant and active in deep ocean environments, they were not expected to contribute significantly to gene expression in this shallow coastal water system; however, the *N. maritimus* taxonomic bin was the second largest in the metatranscriptome (Hollibaugh *et al.*, in press). The oligotrophic marine gammaproteobacteria clades, which are usually in low abundance in oceanic 16S rRNA libraries ($< 3\%$; Cho and Giovannoni, 2004), were also unexpectedly well represented in the metatranscriptome (Figure 6; 8% of transcripts), and transcripts binning to three genes from a gammaproteobacteria prophage (3% of transcripts) may indicate an ongoing infection of these oligotrophic marine gammaproteobacteria populations. Eukaryotic transcripts composed 6% of the total, with those binning to *Ostreococcus* spp. particularly well represented (20% of eukaryotic hits).

Copy numbers of transcripts representing 82 genes diagnostic for P, N and S cycling were determined simultaneously from the metatranscriptomic data (Figure 4). For P transformations, annotations suggest bacterioplankton were transporting phosphate by both high- and low-affinity transporters. The expression of low-affinity transporters, along with polyphosphate storage genes, is consistent with elevated phosphate concentrations ($1.2\ \mu\text{M}$) at the time of sampling, which is typical of late summer in this coastal ocean (Figure 4). Expression patterns also indicated ongoing use of organic P, including phosphoesters (by *phoX*, *phoD*

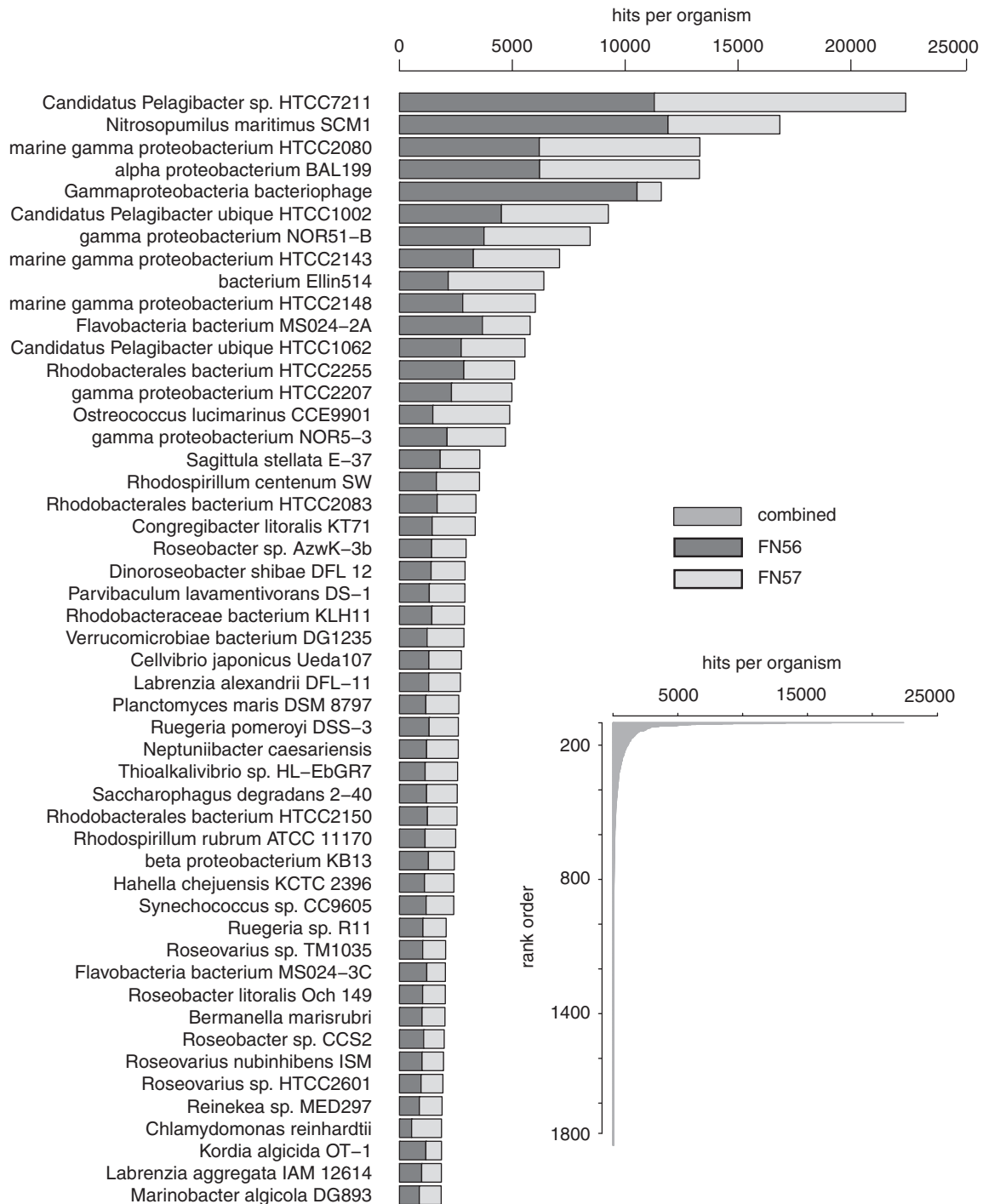


Figure 6 Rank-order abundance of taxonomic bins (species or strain level). Main figure: top 50 taxonomic annotation bins; inset: all 1909 taxonomic annotation bins.

and *phoA*) and phosphonates (although transcripts for the canonical C-P lyase pathway were near the limit of detection). For N transformations, sampling occurred during a local ammonia peak (2.6 μM ; Figure 4), and transcripts related to the uptake and oxidation of ammonia (*amt*, *amoA*, *B* and *C*) were orders of magnitude higher in abundance than genes mediating nitrate or nitrite processing (for example,

nar, *nap* and *nir* genes) (Figure 4). Transcripts for urea metabolism, the only representative of dissolved organic N use included in the analysis, made up the second most abundant group of N-related sequences (Figure 4). Nitrogen is often the limiting nutrient (or co-limiting with carbon; Pomeroy *et al.*, 2000) to microbial activity in this ecosystem, and dissolved organic N is 2- to 200-fold

higher in concentration than inorganic N. For S transformations, gene expression suggested substantial use of reduced S compounds typically found in high concentrations in marsh-dominated coastal systems (Kiene and Capone, 1988; Pakulski and Kiene, 1992). Transcripts were found for metabolism of dimethylsulfoniopropionate (*dmdA*, *dddP* and *dddD*), as well as oxidation of sulfide/thiosulfate (*sox* genes) (Figure 4). This broad inventory of P, N and S cycle transcripts represents an absolute benchmark against which time-series and experimentally manipulated transcriptomes in this ecosystem can be compared.

Conclusions

Addition of an internal mRNA standard provides a significant advantage in metatranscriptomics protocols as it allows estimation of the fraction of the microbial transcriptome captured in the sequence library, as well as the absolute quantification of transcript copy number in the environment (Figures 1 and 4). Although RT-qPCR approaches can also provide absolute transcript numbers, often with greater sensitivity (Church *et al.*, 2010), they are currently limited to a handful of functional genes at a time. Furthermore, the high microdiversity found in many natural gene populations (Figure 3) makes primer design challenging (Varaljay *et al.*, 2010), and likely results in RT-qPCR only quantifying a subset of the total functional gene population. Multiple internal standards that vary in length and concentration (van de Peppel *et al.*, 2003) will allow for more robust calculations of sequencing depth in future studies, and better position 'omics' data for integration with biogeochemical rate measurements.

As low-count transcript categories are difficult to resolve statistically, library size had a critical effect on comparative metatranscriptomic analyses. Many of the biogeochemically diagnostic transcripts detected in our libraries would have been detected in ones that were one-fourth or one-tenth the size, but these theoretically smaller libraries resulted in a decreased ability to statistically differentiate between samples. Typical library sizes for metatranscriptomes (10^5 – 10^6 sequence reads) are therefore sufficient for descriptive studies, but significant gains in comparative analyses of biogeochemically informative gene expression patterns will require a greater sequencing investment. Indeed, 54% of the 46 detected steps in the marine N, P and S cycles would require at least a twofold difference in copy number between samples to meet statistical criteria for hypothesis testing (that is, $P < 0.05$). Although a twofold change in transcript abundance is an appropriate minimum criterion for expression studies of clonal bacterial cultures in synchronized growth (Bürgmann *et al.*, 2007), it may fail to catch smaller expression differences among complex microbial communities that are ecologically relevant.

Despite a sample-sequencing depth of only 1 in 10^7 transcripts, the libraries provided remarkable

insights into gene expression in a marine microbial community, including evidence for active microbes not known previously to have a major role in the ecosystem and quantification of transcripts for scores of steps in marine elemental cycles. These data establish the foundation for comparative assessments of diel, seasonal and annual changes in microbial gene expression that will provide insights into the regulation of biogeochemical processes in the coastal ocean.

Acknowledgements

We thank R Newton for assistance with sample collection and bioinformatics analysis, L Tomsho and S Schuster at Penn State University for 454 sequencing expertise, JT Hollibaugh for comments and discussion on the article and S Rathbun for helpful discussions on statistical methods. Nutrient data were provided by K Hunter and S Joye through the Georgia Coastal Ecosystems Long Term Ecological Research program (OCE-0620959). This project was supported by funding from the Gordon and Betty Moore Foundation and the National Science Foundation Microbial Observatories Program (MCB-0702125).

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