

ORIGINAL ARTICLE

Bacterial transcriptome remodeling during sequential co-culture with a marine dinoflagellate and diatom

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In their role as primary producers, marine phytoplankton modulate heterotrophic bacterial activities through differences in the types and amounts of organic matter they release. This study investigates the transcriptional response of bacterium *Ruegeria pomeroyi*, a member of the Roseobacter clade known to affiliate with diverse phytoplankton groups in the ocean, during a shift in phytoplankton taxonomy. The bacterium was initially introduced into a culture of the dinoflagellate *Alexandrium tamarense*, and then experienced a change in phytoplankton community composition as the diatom *Thalassiosira pseudonana* gradually outcompeted the dinoflagellate. Samples were taken throughout the 30-day experiment to track shifts in bacterial gene expression informative of metabolic and ecological interactions. Transcriptome data indicate fundamental differences in the exometabolites released by the two phytoplankton. During growth with the dinoflagellate, gene expression patterns indicated that the main sources of carbon and energy for *R. pomeroyi* were dimethylsulfoniopropionate (DMSP), taurine, methylated amines, and polyamines. During growth with the diatom, dihydroxypropanesulfonate (DHPS), xylose, ectoine, and glycolate instead appeared to fuel the bulk of bacterial metabolism. Expression patterns of genes for quorum sensing, gene transfer agent, and motility suggest that bacterial processes related to cell communication and signaling differed depending on which phytoplankton species dominated the co-culture. A remodeling of the *R. pomeroyi* transcriptome implicating more than a quarter of the genome occurred through the change in phytoplankton regime.

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Introduction

Marine bacteria participate in the biogeochemical cycling of all major elements. They are the primary degraders of marine dissolved organic matter (DOM), a pool of carbon equivalent in mass to the atmospheric CO₂ reservoir (Hansell, 2013). Phytoplankton constitute the main source of labile carbon for bacteria in the surface ocean via the metabolites they release into the DOM pool, either through exudation or death-related processes such as senescence, predation, or viral lysis (Azam *et al.*, 1983).

Diatoms and dinoflagellates are common phytoplankton groups in the ocean (Hasle, 1976; Bathmann *et al.*, 1997; Taylor *et al.*, 2008), and mixed assemblages or successions of taxa from these groups have been observed in coastal regions and upwelling systems (Seeyave *et al.*, 2009; Ryan *et al.*, 2014; Varaljay *et al.*, 2015). One consequence

of fluctuations in primary producer composition is that the supply of labile compounds to heterotrophic bacteria is likely to vary (Becker *et al.*, 2014). Recent work shows that marine bacteria possess strategies to efficiently detect and utilize phytoplankton metabolites by rapidly congregating near particles, including phytoplankton cells (Smriga *et al.*, 2016). Active movement to the enriched phycospheres associated with phytoplankton (Stocker *et al.*, 2008) allows bacteria access to a wide variety of metabolites, a potentially effective strategy for growth in a diverse and patchy chemical environment.

Bacteria may also have a role in shaping the availability of labile organic matter, influencing metabolite release through excretion of diffusible compounds required by phytoplankton; examples of such compounds include vitamins and enzymes (Croft *et al.*, 2005; Grossart and Simon, 2007; Biller *et al.*, 2016). In cases where interactions are species-specific, bacteria are likely to require repertoires of physiological and ecological strategies unique to each primary producer (Sison-Mangus *et al.*, 2014). These interactions may have important consequences for global-scale carbon fluxes (Harvey *et al.*, 2016; Segev *et al.*, 2016).

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The Roseobacter clade is a marine alphaproteobacterial group whose members are often found associated with phytoplankton (González *et al.*, 2000). Diatom and dinoflagellate blooms in the ocean are usually associated with high Roseobacter abundance, and laboratory cultures of these phytoplankton groups are frequently populated with Roseobacter strains (Lafay *et al.*, 1995; Schäfer *et al.*, 2002; Jasti *et al.*, 2005; Amin *et al.*, 2012). One hypothesis for why roseobacters co-occur with phytoplankton is that their typically large genomes encode high metabolic versatility and sophisticated regulatory capabilities (Newton *et al.*, 2010) allowing them to take advantage of increased substrate concentrations in phycospheres (Luo and Moran, 2014). The observation that individual Roseobacter strains have different effects on phytoplankton growth and metabolism supports the hypothesis that specific mechanisms exist for recognition and interaction (Seyedsayamdost *et al.*, 2011; Amin *et al.*, 2015), and these could explain the success of roseobacters in phytoplankton blooms.

In this study, we established a three-member model system to examine transcriptional changes in bacteria in response to shifts in the dominant primary producer. The bacterium selected for the model system is *Ruegeria pomeroyi* DSS-3, a member of the Roseobacter clade for which a genome sequence (Moran *et al.*, 2004) and genetic system (Howard *et al.*, 2006) is available. The two other members are the dinoflagellate *Alexandrium tamarense* CCMP1771 and the diatom *Thalassiosira pseudonana* CCMP1335, two phytoplankton strains that are well characterized and readily grow in culture (Armbrust *et al.*, 2004; Moustafa *et al.*, 2010). In the model system, bacterial carbon and nitrogen requirements were satisfied exclusively by metabolites provided first by *A. tamarense*, and subsequently by *T. pseudonana* as it outcompeted the dinoflagellate. Here we report changes in the bacterial transcriptome that address three questions: (1) How plastic is gene expression by *R. pomeroyi* when associating with different phytoplankton taxa? (2) What different metabolites are released from the model dinoflagellate and diatom? (3) Do transcriptional responses suggest a role for interactions or signaling between the bacterium and primary producers?

Materials and Methods

Experimental set up

Axenic phytoplankton cultures from the National Center for Marine Algae were maintained without antibiotic addition and examined regularly by microscopy and plating to check for bacterial contamination. Six 20-l LDPE cubitainers (Reliance, Winnipeg, Canada) were filled with 18 l of autoclaved, 0.2 µm-filtered Guillard's f/2 medium with vitamin B₁₂ added at f/50 concentration. Two

liters of an axenic dinoflagellate *Alexandrium tamarense* CCMP1771 culture grown to exponential phase in the same medium were inoculated into each cubitainer. Cubitainers were maintained at 18 °C with a 16–8 h light-dark cycle under ~160 µmol photons m⁻² s⁻¹. After seven days, three cultures were inoculated with *Ruegeria pomeroyi* DSS-3 while the other remained axenic. Bacteria had been grown overnight in ½YTSS medium, harvested in exponential phase, washed five times with sterile phytoplankton medium, and inoculated at ≤10⁶ cells ml⁻¹. Five days after bacterial inoculation, 800 µl of an axenic culture of diatom *T. pseudonana* CCMP1335 prepared as for *A. tamarense* were inoculated into all six cubitainers, and silicate was added (100 µM final concentration). The order in which the phytoplankton were introduced was determined based on *T. pseudonana*'s ability to outcompete *A. tamarense* under the culture conditions used. Cubitainers were maintained for 37 days.

Sampling and monitoring

To monitor growth, triplicate samples of 0.6 ml were mixed with 0.6 ml of glutaraldehyde (2% final concentration) and stored at -80 °C. Once thawed, 500 µl with 5 µm fluorescent beads added (Spherotech, Lake Forest, IL, USA) were stained for 20 min with SYBR Green I (0.75X recommended concentration, Life Technologies, Waltham, MA, USA) and analyzed on a CyAn instrument (Beckman Coulter, Brea, CA, USA) with a 488 nm laser. SYBR Green fluorescence (bacteria) was detected using a FL1–530/30 bandpass filter and chlorophyll *a* fluorescence (phytoplankton) using a FL4–680/30 bandpass filter. Data were analyzed in FlowJo. Bacterial contamination was checked by streaking onto ½YTSS plates. No-bacteria phytoplankton controls were checked for any growth, while experimental cubitainers were checked for growth of bacteria other than *R. pomeroyi*. On days 7, 9, 12, 15, 18, 23, 30, and 37, 1-liter samples were collected for RNA sequencing and chemical analysis.

Chemical analyses

Duplicate 6-ml samples were collected from cubitainers, filtered through 0.45-µm hydrophilic polyethersulfone Acrodisc Supor membranes (Pall Life Sciences, Port Washington, NY, USA), and stored at -20 °C. Samples were analyzed for dissolved inorganic nitrogen (nitrate+nitrite) (APHA Standard Method 4500-NO₃ F), ammonium (4500-NH₃ G), dissolved inorganic phosphorus (4500-PF), and silicate (spectrophotometric silicomolybdate assays). For DOC, triplicate samples filtered through ashed Whatman GF/F filters were combusted on a Shimadzu TOC-5000A Analyzer. Analyses were performed at the Center for Applied Isotope Studies (University of Georgia, Athens, GA, USA).

Bacterial RNA extraction and sequencing

One liter samples were filtered through 2- μm polycarbonate (PC) membranes to remove eukaryotic cells, and then through 0.2- μm PC membranes to collect bacterial cells. Filters were flash-frozen in liquid nitrogen and stored at -80°C . For RNA extraction, filters pieces were incubated at 37°C for 1 h in TE buffer, SDS (0.6% final concentration), and proteinase K (120 ng μl^{-1} final concentration, Qiagen, Hilden, Germany). Acid phenol:chloroform:isoamylalcohol was added (1:1 volume) and samples were shaken, incubated for 15 min, and centrifuged at 4°C for 15 min for collection of the supernatant. The process was repeated and the supernatant mixed 1:1 vol:vol with isopropanol, vortexed, incubated for 15 min, centrifuged at 4°C for 15 min, washed twice with 75% ethanol, and dried and resuspended in RNase-free water. RNA was precipitated with sodium acetate (0.3 M final concentration) and three volumes of 100% ethanol, and incubated overnight at -20°C . Pellets were washed twice with 75% ethanol, centrifuged, dried, and resuspended in RNase-free water.

Potential traces of DNA were removed using the Turbo DNA-free kit (Invitrogen, Waltham, MA, USA). Samples were tested for residual DNA by a 40-cycle PCR targeting the 16S rRNA gene of *R. pomeroyi*. Samples were depleted of rRNA using custom probes for small and large subunit rRNA genes from all three microbes (Stewart *et al.*, 2010). Libraries were prepared for two replicate cubitainers at eight time points (16 samples) using the KAPA Stranded mRNA-Seq Kit (Kapa Biosystems, Wilmington, MA, USA) at the Georgia Genomics Facility (University of Georgia) and sequenced on a HiSeq Illumina 2500 at the Hudson Alpha Institute for Biotechnology (AL, USA).

Bioinformatic analysis

Quality control was performed on 249 million 50-bp reads (10 ± 2 million reads per sample; Supplementary Table S1) using the FASTX toolkit, imposing a minimum quality score of 20 over 80% of read length. Reads aligning to an in-house rRNA database were removed (blastn, score cutoff ≥ 50). Remaining reads were mapped to the *R. pomeroyi* genome (Bowtie 2; Langmead and Salzberg, 2012) and counted (HTSeq; Anders *et al.*, 2015), conserving strand information and removing reads that mapped to more than one location (Supplementary Table S1). Counts were converted to transcripts per million (TPM, Supplementary Table S2) and data deposited in the NCBI BioProject database under accession PRJNA381627.

Genes with differential expression between selected time points were determined with DESeq2 (Love *et al.*, 2014), and linear relationships of gene expression with phytoplankton abundance with Pearson correlations. Self-organizing maps (SOM) were built with the Kohonen package in R (Wehrens

and Buydens, 2007) using expression of each gene (in TPM) normalized by dividing by the median expression for that gene across the 8 time points followed by a \log_2 transformation.

DHPS transporter mutant

A deletion mutant of DHPS transporter operon *hpsKLM* was constructed by homologous recombination (Reisch *et al.*, 2011). Regions up- and downstream of the target region (~1 kb) and *tetAR* were cloned into pCR2.1 using sequence and ligation-independent cloning (SLIC) (Li and Elledge, 2007) and electroporated into competent *R. pomeroyi* cells. Recombined clones were selected on tetracycline-amended medium. Triplicate wild type and Δ *hpsKLM* cultures were incubated at 30°C in marine basal medium amended with DHPS, glucose, xylose, or acetate as a sole carbon source; or in co-cultures with *T. pseudonana*. Growth was assessed with OD measurements over 4 days (pure cultures) or on $\frac{1}{2}$ YTSS plates, amended with tetracycline in the case of the deletion mutant (co-cultures).

Xylose transporter expression

Triplicate *R. pomeroyi* cultures were grown on each of four substrates at 30 mM carbon (5 mM glucose, 15 mM acetate, 10 mM pyruvate, 6 mM xylose). Cells were collected in early exponential phase, RNA extracted using the RNeasy mini extraction kit (Qiagen), and DNA removed as described above. *xylG* (SPO0863) transcripts were assayed by RT-qPCR on an iCycler iQ5 (Bio-Rad, Hercules, CA, USA) using the iTaq SYBR Green one-step kit with a six-fold serially-diluted standard (10^2 to 10^7 copies per reaction). Negative controls had no reverse transcriptase addition.

Results and Discussion

In this three-member co-culture system, the phytoplankton species were the sole source of carbon and usable nitrogen for marine bacterium *Ruegeria pomeroyi*, since no exogenous carbon source was added and *R. pomeroyi* cannot assimilate nitrate (Moran *et al.*, 2004). Over the 37 days of the experiment, there was a progression from a dinoflagellate-dominated to a diatom-dominated system. *Alexandrium tamarense* was the only phytoplankton species present until day 12 when *Thalassiosira pseudonana* was inoculated into the cultures. At day 15 the diatoms became numerically dominant and remained so until the end of the experiment. During the final stage, dinoflagellate cells decreased dramatically, and they were no longer detectable at day 30 (Figure 1). A biovolume-based estimate accounting for size differences showed nearly equivalent biovolume for each species at their peak abundance, with the switch in biovolume dominance occurring at day 19

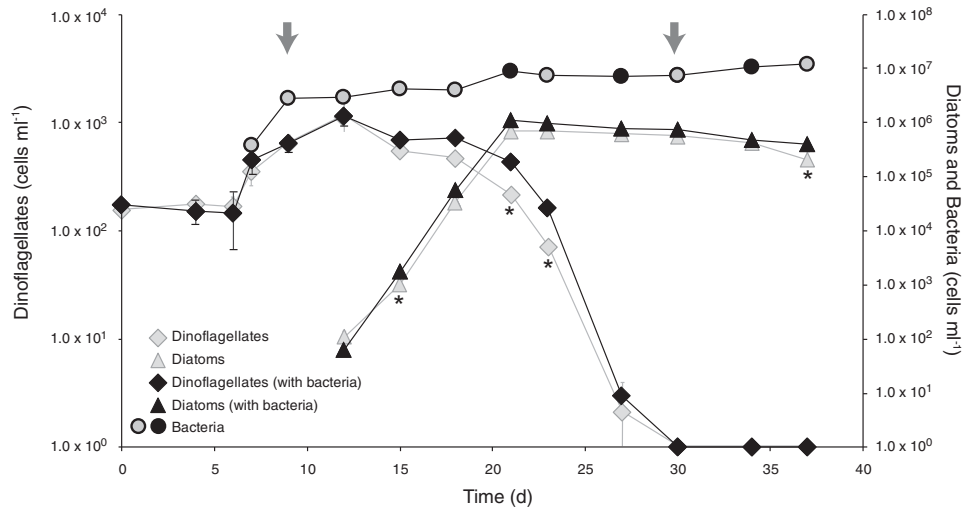


Figure 1 Cell counts of dinoflagellates (diamonds), diatoms (triangles), and bacteria (circles) as determined by flow cytometry. For phytoplankton, the gray diamonds and triangles correspond to the cell counts measured in no-bacteria controls. For bacteria, gray inner circles indicate RNaseq sample collection. Arrows identify the two time points tested for differential expression comparisons. Asterisks indicate significant differences in phytoplankton cell numbers between co-cultures and no-bacteria controls. Mean value \pm s.d. for the triplicate cultures are shown.

(Supplementary Figure S1). Chemical analysis indicated that the co-cultures were never depleted of nitrate or phosphate, although drawdown was evident (Supplementary Table S3). Ammonium remained undetectable, indicating that if any ammonium was produced, it was rapidly utilized. The phytoplankton sustained the growth of *R. pomeroyi* throughout the experiment, with a \sim 30-fold increase in bacterial counts between inoculation and day 37 (Figure 1). On the basis of the no-bacteria phytoplankton controls, bacterial presence did not affect maximum phytoplankton abundance but may have extended the duration of the stationary phase for both species (Figure 1, *t* test, $P \leq 0.02$).

Gene expression clusters

The *R. pomeroyi* transcriptome changed dramatically above the backdrop of shifting phytoplankton dominance. In a comparison of day 9, when dinoflagellates were the only species present in the co-culture, with day 30, when only diatoms were detectable, 1,179 genes in the *R. pomeroyi* genome (27%) were differentially expressed (DESeq2, $P \leq 0.05$). All 4,371 *R. pomeroyi* genes were sorted into clusters based on expression patterns over time using a self-organizing map algorithm (Kohonen, 2001; Wehrens and Buydens, 2007) (Supplementary Figure S2). Of the nine delimited clusters, Clusters A and B had patterns that broadly followed dinoflagellate abundance and Cluster I had patterns that followed diatom abundance (Figure 2, Supplementary Figure S3). Positive linear relationships with dinoflagellate cell numbers were found for 31% of genes in Cluster A and 1% in Cluster B, and with diatom cell numbers for 71% in Cluster I ($P \leq 0.05$, Figure 2). Further, many genes in Clusters A and B had significantly higher expression at dinoflagellate-dominated day 9 time point versus diatom-dominated day 30 time point

(80% and 49%), while almost all genes in Cluster I had significantly higher expression at day 30 versus day 9 (95%, Figure 2). Most of the other clusters contained few or no genes having positive linear relationships with phytoplankton abundance and few genes with differential expression between day 9 and 30 (Figure 2).

We also examined 86 bacterial genes whose expression is expected to correlate with cell activity. Assignment of these genes into the putative dinoflagellate or diatom clusters could indicate a growth signal rather than a specific phytoplankton response, since they encode intrinsic processes involved in protein metabolism and secretion, nucleic acid metabolism, and cell cycle and division (Gerdes *et al.*, 2003). Most of the activity-related genes grouped into Cluster D (49 genes), and to a lesser extent in Cluster H (17 genes) and Cluster C (13 genes) (Supplementary Figure S4). Only 1 grouped into the three phytoplankton-related clusters combined, indicating that expression of genes in Clusters A, B, and I does not simply track with bacterial growth rate or activity. Based on these multiple assessments of expression patterns, we subsequently focused analysis on the three gene clusters with clear relationships to dinoflagellate (Clusters A and B) or diatom (Cluster I) dominance in the co-cultures. In this evaluation, we concentrated on biogeochemically-relevant genes whose functions have been experimentally verified in *R. pomeroyi* in order to minimize speculation built on automated gene function assignment.

Metabolite utilization during growth with dinoflagellates

Several genes involved in degradation of dimethylsulfoniopropionate (DMSP), a phytoplankton osmolyte used by marine bacteria as a source

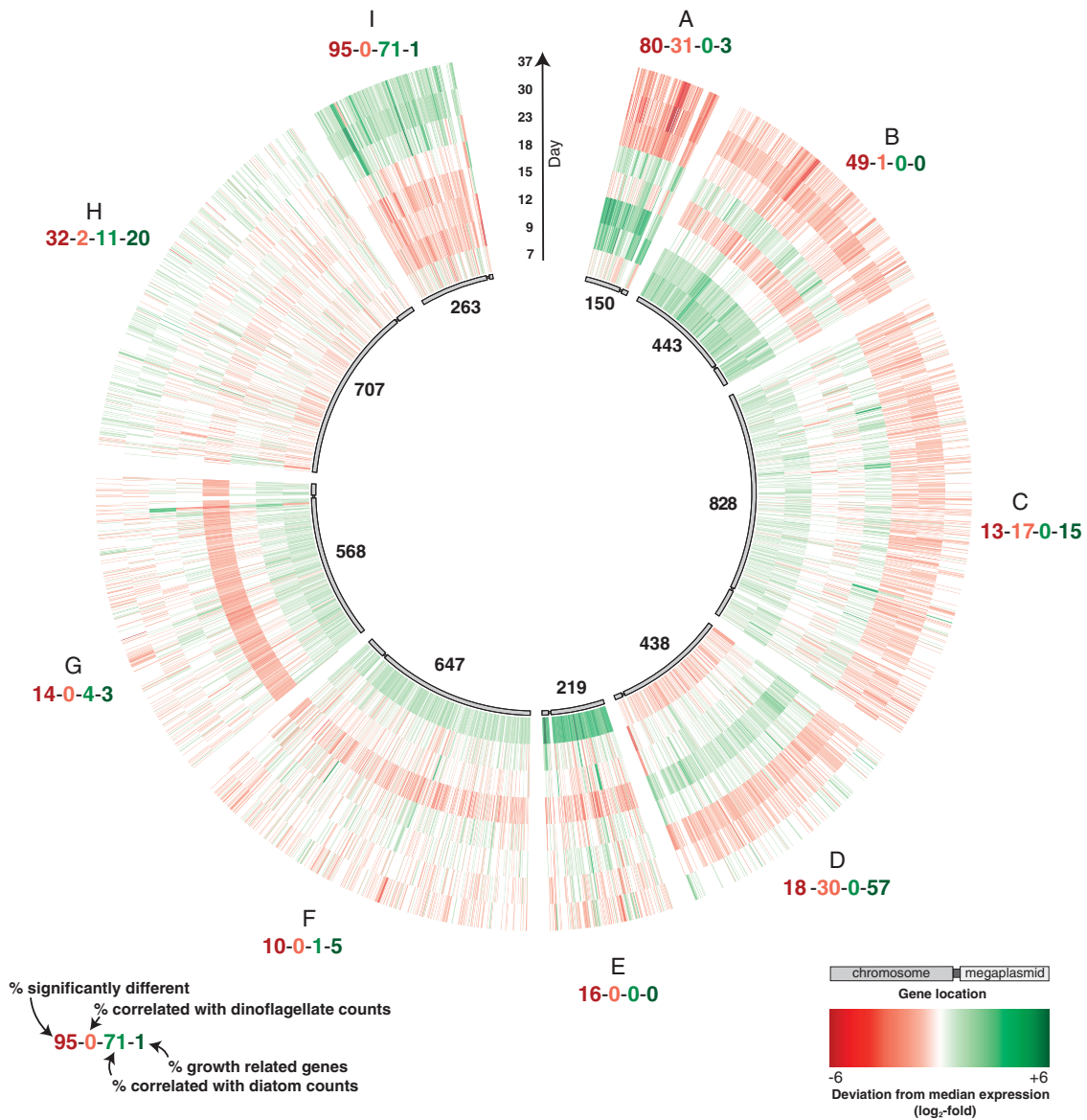


Figure 2 Average normalized expression of *R. pomeroyi* genes from two replicate co-cultures organized into nine expression pattern clusters. The number of genes in each cluster is indicated in the inner circle. Each concentric circle represents one time point.

of organic carbon and sulfur, have been experimentally verified in *R. pomeroyi* (Howard *et al.*, 2006; Reisch *et al.*, 2008, 2011, Todd *et al.*, 2012a, b). The clustering algorithm classified several DMSP-associated genes into dinoflagellate-related Cluster A (Figures 3 and 4), including genes responsible for the committed steps of the DMSP demethylation pathway (*dmdA*) and cleavage pathway (*dddW*). In fact, *dddW* was one of the most differentially expressed genes in the day 9 versus day 30 comparison (72 fold-difference, Table 1). The high expression levels of DMSP degradation genes during *R. pomeroyi* growth with *A. tamarensis* but not *T. pseudonana* accords with findings that dinoflagellates typically produce high levels of DMSP, while diatoms do not (Keller, 1989; Keller *et al.*, 1999; Sunda and Hardison, 2008).

The genetic basis of transport and catabolism of taurine and *N*-acetyltaurine as well as their inducible expression have been verified experimentally in *R. pomeroyi* (González *et al.*, 2003; Gorzyska *et al.*, 2006; Durham *et al.*, 2015). Gene expression patterns grouped most metabolic genes from these pathways into dinoflagellate-related Clusters A and B, and expression differences were as high as 28-fold between day 9 and day 30 (Table 1, Supplementary Table S4). These included *xsc*, *tpa*, and *pta*, encoding a shared lower pathway for metabolism of both compounds, and *naaC'*, encoding a transporter component for *N*-acetyltaurine (Figures 3 and 4). The ability to use taurine appears to be widespread among marine bacteria, while *N*-acetyltaurine use seems restricted to only a few members of the Roseobacter (Figure 5; Durham *et al.*, 2015) and Rhodospirillales clades based on

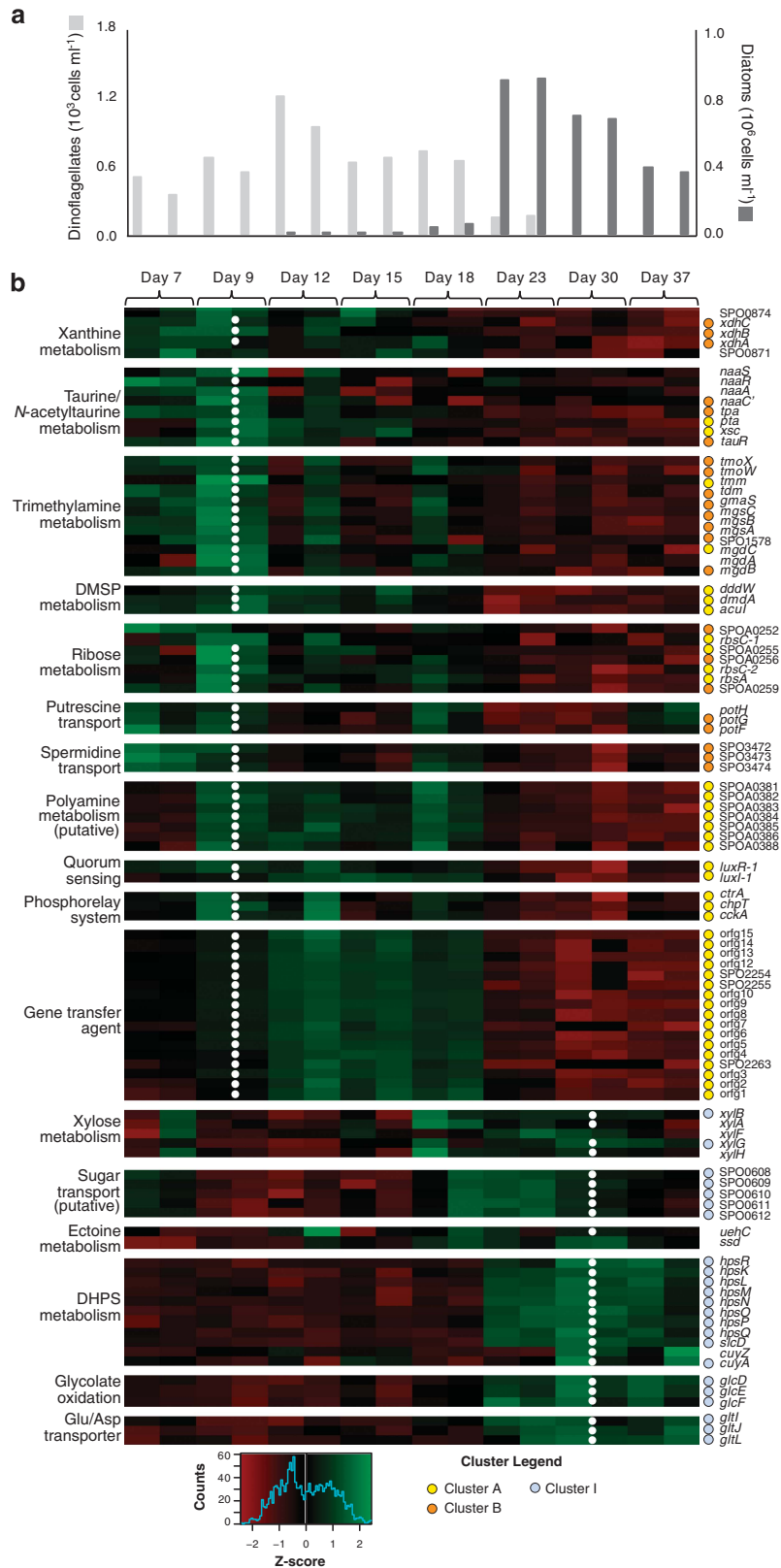


Figure 3 (a) Cell numbers of dinoflagellates (light grey) and diatoms (dark grey). (b) Expression patterns of genes involved in key pathways. Cluster assignments are indicated with colored dots. Normalized expression was scaled by row, and both replicates are shown for each time point. White dots indicate significantly higher expression on day 9 or day 30.

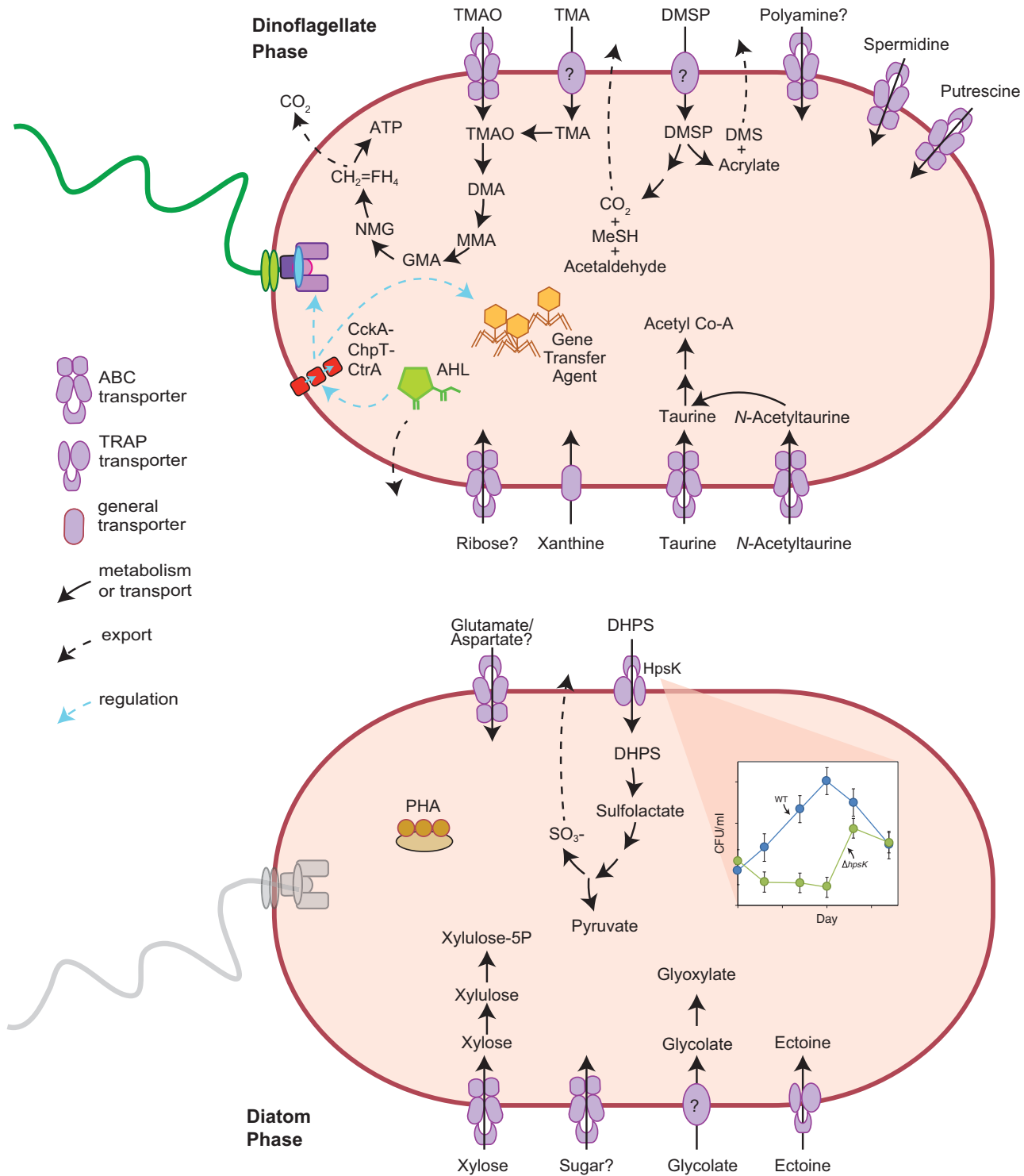


Figure 4 Selected cell functions differentially regulated by *R. pomeroyi* during sequential co-culture with dinoflagellates (top) and diatoms (bottom). CH₂=FH₄, 5,10-methylene tetrahydrofolate; AHL, acylhomoserine lactone; DHPS, dihydroxypropanesulfonate; DMA, dimethylamine; DMS, dimethylsulfide; DMSP, dimethylsulfoniopropionate; GMA, gamma-glutamylmethanimide; MeSH, methanethiol; MMA, monomethylamine; NMG, N-methylglutamate; PHA, polyhydroxyalkanoates; TMA, trimethylamine; TMAO, trimethylamine N-oxide.

the diagnostic gene *naaS* (the amidohydrolase converting N-acetyltaurine to taurine). Transporters for taurine (*tauABC*) and N-acetyltaurine (*naaABB'C*) had more consistent levels of expression throughout the co-culture than catabolic

genes. While both phytoplankton species may release these metabolites (Durham *et al.*, 2015), gene expression patterns predict they are relatively more important in the *A. tamarensis* exometabolome.

Table 1 *Ruegeria pomeroyi* genes with the highest fold-differences in expression at dinoflagellate-dominated Day 9 (positive fold-change values) and at diatom-dominated Day 30 (negative fold-change values)

Gene Name	Locus Tag	Annotation	Category	SOM Cluster	Fold Change
<i>tmm</i>	SPO2508	hypothetical protein		A	151.6
	SPO1551	TMM monooxygenase	Methylamines	A	124.6
	SPO2259	head-tail adaptor, putative	GTA	A	87.8
<i>dddW</i>	SPO0021	Hpt domain protein		A	73.2
	SPO0202	hypothetical protein		A	72.7
	SPO0453	DMSP lyase	DMSP	A	71.8
	SPO2260	hypothetical protein	GTA	A	59.6
	SPO1554	ammonium transporter family		A	59.6
	SPO2261	major capsid protein, HK97 family	GTA	A	59.5
	SPO0931	response regulator		A	54.9
<i>fliN</i>	SPOA0268	IclR family regulator		A	51.9
	SPO0180	flagellar hook-basal body protein	Motility	A	50.9
	SPO0197	flagellar motor switch protein	Motility	A	49.3
	SPOA0269	hypothetical protein		A	48.3
	SPOA0270	hypothetical protein		A	47.2
<i>flgC</i>	SPOA0271	methylamine utilization MauG		A	45.3
	SPO2263	hypothetical protein	GTA	A	44.5
	SPO0181	flagellar basal-body rod protein	Motility	A	39.3
	SPOA0381	polyamine transporter, periplasmic	Polyamines	A	39.1
<i>fliF</i>	SPOA0382	polyamine transporter, ATP	Polyamines	A	38.3
	SPOA0383	polyamine transporter, permease	Polyamines	A	37.1
	SPO0199	flagellar M-ring protein	Motility	A	36.6
	SPO2256	hypothetical protein	GTA	A	34.7
	SPOA0384	polyamine transporter, permease	Polyamines	A	33.2
	SPO2264	portal protein, HK97 family	GTA	A	31.4
<i>flgB</i>	SPO2258	hypothetical protein	GTA	A	30.6
	SPO0930	hypothetical protein		A	30.2
	SPO0182	flagellar basal-body rod protein;	Motility	A	30.1
<i>xsc</i>	SPO2313	hypothetical protein		A	29.4
<i>lpxC</i>	SPO3561	sulfoacetaldehyde acetyltransferase	Taurine	A	28.4
	SPO1205	UDP-3-0-acyl NAG deacetylase		I	-27.5
<i>slcD</i>	SPO1114	TRAP transporter, DctM		I	-27.9
	SPO2269	hypothetical protein		I	-28.2
	SPOA0369	copper resistance protein B		I	-28.6
	SPO1112	TRAP transporter, periplasmic		I	-29.3
	SPO0598	Sulfolactate dehydrogenase	DHPS	I	-29.9
	SPO2180	hypothetical protein		I	-33.7
	SPO1372	hypothetical protein		I	-33.7
	SPO3301	LysR family regulator		I	-34.1
	SPO2657	D-cysteine desulfhydrase		I	-34.6
	SPO0597	UspA stress protein	DHPS	I	-34.7
<i>hpsQ</i>	SPO0584	aspartate aminotransferase		I	-38.9
<i>aspC-1</i>	SPO0595	R or S-DHPS-2-dehydrogenase	DHPS	I	-40.3
<i>hpsO</i>	SPO2914	merR family regulator		I	-41.8
	SPO0636	EF hand domain protein		I	-42.5
	SPO0592	DHPS TRAP transporter	DHPS	I	-45.9
<i>hpsL</i>	SPO0793	Cu(I)-responsive regulator		I	-46.7
<i>cueR</i>	SPO3479	glycolate oxidase	Glycolate	I	-47.8
<i>glcE</i>	SPOA0359	cytochrome c family protein		I	-52.1
	SPO0794	copper-translocating ATPase		I	-55.6
	SPO0594	DHPS-3-dehydrogenase	DHPS	I	-57.7
	SPOA0363	copper-transporting P-type ATPase ActP		I	-60.6
	SPOA0368	hypothetical protein		D	-66.9
<i>phaP</i>	SPO1293	phasin, PhaP	C Storage	I	-73
<i>hpsP</i>	SPO0596	S or R-DHPS-2-dehydrogenase	DHPS	I	-75.1
<i>copA</i>	SPOA0370	copper resistance protein A		I	-79.6
	SPOA0371	hypothetical protein		I	-115.3
<i>hpsK</i>	SPO0591	DHPS TRAP transporter	DHPS	I	-117.2
	SPO2237	hypothetical protein		I	-347.7
	SPO2236	hypothetical protein		I	-1063.3

Day 9 versus Day 30 comparisons were significantly different (DESeq2, $P < 0.001$) for all genes.

A transport and catabolic pathway for the purine xanthine has been proposed for *R. pomeroyi* that entails conversion to glyoxylate and ammonium (Cunliffe, 2016). Genes encoding the putative

xanthine dehydrogenase (*xdhA,B,C*) were assigned to dinoflagellate-related Cluster B (Figure 3, Supplementary Table S4). Putative xanthine permease (SPO0874) and the catabolic gene

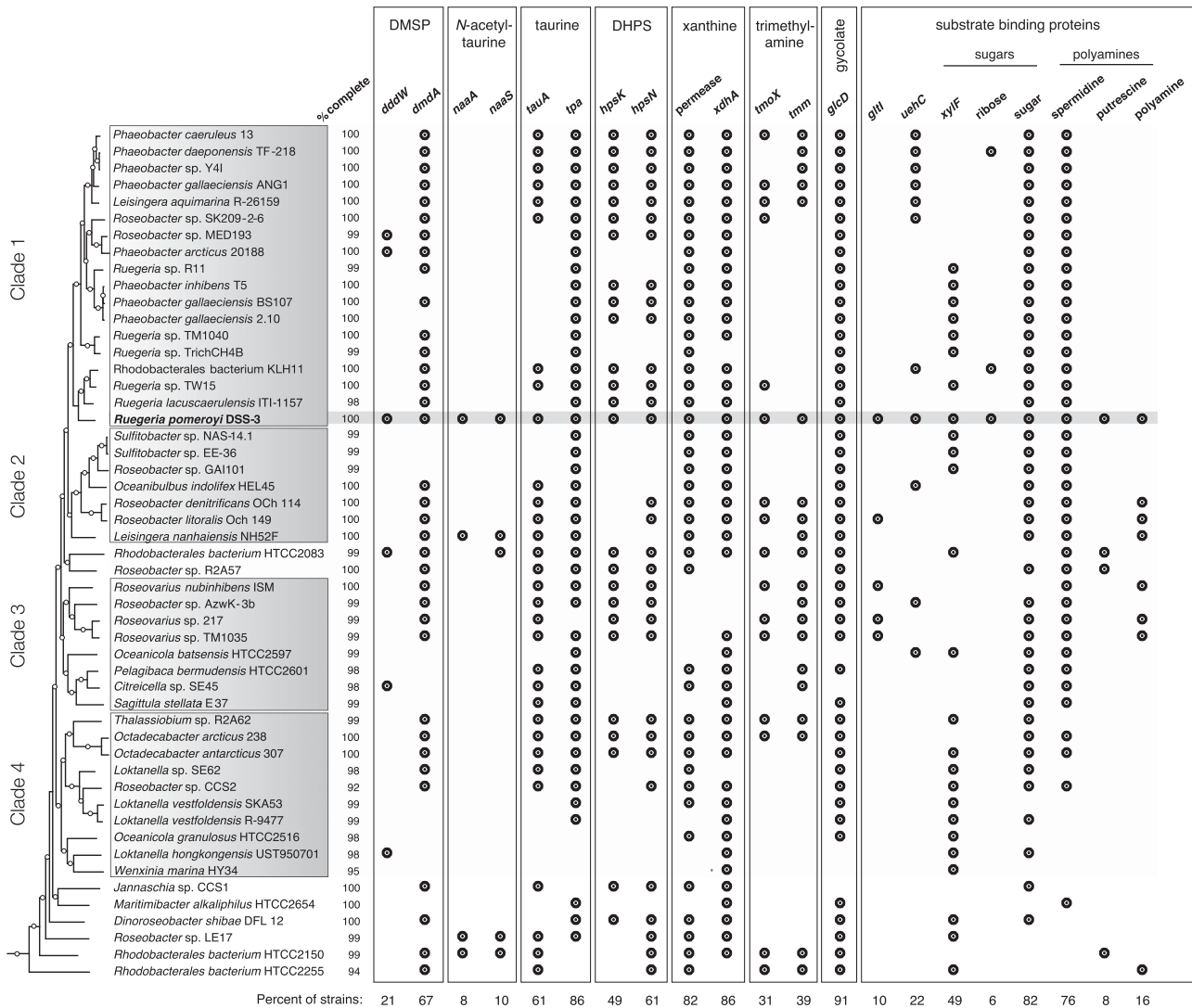


Figure 5 Genome content of *R. pomeroyi* and 50 other Roseobacter clade members. For metabolic genes, analysis focused on enzymes responsible for the first degradation step. For transporters, analysis focused on the periplasmic substrate-binding protein. Genomes were analyzed at roseobase.org. The phylogenetic tree is based on Luo and Moran (2014), with white circles marking nodes with bootstrap values $\geq 50\%$. Substrate binding protein genes: *ghlI*, putative glutamate/aspartate; *uehC*, ectoine; *xylF*, putative xylose.

5-hydroxyisourate hydrolase had expression patterns similar to that of xanthine dehydrogenase (Figure 3). Transcripts for xanthine permeases have been observed previously in marine bacterioplankton communities (Poretsky *et al.*, 2010).

While *R. pomeroyi* cannot grow on trimethylamine (TMA) as a sole carbon source, the substrate induces activities of key enzymes needed for TMA catabolism (TMA monooxygenase, dimethylamine monooxygenase, N-methylglutamate dehydrogenase; Chen *et al.*, 2011) and provides ammonium and energy for the bacterium, with positive effects on biomass production (Lidbury *et al.*, 2015). In the co-culture, multiple genes involved in TMA utilization grouped into dinoflagellate Cluster A (*tmm*, *mgdC*) or Cluster B (*tmoX*, *W*, *tdm*, *gmaS*, *mgsA*, *B*, *C*, *mgdB* and SPO1578) (Figure 3, Supplementary Table S4). The gene encoding the first step in trimethylamine catabolism (*tmm*, mediating oxidation of

trimethylamine to trimethylamine *N*-oxide; Figure 4) was one of the most differentially expressed in the day 9 versus day 30 comparison (125-fold higher on day 9; Table 1).

The genome of *R. pomeroyi* encodes six operons annotated as transporters of polyamines (Moran *et al.*, 2007), organic nitrogen compounds found in phytoplankton metabolomes and marine DOC during phytoplankton blooms (Lee and Jørgensen, 1995; Nishibori *et al.*, 2001). Three of these had expression patterns linked to dinoflagellate abundance. Transporter system SPOA0381-0384 (Cluster A) was the most highly upregulated of the polyamine transporters, with expression differences of up to 39-fold between day 9 and 30 (Figure 3, Table 1). The other two (SPO3467-3469 and SPO3472-3474; Cluster B) both had ~4-fold higher expression on day 9 (Supplementary Table S4). A previous study of *R. pomeroyi* gene transcription patterns during

growth on five polyamines predicted that SPO3467-3469 transports spermidine and SPO3472-3474 transports putrescine, but there was no expression response for SPOA0381-30384 to the polyamines tested (Mou *et al.*, 2010). *A. tamarense* has been shown to release polyamines (Nishibori and Nishio, 1997), and along with *N*-acetyltaurine and xanthine, these represent sources of nitrogen to *R. pomeroyi* in the co-culture. A peak in expression of the putrescine transporter on the last day of the co-cultures indicates decaying diatoms might also release this compound, nuancing the otherwise strong dinoflagellate polyamine signal (Figure 3).

For a number of genes upregulated on day 9 compared to day 30, functional annotations were assigned during genome annotation (Moran *et al.*, 2004, 2007; Rivers *et al.*, 2014) but have not yet been tested experimentally. Six genes encoding a putative ribose transporter and two associated transcriptional regulators were classified into Clusters A and B (Figure 3, Supplementary Table S5). Nineteen other transporter systems were grouped into Cluster B, including four with annotations suggesting opine transport, ten with annotations for amino acids or (di)peptides, and one for an unidentified sulfonate compound (Supplementary Table S5).

Metabolite utilization during growth with diatoms

Expression patterns indicated a different suite of metabolites supporting *R. pomeroyi* during the diatom-dominated phase of the co-culture. The genetic basis of dihydroxypropanesulfonate (DHPS) uptake (*hpsKLM*), catabolism (*hpsOPN*, *slcD*), and pathway regulation (*hpsR*) has been established for *R. pomeroyi* (Denger *et al.*, 2009; Mayer *et al.*, 2010). These genes grouped into diatom-related Cluster I (Figure 3, Table 1). Downstream gene converting cysteate to pyruvate (*cuyA*) also grouped into diatom-related Cluster I, and the gene exporting the resulting sulfite (*cuyZ*) (Denger *et al.*, 2006) exhibited similar expression patterns to *cuyA* (Figure 3). On the basis of the strong positive correlation with diatom abundance ($0.72 \leq \text{Pearson } r \leq 0.90$, $n = 8$), the extent of upregulation (50- to 117-fold), and a previous finding that *T. pseudonana* releases DHPS (Durham *et al.*, 2015), this sulfonate appears to one of the most important bacterial substrates provided by the diatoms (Table 1).

To test this hypothesis, we constructed an *R. pomeroyi* mutant in which the DHPS transporter (*hpsK,L,M*) was deleted. The mutant was unable to grow at the expense of DHPS, but grew at comparable rates to the wild type on other substrates (Supplementary Figure S5). Following inoculation into a co-culture with *T. pseudonana*, cell numbers were compared for mutant and wild-type *R. pomeroyi* over a period of 18 days. Lower cell numbers were observed for the $\Delta hpsKLM$ strain compared to the wild type by as much as six-fold (a difference of 1.2×10^6 cells ml^{-1} after 10 days) suggesting that

DHPS supports the majority of bacterial biomass during co-growth (Figure 4, Supplementary Figure S5). Mutant cells recovered to similar levels as the culture aged, possibly due to a shift in the composition of organic matter released from the diatoms.

The three genes encoding subunits of glycolate oxidase (*glcDEF*) grouped in Cluster I, with 16- to 48-fold higher expression at day 30 compared to day 9 (Table 1, Supplementary Table S4) and strong positive correlations with diatom abundance (Pearson $r \geq 0.7$). Glycolate is a product of phytoplankton photorespiration, with excretion rate dependent on nutrient and light conditions and varying by species (Hellebust, 1965). The *T. pseudonana* strain used here was previously shown to release glycolate, particularly when nitrate was the dominant form of available nitrogen (Parker and Armbrust, 2005), as was the case in the co-cultures (Supplementary Table S3). Once released, the metabolite is thought to be used by marine bacteria primarily for energy production (Wright and Shah, 1977; Lau and Armbrust, 2006).

A component of a putative xylose transporter system was assigned to expression Cluster I (Figure 3, Supplementary Table S4), but the function of this transporter had not yet been verified. We therefore grew *R. pomeroyi* on xylose and three other substrates (glucose, acetate, pyruvate) as sole carbon and energy sources and compared transcription levels under each condition. As was found in a proteomic analysis of an orthologous transporter in *Roseobacter* member *Phaeobacter inhibens* (Wiegmann *et al.*, 2014), both xylose and glucose induced expression of the transporter but xylose had a higher signal (five-fold higher in the case of *R. pomeroyi*; Supplementary Figure S6). The placement of *xylB* (xylulokinase) in Cluster I, along with similar expression patterns for xylose catabolism genes such as *xylA* (xylose isomerase, Figure 3) provided additional transcriptional evidence that xylose is a component of the diatom's exometabolome.

Ectoine is a nitrogen-rich organic osmolyte whose transport and catabolic genes have been experimentally verified in *R. pomeroyi* (Schulz *et al.*, 2017). The substrate binding protein for the ectoine TRAP transporter had 5-fold higher expression on day 30 compared to day 9 and, along with ectoine catabolic genes, its expression pattern was consistent with a diatom-derived source of carbon and nitrogen (Figure 3).

Genes with unverified substrate annotations that were upregulated during the diatom phase involved those encoding a putative amino acid transporter potentially involved in the uptake of glutamate or aspartate (putative *gltIJL*, Cluster I, Figure 3, Supplementary Table S5), and two ABC transporters annotated as sugar uptake systems (Supplementary Table S5). The remaining diatom-related transporters with unverified function included two annotated

for amino acid uptake (polar and branched chain) and two for carboxylic acid uptake. Of the 94 *R. pomeroyi* transporter systems predicted to function in organic compound uptake, 26 were assigned to dinoflagellate-related clusters and 10 to diatom-related clusters (Supplementary Table S5).

Evidence for signaling-driven processes

The expression patterns of genes predicted to be involved in cell–cell communication and signaling suggests that *R. pomeroyi* may be regulating transcription in response to *A. tamarensis*. The bacterial genome encodes two experimentally verified quorum-sensing systems consisting of *luxI/luxR* pairs that are controlled by acylhomoserine lactone (AHL)-type signal molecules (Moran *et al.*, 2004). The *luxR-1/luxI-1* system was recruited into Cluster A with 26-fold (*luxR-1*) and 14-fold (*luxI-1*) higher expression on day 9 compared to day 30; *luxI-1* expression was also positively correlated with dinoflagellate abundance (Pearson $r=0.75$, Supplementary Table S4). Quorum sensing was initially described as a regulation system driven by bacterial density, but there is growing evidence for triggers other than cell numbers, some of which include eukaryote metabolites (Miller and Bassler, 2001; Williams, 2007; Natrah *et al.*, 2011). In our co-culture experiment, quorum sensing expression was decoupled from bacterial abundance. AHL production in *R. pomeroyi* has been shown to be induced by DMSP (Johnson *et al.*, 2016), and it is hypothesized that DMSP signals the presence of phytoplankton and stimulates rearrangement of the bacterium's metabolism (Bürgmann *et al.*, 2007; Johnson *et al.*, 2016).

In *Rhodobacter capsulatus* and Roseobacter strains *Ruegeria* TM1040 and KLH11, quorum sensing was shown to induce a phosphorelay system mediated by the *cckA-chpT-ctrA* genes, which in turn induces expression of motility (Miller and Belas, 2006; Leung *et al.*, 2012; Mercer *et al.*, 2012; Zan *et al.*, 2013). In *Ruegeria* sp. TM1040 this regulation cascade is involved in the bacterium's symbiont-like relationship with dinoflagellate *Pfiesteria piscicida* (Miller and Belas, 2006), while in *Ruegeria* sp. KLH11 it is implicated in a symbiosis with a marine sponge (Zan *et al.*, 2013). In *R. capsulatus*, the phosphorelay system is responsible for the expression of the transduction-like gene transfer agent (GTA), a 15-gene cassette that transfers random 4–5 kb genome fragments between cells (Lang and Beatty, 2002). In *R. pomeroyi*, the function of GTA in gene transfer has also been demonstrated experimentally (Biers *et al.*, 2008). In the co-culture, quorum sensing genes, the *cckA-chpT-ctrA* phosphorelay genes, flagellar assembly genes, and GTA genes were all assigned to dinoflagellate-related Cluster A and had higher expression at day 9 versus day 30 by 2- to 88-fold (Figure 3, Table 1, Supplementary Table S4). The classification of these

genes into the same expression cluster suggests that a similar regulation cascade could exist in *R. pomeroyi* (Figure 4), and interactions between *R. pomeroyi* and *A. tamarensis* might be influenced at the level of the *luxR-1/luxI-1* quorum sensing system. None of the genes linked in this regulatory cascade had expression patterns that tracked with *T. pseudonana* abundance.

Conclusions

The abundance and taxonomy of phytoplankton, the chief producers of labile substrates for marine bacteria, vary on scales of days to weeks in the surface ocean (Teeling *et al.*, 2012; Needham and Fuhrman, 2016) and provide heterotrophic bacteria with a dynamic repertoire of organic substrates. Roseobacter member *R. pomeroyi* changed expression of 27% of its genes when co-grown with a dinoflagellate versus diatom, with many expression shifts linked to non-overlapping suites of metabolites fulfilling bacterial C and N requirements. Further development of chemical methods for identifying low-concentration plankton-derived metabolites in seawater (Moran *et al.*, 2016), in concert with the mining of biological signals through genomics, transcriptomics, and proteomics, will continue to improve our understanding of the flux and fate of marine carbon.

Other members of the Roseobacter clade may have similar metabolic links to phytoplankton exudates. For example, genes for use of putative dinoflagellate metabolites DMSP, taurine, xanthine, and spermidine were found in over half of 50 Roseobacter isolates surveyed, as were the genes for use of putative diatom metabolites DHPS, xylose, and glycolate (Figure 5). These metabolites may support a non-trivial component of Roseobacter heterotrophy in the surface ocean. Other genetic capabilities were infrequent within the clade, including *N*-acetyltaurine and trimethylamine utilization, and these may be examples of currencies of more specific relationships between individual roseobacters and phytoplankton species. Further, metabolites not used as substrates by *R. pomeroyi* would have escaped detection in our experimental system. Thus the surprising diversity and divergence of labile metabolites produced from two phytoplankton strains, each representing just one member of extensive marine groups, and detected by just one bacterial strain highlights the need for better understanding of the compounds that sustain bacterial heterotrophy in the ocean.

The capacity for extensive bacterial transcriptome remodeling observed in response to a shifting mixture of metabolites may be a requirement for successful interactions with diverse phytoplankton. The evidence for a possible recognition response by *R. pomeroyi* during co-growth with the dinoflagellate further suggests that we understand only the tip of an intricate iceberg of autotroph-heterotroph

interactions that influence carbon flux through surface ocean plankton.

Conflict of Interest

The authors declare no conflict of interest.

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