

ORIGINAL ARTICLE

Genomic island genes in a coastal marine *Synechococcus* strain confer enhanced tolerance to copper and oxidative stress

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Highly variable regions called genomic islands are found in the genomes of marine picocyanobacteria, and have been predicted to be involved in niche adaptation and the ecological success of these microbes. These picocyanobacteria are typically highly sensitive to copper stress and thus, increased copper tolerance could confer a selective advantage under some conditions seen in the marine environment. Through targeted gene inactivation of genomic island genes that were known to be upregulated in response to copper stress in *Synechococcus* sp. strain CC9311, we found two genes (*sync_1495* and *sync_1217*) conferred tolerance to both methyl viologen and copper stress in culture. The prevalence of one gene, *sync_1495*, was then investigated in natural samples, and had a predictable temporal variability in abundance at a coastal monitoring site with higher abundance in winter months. Together, this shows that genomic island genes can confer an adaptive advantage to specific stresses in marine *Synechococcus*, and may help structure their population diversity.

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Introduction

Marine picocyanobacteria including *Synechococcus* and its sister taxon *Prochlorococcus*, are a highly diverse group that are found throughout the world oceans, and contribute significantly to global primary production (Li, 1994; Richardson and Jackson 2007). Multiple *Synechococcus* clades or 'species' have been identified using various phylogenetic marker genes, and many of these clades are found to coexist in a single location (Ferris and Palenik 1998; Rocap *et al.*, 2002; Zwirgmaier *et al.*, 2007). The physiological and ecological traits associated with this diversity are not completely understood. With the advent of genome sequencing and the ability to compare whole genomes of these related species, it was discovered that there are a large number of horizontally transferred genes only found in individual strains, and that these genes are concentrated in highly variable regions of the genome called 'genomic islands' (Palenik *et al.*, 2003; Coleman *et al.*, 2006; Palenik *et al.*, 2006). These regions appear to evolve quite rapidly via horizontal gene

transfer as evidenced in part by a *Synechococcus*-enriched metagenome from coastal California waters that showed high recruitment to sequenced reference genomes, except in these island regions (Palenik *et al.*, 2009).

Genomic islands have been found in a wide range of bacterial phyla, and in many cases carry genes known to be involved in niche differentiation, such as nitrogen fixation genes in the symbiotic *Rhizobiaceae* species (Sullivan and Ronson 1998). These acquisitions are considered by some to drive the evolution of bacterial species (Gogarten *et al.*, 2002) and it was predicted that these islands might be a driving force behind niche differentiation in picocyanobacteria (Coleman *et al.*, 2006). However, while some predictions can be made based on known gene function, these islands are characterized by their high number of novel genes of unknown function making it difficult to fully assess their ecological role (Hsiao *et al.*, 2005).

Several studies of phosphate acquisition genes in *Prochlorococcus* have found evidence that certain phosphate nutrition-associated genes are only seen in ocean regions with low phosphate levels and can be found in many phylogenetically distinct clades, indicating that these genes are likely to be laterally transferred at a high rate dependent on environmental conditions (Martiny *et al.*, 2006, 2009a). Given that many *Prochlorococcus* strains have

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small, streamlined genomes, acquisition and retention of new genetic material is most likely under negative selective pressure (Garcia-Fernandez *et al.*, 2004), and island genes likely provide a positive selection advantage. Moreover, metatranscriptomic analyses have found that some of these variable genes in *Prochlorococcus* are very highly expressed and thus, may be important to cellular processes in specific strains, despite their low abundance in the metagenomic data sets (Frias-Lopez *et al.*, 2008). Additionally, a recent study by Avrani *et al.* (2011) found that one island in *Prochlorococcus* is involved in phage-resistance. These studies have been key to our understanding of the role of these islands in marine picocyanobacterial ecology, but targeted gene inactivation of genomic island genes with a predicted function to prove they confer an adaptive response has not been done to our knowledge.

Marine picocyanobacteria are known to be the most sensitive of the major marine phytoplankton groups to copper stress (Brand *et al.*, 1986) and given our knowledge of the temporal and spatial variability of copper in the marine environment (Coale and Bruland 1990; Moffett *et al.*, 1997) it seems likely that tolerance to elevated copper levels would confer an ecological advantage. Indeed, copper has been suggested to affect marine picocyanobacterial distribution (Mann *et al.*, 2002), and marine *Synechococcus* strains have been shown to produce strong extracellular binding ligands in response to excess copper (Moffett and Brand, 1996). Copper toxicity is thought to result from a number of different mechanisms including the production of hydroxyl radicals, the blocking of essential metal binding sites and inhibition of electron transport in photosystem II (Baron *et al.*, 1995; Sunda and Huntsman, 1998; Pinto *et al.*, 2003). Much of the characterized copper stress response can consequently overlap with the oxidative stress response. In addition, copper induces an osmotic shock-like response in *Synechococcus* (Stuart *et al.*, 2009).

In a previous study done on the transcriptional response to copper stress, we found that the coastal *Synechococcus* sp. strain CC9311 (CC9311) had a response distinct from its open ocean cousin (*Synechococcus* sp. strain WH8102), in that a large percentage of the genes it upregulated came from putative horizontally transferred genes (Stuart *et al.*, 2009). In particular, two of the most highly upregulated putative operons were from two genomic islands in this strain. The most highly expressed putative operon contains four genes that are predicted to be a thioredoxin-like protein (sync_1496), a cytochrome c biogenesis protein-like (sync_1495) an extracytoplasmic function sigma-70 factor (sync_1494) and a potential anti-sigma factor (sync_1493). It is on an 83 kb island, the largest island predicted in the genome, with 78 genes, many of which are metal-related (Dufresne *et al.*, 2008).

Bacterial copper resistance has been best characterized in *Escherichia coli* (for review see (Rensing and Grass 2003)) and the gene sync_1495 has predicted transmembrane segments and a conserved domain, DsbD, that has been characterized in *E. coli*. In *E. coli*, DsbD is a inner membrane-bound protein that transfers electrons to DsbC, a disulfide isomerase that repairs nonnative disulfide bonds in the periplasm (Rietsch *et al.*, 1997) and DsbE, a thioredoxin protein essential to cytochrome c maturation (Stirnemann *et al.*, 2005). The *DsbD* knockout is extremely copper sensitive, which is not surprising, since copper as a redox metal catalyzes disulfide bonds quite readily (Katzen and Beckwith, 2003), and the entire pathway is thought to be involved in copper resistance and potentially other oxidative stressors (Hiniker *et al.*, 2005). Indeed, in *Synechocystis* PCC6803 the orthologous operon (sll0685-sll0688) is transcriptionally upregulated in response to hydrogen peroxide stress, osmotic stress and DCMU ((3-(3,4-dichlorophenyl)-1,1-dimethylurea)) (Singh *et al.*, 2010). The best available evidence thus, predicts that the sync_1496–93 putative operon in CC9311 might use a similar mechanism of disulfide repair to confer increased copper and oxidative stress tolerance.

The second most upregulated putative operon from the CC9311 copper stress studies includes an ABC transporter (sync_1214–1215), a HlyD secretion protein (sync_1216) and a 6.5 kb RTX toxin domain protein (sync_1217). This operon is on a much smaller 14 kb island with only eight genes, of which the four described above were induced in response to copper (Dufresne *et al.*, 2008; Stuart *et al.*, 2009). RTX toxin-domain proteins have a range of functions from enzymatic toxins to protective S-layer proteins, but most are secreted (Linhartova *et al.*, 2010). An S-layer-localized RTX toxin domain protein in *Synechocystis* PCC6803 (sll1951) was found to be important in resistance to heavy metals, antibiotics and osmotic stress suggesting a role as general barrier against these stressors (Sakiyama *et al.*, 2011). Although not an ortholog, we predicted that the sync_1214–1217 operon may secrete the RTX toxin-domain protein sync_1217, given the ABC transporter that is part of the putative operon, to function as an outer membrane barrier.

Using bioinformatic searches, we found that close homologs of these two copper-regulated operons were not abundant in marine metagenomic data sets, which was expected given the low recruitment to these genomic island regions in metagenomic data sets shown in Palenik *et al.* (2009). By using targeted gene inactivations of predicted copper tolerance genes, as well as investigation of the temporal and spatial distribution of one of these genes in the marine environment with quantitative PCR (qPCR), we have sought to better understand the functional and ecological role of genomic island genes in *Synechococcus*, using copper and oxidative stress tolerance in the coastal strain CC9311 as a model.

Materials and methods

Cell culture and growth assays

1L batch cultures of CC9311, sync1495- and sync1217- were grown in chelexed synthetic ocean water on stir plates as previously described in Stuart *et al.* (2009). For mutant cultures, spectinomycin and streptomycin were both added at $2\ \mu\text{g ml}^{-1}$ concentration at the time of inoculation, to maintain the insertion. For growth assays, cultures were grown to early log phase, and 50 ml were aliquoted in triplicate into acid-washed glass flasks and grown with shaking at $23\ ^\circ\text{C}$. Cultures were given 24 h in these conditions to adjust to the transfer then CuEDTA was added in triplicate at $10\ \mu\text{M}$, $4\ \mu\text{M}$, $1\ \mu\text{M}$, $0.4\ \mu\text{M}$ for the copper stress, while sterile milli-Q water (Millipore, Billerica, MA, USA) was added to the control. Using the MINEQL algorithm (Westall *et al.*, 1976), we calculated that this corresponds to equilibrium free copper levels equivalent to pCu 10.1, pCu 10.5, pCu 11.1 and pCu 11.5. For methyl viologen stress, methyl viologen (Sigma, St. Louis, MO, USA) was added at 500 nM, 100 nM, 50 nM and 5 nM. For AlexaFluor-488 C₅ maleimide (Life Technologies, Carlsbad, CA, USA) treatment it was added at $1\ \mu\text{M}$. Flow cytometry samples were collected daily for up to 7 days and were fixed and analyzed as described in Stuart *et al.*, 2009.

RNA extraction and quantitative reverse transcription-PCR

Cells were harvested and RNA extracted as described in Stuart *et al.*, 2009. Briefly, cells were re-suspended in Trizol Reagent (Life Technologies, Carlsbad, CA, USA), heated, extracted according to the manufacturers specifications, and then purified and concentrated on a glass fiber column. The following modifications were made to the original procedure: the $65\ ^\circ\text{C}$ incubation was for 30 min instead of 60 min, a Zymo PCR inhibitor removal kit (Zymo, Irvine, CA, USA) was used to purify RNA instead of the Qiagen RNeasy kit (Qiagen, Venlo, Netherlands). Semiquantitative PCR was performed as described in (Stuart *et al.*, 2009). Briefly, complimentary DNA was generated using a commercially available kit and a set amount of total starting RNA. Control reactions with no reverse transcriptase were also included. QPCR reactions included both a negative control and the complimentary DNA control as well as a 5 point standard curve. The following changes to Stuart, 2009 protocol were made: complimentary DNA was generated using 250 ng of total RNA using Ambion's Retroscript kit (Life Technologies, Carlsbad, CA, USA) according to the manufacturer's specifications. QPCR was performed on a Qiagen RotorGene-Q with Promega GoTaq qPCR mastermix according to the manufacturer's specifications. *Rpoc1* was used as a house-keeping gene, because its expression was high and

did not change significantly in any of the expression microarrays datasets available for strain CC9311 or strain WH8102 (Stuart *et al.*, 2009; Tai *et al.*, 2009; Tetu *et al.*, 2009).

Insertional inactivation of sync_1495 and sync_1217

Gene inactivations were conducted as described in Brahamsha, 1996 (Brahamsha 1996a, b). *Synechococcus* sp. strain sync1495- and sync1217-, in which the genomic gene sync_1495 or sync_1217 is interrupted by a duplication, and insertion of a non-replicating plasmid was constructed as follows. First, the omega cassette conferring spectinomycin/streptomycin resistance was cut from the pDW9 vector (Golden and Wiest 1988) using *Bam*HI and cloned into the *Bam*HI site of pBR322 (New England Biolabs, Ipswich, MA, USA) resulting in plasmid pBR322_Ω. A 399 bp segment of sync_1495 that is completely internal to the gene was amplified from CC9311 genomic DNA using primers 1495mutfwd and 1495mutrvs (Supplementary Table S1). A 500 bp segment of sync_1217 that is completely internal to the gene was also amplified from CC9311 genomic DNA using primers 1217mutfwd and 1217mutrvs. The amplified fragments were ligated separately into pCR2.1-TOPO (Life Technologies, Carlsbad, CA, USA) according to the manufacturer's specifications resulting in plasmids TOPO_1495 and TOPO_1217, respectively, and insertion of the correct fragment was verified by sequencing. The 407 bp *Eco*RI fragment from TOPO_1495 was ligated into the *Eco*RI site of pBR322_Ω resulting in plasmid pBR322_Ω_1495. The 507 bp *Eco*RI fragment from TOPO_1217 was ligated into the *Eco*RI site of pBR322_Ω resulting in plasmid pBR322_Ω_1217. Both plasmids were then separately introduced via conjugation into *Synechococcus* sp. strain CC9311 as described in Brahamsha, 1996 (Brahamsha, 1996b), with the following modifications: SN pour plates contained 0.4% (wt/vol) SeaPlaque agar (Lonza, Koeln, Germany) and $2\ \mu\text{g ml}^{-1}$ each of spectinomycin and streptomycin. Exconjugants were screened by PCR to verify that no wild-type copy of the gene remained and that insertion had occurred at the predicted region (Supplementary Figure S1 and Supplementary Table S1).

CAMERA search, environmental clone library construction and qPCR

Bioinformatic searches for sync_1495 homologs were performed on the CAMERA website (Sun *et al.*, 2011) by first performing a tBLASTn (with an e-value cutoff of 10^{-6}) of both the 'All metagenomic open reading frames' and the 'All metagenomic 454 Reads' databases and combined the hits. A reciprocal BLASTx using this combined list of hits was then performed to the non redundant database to assess the similarity of the hits to

sync_1495. Sequences that did not hit sync_1495 as a top hit were excluded.

To assess the abundance of this gene, primers to sync_1495 were designed with several considerations in mind. First, primers should amplify only close homologs of sync_1495, including all Clade I *Synechococcus* members with the gene or potentially very recent horizontally transferred versions, and excluding other *DsbD* genes. At the nucleotide level this gene is almost unique in the non redundant database, with no close BLASTn hits, which made it simple to exclude known *DsbD* genes. At the time of primer design, we did not have other Clade I versions of the gene to align to so we chose a region within the predicted DsbD domain, which extends from 198 to 811bp, and would be less likely to have divergence within the clade. We also wanted to amplify as large a region as possible given qPCR limitations (which optimally amplifies 50–150 bp) in order to cover a sufficient portion of the gene for TOPO-cloning verification. In consequence, the 254 bp amplified region chosen covers from 549 – 802 bp of the gene and the primers had no close BLASTn hits in the database (Supplementary Table S1). DNA from Tai *et al.*, 2009 (Tai and Palenik 2009) was used first to generate a clone library from one sample (14 May 2006) using TOPO TA Cloning Kit (Life Technologies, Carlsbad, CA, USA) according to the manufacturer's specifications. Five clones were picked and sequenced using the forward primer included in the kit. QPCR was performed on a Stratagene Mx300P using Promega GoTaq qPCR master mix (Promega, Madison, WI, USA) according to the manufacturer's specifications. QPCR samples included DNA from monthly samples from 2005 (no February or August samples due to sample loss) and 2007 from Tai *et al.*, 2009 (Tai and Palenik 2009) and DNA from depth samples collected in July, 2007 as described in Tai *et al.*, 2011. QPCR primers used on above samples were for sync_1495 and *rpoc* Clade I (Supplementary Table S1).

Results

Expression of sync_1495 and sync_1217 in wild type in response to stress

As the response to copper stress often shares similarities to the oxidative stress response, we wanted to determine whether copper-responsive genes could be induced by other sources of oxidative stress. We also wanted to investigate the timing of upregulation of copper responsive genes to confirm their importance in copper stress response. The first two genes in putative operon sync_1496 – 1493 (Figure 1a) had very high induction in response to copper (6.3 and 6.1 log₂ fold changes, respectively) more than double the log₂ fold changes of other induced genes (Stuart *et al.*, 2009), so we chose one, sync_1495, to further investigate the

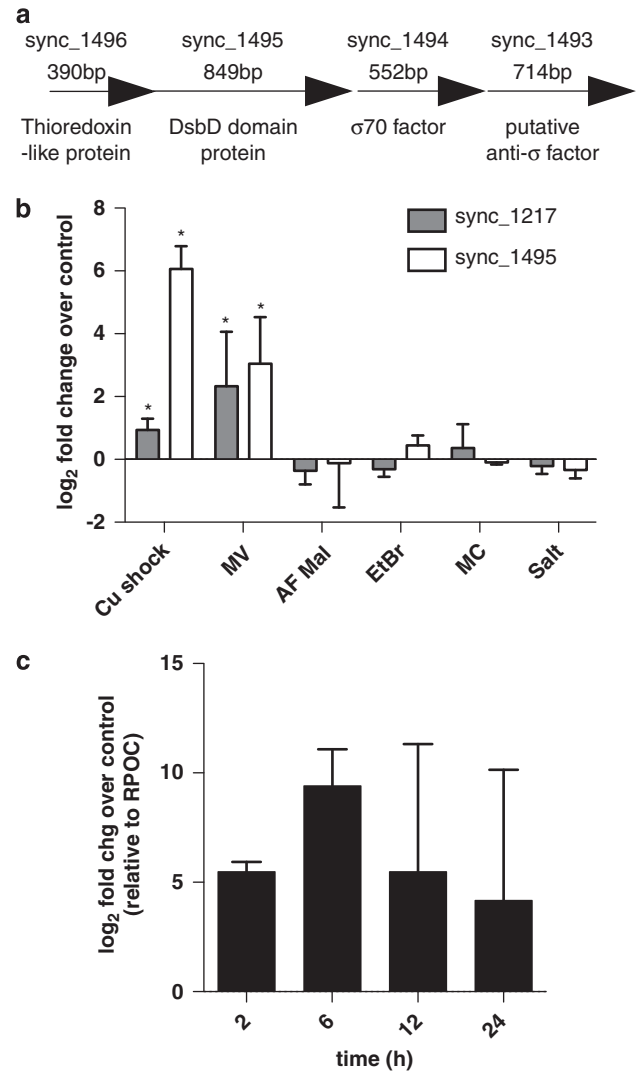


Figure 1 Sync_1495 gene expression in response to stressors and over time. (a) is a schematic of the operon sync_1496 – 1493. (b) is gene expression in response to various stressors. Log₂ fold change is calculated relative to a control treatment. Error bars represent one s.d. between three biological replicates. ‘Cu’ is quantitative reverse transcription (RT)-PCR after 2 h of copper addition (pCu10.1, 10 μM CuEDTA), ‘MV’ is qRT-PCR of methyl viologen treatment 2 h after addition (100 nM). ‘AF Mal’ is qRT-PCR AlexaFluor-488 C₅ maleimide 2 h after addition (1 μM). ‘EtBR’, ‘MC’ and ‘salt’ are microarray expression data (S Tetu, D Johnson, K Phillippy, R Stuart, C Dupont, K Hassan, B Palenik and I Paulsen, unpublished data). (c) is gene expression of sync_1495 over 24 h after copper stress (pCu10.1, 10 μM CuEDTA).

specificity of this response. Sync_1495 was chosen because it is the longer of the two genes, which makes it a more feasible target for eventual genetic inactivation, which is discussed below. In quantitative reverse transcription-PCR (QRT-PCR) experiments sync_1495 was also significantly induced by methyl viologen (Figure 1b), which is a mild oxidant that subsequently reduces oxygen and generates superoxide anions (Imlay, 2008). The gene was not induced by AlexaFluor-488 C₅ maleimide, which binds to surface free thiols, and in microarray experiments (S Tetu, D Johnson, K Phillippy, R

Stuart, C Dupont, K Hassan, B Palenik and I Paulsen, unpublished data), the gene was also not significantly induced by ethidium bromide, mitomycin C, or salt shock (Figure 1b). Induction in response to copper stress was significant at 2 h and 6 h after treatment (one sample *t*-test of \log_2 fold changes, *P*-values of 0.0022 and 0.0103, respectively), but began to decrease by 12 h with more variation between the three biological replicates (Figure 1c).

In addition, *sync_1217* was induced by methyl viologen with a 2.3 \log_2 fold change (± 1.7), but not by ethidium bromide, mitomycin C, or salt shock (S Tetu, D Johnson, K Phillippy, R Stuart, C Dupont, K Hassan, B Palenik and I Paulsen, unpublished data). These results provide evidence that these acquired genes are responsive to methyl viologen stress and copper stress, but not other tested stressors. Methyl viologen and copper are similar as known inducers of oxidative stress-like responses.

Mutant sensitivity to methyl viologen and copper stress
Insertional inactivations of *sync_1495* (designated strain 'sync1495-') and *sync_1217* (designated strain 'sync1217-') were validated using PCR (Supplementary Figure S1). Growth assays with methyl viologen additions showed that both *sync1495-* and *sync1217-* exhibited significant sensitivity to methyl viologen stress relative to the wild type. By 4 days after methyl viologen treatment at 50 nM, the wild type showed no difference in growth rate from the control treatment, whereas the mutants exhibited significantly slower growth rates (Figure 2, Supplementary Figure S2, Table 1).

The response to copper stress in both mutants was slightly more complicated, as copper is both a micronutrient and a toxicant so growth stimulation is possible with copper additions. This can be seen

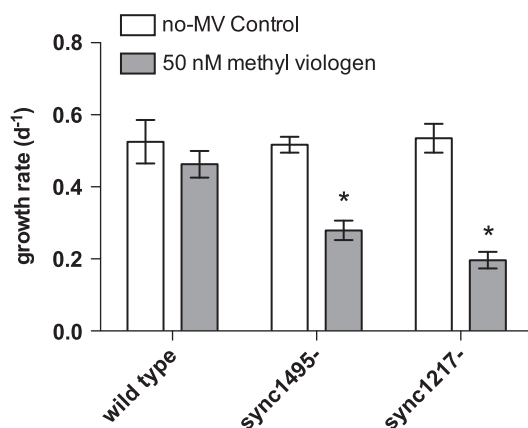


Figure 2 *Synechococcus* CC9311 wild type and mutant methyl viologen growth assays. Growth rates in both wild-type CC9311 ('wt'), *sync1495-* and *sync1217-*. Cells per ml from flow cytometry counts were ln-transformed and a linear regression was performed on each. Bars represent linear regression slope, error bars represent one s.d. between three biological replicates. Star indicates significant difference from no-MV control treatment (one sample *t*-test, *P*<0.05).

in the wild-type growth assays, where in the first several days the lowest copper addition led to higher cell numbers than the control treatment, although the final maximum yields between these two treatments was not significantly different (Figure 3).

This partially positive effect of providing some copper may have complicated the growth rate data, and we were not able to demonstrate a statistically significant difference in mutant growth rates relative to wild-type controls (Figure 3, Table 2). However, both mutants were sensitive to copper stress relative to the wild type as evidenced by the differences in maximum cell yield (Figure 3, Table 2). In the wild type the growth rate and the maximum yield for pCu11.5 (Cu^{2+} ion concentration of $10^{-11.5}$ M; 0.25 μM CuEDTA, lowest copper addition) was not significantly different from the no-Cu control. By contrast, both mutants showed significantly lower maximum yields in the copper treated cultures (Figure 3, Table 2). Additionally, for the next highest copper level, pCu11.1, the wild type showed no difference in either growth rates or maximum yield from the control, whereas *sync1495-* and *sync1217-* had significantly lower maximum cell yields (Table 2).

Under no-copper conditions *sync1217-* had much higher cell yields than the wild type, however, the mean cell size of *sync1217-* over the course of the experiment was significantly smaller than the wild type, indicating that although cell number may be higher, total biomass may be similar between the two (Supplementary Figure S3). *Sync1495-* also had higher maximum cell yields than wild type, although not as high as *sync1217-*, and given that it did not have a smaller average cell size, it may be that *sync1495-* may actually have a higher biomass yield than wild type in the no-copper conditions, indicating that there may be a tradeoff in fitness associated with this operon. Conversely, it is also possible that the spectinomycin/streptomycin resistance cassette provides a slight growth advantage to the mutants.

Downstream gene expression in *sync1495-*

As insertional inactivation can sometimes lead to inactivation of downstream genes, we looked at the expression of downstream genes in the *sync_1495* putative operon (Figure 1a). QRT-PCR of the genes surrounding *sync_1495* in the wild type, revealed that the upstream gene and two downstream genes were highly upregulated in response to 2 h of copper stress (Supplementary Figure S4a). In the mutant, transcripts of these two downstream genes were detectable, but there was no induction of these genes under copper stress relative to the control (Supplementary Figure S4b).

Temporal and spatial distribution of *sync_1495* gene in the marine environment

A BLAST-based search was performed to find homologs of *sync_1495* in marine metagenomic data

Table 1 Growth rates and maximum yield after methyl viologen treatments

Treatment ^a	Wild type ^b	<i>Sync1495</i> - ^b	<i>Sync1217</i> - ^b
0 (no-MV control)	0.5251 ± 0.060 (3.023 ± 0.23)	0.5170 ± 0.022 (8.366 ± 0.518)	0.5353 ± 0.040 (12.193 ± 1.726)
50 nM	0.4627 ± 0.037 (1.303 ± 0.131)	0.2791 ± 0.027 (1.972 ± 0.468)	0.1964 ± 0.023 (1.369 ± 0.154)
5 nM	0.5558 ± 0.0439 (2.806 ± 0.696)	0.4199 ± 0.026 (6.065 ± 0.753)	0.4218 ± 0.039 (5.682 ± 1.037)

^aConcentration methyl viologen (MV) added.

^bGrowth rate (d⁻¹) ± 1 s.d. and in parentheses maximum yield cells per ml (*10⁸) ± 1 s.d.; bold indicates significant difference from no-MV control treatment (*P* < 0.05, one sample *t*-test).

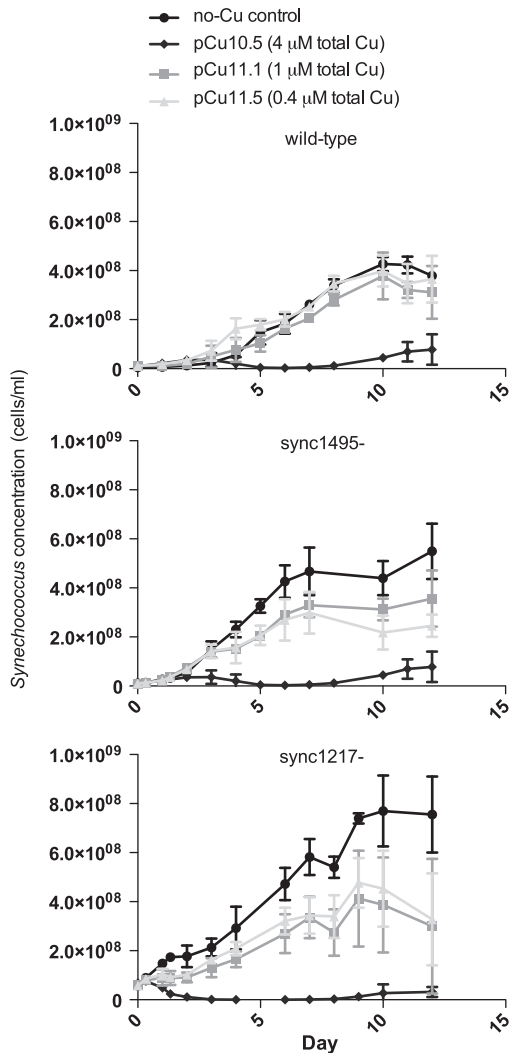


Figure 3 *Synechococcus* CC9311 wild type and mutant copper growth assays. Wild-type CC9311 ('wt'), *sync1495*- and *sync1217*- cells per ml from flow cytometry counts. Error bars represent one s.d. between three biological replicates. Cells/ml do not take into account mean cell size, which was significantly smaller in *sync1217*- (see Supplementary Figure S3).

sets using the CAMERA website (Sun *et al.*, 2011), and in accordance with our predictions, only five hits were found to closely match *sync1495* (Supplementary Table S2).

Primers were designed to amplify 254 bp of the 849 bp gene, *sync1495*, that match the genome

sequence exactly without introduced degeneracy (Supplementary Table S1). The primers were tested for specificity to *sync1495* in one DNA sample extracted from water collected at the Scripps Institution of Oceanography pier by generating a clone library from, which five clones were sequenced. All five clones aligned very closely to *sync1495* with 98.0–100% identity at the nucleotide level (Supplementary Figure S5). There were single nucleotide polymorphisms detected at residues 26, 68, 70, 85, 169, 178, 209 and 226, which resulted in amino acid substitutions in single nucleotide polymorphisms 26, 68 and 209 (Supplementary Figure S5). Using quantitative reverse transcription-PCR, the above primers along with Clade I-specific *rpoc1* primers from Tai *et al.*, 2009 (Tai and Palenik, 2009) were then used on monthly DNA samples from 2005 and 2007 from surface waters off the Scripps Institution of Oceanography pier. *Sync1495* was more abundant in the winter months and decreased to almost undetectable levels in the summer, relative to total Clade I *Synechococcus* abundance (Figure 4). This trend is not correlated to either total *Synechococcus* numbers or Clade I *Synechococcus* abundance (Figure 4, Supplementary Figure S6). We found that abundance of *sync1494*, the gene directly downstream of *sync1495*, was also higher in the winter months (Supplementary Figure S7), indicating that this trend holds for other genes in this putative operon. *Sync1494* was also detected using perfect primers (Supplementary Table S1) and compared with Clade I-specific *rpoc1* abundance.

Additionally, from two sites off the California coast, DNA from several depths was obtained in July of 2007. Site 93.40 (32.51304°N, 118.21386°W), which was closer to the coast, showed an increase in *sync1495* abundance with depth. Site 93.65 (31.6776°N, 119.9025°W), which was further from shore, did not show an increase of the gene with depth in the top 100 m, while copper measurements were not available for this station (Figure 5). Along this transect, total dissolved copper ranged between 0.9 and 2.0 nM, and did not seem to increase consistently with distance from shore and only slightly increased with depth to 250 m (K. Buck and K. Barbeau unpublished data). Free copper and total copper generally have been shown to increase with depth (Coale and Bruland 1990; Moffett and

Table 2 Growth rates and maximum yield after copper treatments

Treatment ^a	Wild type ^b	<i>Sync1495</i> - ^b	<i>Sync1217</i> - ^b
0 (no-Cu control)	0.6656 ± 0.044 (4.269 ± 0.288)	0.7023 ± 0.039 (5.781 ± 0.765)	0.3012 ± 0.028 (8.447 ± 0.870)
pCu10.1	NG (0.195 ± 0.008)	NG (0.206 ± 0.020)	NG (0.642 ± 0.057)
pCu10.5	NG (0.447 ± 0.006)	NG (0.406 ± 0.107)	NG (0.762 ± 0.0186)
pCu11.1	0.4816 ± 0.079 (3.783 ± 0.951)	0.6253 ± 0.057 (3.565 ± 1.145)	0.2305 ± 0.026 (4.129 ± 1.954)
pCu11.5	0.6108 ± 0.051 (3.983 ± 0.633)	0.6278 ± 0.069 (2.985 ± 0.851)	0.2633 ± 0.017 (4.767 ± 1.010)

^aConcentration copper added.

^bGrowth rate (d⁻¹) ± 1 s.d. and in parentheses maximum yield cells per ml (*10⁸) ± 1 s.d.; bold indicates significant difference from no-Cu control treatment (*P* < 0.05, one sample *t*-test); 'NG' indicates no significant increase in cells per ml over experiment.

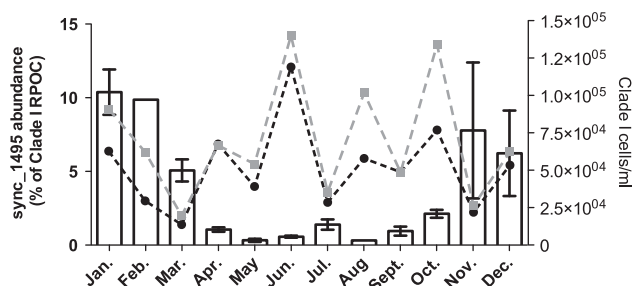


Figure 4 Temporal distribution of *sync_1495* gene over 2 years off the Scripps Institution of Oceanography pier. Bars show qPCR of *sync_1495* as a percentage of total Clade I *Synechococcus* abundance (as determined by Clade I *rpc1* abundance) in surface samples from 2005 and 2007. Error bars represent +/- one s.e.m. between 2 replicate years (2005 and 2007). For Feb. and Aug. months no data was available for 2005 and only 2007 value is shown. Lines show *Synechococcus* abundance data from Tai *et al.*, 2009; black line shows Clade I total abundance from qPCR and gray line shows total *Synechococcus* numbers from flow cytometry.

Dupont 2007), however, marine *Synechococcus* is known to produce strong copper binding ligands in response to excess copper (Moffett and Brand 1996) and could influence free copper levels. At 60 m at site 93.40 the sharp decrease in total *Synechococcus* numbers (Tai *et al.*, 2011) coincides well with a clear increase in Cu²⁺ concentration as found using CLE-ACSV (Figure 5, Supplementary Figure S8) (Buck and Bruland 2005; Buck *et al.*, 2010)), and the largest positive difference between total dissolved copper and free Cu²⁺ occurs at the depth (30 m) with maximal *Synechococcus* numbers. Accordingly, we would predict that site 93.65, which has two to four times higher abundance of *Synechococcus*, might have lower free copper levels due to high levels of organic ligand.

Discussion

Highly variable genomic regions in picocyanobacteria have been implicated in picocyanobacterial adaptation to various ecological niches by conferring advantage, be it nutrient acquisition or some other stressor (Coleman *et al.*, 2006; Martiny *et al.*, 2009a). Identifying and characterizing an advantage can best be done in culture by knocking out these genes, and once this has been done it permits further

analysis of environmental data to better understand the ecological relevance of these genes. The present data show that both *sync_1495* and *sync_1217* confer methyl viologen and copper tolerance to CC9311, suggesting that both genes have a role in an oxidative stress-like response, as both copper and methyl viologen can act as electron oxidants. Both are upregulated in the wild type in response to methyl viologen and copper, but not other stressors, again helping to confirm this role. The increase in the expression of *sync_1495* is sustained over at least 6 h after copper stress in wild type, indicating it may be necessary for more than just early shock response.

As might be expected for genes that are not part of the *Synechococcus* core genome, the stress tolerance is manifested as a decrease in growth rate in response to methyl viologen in the mutants at levels that elicited no effect in the wild type. In response to copper the effect is slightly more complex. However, although the growth rate differences are not as large as for the methyl viologen treatments, comparison of the maximum cell yields clearly indicate that both mutants are impaired by copper levels that have no effect on wild-type cell yields.

Interestingly, the closest homologs to both these operons are found in estuarine cyanobacteria (*Synechococcus* WH5701 gene locus WH5701_04100 and *Cyanobium* PCC7001 gene locus CPCC7001_837), suggesting the possibility of gene flow between coastal marine and estuarine populations of cyanobacteria. On the basis of GC content, the genes in these two estuarine cyanobacteria do not themselves appear to have been laterally acquired. Indeed, the levels of copper, and other heavy metals, can be much higher in estuarine environments due to both natural and anthropogenic effects (for review of San Francisco Bay see (Buck *et al.*, 2007)), and that these groups would have extra copper resistance mechanisms is logical.

Although the exact mechanisms of action of each of these genes has yet to be confirmed, we can make informed predictions based on both their phenotypes and what we know of homologous genes. The cellular localization of *sync_1217* in strain CC9311 is not known, but we have some initial indications that a large protein of this size (the largest predicted protein in the genome) can be found in the outer

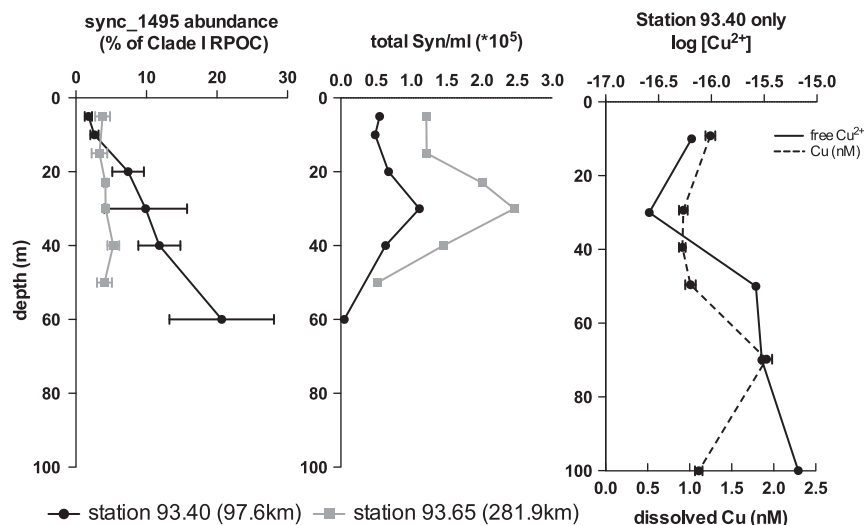


Figure 5 Spatial distribution of *sync_1495* gene from two stations in the Southern California Bight. Numbers in parentheses indicate distance from shore. Left vertical profile shows qPCR of *sync_1495* as a percentage of total Clade I *Synechococcus* abundance (as determined by Clade I *rpoC1* abundance). Error bars represent one s.d. between three technical replicates. Middle profile shows *Synechococcus* counts from flow cytometry taken from Tai *et al.*, 2011. Right profile shows both total dissolved copper concentration (dashed line) and free copper ($-\log[\text{Cu}^{2+}]$) concentrations (solid line). $-\log[\text{Cu}^{2+}]$ measurements from copper speciation using CLE-ACSV at 25 μM SA (Buck *et al.*, 2010).

membrane (R. Stuart, B. Brahmsha, unpublished data). A recent review of RTX domain genes surveyed all available genomes, and cyanobacteria tend to have a relatively high number of these genes most of which are of unknown function (Linhartova *et al.*, 2010). Three that have been characterized in cyanobacteria are involved in motility including SwmA from *Synechococcus* WH8102 (swimming), oscillin from *Phormidium uncinatum* (gliding) and a hemolysin-like protein in *Synechocystis* PCC6803 that is involved in motility through an unknown mechanism, but CC9311 is nonmotile. In *E.coli*, a Type I secretion system secretes HlyA, an RTX hemolysin protein and the operon, HlyCABD, includes an ABC transporter (HlyB), an membrane fusion protein (HlyD) that is involved in the folding of HlyA and a toxin activation protein (HlyC). The *sync_1217* putative operon does include both an ABC transporter and a HlyD homolog, which strongly suggests that *sync_1217* is secreted. The S-layer RTX protein that has been best characterized is RsaA in *Caulobacter crescentus* that is thought to help form a selective porosity barrier that may prevent predation (Linhartova *et al.*, 2010), and it could be that *sync_1217* has a similar function, acting as an outer membrane barrier to toxins.

Although *sync_1495* was also sensitive to copper and methyl viologen, it did not have a smaller average cell size than wild type and indeed had higher maximum cell yields than wild type in some cases. This indicates that the activity of the proteins in the operon may have a cost in terms of cell growth under control conditions even, while it provides an advantage under copper and oxidative stress conditions. Copper induction of the downstream genes, the sigma factor and putative anti-sigma factor

(*sync_1494* and *sync_1493*) were also inactivated in this mutant, although the upstream thioredoxin-like protein (*sync_1496*) was still copper inducible. As noted in the introduction *sync_1495* has a domain similar to DsbD in *E.coli*, which is involved in disulfide bond isomerization. Additionally, the DsbD superfamily is present in many bacterial groups and the DsbD homolog in *Caulobacter crescentus*, ScsB, acts on a different set of substrates from *E.coli*, including a novel cell envelope reductive pathway providing electrons to a thioredoxin-like protein, which then reduces a novel peroxiredoxin that acts in cell envelope to directly scavenge peroxides (Cho *et al.*, 2012). This suggests that this superfamily may be involved in several mechanisms of reduction in the cell envelope including disulfide isomerization as in *E.coli*, but also potentially directly scavenging of reactive oxygen species produced by copper or methyl viologen in the periplasm.

On the basis of qPCR, *sync_1495* was relatively more abundant within the clade I species in the winter months in 2005 and 2007 off the Scripps Institution of Oceanography pier. The primers used were not degenerate and matched the CC9311 version of *sync_1495* and although we know little about the rate of transfer of these genomic island genes, unless it is very rapid, it is likely that these primers only amplify versions of this gene that are in strains closely related to CC9311, that is, the clade I species. Our data thus show that the strain(s) that includes *sync_1495* is under a different selective pressure than the clade I species as a whole. The surface waters off the pier in the winter months are characterized by lower water temperature, a higher degree of mixing and more runoff from rains. The

latter two of these could lead to episodic pulses of higher copper in surface waters, and although we have no direct measurements of seasonal copper at this location, it is well-known that runoff does tend to increase copper levels in San Diego Bay, and the rainy winter months have significantly higher copper levels (Blake *et al.*, 2004; Buck *et al.*, 2007). Additionally, it could be that copper is also upwelled seasonally (Takesue and van Geen 2002).

At one site in the Southern California Bight the gene also appeared to increase with depth, but more samples are needed to explain the depth distribution pattern seen as the two sites sampled were quite different. In Tai *et al.* (2011) distribution of several subclades from clades I and IV were quantified at these sites and it was found that at site 93.40 (and other coastal stations) the main clade I groups increased with depth, whereas the dominant clade IV group decreased with depth. In contrast, at site 93.65 most of the groups increased with depth and thus, at depth these sites had quite different relative abundances of the various groups. Moreover, at site 93.40 where sync_1495 was found to increase with depth, Cu²⁺ did increase between 30 m and 70 m. Although copper measurements were not available for site 93.65 it did have much higher abundances of *Synechococcus*, which could mean lower free copper levels.

Although it is possible that copper is driving these trends in sync_1495 abundance, it is also possible that some other variable that increases in the winter months and causes increased envelope or oxidative stress is the driver of this trend. Increased runoff, a deeper mixed layer and decreased temperature can lead to an increase in many environmental stressors (that is, anthropogenic toxins, fecal bacteria (Dwight *et al.*, 2002)) and an increase in dynamic conditions. So while copper might not be the driver of this trend, the more important point is that a genomic island gene that is known to confer stress tolerance in cultures is more abundant in the winter months, when many different environmental stressors are increasing, irrespective of seasonal variability in the main group, in which the gene has been found.

Most intraclade (intraspecies) strain variability in *Synechococcus* appears to lie within genomic islands (Palenik *et al.*, 2009) and accordingly it is likely that the island(s) carried by the CC9311 strain and closely related strains are the drivers of the seasonal variability in relative abundance seen in this one gene, sync_1495, involved in copper/oxidative stress response. In *Prochlorococcus*, macronutrient levels in the ocean are predicted to drive genomic content and ecotype formation, potentially through lateral gene transfer (Coleman *et al.*, 2006; Martiny *et al.*, 2006, 2009a, 2009b). Although copper and oxidative stress response genes may be less recognizable than nutrient acquisition genes, our data predicts that this stress response differentiation can also influence genomic content and may be an additional mechanism

driving population structure in picocyanobacteria. A collection of strains with different genomic islands including ones conferring tolerance to these stresses could contribute to the overall fitness of *Synechococcus* in dynamic coastal conditions.

Conflict of Interest

The authors declare no conflict of interest.

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