

ORIGINAL ARTICLE

Coral-mucus-associated *Vibrio* integrons in the Great Barrier Reef: genomic hotspots for environmental adaptation

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Integron cassette arrays in a dozen cultivars of the most prevalent group of *Vibrio* isolates obtained from mucus expelled by a scleractinian coral (*Pocillopora damicornis*) colony living on the Great Barrier Reef were sequenced and compared. Although all cultivars showed >99% identity across *recA*, *pyrH* and *rpoB* genes, no two had more than 10% of their integron-associated gene cassettes in common, and some individuals shared cassettes exclusively with distantly-related members of the genus. Of cassettes shared within the population, a number appear to have been transferred between *Vibrio* isolates, as assessed by phylogenetic analysis. Prominent among the mucus *Vibrio* cassettes with potentially inferable functions are acetyltransferases, some with close similarity to known antibiotic-resistance determinants. A subset of these potential resistance cassettes were shared exclusively between the mucus *Vibrio* cultivars, *Vibrio* coral pathogens and human pathogens, thus illustrating a direct link between these microbial niches through exchange of integron-associated gene cassettes.

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Introduction

Corals are reef-building animals that depend on phototrophic dinoflagellate endosymbionts in the genus *Symbiodinium* (commonly called zooxanthellae) for much of their fixed carbon and molecular oxygen (Rosenberg *et al.*, 2007). The importance of this symbiosis is perhaps most apparent in coral bleaching, which generally entails loss of zooxanthellae (Glynn, 1993). Severe bleaching can result in significant coral mortality (Hoegh-Guldberg, 1999). Recently, a complex role in coral health has also been claimed for the prokaryotic component of the coral microbiome, itself complex and diverse (Rohwer *et al.*, 2001, 2002; Pantos *et al.*, 2003; Bourne and Munn, 2005; Koren and Rosenberg, 2008). This diversity extends to location-specific differences but healthy coral includes a Vibrionaceae component (Kvennefors *et al.*, 2010). Conversely, nitrogen-fixing bacteria (Lesser *et al.*, 2004; Olson *et al.*, 2009) and chitin degraders (Ducklow

and Mitchel, 1979) thought to supply metabolic intermediates from these pathways to coral, pathogens that can cause bleaching (Kushmaro *et al.*, 1996, 1997; Ben-Haim and Rosenberg, 2002; Ben-Haim *et al.*, 2003) and commensals that might serve as ‘probiotics’ (Reshef *et al.*, 2006) defending the coral from pathogens by production of antimicrobial compounds (Koh, 1997; Castillo *et al.*, 2001; Ritchie, 2006). Overall it is clear that coral/bacterial associations can be either detrimental or beneficial to the former.

Unlike vertebrate animals, invertebrates do not accumulate an adaptive immune response to infection, but rely on a first-line innate identification of pathogens, localized inflammation of the injured area, and the synthesis of specific antimicrobial compounds at the cell surface or within cells to target the microbe (Fearon and Locksley, 1996). In marine invertebrates such endogenous antimicrobial effectors include glycosides, brominated phenols, polyphenolics, polyketides, ribosomal and nonribosomal peptides, alkaloids, fatty acids and terpenoids (Blunt *et al.*, 2004).

The coral probiotic hypothesis holds that the coral prokaryotic microbiome is also adaptive for the coral, providing resistance to pathogen-mediated bleaching. (Reshef *et al.*, 2006). Studies of coral

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disease caused by *Vibrio corallilyticus* in *Pocillopora damicornis* (Geffen and Rosenberg, 2005) show that resistance to the pathogen is mediated by the coral's mucus, a protective layer that coats all types of coral and is densely colonized with bacteria (Brown and Bythell, 2005; Rosenberg *et al.*, 2007), a significant fraction of which produce antimicrobials (Koh, 1997; Castillo *et al.*, 2001; Ritchie, 2006). Moreover, corals once susceptible to bleaching caused by a specific bacterial pathogen can become immune to it, a phenomenon called 'experience-mediated tolerance'. For instance, Rosenberg and colleagues demonstrated that the coral *Oculina patagonica*, once strongly affected by bleaching events caused by *Vibrio shiloi*, can become resistant to reinfection by this bacterium (Reshef *et al.*, 2006). Similarly, immunity to *Aurantimonas coralicida*, the putative bleaching agent responsible for the 1995 white plague outbreak in corals situated in the Florida Keys (Denner *et al.*, 2003) has been observed. The coral probiotic hypothesis suggests that such instances of acquired immunity reflect changing patterns of antimicrobial production by commensal nonpathogenic bacteria (Brown *et al.*, 2000; Rosenberg *et al.*, 2007).

Pathogens may in turn develop antimicrobial resistance, recruiting resistance determinants from a 'resistome' (D'Costa *et al.*, 2006) through lateral gene transfer (LGT). One genetic element that facilitates this evolutionary arms race is the integron. This system comprises an integrase gene (*intI*) and an associated integration site (*attI*), at which the integrase protein (IntI) can catalyze gene cassette insertion or removal (Stokes and Hall, 1989). Gene cassettes are small mobile elements, usually composed of a single gene and a recombination site (*attC*). There are good reasons to suppose that integrons might have a significant role in coral bleaching and resistance. First, integrons are known to be crucial in the spread of antibiotic resistance in many human and animal pathogens (Hall *et al.*, 1999). Second, integrons are especially prominent and well studied chromosomal elements in pathogenic and environmental *Vibrio*: indeed all sequenced *Vibrio* have at least one such integron array, often bearing more than 100 cassettes (reviewed in (Stokes and Hall, 1989; Rowe-Magnus *et al.*, 2003; Wright, 2007)). Third, many species of *Vibrio*, such as *V. shiloi* and *V. corallilyticus*, are known or suspected coral pathogens and a role for *Vibrio* in bleaching can be inferred from an increasing proportion of *Vibrio* 16S rRNA genes sequenced during bleaching events (Bourne *et al.*, 2008). But, fourth, there are also rich commensal vibrio populations associated with healthy corals (Arboleda and Reichardt, 2009; Kvennefors *et al.*, 2010).

Here, we describe the diversity and function of encoded genes within gene cassettes associated with chromosomal integrons of *Vibrio* isolates cultivated from mucus expelled by a healthy *Pocillopora damicornis* colony living in the Great Barrier Reef

off the coast of Australia. Integron sequences were obtained from twelve cultivars, which collectively share a pairwise 99% average nucleotide identity in housekeeping genes. We show that integron arrays in these coral *Vibrio* are extraordinarily dynamic (compared even with arrays in other members of the genus) and that this dynamic nature likely reflects LGT. Antibiotic resistance might be inferred as prominent function of these cassettes. We also show that, although the gene cassette repertoires of (a) coral mucus vibrios, (b) *Vibrio cholerae* isolates, (c) other marine vibrio and (d) non-vibrio from other (especially polluted) environmental sites are clearly distinguishable, cassettes are shared and presumably exchanged between them.

Materials and methods

Sample collection and cultivation

Coral mucus samples were collected from a physiologically stressed scleractinian coral, *Pocillopora damicornis*, sampled from Davies Reef (Latitude 18°50.9'S/Longitude 147°41'E) within the Great Barrier Reef Marine Park off the coast of Queensland, Australia in March, 2006. Mucus was obtained by exposure of the colony to the air inducing excessive mucus production (milking), which was collected in sterile tubes. This coral mucus was plated on both Marine Agar 2216 (Difco Laboratories, Detroit, MI, USA) and thiosulfate citrate bile sucrose media (McLaughlin, 1995). Plates were incubated over night at 37 °C and then colonies randomly picked and re-streaked three times for pure cultures.

Gene amplification, cloning and sequencing

PCRs were carried out in a final volume of 25 µl containing 1–5 ng of template DNA, 1.0 mM of each primer and 12.5 µl of PCR Master Mix (PROMEGA, Alexandria, New South Wales, Australia). The reactions were performed with an initial denaturation step at 94 °C for 2 min, followed by 30 cycles with a denaturation at 94 °C for 30 s, primer annealing at 55 °C for 30 s and primer extension at 72 °C for 1 min. PCR products were gel purified with the MinElute kit (QIAGEN, Doncaster, Victoria, Australia) and cloned in TopoTA (INVITROGEN, Mulgrave, Victoria, Australia). Clones were sequenced from both strands.

Colony screening for IntI by PCR

Pure colonies were picked and used as template DNA in PCR reactions specific for the *intI* gene. This reaction was performed using class 1-specific primers HS463A and HS464 (Gillings *et al.*, 2009). Colonies were also screened for gene cassettes with degenerate primers (HS287 and HS286) targeting *attC* regions (Stokes *et al.*, 2001). PCR products by these reactions were cloned and sequenced as described above.

Fosmid library construction, screening and sequencing of Vibrio cultivars

Fosmid libraries were constructed from genomic DNA using the EPIFOS kit (EPICENTRE, Gymea, New South Wales, Australia). Purified genomic DNA was run on a low-melt 1% agarose gel (AMRESCO, Gymea, New South Wales, Australia) in a pulsed-field gel electrophoresis apparatus (Bio-Rad, Gladesville, New South Wales, Australia) and the DNA of ~40 kb was purified from the agarose, ligated to the fosmid vector, packaged in phage capsids and used to infect *E. coli* as described in the EPIFOS kit manual. A total of 480 colonies from each library were picked from agar plates and used to inoculate 96-well blocks containing 1 ml of LB broth with 12.5 µg ml⁻¹ of chloramphenicol in each well. These cultures were grown overnight and glycerol stocks were made by mixing 140 µl of culture from each well with 60 µl of 50% glycerol in a 96-well plate. PCR screening was done using primers targeting the *intI* gene as described above. Clones positive for the *intI* gene were re-streaked onto LB agar plates containing 12.5 µg ml⁻¹ of chloramphenicol, which were used to inoculate a liquid culture for extraction of pure fosmid DNA. These clones were used for shotgun library construction, sequence and assembly performed by MacroGen Inc. (Seoul, South Korea).

Housekeeping-gene sequence acquisition from Vibrio cultivars and genomes

In the case of *Vibrio* cultivars, housekeeping-gene sequences were amplified using *recA*, *pyrH* and *rpoB* primers (Thompson *et al.*, 2005) and sequenced as described above. The same sequences were retrieved from whole *Vibrio* genome sequences stored in the public database NCBI.

Taxonomic assignment of Vibrio cultivars by recA phylogeny

recA DNA sequences were amplified from 128 *Vibrio* coral mucus cultivars and combined with 210 reference sequences collected from the NCBI database. These were aligned together in MUSCLE (Edgar, 2004) and manually edited. The final alignment of 725 nucleotide positions was used as an input file for the Maximum Likelihood-based (ML) phylogenetic program PHYML (Guindon and Gascuel, 2003). The general time reversible nucleotide substitution model was implemented with the proportion of invariable sites and the gamma parameter of across-site rate variation (using four categories) estimated from the dataset. Bootstrap support values were calculated with the same parameters (100 replicates).

Vibrio concatenated gene phylogeny

The phylogenetic tree of coral mucus *Vibrio* isolates whose integrons were sequenced, in addition to reference strains of known taxonomic affiliation,

was reconstructed using PHYML (Guindon and Gascuel, 2003) with the general time reversible nucleotide substitution model, the proportion of invariable sites and the gamma parameter of across-site rate variation (using four rate categories) estimated from the dataset. Bootstrap support values were calculated with the same parameters (100 replicates). The sequence dataset used for the analysis is a concatenation of the *recA*, *pyrH* and *rpoB* genes amplified from *Vibrio* coral mucus isolates and retrieved from reference strains publicly available at NCBI. These were aligned using MUSCLE (Edgar, 2004) and manually edited to a final length of 2144 nucleotide positions.

Integron annotation

Assembled fosmid arrays encoding integrons were uploaded into the integron database: Annotation of Cassette and Integron Data (ACID) (Joss *et al.*, 2009). ACID contains publicly available integron sequences and users can annotate integron-related components including and associated cassette open reading frames (ORFs).

Cassette *attC* sites were identified using a string-search-scoring method. The total score is composed of several individual categories that reflect *attC* structural components (Bouvier *et al.*, 2009). Specifically, one point per match between inverse core/core sites of the *attC* site was assigned. The total possible score of this property referred to as R, is four as described by Joss *et al.* (2009). Briefly, one point is scored per match between the inner repeats of the *attC* site. These base-pairing events were weighted to make them more meaningful. Specifically, potential secondary structures of this simple site were considered as two 3 bp regions. These properties, referred to as L_A and L_B, when multiplied, produce a maximum score of 9. These are multiplied so that pairing is required in each of these two domains to produce a score. For example, L_A = 3 and L_B = 0 would produce a score of 0 (3 × 0). However, if each L_A and L_B had two base-pairing events, the score would be four. This weighting strategy ensures that the extra-helical base essential for IntI-*attC* interaction is considered. In addition, one point is scored for the presence of the typical extra-helical base, defined as E. Finally, one point is awarded for each mismatch in the 2 bp region located upstream from R', referred to as site S. These mismatches produce the bulge necessary for IntI catalysis of the hairpinned *attC* substrate (Mazel, 2006). Final query scores are divided by the maximum possible score and the cut off was set to 75%, which results in the overall lowest rate of false identifications (Joss *et al.*, 2009).

Cassettes identified by ACID were extracted from this database and uploaded to the Meta Genome Rapid Annotation using Subsystem Technology (MG-RAST (<http://metagenomics.nmpdr.org>)) server for functional assignment. MG-RAST was built as a modified version of the RAST server (Aziz *et al.*,

2008) and is a high-throughput pipeline and provides automated functional assignments of sequences by comparing both protein and nucleotide databases. Finally, given that cassettes are mobile, the Basic Local Alignment Search Tool (BLASTx) algorithm (Altschul *et al.*, 1990) was used to taxonomically define cassettes with the E-value threshold set to 10^{-5} . These results were graphically illustrated using the software package MEGAN (Huson *et al.*, 2007).

Cassette ecology

Cassette sequences were extracted from ACID in both nucleotide and amino acid FASTA formats, as were *attC* nucleotide sequences. These files were formatted for BLAST analysis with the formatdb freeware package (Altschul *et al.*, 1990). Freely available blastall (Altschul *et al.*, 1990) software was employed for BLAST 'all versus all' analysis for each of the three datasets. This BLAST output was used as an input file for the metagenomic distance-based operational taxonomic units and Richness determination toolbox (Schloss and Handelsman, 2008). Using the furthest neighbor algorithm (Legendre and Legendre, 1998), this software package was used to arbitrarily define cassette family and *attC* cluster operational taxonomic units as two or more cassette or *attC* nucleotide sequences producing a pairwise BLAST Score Ratio (BSR) (Rasko *et al.*, 2005) of >0.70 .

When analyzing these estimates with greater scrutiny, however, it was clear that the estimate continued to grow. Specifically, the range of the Chao1 95% confidence intervals was analyzed as a function of sampling effort for these gene cassettes as suggested in (Schloss and Handelsman, 2005). There was a positive correlation with sequencing effort ($R^2 = 0.722$) for these cassettes suggesting that the uncertainty of the Chao1 estimate increases with additional sampling and thus the 95% confidence intervals is artificially low for cassette sequence data and therefore more sampling of this gene pool is required for more accurate richness estimates (data not shown).

Operational taxonomic unit clustering

The relationship of shared cassettes or *attC* sites was visualized with Cytoscape 2.6 network visualization software (Shannon *et al.*, 2003). The cassette/*attC* networks are based on the families/clusters defined above. The network was generated using a spring-embedded algorithm based on a 'force-directed' paradigm (Kamada and Kawai, 1989). Network nodes are treated like physical objects that repel each other, and the connections between nodes attract their end points such that nodes sharing more connections will be placed more closely (relative to others sharing less) in three-dimensional space.

Statistical support for *attC* clustering

The UniFrac implemented G-test (Lozupone *et al.*, 2006) was employed to assess whether or not the

observed counts of *attCs* are distributed evenly across *Vibrio* lineages. The input data was in the form of an ML tree calculated using RAxML (Stamatakis, 2006). The best scoring tree of selected *attC* nucleotide sequences was obtained from 200 tree-building iterations using the GAMMA + P-Invar model of rate heterogeneity with an ML estimate of the alpha parameter.

Cassette phylogeny

Nucleotide alignments for each cassette family were subjected to ML-based phylogeny with PHYML (Guindon and Gascuel, 2003). The general time reversible nucleotide substitution model was implemented with the proportion of invariable sites and the gamma parameter of across-site rate variation (using four categories) estimated from the dataset. Bootstrap-support values were calculated with the same parameters (100 replicates).

Vibrio cassette dendrogram assembly

Selected *Vibrio* spp. were clustered according to the heuristic tree searching algorithm available in PAUP* (Swofford, 2003) with the search criterion set to distance (UPGMA) and the data (presence-absence) originating from a matrix of shared cassette families with >0.70 BSR values. Bootstrap support values were calculated with the same parameters (100 replicates). The cassette families themselves were clustered according to shared *Vibrio* spp. using the same algorithm and therefore the result is a two-dimensional clustering of shared cassette families.

Results

Pocillopora damicornis mucus contains a variety of *Vibrio* species

A total of 128 *Vibrio* cultivars were isolated on rich marine and *Vibrio*-selective growth media from *Pocillopora damicornis* mucus (see Materials and methods) and the complete cassette arrays from 12 of these isolates were sequenced and are shown diagrammatically in Figure 1. The taxonomic distribution of the 128 *Vibrio* isolates, according to their PCR-amplified *recA* nucleotide sequences, was determined by maximum-likelihood phylogeny (Guindon and Gascuel, 2003). Figure 2 shows that a diversity of bacteria from the genus *Vibrio* was recovered, including many whose *recA* nucleotide sequences grouped with reference sequences from the completed genomes of *V. fischeri*, *P. eurosenbergii*, *V. tubiashi*, *V. fortis* and *V. coralliilyticus*. In addition to these, the clade most represented in this dataset of cultivars (referred to as Clade 1) also contained *V. alginolyticus*, *V. harveyi*, *V. campbellii* and *V. rotiferianus recA* reference sequences.

Most of the *Vibrio* recovered in this study were part of a single monophyletic group (Clade 1, Figure 2). Thus, to examine the tempo and mode

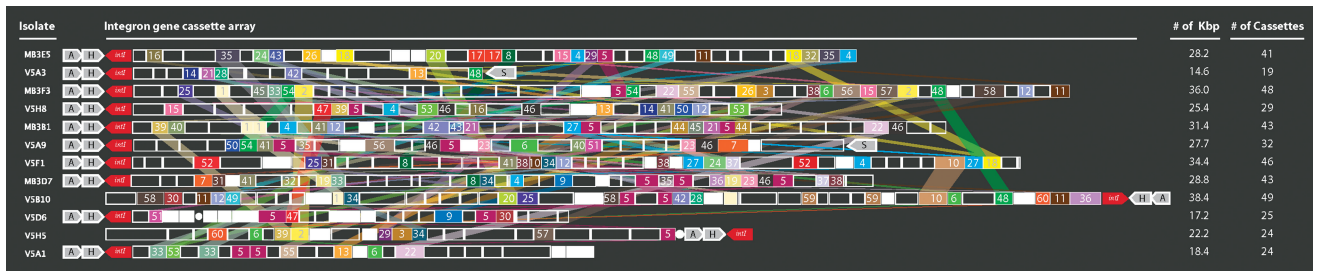


Figure 1 Integron gene-cassette arrays and array dynamics of *Vibrio* isolates cultured from coral mucus. Integron gene-cassette arrays are drawn to scale with each cassette boxed. Dark gray cassettes represent those that have no known homologs. The light gray ORFs that flank some of the integrons correspond to putative acetyltransferase (A), hypothetical unknown (H) and sodium-solute symporter (S) proteins. IntI ORFs are shown in red, whereas the white cassettes are those that have pairwise nucleotide sequence BSR values >0.70 to cassettes encoded by other more evolutionary distant *Vibrio* spp. not accounted for in this figure (none of these are like any other in this figure). White circles represent sequence gaps ~ 10 kbp for V5H5 and an unknown value for V5D6. Cassettes that are shared between the different *Vibrio* isolates cultured from coral mucus are colored accordingly and are assigned a family number that is consistent with how cassette families are named in this study. Colored bars connecting cassettes common to different isolates indicate the dynamic nature of integrons between closely related bacteria.

of cassette sharing within a population of closely related *Vibrio* living in the same environment, the 12 isolates for which integron cassette arrays were sequenced all came from Clade 1.

Mucus Vibrio-cassette arrays are highly dynamic

The 12 isolates selected had a nucleotide identity of at least 99% across 2144 nucleotide positions sequenced for the *recA*, *pyrH* and *rpoB* genes, placing them well within even the most stringent 'species' definitions (Doolittle and Zhaxybayeva, 2009). Despite this, the number of shared cassettes encoded in their integron gene cassette arrays ranges from only ~ 1 –10%. Synteny maps show that even cassettes shared between arrays have shuffled to randomness in position in the time required for 1% genomic sequence divergence to accumulate in housekeeping genes (Figure 1).

In all, 10 of the integrons from these coral mucus-associated *Vibrio* isolates are downstream from ORF H, encoding a conserved hypothetical protein of unknown function (Figure 1). The other two, while also associated with this same gene, had undergone genomic rearrangements. In one instance (V5B10), the *intI* is downstream from ORF H, but the associated array appears inverted relative to the *intI* gene. In the second, V5H5, the array is not only inverted but interrupted by ~ 10 kbp of genomic sequence encoding ORFs without any flanking *attC* sequences. Interestingly, *Vibrio* species, *V. parahaemolyticus* RIMD 2210633 and *V. sp.* Ex25, both had their integrons inserted adjacent to ORF H, whereas the integrons associated with the two *V. vulnificus* strains (YJ016 and CMCP6) and all *V. cholerae* isolates were at a different site (downstream from the LSU ribosomal protein L20p).

Mucus Vibrio-cassette arrays evolve more rapidly than those of V. cholerae

To assess whether the mucus *Vibrio*-cassette arrays are evolving more rapidly than the core genome

in terms of structure and composition, available chromosomal integrons from any member of the *Vibrio* genus were compared in a pairwise manner, correlating similarity in cassette array composition to divergence in sequence of the three housekeeping genes. These analyses revealed only a weak positive correlation overall ($R^2=0.3$) between the percent of gene cassette families shared between any two *Vibrio* isolates and their evolutionary relatedness as defined by percent nucleotide identity of concatenated housekeeping genes *recA*, *pyrH* and *rpoA* (Figure 3a), although *V. cholerae* isolates (upper cluster, Figure 3a) appeared different in this regard. Indeed, the same analysis focused at a finer evolutionary scale revealed a distinction between *V. cholerae* and coral mucus isolates specifically (Figure 3b). Six strains of the former (Thompson *et al.*, 2004), showed a strong correlation of shared gene-cassette content and evolutionary relatedness ($R^2=0.6$) compared with the genus level analysis (Figure 3b). Conversely, analysis of Clade 1 coral mucus *Vibrio* cultivars (Figure 1), together with the six most closely related strains of the *V. harveyi* species group, showed no correlation between shared cassettes and evolutionary relatedness ($R^2=0$; Figure 3b).

Possible LGT among mucus *Vibrio* cassettes can be examined phylogenetically. In total, 404 cassettes were sequenced from these twelve *Vibrio* isolates, with sixty instances of related cassettes occurring more than once (that is, having BSR values >0.70) corresponding to 70% of amino acid sequence identity corrected for length). Phylogenetic analysis (Guindon and Gascuel, 2003) was performed on the basis of nucleotide sequences of all eleven individual cassette families that had four or more members (Supplementary Figure S1). Five of these trees presented conflicting topologies. Although it is possible that these trees reflect gene duplication and differential loss of integron gene cassettes, LGT between members of this *Vibrio* population seemed the more

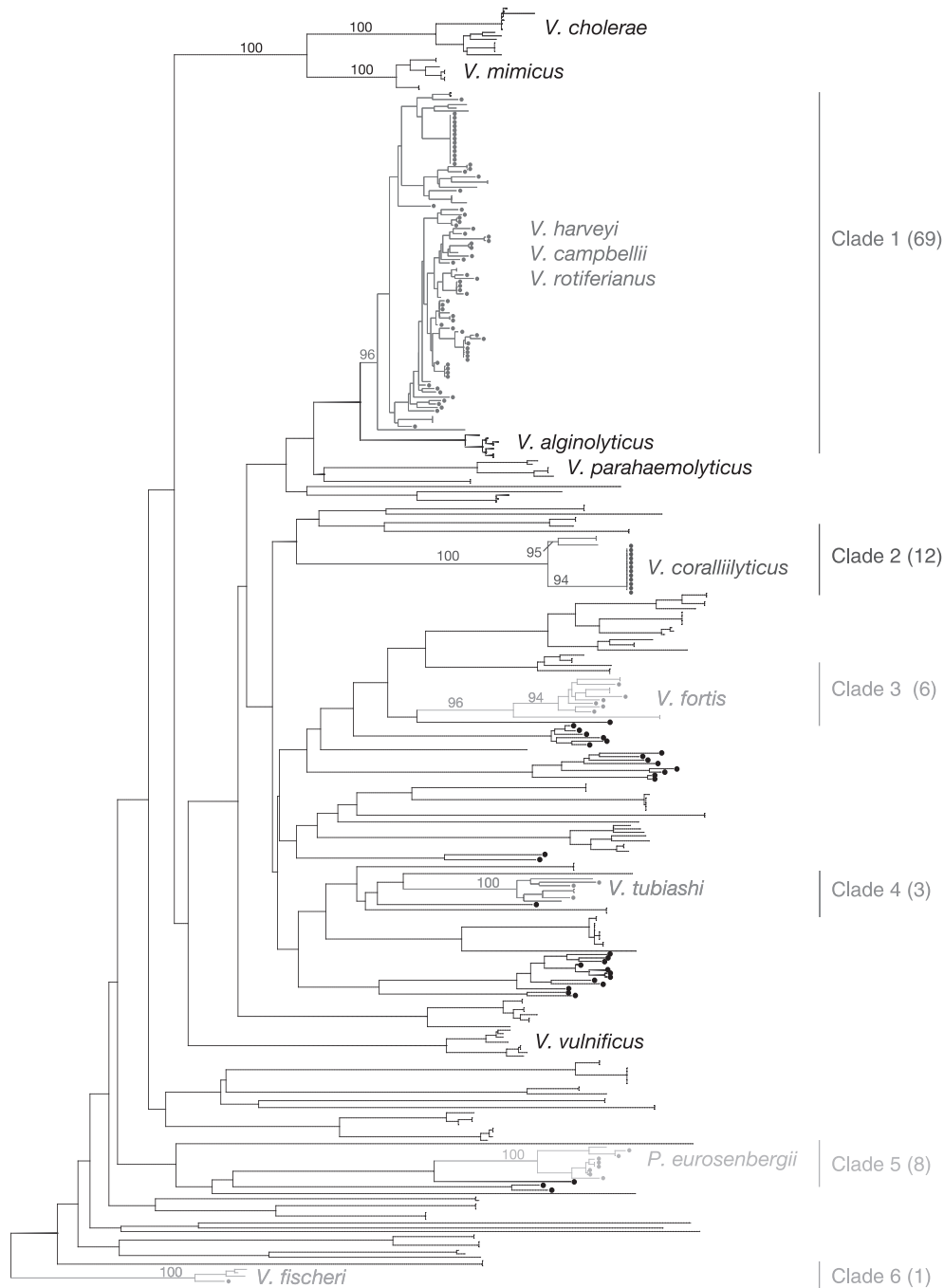


Figure 2 Taxonomic distribution of *recA* nucleotide sequences amplified from cultivars isolated from coral mucus samples. Circles at the tips of branches represent the 128 isolates cultured from mucus samples collected from a *Pocillopora damicornis* colony living in the Great Barrier Reef.

parsimonious explanation. Many *Vibrio* species are transformation competent, and high rates of transformation have been observed in marine populations (Meibom *et al.*, 2005). Phylogenetic analyses of the *attC* sites linking cassettes in these strains (data not shown) reveal little of the strain-specific signal that might be expected if within-strain integrase catalyzed rearrangements, rather than between-strain LGT, were responsible for the shuffling shown in Figure 1.

Resistance to antimicrobials may be an important function of mucus Vibrio gene cassettes

If natural antimicrobials have a determining role in coral microbiome ecology (Shnit-Orland and Kushmaro, 2009), we would expect a significant fraction of antibiotic-resistance (or synthesis) determinants among integron cassettes. Indeed, of the approximately 26% of gene cassettes encoding a protein that can be ascribed a function, nearly 48% were implicated in biochemical processes

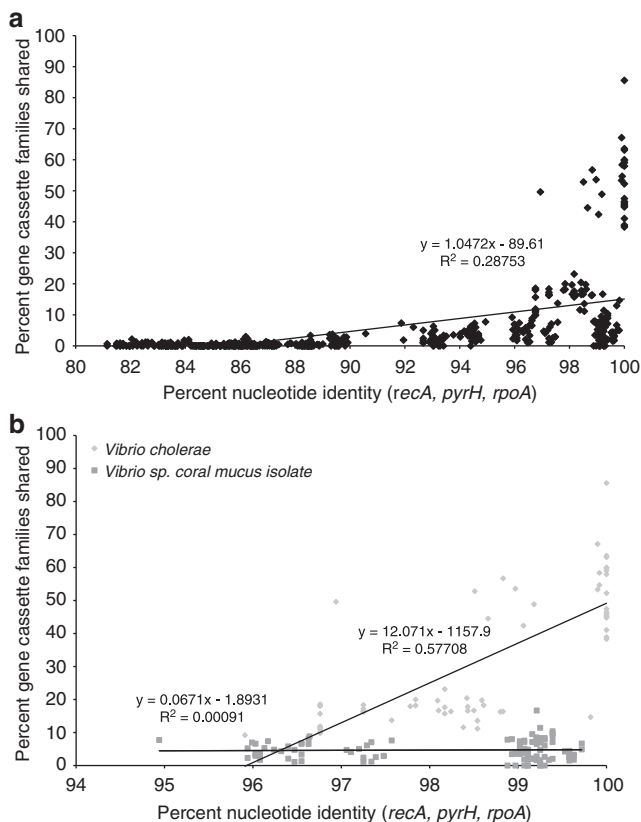


Figure 3 Pairwise comparison of the genetic identity versus integron gene-cassette array conservation in *Vibrio* isolates. (a) All *Vibrio* isolates for which gene cassettes have been sequenced. (b) *Vibrio cholerae* isolates as well as *Vibrio* strains collected from coral mucus and their close relatives. The trend line was obtained through a linear regression analysis of all data points, each point representing a pairwise comparison of isolates based on their combined *recA*, *pyrH* and *rpoA* housekeeping-genes nucleotide identity and the gene-cassette families they share. The R^2 values were calculated using the square of the Pearson product moment correlation coefficient through the given data points.

previously identified in antibiotic resistance (Supplementary Table S2). Most prevalent among these are 39 cassette-encoded acetyltransferases, an enzyme class implicated in resistance to different aminoglycosides (Wohlleben *et al.*, 1989). A total of 12 families were resolved from these acetyltransferase amino acid sequences when clustered at the BSR threshold >0.70 . Of these, four families in particular were highly similar to four different acetyltransferase genes that have been shown to confer resistance to streptothricin and virginiamycin (Supplementary Figure S2) (Liu and Pop, 2009). In addition to these acetyltransferases, eleven cassette-encoded proteins were identified as putative homologs of glyoxalase/bleomycin-resistance proteins implicated in resistance to the glycopeptide bleomycin. This abundance of cassette-encoded acetyltransferases and glyoxalases is of interest given that glycoproteins or ‘mucins’ are the primary constituents of the coral mucus matrix and may have antimicrobial activity (Brown and Bythell, 2005).

Among other cassette-encoded proteins, FtsZ and DNA topoisomerase I were also identified. FtsZ, a bacterial protein essential for cell division (Errington *et al.*, 2003), was recently targeted with a new class of antibiotic (Haydon *et al.*, 2008). Furthermore, DNA topoisomerase I, present in all bacteria (Forterre *et al.*, 2007) and responsible for removal of excess transcriptionally induced negative DNA supercoiling (Viard and de la Tour, 2007) is sensitive to a few known antibiotics. Even though DNA gyrase is considered the primary target of quinolones from *E. coli*, some of these have been shown to inhibit the relaxation activity of *E. coli* topoisomerase I (Tabary *et al.*, 1987; Moreau *et al.*, 1990; Tse-Dinh, 2009).

The mucus cassette repertoire overlaps with that of other environmental Vibrio, and in particular with those of pathogens

To assess whether cassettes obtained from our *Vibrio* cultivars were also present in known pathogens, they were compared with publicly available cassettes from *V. coralliilyticus* and *V. shiloi* (coral pathogens) as well as *V. cholerae* and *V. vulnificus* (human pathogens). Indeed, a number of the cassettes listed in Supplementary Table S3 were shared (amino acid BSR values >0.70) exclusively between *Vibrio* mucus cultivars, known coral pathogens and known human pathogens including *V. vulnificus* and *V. cholerae*. These cassette-encoded functions include putative resistance genes, specifically glyoxalases/bleomycin resistance, three different acetyltransferases and DinB, the damage-inducible protein, which is triggered during bacterial SOS response and has the potential to increase mutation rate (McKenzie *et al.*, 2001) and, potentially, the rate of cassette rearrangements (Guerin *et al.*, 2009).

To assess the connections between mucus *Vibrio* cassette arrays and cassettes within a larger global cassette metagenome regardless of taxonomic assignment, 10 365 integron gene cassette sequences were extracted from the ACID database (Joss *et al.*, 2009). These cassettes were clustered using the furthest neighbor algorithm (Legendre and Legendre, 1998) at the amino acid BSR threshold >0.70 . Figure 4 displays clusters of organisms (or environments) formed on the basis of their shared cassettes. Three main clusters emerged from this analysis and were named *Vibrio* cassettes, clinical studies and environmental surveys. The ‘clinical studies’ cluster contained the same resistance cassettes conserved in distantly related pathogens (Boucher *et al.*, 2007). For the ‘Environmental Surveys’ cluster, similar environments exhibited comparable cassette profiles (Koenig *et al.*, 2008).

Some interesting subdivisions within and between clusters could be observed (Figure 4). For example, the >2000 cassettes that were sampled from the Halifax Harbour (Canada) and its vicinity form two discrete subclusters based on shared

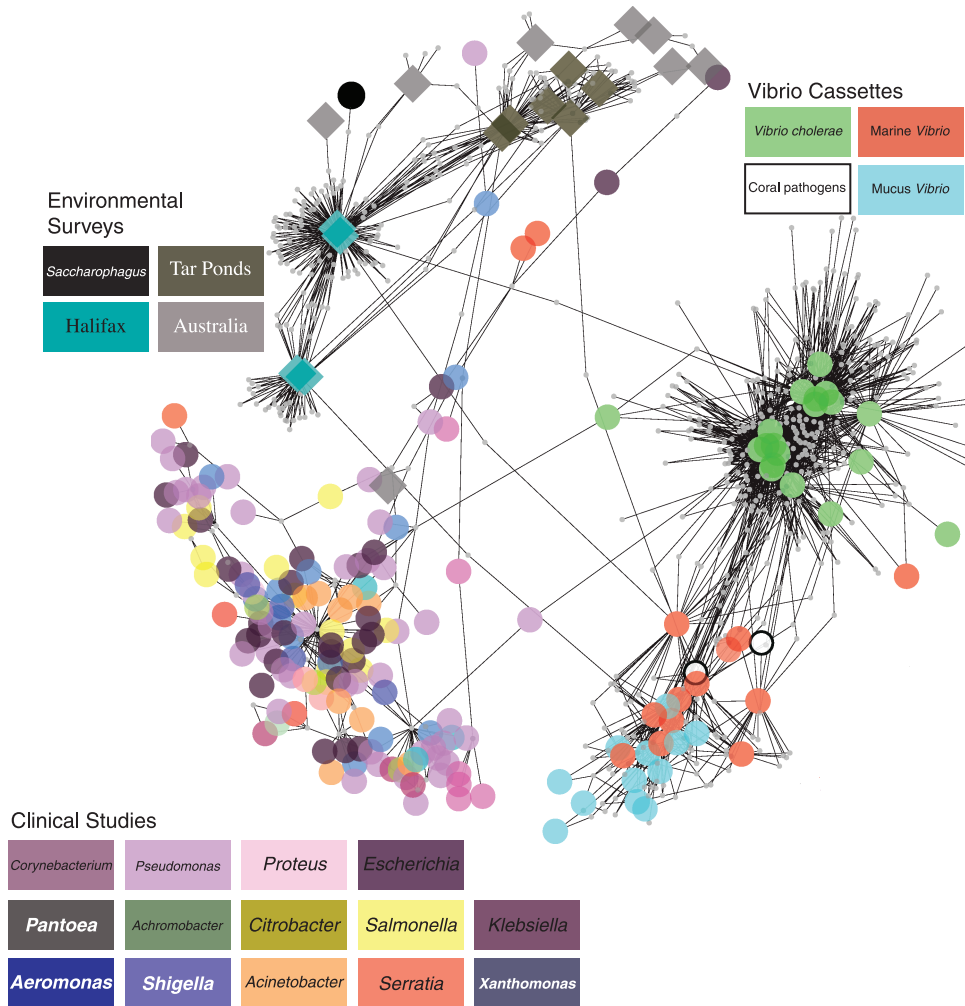


Figure 4 Network clusters based on integron gene cassettes. Cassette sequences were collected from isolates (colored circles) or environment (colored diamonds). The network is based on cassette families (small gray-colored nodes) that have an amino acid BSR thresholds > 0.70.

cassettes, one of these corresponding to the abundance of cassettes that were observed to be present in samples taken near sewage outfalls (Figure 4). These Halifax samples and the Australian samples share cassettes with each other but also with *Pseudomonas*, *Saccharophagus* and *Shigella*, specifically cassettes encoding acetyltransferases and isochorismatases-like proteins (as well as ‘conserved hypothetical’ proteins).

Phylogenetic analysis of cassettes belonging to the ‘*Vibrio* Cassettes’ cluster revealed several instances of trees, topologies of which contradict organismal phylogeny based on *rpoB* gene sequences and supports the notion that cassettes may be laterally transferred between these *Vibrio* species (Supplementary Figure S3). Furthermore, the cassettes observed to connect the ‘*Vibrio* cassettes’ and the ‘clinical studies’ clusters encode β -lactamases, chloramphenicol resistance and aminoglycoside resistance genes. Cassette-encoded amino acid sequences obtained from resistance genes were

subjected to phylogenetic analysis (Guindon and Gascuel, 2003) and produced tree topologies supporting the idea that these cassettes have been exchanged recently between *Vibrio cholerae* and other pathogens (Supplementary Figure S4).

Discussion

Cassette arrays in vibrios are generally understood to evolve rapidly in relation to chromosomal genes (Labbate *et al.*, 2007). This study has reinforced that observation but, remarkably, has also revealed that the integrons of coral mucus *Vibrio* are particularly strong evolutionary hotspots in their genomes, with a very high flux of mobile genes coming in and going out that exceeds even that seen in more free-living vibrios. This was obvious from the pairwise comparisons correlating cassette diversity with clonal divergence (Figure 3). The clear implication of this analysis is that coral mucus isolates within

a single community are rapidly exchanging gene cassettes by LGT, whereas *V. cholerae* isolates, from different hosts and outbreaks, have more independent evolutionary trajectories. Apart from the fact that individuals in coral mucus are in close proximity, what would be the advantage of this? The diversity of cassette genes within coral mucus may represent a shared resource functioning cooperatively to help maintain the health of both the microbial and coral community. The ability to use cassette gene products as a shared resource potentially has many advantages in such a community. For example, it may allow the cooperative generation of a greater array of defense products than would be the case if all individuals had to carry the same extensive genetic burden. The fact that the coral mucus bacterial community is a dense and physically limited environment means that potentially all members could benefit from the production of adaptive diffusible gene products, even if they are only produced by a small subset of the population. This could, in theory, extend to products of complex genetic pathways where individual genes in a pathway are spread across multiple individuals.

As is the case for the majority of cassette products located within chromosomal arrays, the high degree of novelty makes identifying specific functions difficult. However, meta-analysis involving in excess of 10 000 cassettes from different organisms and environment types does suggest exchange between other vibrio types (Figure 4). In part, this is almost certainly because of relatedness of *attC* sites and IntI integrase preferences but more specific host-pathogen interactions may also be important. Known ecological associations of the characterized *Vibrio* species that have close relatives among our coral mucus cultivars are summarized in Supplementary Table S1. It can be seen that eight of the nine *Vibrio* species (Supplementary Table S1) that have close relatives among our coral mucus cultivars have been implicated in the pathogenesis of marine organisms or humans, and seven of these vibrios were isolated from bleached corals (Ritchie *et al.*, 1994; Thompson *et al.*, 2005; Bourne *et al.*, 2008), although only one, *V. coralliilyticus*, has been directly proven to be an etiological agent of bleaching in *P. damicornis* (Ben-Haim and Rosenberg, 2002). *V. coralliilyticus* is also a causative agent of white syndrome disease in other coral species (Bally and Garrabou, 2007; Sussman *et al.*, 2008, 2009; Vezzulli *et al.*, 2010).

A function of some cassette proteins may also be for microbial defense—specifically to provide resistance to coral produced antimicrobials. This hypothesis is supported to the extent that, where putative general functions can be ascribed, a significant number relate to either resistance to antimicrobial compounds or to stress adaptation (Supplementary Table S3). These include matches to bleomycin-resistance compounds and acetyltransferases in the case of the former and damage-inducible proteins in the case of the latter.

Evidence continues to emerge that integron-encoded cassette products are likely to be crucial to adaptation of microbes to stressful environments. The outstanding example of this is the contribution mobile cassette products make to assisting human pathogens become multi-drug resistant (Mazel, 2006; Boucher *et al.*, 2007). For bacteria, intimate association with coral mucus may be an analogous type of stress. If so, the recent observation that stress increases IntI integrase activity thereby increasing rates of cassette rearrangement may also partly explain the increase in array diversity seen here (Guerin *et al.*, 2009). As noted above, the association between coral and vibrio may also be potentially mutually beneficial if cassette proteins contribute to coral maintenance in ways that are yet to be elucidated.

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