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Ecological Populations of Bacteria Act as Socially Cohesive Units of Antibiotic Production and Resistance

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Abstract :

In animals and plants, social structure can reduce conflict within populations and bias aggression toward competing populations; however, for bacteria in the wild it remains unknown whether such population-level organization exists. Here, we show that environmental bacteria are organized into socially cohesive units in which antagonism occurs between rather than within ecologically defined populations. By screening approximately 35,000 possible mutual interactions among Vibrionaceae isolates from the ocean, we show that genotypic clusters known to have cohesive habitat association also act as units in terms of antibiotic production and resistance. Genetic analyses show that within populations, broad-range antibiotics are produced by few genotypes, whereas all others are resistant, suggesting cooperation between conspecifics. Natural antibiotics may thus mediate competition between populations rather than solely increase the success of individuals.

Main Text:

The ratio of intra- versus interspecific competition is a key element regulating populations and determining their success within diverse communities. It is especially important in structured animal and plant populations, where closely related individuals live in patches and encounter each other often (1). In these cases, modulation of intra-specific antagonism or cooperation can mitigate the detrimental effects of niche overlap. However, for bacteria in the wild, it has been postulated that populations merely represent loose assemblages of individuals driven by ecological opportunity (2, 3). The reasons given include high dispersal rates and rapid horizontal gene transfer (HGT), which both can rapidly erode population structure by mixing unrelated individuals and introducing novel, potentially advantageous genes to their genomes. This may initiate a dynamic process of rapid, but locally and/or temporarily limited expansion of

individuals (clones). A classical example of such interactions is interference competition via colicin-type bacteriocins (4, 5), which are almost always encoded by plasmids and are able to kill closely-related competitors in a highly specific manner. In these cases, population dynamics are primarily driven by the cyclic invasion of antibiotic production and resistance genes. Similarly, a recent high-throughput screen of mutual interactions among soil isolates indicated changing types of interactions occur over relatively short evolutionary distances. This was interpreted as short-lived dynamics of gene gain and loss, where antibiotic-production selects resistance, which subsequently promotes loss of production and reversion to sensitivity (6). In contrast to this gene-centric view of bacterial population dynamics, recent fine-scale environmental mapping of bacterial diversity has suggested that population structure may exist beyond individual clones. Such ecologically defined populations consist of phylogenetic clusters of closely related but non-clonal individuals, which share common ecological associations (7, 8). However, it remains unknown whether individuals within such populations interact sufficiently strongly to allow for the development of cohesive population-level social organization akin to structured animal and plant populations.

Here, we asked whether ecologically defined populations show social cohesion beyond association with similar sets of resources. We reasoned that an obvious and important test case would be interference competition mediated by antibiotic production. This required mapping the network of potential antagonistic interactions between bacteria onto their fine-scale genotypic structure in the environment. Thus far, such an exercise has been impeded by the lack of data on genotypically and ecologically delineated natural microbial populations. In recent years, however, we have taken bacteria of the family *Vibrionaceae* as a model for the ecology and

evolution of bacterial populations (8-10). These populations were originally identified using an unsupervised machine learning approach that combines genetic similarity and micro-habitat specialization to cluster clades with cohesive ecology (8) in a way that is agnostic to any preconceived notion of species. The populations thus identified consist of groups of individuals coexisting in micro-habitats and closely related in protein housekeeping gene sequences (genotypic clusters). Many of these clusters cannot be differentiated by the most commonly used marker for phylogenetic classification of microbes, the rRNA gene, suggesting recent evolutionary age (8). Importantly, although these genotypic clusters are distinguished by their propensity to occur as free-living or associated with different types of suspended organic particles and zooplankton (8, 11), they co-occur in the guts and on the surfaces of larger marine animals (11). Because of their genotypic cohesion and differential environmental distributions, these clusters are hypothesized to represent natural populations and provide a platform to inquire whether factors beyond similarity in environmental association enforce population structure.

We first determined the potential for interference competition among individuals from different populations by measuring antagonistic interactions in an all-against-all framework using 185 *Vibrio* isolates that have been characterized to high genetic resolution by sequencing of several protein-coding, housekeeping genes (table S2). We used the classic Burkholder plate assay (12, 13), which allows testing for local growth inhibition between bacteria co-plated on nutrient agar (14). Testing inhibition in this way provides somewhat realistic conditions for ocean bacteria, since interference competition is most likely to occur among individuals co-existing on particle surfaces or in the guts of animals, where local population density can be high (13). After screening ~35,000 possible interaction pairs, we obtained a large network containing 830

antagonistic interactions between naturally co-occurring *Vibrio* strains (individual genotypes). Replication of the antagonism assay for a selected number of strains showed this data to be very robust, with more than 97% replicability of interactions (14). The data show that nearly half (44%) of the strains were able to inhibit at least one other strain, while 86% were inhibited by at least one strain. A few (<5%) ‘super-killer’ strains were able to inhibit more than 25% of all other strains in the collection (Fig. 1A). Using a membrane diffusion method (14), we estimate that 96% of the antagonistic interactions are mediated by small molecules and not by proteins such as bacteriocins.

Mapping the network of antagonistic interactions onto the fine scale genotypic structure of the tested strains shows that the potential for interference competition is much lower within natural populations than between them. This is expressed by the conditional probability of observing an antagonistic interaction as a function of genetic distance, $P(A|d)$ (Fig. 1b), where distances were computed based on a concatenated alignment of six housekeeping genes. $P(A|d)$ has a sigmoidal shape, with a 4-5 fold reduction in the probability of observing antagonism over relatively short genetic distances. Despite the strong influence that super-killer strains have on these data, this trend holds when considering only narrow range antagonists with ≤ 5 inhibited strains (Figure S1), and the bias to long-distance killing is statistically significant ($p = 0.001$) even when controlling for phylogenetic structure and killing activity per strain (14). The sharp increase in $P(A|d)$ at a threshold distance indicates that a natural genetic boundary exists for interference competition. Remarkably, this boundary coincides almost exactly with the average value of d between strains in different populations, d_{pop} , that is, the average genetic boundary between

ecologically cohesive genotypic clusters determined in previous studies (8, 10) (Fig 1C). This means that antagonism occurs mostly between rather than within natural *Vibrio* populations.

The observed low antagonism within populations is not a result of resistance between near-clonal strains as would be expected from dynamics of clonal expansion followed by gradual gene loss (6). Although the *Vibrio* populations consist of isolates with high sequence similarity in the set of shared genes, there is significant gene content diversity between strains. In 41 sequenced genomes representing 10 *Vibrio* populations (table S4), we find that although populations are clustered by gene content, the average percentage of shared genes between genomes at distances $< d_{pop}$ is only 72% (Figure S2). Moreover, these genomes are highly recombinogenic and show no evidence of a clonal origin (15). This implies that the pattern of low intra-population antagonism is not likely to be explained by simple vertical inheritance and gene loss; rather, this pattern is generated and maintained in a regime of fast allelic turnover and potential for losing and acquiring new genes.

To further explore whether antibiotic production might have co-evolved with populations or was horizontally acquired, we increased the isolate sampling around the most prolific super-killer in our collection, strain 12B09, belonging to a population of *V. ordalii*. We added a tight cluster of 29 highly related co-isolates ($d < 0.01$), which we used to study the population genomics of the super-killer phenotype. Using random transposon mutagenesis, we identified the genetic basis of antibiosis in 12B09 and studied its evolution using whole genome sequences of both producers and non-producers. This genomic approach was complemented with chemical screening and identification of active compounds.

By screening a library of 20,000 12B09 transposon mutants against a sensitive indicator strain, we identified 10 mutants with no antagonistic activity, which all had transposon insertions within a hybrid polyketide synthase (PKS) / non-ribosomal peptide (NRP) gene cluster (Figure 2A). A genetic knock-out of the central NRP biosynthesis gene shows complete loss of activity, demonstrating a single specific antibiotic biosynthesis cluster is responsible for the antagonistic activity. This is consistent with results obtained from screening a chemical extract from cell-free 12B09 supernatant separated by high performance liquid chromatography, showing that 100% of the activity could be accounted for by a single peak. Accordingly, this peak was absent from the knockout mutant 12B09-HW44, which had no antibiotic activity (Figures S4-5). Genes in the PKS-NRP cluster possess sequence similarity to cyclic lipopeptide antibiotic synthases, which typically cause membrane depolarization or pore formation, triggering cell lysis (16).

Using the cluster of 29 highly related strains from the expanded *V. ordalii* population, we obtained a high-resolution all-against-all antagonism network comprising 91 strains subdivided roughly equally across three populations of *V. ordalii*, *V. crassostreae* and *V. tasmaniensis* (table S3). This network revealed that the super-killer phenotype is present in a small fraction (5 out of 29) of the highly related strains within *V. ordalii* (Figure 2B), and was absent in any of the other populations. Moreover, all *V. ordalii* strains were resistant, confirming our previous result at a much higher genetic resolution level. A PCR screen with multiple specific primers diagnostic for the PKS-NRP biosynthetic cluster confirmed that this specific pathway is found in all of the five super-killers identified in the plate assays, and not in any of the other *V. ordalii* strains (14). This was consistent with data from the 185 strain network (Fig. 1A) showing that individuals resistant to each other did not antagonize the same set of strains (Pearson's $\phi = 0.04$). Overall, this result

shows that the super-killer phenotype in the population is caused by differential presence/absence of genes, not by transcriptional regulation or silencing mutations.

The study of sequenced genomes from two super-killers and three resistant conspecifics confirmed our prediction that recent horizontal gene transfer mediated the acquisition of production and resistance genes in *V. ordalii*. Different lines of evidence supported this. First, a whole genome phylogeny of the sequenced strains (Figure S3) showed that super-killers share a recent common ancestor, suggesting that the gene cluster is not ancestral to the population but that it was acquired in a single recent event. Second, BLAST (17) searches against public databases of fully sequenced genomes identified the antibiotic cluster only in two previously sequenced *Vibrio* isolates from the Pacific Ocean with very low genomic similarity to *V. ordalii*: the shellfish pathogen *V. tubiashii*, and the particle attached SWAT-3 (14, 18). Third, a whole genome alignment of 12B09 and SWAT-3 revealed a collinear and conserved fragment of 16.3Kb containing only the antimicrobial cluster, which indicated recent acquisition. Moreover, in SWAT-3 the antimicrobial peptide cluster is flanked by a large arrangement of transposases and integrases (Fig 1B), suggesting that the cluster comprises a mobile element, which was recently acquired in different *Vibrio* populations across distant regions of the ocean. Importantly, the resistance factors did not appear to be coded within the same mobile cluster, since none of the genes in the cluster were present in the resistant but non-producing *V. ordalii* strains. This suggested that these genes for antimicrobial production are unlinked from their resistance factors and can only invade in populations where some individuals carry pre-adaptations that enable them to survive the acquisition of antimicrobial production genes.

Contrary to the widespread idea that bacterial populations are driven by gene centric and selfish dynamics, we have shown that ecological populations defined by common microhabitat association also represent socially interacting units. Although it remains unknown how widespread the observed phenomenon is, low frequency of antagonism within short genetic distances has also recently been observed among *Streptomyces* isolates (6). Our results indicate that, similar to the case of marine vibrios, this pattern could reflect the ecological and genetic boundaries of structured populations and not a transient gene-centric dynamic (6). The fact that each antibiotic is produced by only a small fraction of the population while the rest is resistant supports the hypothesis that antibiotics can constitute public goods within populations, benefiting non-producing but resistant conspecifics. Importantly, such social structure mediates competition between ecological populations rather than benefiting only the carrier of the antibiotic production gene. Finally these result suggests that the ecological population structure of bacteria in the wild is much stronger than previously assumed.

References and Notes

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19. **Acknowledgements:** We thank W. Hanage and E. Alm for valuable suggestions. Funding was provided by the Moore Foundation, the Broad Institute's SPARC program, NSF grant DEB 0821391 and the NSF sponsored Woods Hole Center for Oceans and Human Health. Support for O.X.C. was provided by the Netherlands Organisation for Scientific Research. Whole Genome Shotgun projects have been deposited at DDBJ/EMBL/GenBank under the accessions AJWN000000000, AJYD000000000-AJYZ000000000 and AJZA000000000-AJZQ000000000 (table S4).

Supporting Online Material

Materials and Methods.

Fig. S1 to S5.

Tables S1 to S4.

References.

Figures

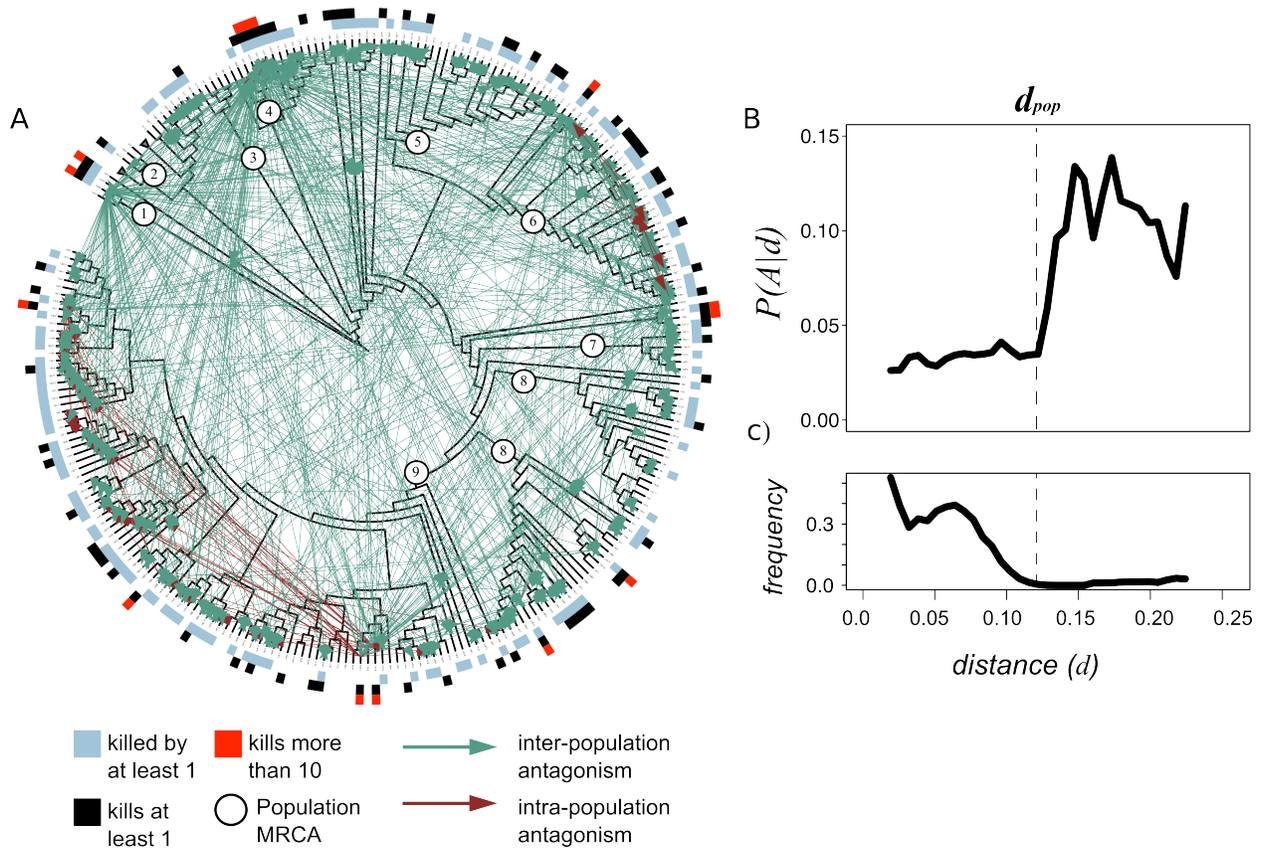


Fig. 1. Distribution of antagonistic interactions in relationship to *Vibrio* phylogeny and genetic distance. **(A)** Phylogeny of *Vibrio* isolates based on 6 housekeeping genes with outer, colored rings highlighting antagonists and sensitive strains. Green arrows connect antagonists to sensitive strains. Circles identify the most recent common ancestor (MRCA) of previously identified ecologically cohesive populations: 1, *V. ordalii*; 2, *V. fischeri*; 3, *V. breoganii*; 4, *V. alginolyticus*; 5, *V. sp. F12*; 6, *V. crassostreae*; 7, *V. cyclotrophicus*; 8, *V. tasmaniensis*; 9, *V. splendidus*. **(B)** The conditional probability of antagonism as a function of genetic distance, $P(A|d)$, shows that antagonistic interactions occur mostly between strains whose genetic distance exceeds a critical threshold. This threshold coincides with the average distance between previously defined populations (dashed line). **(C)** Frequency distribution of within-population genetic distances, showing that the transition point for $P(A|d)$ matched the average boundary of populations.

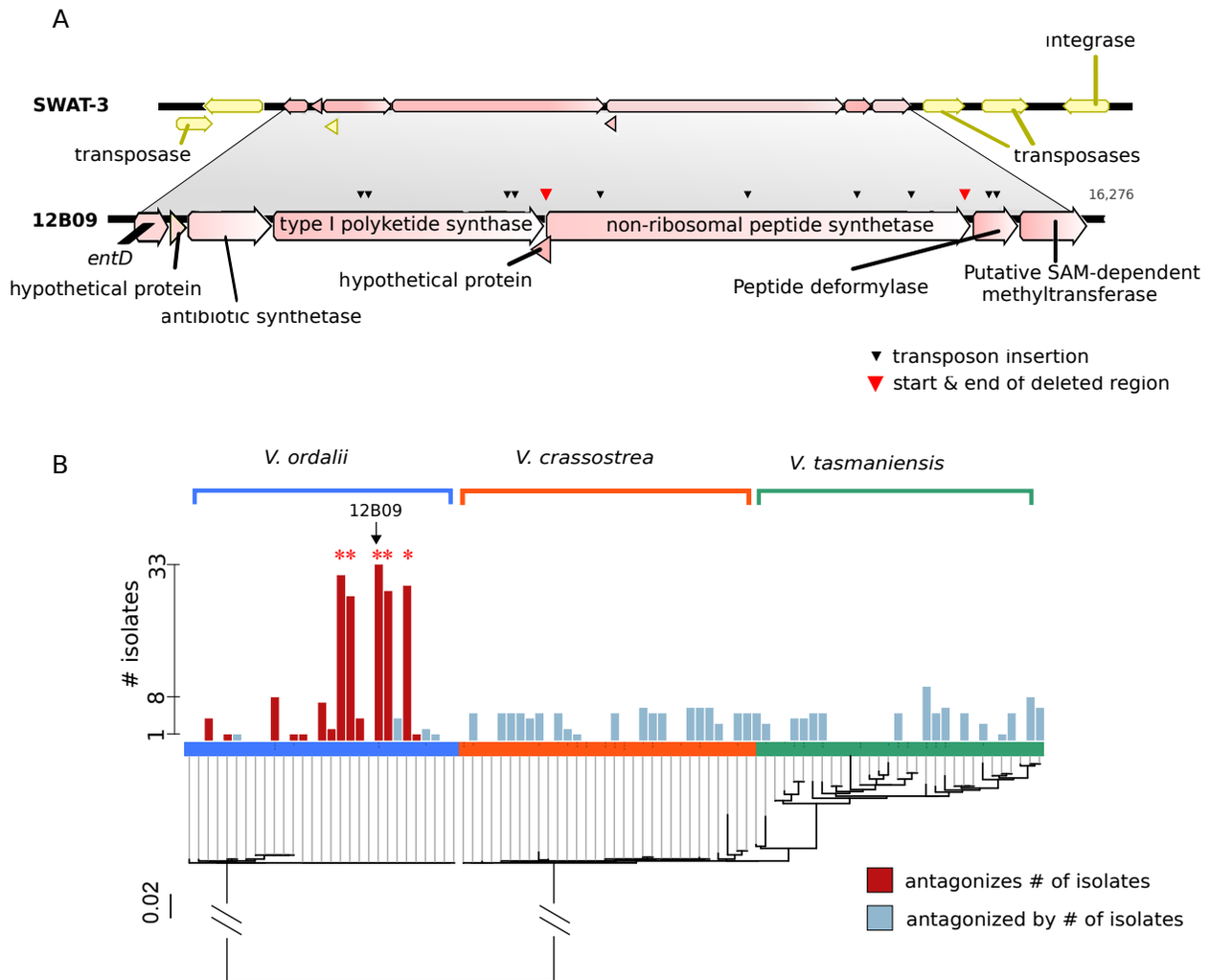


Fig. 2. Antimicrobial peptide cluster and its distribution in the *V. ordalii* population **(A)** Alignment of NRP cluster from 12B09 showing transposon insertions and knocked-out gene, and homologous cluster from environmental isolate SWAT-3. The figure shows that in SWAT-3, a nearly identical cluster is flanked by transposases and integrases, consistent with the idea that the cluster comprises a mobile element. **(B)** Antibiotic activity of *V. ordalii* isolates against *V. ordalii*, *V. crassostreae* and *V. tasmaniensis* populations. The phylogeny is based on the *hsp60* genetic marker and the bars next to each isolate indicate the number of isolates antagonized (red) and the number of *V. ordalii* isolates that antagonized the isolate (light blue). The red stars mark the five isolates where the NRP gene cluster was detected by PCR.

Supplementary Online Material

Ecological populations of bacteria act as socially cohesive units of antibiotic production and resistance

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Materials and Methods

Figs. S1 to S5

Tables S1 to S4

References

Materials and Methods

Antagonism assay and robustness of data.

Strains were grown overnight 36-48 hours prior to the time of assay in brain heart infusion (Difco) broth + 1.4% NaCl. To create a lawn of a potentially sensitive strain, 300 uL of the culture was spread on agar plates. Subsequently, potential killers were stamped onto the lawn from 96-well microtiter plates using a replicator. Zones of clearing were recorded between 18 and 24 hours after incubation at 25°C. To infer expected false positive (FP) and false negative (FN) rates, two strains were selected and their sensitivity tested against all other strains in 5 replicates. True positives are assumed to have replicability $\geq 3/5$, and otherwise they are assumed to be true negatives. Using this assumption the expected FP rate is $1.7e-4$ and the expected FN rate 0.025. In addition, a subset of 116 interactions between 130 different strains were re-tested for antagonism using 2KDa membranes to separate antibiotic producing and sensitive strains. This size-exclusion method allowed screening for antibiotics $< 2\text{KDa}$, and not proteins, which are normally in the range of $> 30\text{KDa}$. Overall, 111 out of the previously scored 116 interactions (96%) were replicated, highlighting not only the high replicability of our assay but also that the vast majority of interactions are mediated by small molecules.

Statistical significance of long distance inhibition.

We used a randomization test to measure the statistical significance of the bias towards long distance killing. We randomized the network of antagonistic interactions keeping the out-degree per node (number of killed strains) constant. This controls simultaneously for the position of killers in the phylogeny as well as for their killing activity. We measured the bias towards long distance killing by counting the number of killed strains at distance $> d_{pop}$ and found that only 1 out of 1,000 randomized networks had a bias equal or larger than the one observed in the data (non parametric p-value = 0.001).

Mapping of strains to characterized genotypic clusters

To map isolates to previously characterized populations, we created phylogenetic trees combining all sequences (with and without population assignment). To calculate the confidence of the population mapping, 100 bootstrap replicates were generated with PHYML (1). Sequences were assigned to the best matching population with at least 80% bootstrap confidence.

Measuring genetic distance.

To accurately measure evolutionary distances we used a concatenated alignment of 6 housekeeping loci: hsp60, adk, mdh, gyrB, pgi and recA, available for 166 strains. Distances were calculated using the F84 model in the software DNADIST, available in the PHYLIP package (2).

Measuring gene content overlap within populations

To measure gene content overlap we first clustered protein-coding sequences from 42 sequenced genomes representing 10 different populations using the method defined in OrthoMCL 1.4 (3) with 50% aminoacid identity cutoff and otherwise default parameters. This resulted in 15,190

clusters observed in at least one of the 42 strains. We calculated the percentage shared gene content between pairs of genomes using the Jaccard index $|A \cap B| / |A \cup B|$, where A and B are the sets of gene families present in the two genomes being compared.

Identification of antimicrobial peptide cluster.

The *E. coli* DH5 λ pir donor strain (4) was used to conjugate the mariner-derived transposon vector pSCCAT into *V. ordalii* 12B09. For conjugation, recipient and donor were grown overnight at room temperature in 5 ml tryptic soy broth (TSB, Difco) and 5 ml LB + 150 μ g chloramphenicol (Cm) + 250 μ M diaminopimelic acid (DAP), respectively. Cells of each type were harvested, mixed and plated as a mating spot on TSB plates + 250 μ M DAP. Mating spots were incubated overnight at room temperature and suspended in 20 ml TSB. 100 μ l aliquots were spread on TSB plates + 12.5 μ g/ml Cm and grown overnight at room temperature. 12B09 mutants were replica plated onto the sensitive strain ZF91 (*V. crassostreae*) and screened for loss of antagonism phenotype. Transposon insertion sites were identified by bubble PCR (Polymerase Chain Reaction). A deletion mutant was constructed using a suicide vector for construction of *Vibrio* gene deletion mutants.

Conjugation and Transposon Mutagenesis

The mariner derived transposon vector pSC-CAT was constructed from pSC180 by exchanging the kanamycin gene with the chloramphenicol acetyl transferase gene. The *E. coli* DH5 λ pir donor strain was used to conjugate pSC-CAT into *V. ordalii* 12B09 (4). For conjugation, the donor and recipient were grown overnight at 37°C in 5 ml LB + 150 μ g chloramphenicol (Cm) + 250 μ M diaminopimelic acid (DAP) and at 21°C in 5 ml tryptic soy broth (TSB), respectively. One ml of each cell type was centrifuged for 5 minutes at 13,000 rpm and resuspended in 1 ml TSB. 100 μ l of the recipient and donor was combined, centrifuged for 5 minutes at 13,000 rpm, resuspended in 10 μ l TSB, spotted onto TSB + 250 μ M DAP and incubated overnight at room temperature. The spot was suspended in 20 ml TSB. 100 μ l aliquots were spread plated on TSB + 12.5 μ g/ml Cm and grown overnight at room temperature. 12B09 mutants were replica plated to the sensitive strain ZF91 and screened for loss of antagonism phenotype. Transposon insertion sites were identified by bubble PCR.

DNA isolation and bubble PCR

Mutants with loss of antagonism phenotype were grown in 3 ml TSB + 12.5 μ g/ml Cm overnight at room temperature. For genomic DNA isolation, centrifugation of 1.5 ml cells was performed and pellets were resuspended in 300 μ l cell lysis solution (Qiagen, catalog # 158908). After incubation at 80°C for 5 minutes, 1.5 μ l of 4 mg/ml of RNase A was added and incubated at 37°C for 30 minutes. 100 μ l of protein precipitation solution (Qiagen, catalog # 158912) was added and vortex. After centrifugation, 300 μ l isopropyl alcohol was added to the supernatant. DNA was pelleted by centrifugation. The pellet was rinsed with 70% ethanol, dried and resuspended in 100 μ l DNA hydration solution (Qiagen catalogue # 158916).

Bubble PCR (BPCR) was used to identify transposon insertion in the genome of mutants showing loss of antagonism phenotype. Two μ g of genomic DNA was digested with restriction

enzyme *PvuII*, *SacI*, *SmaI*, or *SspI*. Digested DNA was ligated to 4 μ M of annealed bubble PCR primers BPHI and BPHII using T4 DNA ligase. BPCR I was performed using 2 μ l of digested DNA and 5 μ M primers 224 and pSCCAT2067 which anneal to the ligated primer BPHI and the pSC-CAT vector, respectively. BPCR I was performed at 92°C denaturing for 10 s, 60°C annealing for 60 s, and elongation at 72°C for 90 s and repeated 19 times. BPCR I products were diluted 1:25 and 2 μ l of DNA was used to perform BPCR II with 5 μ M primers 224 and pSCCAT2036. BPCR reaction II was performed at 92°C denaturing for 10 s, 55°C annealing for 30 s, 72°C elongation for 90 s and was repeated 34 times. BPCR II products were sequenced by GENEWIZ, Inc using primers 224, pSCCAT2067, and pSCCAT3036.

Construction of pJC4

Plasmid pJC4 was constructed from pMW91 (4) and used to create *Vibrio* gene deletion mutants. pJC4 is a R6K based suicide vector containing *sacB* gene for counter-selection of allelic exchange. Briefly, pMW91 was digested with *ScaI* to disrupt the beta lactamase gene. The chloramphenicol acetyl transferase gene was amplified from pACYC184 and cloned into the *ScaI* site giving pJC4.

Deletion of non-ribosomal peptide (NRP) biosynthesis gene

The NRP gene was targeted for deletion by creating a fusion PCR fragment that lacked the NRP gene as shown in (5). Briefly, 750 base pairs upstream and downstream of the NRP gene were amplified using the primer pairs 12B09 2900F - 12B09 3683R and 12B09 9848F - 12B09 10593R, respectively. Primers 12B09 2900F and 12B09 10593R have a complementary tail so their respective amplicons can anneal and produce a single template generating a fusion Δ NRP PCR product. For fusion PCR, the 1.5 kb Δ NRP product was generated using the 12B09 2900F and 12B09 10593R primers with one μ l of the up- and downstream products as the DNA template. The fusion PCR product was digested with *SpeI* and cloned into the suicide plasmid pJC4 resulting in pHW44. pHW44 was conjugated to 12B09 as described above. Homologous recombination between fusion PCR product and the 12B09 chromosome resulted in chloramphenicol resistant transconjugants. Transconjugants were grown overnight in TSB + 2%NaCl. A serial dilution was performed on LB plates without salt + 5% sucrose. Loss of the NRP gene was confirmed by sequencing using primers 12B09 2900F and 12B09 10593R.

Table S1. Primers used in this study.

Primer	Sequence
BPHI	5' CAA GGA GAGGACGCT GTC TGT CGA AGG TAA GGA ACG GAC GAG AGA AGG GAG AG 3'
BPHII	5' CTC TCC CTT CTC GAA TCG TAA CCG TTC GTA CGA GAA TCG CTG TCC TCT CCT TG 3'
224	5' CGA ATC GTA ACC GTT CGT ACG AGA ATC GCT 3'
pSCCAT2067	5' GGC CGA AAT CGG CAA AAT CC 3'
pSCCAT2036	5' AAA GAA TAG ACC GAG ATA GG 3'
12B09 2900F	5' GGA CTA GTA CGC GAC ACT CAA AGC CAG C 3'

12B09 3683R	5' GGT GGT CCT CAA AGA AAT TAA CGA GTT CTA AAT ATT CGA CTA AGT G 3'
12B09 9848F	5' TCG TTA ATT TCT TTG AGG ACC ACC AAA CAA CAA TGG CTA TCA GAC C 3'
12B09 10593R	5' GGA CTA GTT ACA AAC CCG GAC CGC AGG C 3'

Extraction of antibiotic from *Vibrio ordalii* 12B09

Strains of 12B09 (either the wt or HW44) were inoculated from overnight starter cultures and grown in 1L of Difco 2216 marine broth in a 2.8 L Fernbach flask with 150 RPM shaking at 25 C° for 24 hours. Subsequently, 20 g of sterile XAD-7 resin was added with shaking at 25 C° for 16 hours. The XAD-7 resin was harvested by filtration through glass wool and rinsed with 1 L of distilled water and dried under vacuum filtration. The resin was eluted thrice with 100 mL of optime grade methanol (Fisher Scientific, Valencia CA.), which was subsequently reduced to dryness using rotary evaporation. Crude extracts were prepared by resuspension in DMSO to a final concentration of 50 mg/mL. These were filtered through a 0.45 µm glass fiber filter and stored at -20 C° until further analyses. Crude bioactivities were compared using quantitative dilution and employing a disk-diffusion method measuring inhibition zone on solid medium as described in methods. The crude extract was found to account for 100% of the culture activity based on cell counts of cultures of 12B09wt. Biological testing was performed in biological triplicates in the testing of 12B09-HW44.

Large-scale normal phase flash chromatography

Crude extract paste material from 10 replicate 1 L cultures of 12B09wt was obtained as outlined above and resuspended in 20 mL 1:1:1 trimethylpentane (TMP):ethylacetate:methanol. The crude resuspension was mixed with 50 g of silica gel mesh 230-400 (Whatman, Piscataway, NJ), dried using rotary evaporation and loaded onto a 300 ml bed volume of silica gel in a 600 ml vacuum funnel. Solvents of increasing polarity were added and collected as separate fractions of one bed volume of TMP, 1:1 TMP:ethylacetate, ethylacetate, 1:1 ethylacetate:acetone, acetone, and methanol. All fractions were evaporated to dryness and tested by a disk-diffusion assay. The acetone fraction was found to account for 100% of the antibiotic activity.

Supporting Figures

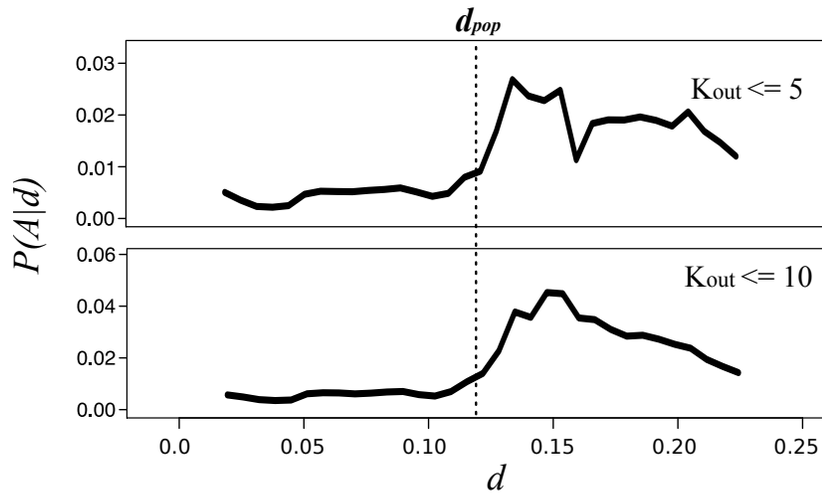


Figure S1. Bias towards long distance antagonism was not driven by broad-range killers. $P(A|d)$ represents the probability of antagonism given the genetic distance, d , and d_{pop} marks the boundary between ecological populations as described in the main text. In the upper panel ($K_{out} \leq 5$), strains inhibiting more than 5 other strains have been removed from the network along with their interactions. In the lower panel ($K_{out} \leq 10$), the maximum allowed number of inhibited strains is 10. The figures show that the trend depicted in Figure 1 of the main text holds even when focusing on highly specific antagonisms.

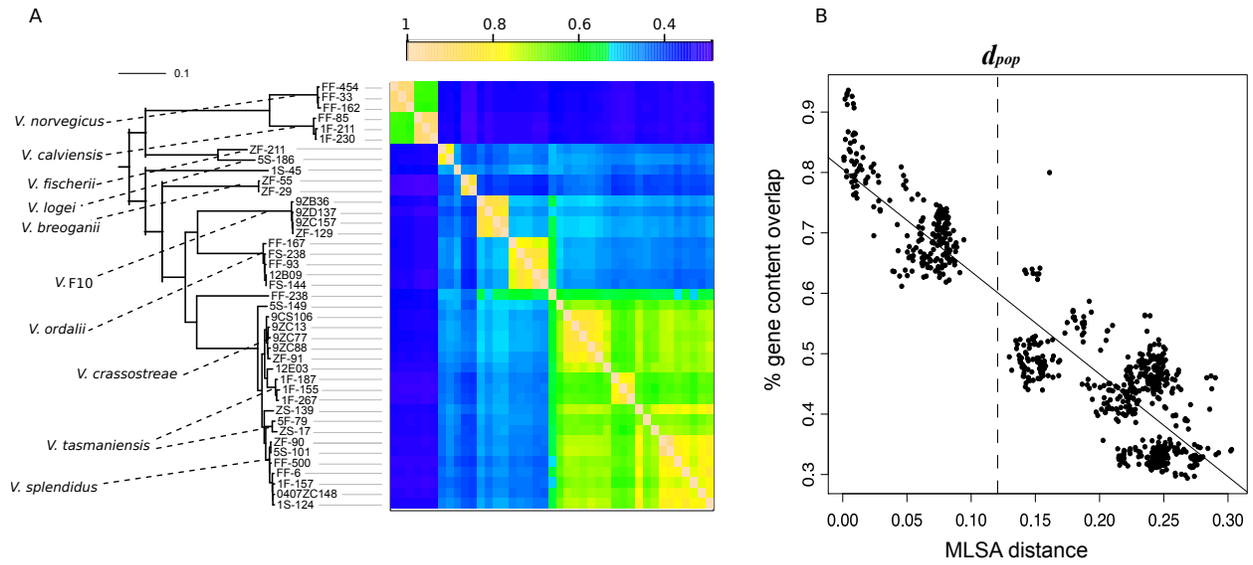


Figure S2. High variability in gene content despite overall coherence of phylogenomic signal. **A)** *Vibrio* phylogeny of 41 sequenced genomes from 10 populations based on 40 gene families present in 97.5% of the sequenced strains. Populations form tight clusters based on housekeeping genes and can also be clustered based on gene content. **B)** The figure shows the overlap in gene content as a function of the genetic distance calculated from 6 housekeeping genes as described in the main text. The average gene content overlap is 72% below d_{pop} . Each data point represents pairwise comparisons between the 41 sequenced strains.

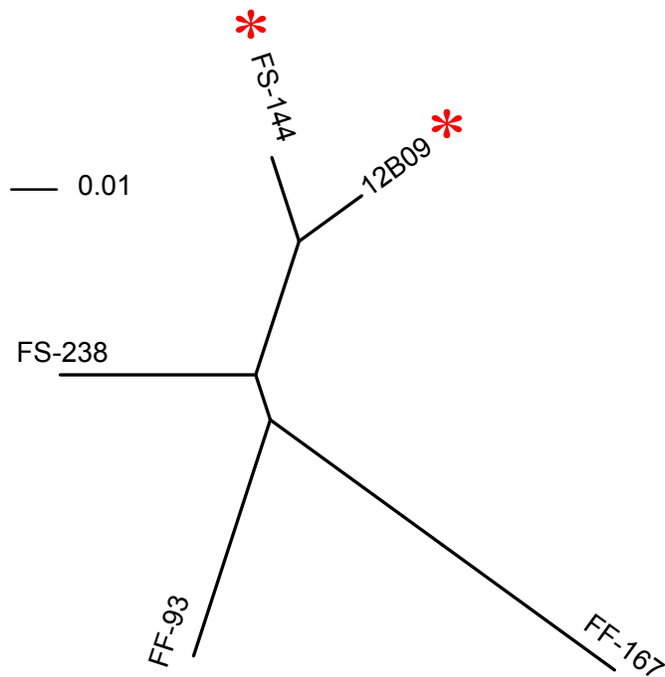


Figure S3. Whole genome tree of five sequenced *V. ordalii* strains shows that super-killers 12B09 and FS-144 have recently diverged, suggesting that the acquisition of the antimicrobial peptide was a single recent event in their common ancestor. Whole genome alignment tree were generated with Mauve (6). Red asterisks indicate super-killer strains carrying the antibiotic cluster.

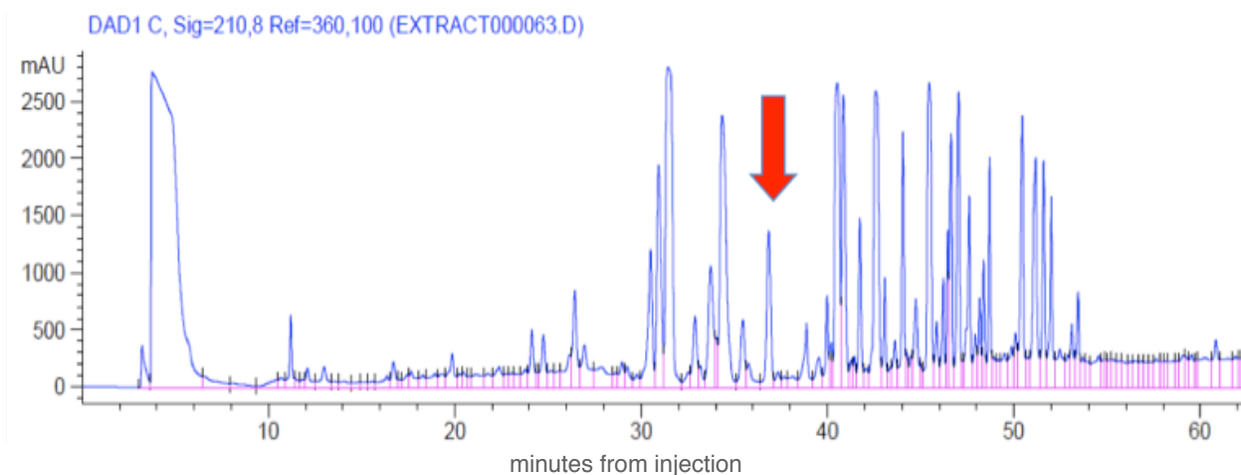


Figure S4. Identification of active fraction from 12B09 by chromatography. The figure shows the reverse-phase HPLC-Diode array detection (at 210nm) chromatogram of a flash chromatography (100% acetone elute from silica gel) run with a gradient of 0-30% acetonitrile:water, 0.4mL/min. The compound marked by red arrow eluting at 37 min (18% acetonitrile:water) was responsible for 100% of the antibiotic activity in the 12B09 antibiotic assay.

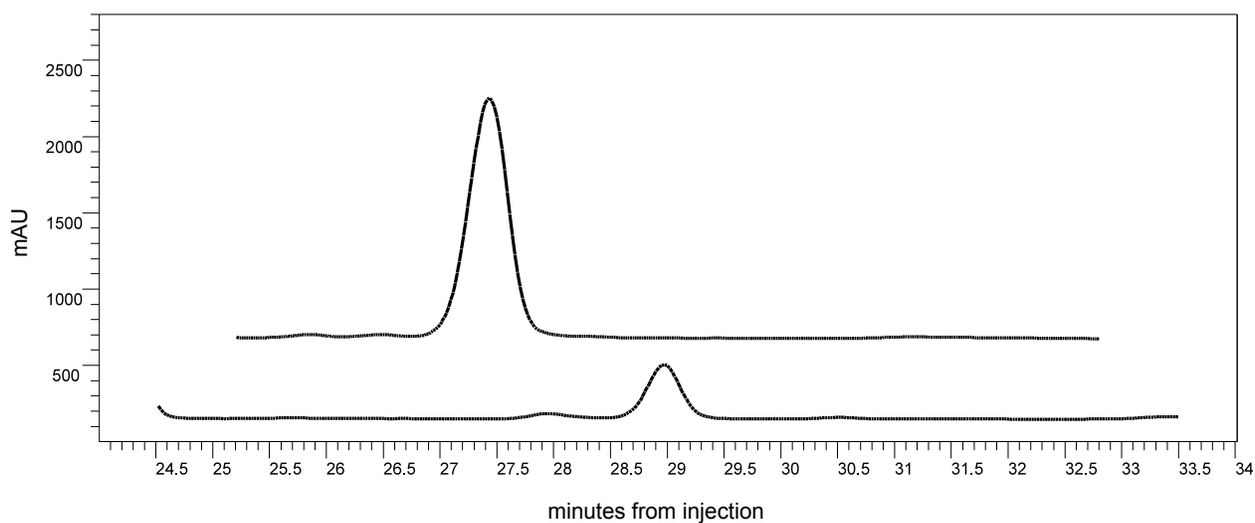


Figure S5: Lack of active compound in knock-out mutant 12B09-HW44. The figure shows a reverse-phase HPLC-DAD analysis (monitored at 300 nm) employing a 20% acetonitrile:water isocratic solvent system at a at 0.4 mL/min flow rate to compare the transposon knock-out mutant HW44 (bottom trace) with 12B09 wt (top trace). The peak at 27.4 minutes accounts for 100% of antibiotic activity and is completely absent in the HW44 trace. The appearance of the small peak at the 29 minute retention time is a malformed product which was found to have no antibiotic activity by peak-based collection and bioassay.

Table S2. List of 185 isolates from the all-against-all network with their corresponding out-degree (number of isolates antagonized) and in-degree (number of isolates that antagonize it).

isolates name	out-degree	in-degree
12A11	0	1
13B11	77	5
12C02	0	3
12A08	2	2
12G02	33	11
1C04	2	0
12G06	0	6
1C02	3	0
13A01	0	8
13A11	1	5
14C02	0	1
14B06	1	4
13E02	0	7
13F02	0	7
14H08	0	2
1A12	0	4
13B10	3	6
14C08	2	1
12F11	1	15
12G11	5	2
13E09	1	0
1B09	0	5
1B01	0	4
1B11	0	6
12G04	0	9
12G12	107	1
12G08	0	7
12B11	3	0
12B04	2	6
13B03	0	1
13D07	0	4
12A10	8	1
12H04	2	4
13E05	1	6
13H05	0	3
12E12	1	4
14D03	0	5
14C03	1	4
14B10	0	4

isolates name	out-degree	in-degree
14D12	0	11
14H05	0	4
14G04	0	6
14D07	44	2
14A03	1	5
1C03	0	8
1C09	1	4
13A03	0	9
13C01	0	4
12D12	1	2
12B05	10	0
12H10	1	0
13G02	2	6
12A09	4	0
12B10	18	3
13H06	0	3
12F12	0	9
14F08	0	3
14F05	0	4
14E05	1	3
14G03	0	4
14E06	0	6
14G09	0	3
12F08	12	9
12F03	3	10
12E06	7	7
12C11	1	5
13E07	1	0
14E02	4	0
1C01	1	2
1D01	4	1
12F05	6	0
1A05	0	1
14H11	0	5
13C05	1	3
12H09	0	6
13A10	0	14
12D11	1	4
12C05	0	3
12D09	0	4
14E01	0	3
14A06	1	4

isolates name	out-degree	in-degree
13G08	0	4
1D08	0	4
13G12	0	3
14C04	55	0
14E03	0	3
12G09	0	6
13H11	0	4
13D01	0	1
1E06	0	3
1D10	0	4
14B11	0	3
12D07	0	6
14H09	0	3
14G06	0	11
1A11	0	5
14D09	0	4
14B12	0	9
12H05	0	7
14C01	0	1
13D12	0	7
13D09	1	8
1D02	0	3
13E06	0	18
12E04	1	0
13C04	0	4
12H11	3	0
12A01	4	14
12C09	2	0
13G10	0	1
1B07	1	8
14A07	0	7
12D08	4	1
13C12	0	3
14C10	2	0
13B01	1	7
14H02	0	5
14C11	0	15
1B03	0	5
13E04	1	6
13F08	0	3
12A06	18	0
1A09	0	1

isolates name	out-degree	in-degree
14B08	0	6
12G05	0	10
12G01	42	28
12G07	1	6
12G03	0	2
12B09	110	3
12F02	1	4
12E02	1	6
14F04	7	0
1A03	0	6
1C07	0	11
1B02	0	5
13A04	0	3
13F04	1	0
1B10	0	9
14A11	13	5
14F06	0	3
14C05	0	2
13H04	1	4
14A10	1	1
14C06	1	1
12E10	0	4
14E04	2	6
13B07	0	10
12H02	0	4
13A02	0	2
14G12	5	1
13E10	4	0
12G10	0	6
14F11	0	3
13A08	0	9
1E01	0	5
1D09	0	7
12D05	2	0
13H12	0	3
13H02	0	2
13B12	5	9
14F09	0	3
1D07	0	7
12F04	31	5
14F10	0	4
14E07	1	4

isolates name	out-degree	in-degree
14C07	38	0
13G07	0	6
12H01	0	5
14C09	61	0
14A04	0	5
1C05	0	1
13F11	0	2
12B08	17	0
13D11	1	1
14H06	0	9
14A12	1	0
12B02	0	12
13D03	2	0
13C10	1	2
7G11	4	0
1D12	0	3
12F10	0	11
12F06	1	9
12E11	0	7
12E05	1	7

Table S3. List of 91 isolates from the *V. ordalii*-tasmaniensis-crassostreae network with their corresponding out-degree (number of isolates antagonized) and in-degree (number of isolates that antagonize it). The field “in-degree (from ordalii)” shows the number of ordalii isolates that contribute to the total in-degree (Fig 2 of the main text).

isolate name	population	out-degree	in-degree	in-degree (from ordalii)
FAL5122	<i>V. crassostreae</i>	0	5	5
FALZ15	<i>V. crassostreae</i>	0	7	5
FALZ91	<i>V. crassostreae</i>	0	8	5
12F10	<i>V. crassostreae</i>	0	8	6
14A01	<i>V. crassostreae</i>	10	6	6
FAL5161	<i>V. crassostreae</i>	0	8	5
FALZ17	<i>V. crassostreae</i>	0	5	2
FAL1165	<i>V. crassostreae</i>	0	7	6
12C04	<i>V. crassostreae</i>	9	1	1
FALZ8	<i>V. crassostreae</i>	0	3	0
13E10	<i>V. crassostreae</i>	2	5	5
14E07	<i>V. crassostreae</i>	0	2	0
FALZ219	<i>V. crassostreae</i>	0	5	5
FALZ269	<i>V. crassostreae</i>	0	7	6
12A08	<i>V. crassostreae</i>	0	3	0
13H05	<i>V. crassostreae</i>	0	1	0
12B04	<i>V. crassostreae</i>	0	6	5
FALF372	<i>V. crassostreae</i>	0	4	4
FALZ49	<i>V. crassostreae</i>	0	5	5
14F03	<i>V. crassostreae</i>	0	5	5
13D10	<i>V. crassostreae</i>	0	6	5
SPR1297	<i>V. crassostreae</i>	0	5	5
FALZ33	<i>V. crassostreae</i>	0	1	0
FALZ100	<i>V. crassostreae</i>	0	5	5
13G12	<i>V. crassostreae</i>	0	0	0
14C11	<i>V. crassostreae</i>	0	0	0
FAL1255	<i>V. crassostreae</i>	0	0	0
FALF58	<i>V. crassostreae</i>	0	0	0
FALZ113	<i>V. crassostreae</i>	0	0	0
12E02	<i>V. crassostreae</i>	0	0	0
13H03	<i>V. crassostreae</i>	0	0	0
SPRF143	<i>V. ordalii</i>	28	0	0
SPR1143	<i>V. ordalii</i>	4	0	0
FALF159	<i>V. ordalii</i>	4	0	0
SPR1257	<i>V. ordalii</i>	7	0	0
FALF68	<i>V. ordalii</i>	0	2	2

isolate name	population	out-degree	in-degree	in-degree (from ordalii)
FALF2	V. ordalii	8	0	0
FALF197	V. ordalii	0	1	1
FALF236	V. ordalii	1	0	0
SPRF219	V. ordalii	1	0	0
SPRF203	V. ordalii	0	1	1
FAL1128	V. ordalii	1	0	0
FALF222	V. ordalii	2	0	0
FALF31	V. ordalii	1	0	0
SPRF144	V. ordalii	27	0	0
FALF443	V. ordalii	29	0	0
SPRF148	V. ordalii	0	4	4
FALF495	V. ordalii	31	0	0
12B09	V. ordalii	33	0	0
FALF19	V. ordalii	0	0	0
FALF498	V. ordalii	0	0	0
FALF166	V. ordalii	0	0	0
FALF134	V. ordalii	0	0	0
FALF317	V. ordalii	0	0	0
FALF343	V. ordalii	0	0	0
FALF344	V. ordalii	0	0	0
FALF93	V. ordalii	0	0	0
SPRF217	V. ordalii	0	0	0
SPRF129	V. ordalii	0	0	0
SPRF206	V. ordalii	0	0	0
FALF36	V. tasmaniensis	1	0	0
12E04	V. tasmaniensis	0	4	4
13E05	V. tasmaniensis	0	5	5
12F06	V. tasmaniensis	1	0	0
14G11	V. tasmaniensis	0	5	5
14B04	V. tasmaniensis	0	5	5
13B12	V. tasmaniensis	1	0	0
FALF386	V. tasmaniensis	0	8	8
13B11	V. tasmaniensis	8	3	3
13G10	V. tasmaniensis	0	5	5
FAL1300	V. tasmaniensis	0	4	1
13B07	V. tasmaniensis	0	1	0
FALZ76	V. tasmaniensis	0	3	3
13H11	V. tasmaniensis	0	3	3
14F05	V. tasmaniensis	0	10	10
FALF481	V. tasmaniensis	0	5	5
14B07	V. tasmaniensis	0	6	6

isolate name	population	out-degree	in-degree	in-degree (from ordalii)
14B02	V. tasmaniensis	0	5	5
14G06	V. tasmaniensis	0	5	4
FALF350	V. tasmaniensis	0	6	6
FAL1292	V. tasmaniensis	0	0	0
FAL1293	V. tasmaniensis	0	0	0
FALF233	V. tasmaniensis	0	0	0
FAL5261	V. tasmaniensis	0	0	0
13F02	V. tasmaniensis	0	0	0
14H03	V. tasmaniensis	0	0	0
14H05	V. tasmaniensis	0	0	0
14G12	V. tasmaniensis	0	0	0
12H02	V. tasmaniensis	0	0	0
12H09	V. tasmaniensis	0	0	0
12E09	V. tasmaniensis	0	0	0

Table S4. Accession numbers of 41 genome sequences used in gene content analysis (Fig S2).

Bioproject ID	Accession number	Strain name
PRJNA164629	AJWN000000000	Enterovibrio norvegicus FF-454
PRJNA164633	AJYD000000000	Enterovibrio norvegicus FF-33
PRJNA164635	AJYE000000000	Enterovibrio norvegicus FF-162
PRJNA164637	AJYF000000000	Enterovibrio calviensis FF-85
PRJNA164639	AJYG000000000	Enterovibrio calviensis 1F-211
PRJNA164641	AJYH000000000	Enterovibrio calviensis 1F-230
PRJNA164727	AJYI000000000	Aliivibrio fischeri ZF-211
PRJNA164729	AJYJ000000000	Aliivibrio logei 5S-186
PRJNA164805	AJYK000000000	Vibrio rumoiensis 1S-45
PRJNA164807	AJYL000000000	Vibrio breoganii ZF-55
PRJNA164811	AJYM000000000	Vibrio breoganii ZF-29
PRJNA164821	AJYN000000000	Vibrio genomosp. F10 str. 9ZB36
PRJNA164823	AJYO000000000	Vibrio genomosp. F10 str. 9ZD137
PRJNA164825	AJYP000000000	Vibrio genomosp. F10 str. 9ZC157
PRJNA164829	AJYQ000000000	Vibrio genomosp. F10 str. ZF-129
PRJNA164835	AJYR000000000	Vibrio ordalii FF-167
PRJNA164837	AJYS000000000	Vibrio ordalii FS-238
PRJNA164841	AJYT000000000	Vibrio ordalii FF-93
PRJNA164843	AJYU000000000	Vibrio ordalii FS-144
PRJNA164847	AJYV000000000	Vibrio ordalii 12B09
PRJNA167003	AJYW000000000	Vibrio genomosp. F6 str. FF-238
PRJNA167004	AJYX000000000	Vibrio kanaloae 5S-149
PRJNA167005	AJYY000000000	Vibrio crassostreae 9CS106
PRJNA167006	AJYZ000000000	Vibrio crassostreae 9ZC13
PRJNA167007	AJZA000000000	Vibrio crassostreae 9ZC77
PRJNA167008	AJZB000000000	Vibrio crassostreae 9ZC88
PRJNA167009	AJZC000000000	Vibrio crassostreae ZF-91
PRJNA167010	AJZD000000000	Vibrio splendidus 12E03
PRJNA167011	AJZE000000000	Vibrio splendidus ZS-139
PRJNA167012	AJZF000000000	Vibrio splendidus ZF-90
PRJNA167013	AJZG000000000	Vibrio splendidus 5S-101
PRJNA167014	AJZH000000000	Vibrio splendidus FF-500
PRJNA167015	AJZI000000000	Vibrio splendidus FF-6
PRJNA167016	AJZJ000000000	Vibrio splendidus 1F-157
PRJNA167017	AJZK000000000	Vibrio splendidus 0407ZC148
PRJNA167018	AJZL000000000	Vibrio splendidus 1S-124
PRJNA167019	AJZM000000000	Vibrio tasmaniensis 1F-187
PRJNA167047	AJZN000000000	Vibrio tasmaniensis 1F-155
PRJNA167048	AJZO000000000	Vibrio tasmaniensis 1F-267
PRJNA167049	AJZP000000000	Vibrio tasmaniensis 5F-79

Bioproject ID	Accession number	Strain name
PRJNA167050	AJZQ00000000	Vibrio tasmaniensis ZS-17

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