Bacterial symbionts use a type VI secretion system to eliminate competitors in their natural host

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Intraspecific competition describes the negative interaction that occurs when different populations of the same species attempt to fill the same niche. Such competition is predicted to occur among host-associated bacteria but has been challenging to study in natural biological systems. Although many bioluminescent Vibrio fischeri strains exist in seawater, only a few strains are found in the light-organ crypts of an individual wild-caught Euprymna scolopes squid, suggesting a possible role for intraspecific competition during early colonization. Using a culture-based assay to investigate the interactions of different V. fischeri strains, we found “lethal” and “non-lethal” isolates that could kill or not kill the well-studied light-organ isolate ES114, respectively. The killing phenotype of these lethal strains required a type VI secretion system (T6SS) encoded in a 50-kb genomic island. Multiple lethal and nonlethal strains could be cultured from the light organs of individual wild-caught adult squid. Although lethal strains eliminate nonlethal strains in vitro, two lethal strains could coexist in interspersed microcolonies that formed in a T6SS-dependent manner. This coexistence was destabilized upon physical mixing, resulting in one lethal strain consistently eliminating the other. When juvenile squid were coinoculated with lethal and nonlethal strains, they occupied different crypts, yet they were observed to coexist within crypts when T6SS function was disrupted. These findings, using a combination of natural isolates and experimental approaches in vitro and in the animal host, reveal the importance of T6SS in spatially separating strains during the establishment of host colonization in a natural symbiosis.

Significance

Competition among cooccurring bacteria can change the structure and function of a microbial community. However, little is known about the molecular mechanisms that impact such interactions in vivo. We used the association between bioluminescent bacteria and their squid host to study how environmentally transmitted bacteria compete for a limited number of host colonization sites. Our work suggests Vibrio fischeri use a type VI secretion system, acting as a contact-dependent interbacterial “weapon,” to eliminate competing strains from cooccupying sites in the host. This work illuminates a mechanism by which strain-specific differences drive closely related bacteria to engage in lethal battles as they establish a beneficial symbiosis, revealing how genetic variation among potential colonizers directly impacts the spatial structure of the host-associated population.

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predicted to be translocated, prevents self-intoxication and also protects isogenic cells from growth inhibition or cell lysis. However, if the toxic effector is translocated into a nonisogenic or competitor cell that does not encode the appropriate immunity gene, the competitor cell is inhibited or eliminated. Despite the broad distribution of interbacterial killing mechanisms, few studies have investigated their ecological roles (14, 17, 43, 50, 51). We hypothesize that in addition to encoding necessary host colonization factors, successful colonizers may also employ mechanisms to exclude closely related competitors and occupy a limited number of colonization sites.

To test this hypothesis, a tractable model system is required. The Vibrio–squid symbiosis is a simplified model for studying bacterial colonization in an animal host (52). Euprymna scolopes squid house multiple strains of bioluminescent Vibrio fischeri bacteria in a structure called the light organ (53). Juvenile squid hatch without their bacterial symbionts, which they must acquire from the surrounding seawater (54). Although studies addressing the strain-level diversity of V. fischeri in the seawater are limited (55), it is predicted to be much higher than the diversity found in a naturally colonized squid, which harbors only a few strains per light organ (53). Potential colonizers first aggregate on the surface of the light organ before entering pores that lead to six physically separated crypt spaces, where cells rapidly proliferate (56). Importantly, the passageway leading to the crypt entrance is physically constricted to the size of approximately one bacterial cell, restricting access to a single cell at a time (53). Because on average only a few cells enter each crypt space, the Vibrio–squid symbiosis offers a unique opportunity to answer questions about intraspecific competition that arises during the natural host colonization process.

Previous work suggests that V. fischeri strains employ competitive mechanisms as they colonize the host. For example, pairwise co colonization experiments of juvenile squid using different combinations of light-organ isolates revealed that colonized animals often possess a light-organ population dominated by a single strain, based on counts of colony-forming units (CFUs) (59, 60). Recently, we showed competition that appears to occur within individual crypts of the light organ: when juveniles were exposed to an inoculum containing two different strains, the resulting crypts were physically separated crypt spaces, where cells rapidly proliferate. Despite the broad distribution of interbacterial killing mechanisms, few studies have investigated their ecological roles. We hypothesize that in addition to encoding necessary host colonization factors, successful colonizers may also employ mechanisms to exclude closely related competitors and occupy a limited number of colonization sites.

V. fischeri Isolates Are Capable of Strain-Specific Competitive Interactions. To begin exploring potential competitive interactions among V. fischeri isolates, we first examined how strains coisolated from the same light organ interact with the commonly studied symbiotic isolate, ES114. V. fischeri ES114 was isolated in 1989 from an adult E. scolopes host collected in Kanehoe Bay, Hawaii (62). For more recently isolated strains, we selected six additional strains: three coisolated strains that each came from two different hosts. We chose to examine interactions using two groups of recently coisolated strains because these isolates may have physically encountered one another in aggregates on the light-organ surface or within crypts during the natural host colonization process in wild Euprymna juveniles. Moreover, some microorganisms lose competitive genes when kept in isolation (63); therefore, we reasoned that newly isolated strains may be less likely to have lost important genes that mediate interstrain interactions.

To identify possible competitive behaviors among different symbiotic strains, we used a culture-based coinubcation assay to reflect the close interactions experienced by symbiotic cells in the host. These assays were optimized for V. fischeri and are based on similar coinubcation assays described previously (13, 16). Briefly, strains were transformed with stable plasmids expressing different fluorescent proteins and antibiotic resistance genes. These differentially labeled coinubcated strains could be spatially distinguished within a mixed colony and directly quantified by plating onto selective medium. Based on CFU measurements, each strain was present at equal abundance (1:1 ratio) at the start of the coinubcation. When strain ES114 was coinubcated 1:1 with each of our six V. fischeri isolates, we found there was a strain-specific difference in the ability of ES114 to grow in the presence of another strain (Fig. 1A). ES114 was able to grow in the presence of strains ABM004, EMG003, and FO-A003: both strains were visibly present in the mixed colony after 15 h. In contrast, ES114 was not visibly detectable after 15 h when coinubcated with strains EBS004, FO-A001, and FO-A002, suggesting that the growth of ES114 was inhibited. These findings suggest that certain light-organ isolates are capable of strain-specific competitive interactions, and that the interactions can be captured using an in vitro assay.

ES114 Is Killed in a Contact-Dependent Manner. We first hypothesized that these strain-specific competitive outcomes could be explained by differences in growth rates among our isolates. For example, if EBS004, FO-A001, and FO-A002 grow significantly faster than ES114, then these three strains could competitively exclude ES114 after only 15 h. To test this hypothesis, we quantified the growth rates of ES114 and the competitive strains (EBS004, FO-A001, and FO-A002) by measuring the CFUs of each strain when grown under the same conditions used in the coinubcation assays (SI Appendix, Fig. S1A). The doubling time of ES114 (39 ± 5.8 min per generation) was not significantly different (Student’s t test, P > 0.05) from those of the three competitive strains: EBS004 (36 ± 4.7 min per generation),
FQ-A001 (43 ± 7.6 min per generation), and FQ-A002 (43 ± 10 min per generation), suggesting their ability to outcompete ES114 in our coinubcation assay is not due to differences in growth rate.

To determine whether the competitive strains outcompete ES114 through growth inhibition or physical elimination of ES114 cells, we measured the change in ES114 abundance in coinubcations with our V. fischeri isolates using two methods: measuring the total CFUs of ES114 remaining after 5 h and direct cell counts by flow cytometry. When ES114 was coinubcated with ABM004, EMG003, or FQ-A003, the CFUs of all strains increased after 5 h (Fig. 1B) and were uniformly mixed in colonies after 15 h (Fig. L4), suggesting these strains were able to coexist. However, when ES114 was coinubcated with the three competitive strains, CFU counts for all three competitor strains increased after 5 h, but ES114 CFU counts decreased by ∼3 logs (Fig. 1B). The decrease in ES114 CFUs was confirmed by direct cell counts of GFP-tagged ES114 using a flow cytometer, which showed that >90% of ES114 cells are eliminated after a 5-h coinubcation with competitive strains (SI Appendix, Fig. S1B).

To determine whether the ability of competitive strains to eliminate ES114 is dependent on direct cell–cell contact, or is mediated by a diffusible antimicrobial compound, we performed coinubcations in which a 0.22-μm filter was placed between ES114 and the coinubcated strain. This filter allows for diffusion of antimicrobial molecules, but prevents direct cell–cell contact between strains. With the filter, ES114 was able to grow and comprised at least 50% of the cells after 5 h (SI Appendix, Fig. S1C), suggesting that the filter protected ES114 cells from being killed by the competitive strains. As a positive control for a diffusible antimicrobial, an antibiotic was used in a parallel experiment. In this case, ES114 was killed (SI Appendix, Fig. S1D), showing that the filter did not protect ES114 from killing by a diffusible compound. Taken together, these findings indicate that some V. fischeri strains are capable of contact-dependent killing of ES114. We term isolates with the ability to kill ES114 as “lethal” strains and isolates that did not inhibit the growth of ES114 as “nonlethal.”

The Ability to Kill Is Independent of Strain Phylogeny. Given that our initial assays included only six light-organ isolates, we wondered how prevalent lethal and nonlethal strains are among V. fischeri isolates and if there was any correlation between phylogeny and killing ability. To determine the relative abundance of lethal and nonlethal strains, we repeated the coinubcation assay with ES114 and 32 other differentially tagged V. fischeri strains, isolated primarily from light-organ symbioses, and tested whether they could prevent growth of ES114 in a manner similar to what was observed for lethal strains in Fig. L4. Because some of these strains have slower growth rates than ES114, we modified the assay so that the ES114 target was outnumbered 5:1 by the coinubcated strain. Mixed colonies were imaged for fluorescence after ∼24 h to score for the presence or absence of ES114 (SI Appendix, Fig. S2), which would indicate whether the coinubcated strain is nonlethal or lethal, respectively, as defined above. The presence of the coinubcated strain was also confirmed by fluorescence (SI Appendix, Fig. S2). This lethal or nonlethal behavior was then mapped to a consensus phylogenetic tree built using four concatenated housekeeping genes (60). Of the 32 V. fischeri strains tested, 16 isolates are lethal (little to no ES114 observed) and 16 isolates are nonlethal (ES114 observed) (SI Appendix, Fig. S2). Moreover, the ability to kill was not correlated with phylogeny: lethal strains (black circles in Fig. 2) are found throughout the tree and do not comprise a monophyletic group, suggesting the killing mechanism is not a trait shared only among closely related strains. Finally, when we considered strains coisolated from four different E. scolopes light organs (purple, yellow, blue, and orange in Fig. 2), each set of coisolates consisted of both lethal and nonlethal strains. Taken together, these findings yielded three important observations: (i) both lethal and nonlethal strains are prevalent, (ii) killing does not correlate with phylogeny, and (iii) strains coisolated from the same light organ are a mix of lethal and nonlethal strains.

Whole-Genome Comparison Reveals a Correlation Between Killing and T6SS2. To identify the genetic determinants of interstrain killing, we performed comparative genomics using the complete genomes of two isolates: a nonlethal strain (ES114) and a lethal strain (MJ11) (64–66). We searched the MJ11 genome for genes encoding putative contact-dependent killing mechanisms that are not present in the ES114 genome and found strain-specific differences in T6SS genes.

Like most vibrios, V. fischeri has two chromosomal, and although both of the genomes examined encode a predicted T6SS on chromosome I (T6SS1), strain MJ11 encodes a second predicted T6SS on chromosome II (T6SS2) that is absent in ES114 (Fig. 3 and SI Appendix, Fig. S3). T6SS1 and T6SS2 gene clusters show very different genetic organization and essential T6SS structural proteins share low levels of homology: IcmF_1 and IcmF_2 are 24% identical, and Vasa_1 and Vasa_2 are 34% identical (Fig. 3 and SI Appendix, Table S1). Moreover, when the presence of IcmF_1 and IcmF_2 homologs was mapped onto a phylogenetic tree consisting of multiple Vibrio species, IcmF_1 homologs were restricted to a monophyletic group consisting of V. fischeri isolates and closely related species, while IcmF_2 homologs were found throughout the tree (SI Appendix, Fig. S4). Similar to other organisms encoding multiple T6SSs (67–69), these findings suggest the two T6SS systems in V. fischeri are likely not the result of a recent duplication event. The MJ11 T6SS2 encodes all necessary components for a functional secretion apparatus, as well as three auxiliary gene clusters encoding putative effector/immunity genes (Fig. 3B and SI Appendix, Tables S1 and S3). Further examination of the T6SS2-encoding gene cluster in the MJ11 genome revealed two hallmarks of genomic islands: the 50-kb region is located next to
a tRNA gene (SI Appendix, Table S1) and is flanked by 16-bp direct repeat sequences (Fig. 3B) (70). ES114 contains a single copy of the 16-bp repeat sequence between the conserved flanking genes (Fig. 3B), and also encodes two of the three auxiliary gene clusters found in MJ11 (SI Appendix, Table S3). Moreover, the T6SS2 gene cluster appears to be broadly conserved among other *Vibrio* species that have been isolated from marine and human hosts (SI Appendix, Fig. S4 and Table S2). Taken together, these findings suggest strain MJ11 contains genomic features that may facilitate horizontal transfer and/or loss of the genomic island. Therefore, we were interested in assaying additional strains for presence or absence of the island.

To determine whether other *V. fischeri* strains harbor this genomic island, we performed PCR using primers specific to two essential T6SS2 genes, *icmF-*2 and *vasA-2* (27), as well as the left junction of this genomic island. We obtained products for *icmF-*2, *vasA-2*, and the left junction for all three lethal strains (EBS004, FO-A001, and FO-A002), as well as MJ11, which served as a positive control (SI Appendix, Fig. SS4). In contrast, products from the three nonlethal strains (ABM004, EMG003, and FO-A003), as well as ES114, were only observed when using primers for the *recA* control (SI Appendix, Fig. SS4), suggesting the T6SS2-encoding genomic island is not present in these strains. When we screened the remaining isolates from Fig. 2 using the left junction primers, we did not observe PCR products for most of the nonlethal strains (Fig. 2, white circles, and SI Appendix, Fig. SSB); however, we did observe left junction PCR products for a few nonlethal strains (ES213, MB1382, ANM004, H905), suggesting some strains may encode the genomic island but their T6SS2 is not functional. We observed PCR products for the left junction for all lethal isolates (SI Appendix, Fig. SS8), suggesting the presence of the genomic island in these strains. Moreover, we observed products using primers specific to *recA* and *vasA-1* for all strains (SI Appendix, Fig. SS5). Taken together, these results indicate that although all strains appear to encode T6SS1, the ability to kill is correlated with the presence of genes associated with the T6SS2-encoding genomic island.

**Killing Is Dependent on T6SS2 Function.** To directly test if either T6SS is required for interstrain killing, we introduced a disruption mutation into the *vasA-1* or *vasA-2* gene in the lethal strain, FO-A001. The *vasA* gene is predicted to encode a homolog of TssF, which is an essential inner membrane protein of the structural machinery of T6SSs that is required for sheath assembly in *V. cholerae* and *Serratia marcescens*; disruption of this gene has previously been shown to eliminate T6SS function in these bacteria (71, 72). We next conducted coincubation assays with ES114 and the wild-type, *vasA-1*, or *vasA-2* mutant strains of FO-A001. When incubated with either wild-type FO-A001 or the *vasA-1* mutant, ES114 was killed by 5 h and not visibly detected at 15 h (Fig. 4A). In contrast, when incubated with the *vasA-2* mutant, ES114 was able to grow after 5 h and was visibly detectable in the mixed colony after 15 h (Fig. 4A). When *vasA-2* was complemented in trans using an isopropyl-β-D-thiogalactopyranoside (IPTG)-inducible expression vector, killing was restored in an IPTG-dependent manner (Fig. 4A).

To determine whether the T6SS2 genes encode the proteins required for constructing a T6SS sheath, we next constructed a VipA-2-GFP expression vector to visualize possible T6SS sheaths in wild-type and mutant FO-A001 strains. Previous work has shown that tagging the VipA subunit of a T6SS sheath allows for direct visualization of sheath assembly in *V. cholerae* (73, 74). We moved the VipA-2-GFP expression vector into wild-type, *vasA-1*, and *vasA-2* mutants of FO-A001 and used single-cell fluorescence microscopy to visualize sheath assembly. We observed GFP-tagged sheaths for the VipA-2 fusion in the wild-type and *vasA-1* mutant, but not in the *vasA-2* mutant (Fig. 4B), suggesting that T6SS2 encodes the necessary components to assemble a sheath, which is dependent on the baseplate component VsaA-2.

To determine whether T6SS2 is required for killing in the other coisolated lethal strains, we made *vasA-2* mutations in strains FO-A002 and EBS004. Disruption of *vasA-2* also abrogated killing of ES114 in these strains (Fig. 4C). Moreover, the *vasA-2* mutant strains were able to coexist with their wild-type parent, suggesting the *vasA-2* mutant is immune to the parental strain’s T6SS2-delivered effectors (SI Appendix, Fig. S1E). These findings suggest that interstrain killing among light-organ isolates is mediated by the T6SS on chromosome II.

**Coisolated Strains Are Incompatible.** The findings from our coinoculation assays with ES114 indicate lethal strains can eliminate a nonlethal strain; however, both lethal and nonlethal strains are commonly isolated from the same light organ (Fig. 2). Given this finding, we wondered how coisolated strains might be able to cohabitate the same host. Because light-organ colonization can only occur within a window of time after the animal hatches (75), we considered two alternative hypotheses: (i) the natural host colonization process selects for compatible strains that are immune to each other’s T6SS2 effectors or (ii) coisolated strains are not compatible and their ability to cohabitate the same light organ is dependent on physical separation within the host.

To test whether naturally coisolated strains are compatible, we performed pairwise coinoculation assays using both sets of squid isolates. For squid 1 isolates, we found the lethal strain EBS004 kills both nonlethal coisolates (ABM004 and EMG003) in a T6SS2-dependent manner after 5 h, resulting in no visibly detectable nonlethal strain after 24 h (Fig. S4). When the two nonlethal isolates were coincubated, each strain was able to grow after 5 h, resulting in a well-mixed colony after 24 h (Fig. S4). Similar results were obtained for squid 2 isolates; both lethal strains (FO-A001 and FO-A002) eliminate the nonlethal strain (FO-A003) in a T6SS2-dependent manner (Fig. 5B). These results suggest that nonlethal strains lack the appropriate immunity genes required to coexist with their coisolated lethal strains.

**Fig. 3.** T6SS2 is located on a strain-specific genomic island. T6SSs located on (A) chromosome I and (B) chromosome II of *V. fischeri* strains: MJ11 (lethal), ES114 (nonlethal), both MJ11 and ES114 encode putative T6SS genes on chromosome I (T6SS1). MJ11 carries a 50-kb gene cluster encoding predicted T6SS genes on chromosome II (T6SS2) that is lacking in ES114 (B). Conserved T6SS genes are the same color in T6SS1 and T6SS2 and genes of unknown function are indicated in white. Orange dashed lines indicate a 16-bp sequence (GTTTAAAAAAGCAG/CAA) flanking the MJ11 genomic island and corresponding sequence location in ES114. Black arrows indicate conserved genes flanking the MJ11 genomic island.
When the two lethal strains from squid 2 were coincubated, they maintained a 1:1 ratio and did not increase in CFUs after 5 h, and they were both observed in the mixed colony after 24 h (Fig. 5C). When the FQ-A001 vasA\_2 mutant was coincubated with wild-type FQ-A002, or if wild-type FQ-A001 was coincubated with the FQ-A002 vasA\_2 mutant, the wild-type strain always eliminated the vasA\_2 mutant (Fig. 5C). However, when the two vasA\_2 mutants were coincubated, these strains maintained a 1:1 ratio and increased in CFUs after 5 h and were both visibly detectable after 24 h (Fig. 5C), consistent with what is observed when coincubating nonlethal strains (Fig. 5A). These results suggest that FQ-A001 and FQ-A002 encode unique T6SS2-exported effectors that allow for T6SS2-mediated elimination of the other strain when it lacks a functioning T6SS2 for defense. Moreover, these data suggest that when coincubating strains are incompatible lethal strains, they actively kill one another, which can initially restrict their growth (Fig. 5C).

Strain Compatibility Is only Partly Predicted by T6SS Toxin Genotype.

Next, we expanded our search for compatible strains by examining the predicted toxin/immunity pairs encoded in our strain collection for isolates with available draft genomes. Because MJ11 is the only lethal V. fischeri strain with a completely sequenced genome, we searched draft genomes of strains ES213, MB11B1, MB13B2, MB13B3, MB13B1, KB4B5, MB15A4, ES114, and SR5 for homologs of the putative T6SS toxins found in the MJ11 genome (SI Appendix, Fig. S6). Previous work in V. cholerae indicates that strain compatibility can be inferred based on predicted T6SS toxin/immunity genes: strains that share the same toxin/immunity genes are compatible and those that have different toxin repertoires are not compatible (47). Moreover, recent work has shown that the C-terminal toxin/immunity genes can diversify via allelic exchange, resulting in varying alleles of toxins among closely related strains (76–78).

The MJ11 genome encodes putative T6SS toxins in the primary T6SS2 gene cluster and has one predicted toxin in each of three auxiliary gene clusters (Fig. 3B and SI Appendix, Table S3). We compared the sequences for these toxins across the available 10 draft genomes and found two alleles for auxiliary toxin 1 (A1), two alleles for auxiliary toxin 2 (A2), three alleles for auxiliary toxin 3 (A3), and one allele for the predicted toxin associated with the primary T6SS2 gene cluster and has one predicted toxin in each of three auxiliary gene clusters (Fig. 3B and SI Appendix, Table S3). We compared the sequences for these toxins across the available 10 draft genomes and found two alleles for auxiliary toxin 1 (A1), two alleles for auxiliary toxin 2 (A2), three alleles for auxiliary toxin 3 (A3), and one allele for the predicted toxin associated with the primary T6SS2 gene cluster (P) (SI Appendix, Fig. S6A). In addition to diverse toxin alleles, some strains were missing certain gene clusters altogether. Based on the toxin alleles we...
were able to divide these 10 strains into 6 predicted compatibility groups (SI Appendix, Fig. S6B). Only two groups contained more than one strain: group 1 (ES213, MB11B1, MB13B2, and MB13B3) and group 3 (KB4B5 and MB15A4).

We next performed pairwise coinoculation assays to test whether any strains within these groups were in fact compatible. For group 1, we found that only two sets of strains were able to coexist in our assays: MB13B2 and ES213 (both nonlethal strains) and MB11B1 and MB13B3 (both lethal strains) (SI Appendix, Fig. S6C). Although strains MB11B1 and MB13B3 can kill other group 1 nonlethal strains ES213 and MB13B2, they were able to coexist with one another even if the other strain was lacking a functional T6SS2, suggesting both MB11B1 and MB13B3 contain the same toxin/immunity genes and are indeed compatible (SI Appendix, Fig. S6C). This result is perhaps not surprising given that these two strains are closely related, according to our phylogenetic tree (Fig. 2). For group 3 strains, both MB15A4 and KB4B5 are lethal; however, when the two are coinoculated, MB15A4 eliminates KB4B5, indicating these strains are not compatible. Taken together, these results suggest that predicted T6SS toxin genotypes can only partly predict strain compatibility among V. fischeri strains, and strains appear to diversify their toxins rapidly and are largely incompatible, except for nonlethal strains.

Lethal Strains Spatially Separate in a T6SS2-Dependent Manner. Our initial characterization of strain compatibility indicates that although nonlethal strains can coexist, competitive outcomes of two lethal strains include: (i) one strain killing the other (MB15A4 vs. KB4B5), (ii) stable coexistence of compatible killers (MB11B1 and MB13B3), or (iii) coexistence by mutual killing (FQ-A001 vs. FQ-A002). Because our initial coinoculations with FQ-A001 and FQ-A002 resulted in coexistence that was T6SS2-dependent, we more closely examined the mixed colonies after 24 h. Interestingly, we found these lethal strains coexisted as spatially separated microcolonies that were T6SS2-dependent (Fig. 6 A, i and iii). Although this mechanism of strain separation has been observed by others (31, 79, 80), it is not yet known if this spatial segregation is stable or if it can be destabilized by ecologically relevant conditions.

We next considered whether physical disruption impacts the spatial separation of incompatible lethal strains observed in vitro. Such physical disruption is expected to occur within the light organ, which experiences daily venting of the crypt spaces: a light cue at dawn stimulates the light organ to contract and forcefully vent ~90–95% of the crypt contents through the pores (81, 82). In addition to diluting the bacterial cell numbers, contraction of the light organ to force out its contents could also act to destabilize the spatial structure of symbiotic cells. We wondered whether incompatible lethal strains could continue to coexist as spatially separated microcolonies in our in vitro assay if physical disruption was introduced. We predicted two possible outcomes: (i) both lethal strains would be present after disruption and would again be spatially separated in a T6SS2-dependent manner, or (ii) disruption of the initial spatial separation of strains would allow one lethal strain to dominate and eliminate the other.

To test these predictions, we coinoculated the two lethal strains FQ-A001 and FQ-A002 and physically disrupted the cells after the initial coinoculation. After the initial 24-h coinoculation, both strains grew equally and remained at a 1:1 ratio (Fig. 6B) and spatially separated in a T6SS2-dependent manner (Fig. 6D). We then resuspended these coinoculation spots in 1 mL broth, physically mixed the cells by pipetting up and down, and spotted 10 μL of this cell suspension onto agar plates. When cells were spotted after this mixing step, both strains were present in a 1:1 ratio, yet after an additional 24-h incubation, FQ-A002 was consistently able to eliminate FQ-A001 while the two T6SS2 mutant strains grew equally well and were spatially well-mixed (Fig. 6). Taken together, these findings suggest that lethal coisolated strains do not intermix under dynamic conditions: although T6SS2-mediated killing can initially spatially separate competing strains, this coexistence is not stable in the presence of physical disruption.

T6SS2 Is Necessary for Strain Incompatibility Within the Host. Our in vitro assay indicates that T6SS2 mediates strain incompatibility among light-organ isolates. Specifically, lethal strains (FQ-A001) are able to kill nonlethal strains (ES114) in a T6SS2-dependent
manner. Moreover, previous work has shown that squids exposed to an inoculum of FQ-A001 and ES114 do not exhibit crypt spaces colonized with both strain types (61). To investigate the impact of T6SS2 on the intercellular interactions that *V. fischeri* cells experience in vivo, we turned to the colonization model involving the light organ of *E. scolopes*. The nascent light organ features six crypt spaces that are each independently colonized by *V. fischeri* cells in direct contact with each other (58). Each crypt space has a physical bottleneck that permits entry of only one to two cells at a time, which proliferate into the resulting symbiotic population (53). By labeling two strains with different fluorescent proteins, it is possible to use confocal fluorescence microscopy to identify the strain types of each colonized crypt (Fig. 7 A–C) (61, 83). For example, in the light organ shown in Fig. 3A, which was colonized with either YFP-tagged ES114 and either YFP- or CFP-labeled FQ-A001, the crypts colonized with ES114 and either YFP, CFP, or both YFP and CFP were counted. When the frequency of cocolonized crypts was calculated, the cells of each strain type were mixed throughout the crypt space (Fig. 7F), showing that ES114 and the FQ-A001–derived cells are in direct contact within colonized crypts. In contrast, none of the colonized crypts resulting from animals exposed to wild-type FQ-A001 and ES114 were CFP+ YFP+ (Fig. 7 E and G), which was consistent with our previous report (61). This effect of T6SS2 on crypt colonization frequencies was statistically significant according to a two-proportion z-test (SI Appendix, Table S5). To ensure that our approach was sensitive enough to detect significant differences in crypt colonization frequencies, we performed a power analysis that estimated the effect size to be greater than 0.37 (two-proportion z-test, α = 0.05, power = 0.80) (SI Appendix, Table S5), suggesting our crypt colonization assay sampled a sufficient number of colonization sites (n > 107 per treatment, per trial) to statistically support the conclusion that VasA_2, and thus T6SS2, prevents the establishment of crypts colonized with FQ-A001 and ES114. Taken together, these data suggest that T6SS2 promotes the separation of incompatible strains within the natural host.

**Discussion**

This work demonstrates that the *Vibrio*–squid symbiosis is a valuable system for investigating intraspecific bacterial interactions and dynamics as well as their impact on host colonization. Our evidence suggests these interactions are relevant in a natural host, where they are capable of shaping the spatial distributions of host-associated microbial populations. This study shows that the T6SS prevents coexistence of strains in the context of a natural, beneficial bacteria–host interaction. Moreover, these interactions can be recapitulated using culture-based assays, which provide a powerful system to probe molecular mechanisms of interbacterial competition and their ecological and evolutionary roles.

Based on our results, we propose a model for the role of T6SS2 during host colonization. Juvenile squid hatch without their bacterial symbionts, and their crypts initially are colonized by one to two cells of different genotypes. During their growth, the initial populations within colonized crypt spaces come into contact with each other. In this context, our data suggest that T6SS2 strains begin to kill competitor cells in a contact-dependent manner. By 44-h postinoculation, T6SS2-deficient cells within these initially colonized crypts have been eliminated, resulting in crypts colonized exclusively by the T6SS2+ genotype.

Further investigation should examine the details of when and where competitors use T6SS2 during the first 44 h of host colonization, as well as the role of the host in influencing these interactions. Although this work did not directly test the impact of venting on T6SS2-mediated competition within crypts, our in vitro data suggest that T6SS2 strains begin to kill competitor cells in a contact-dependent manner. By 44-h postinoculation, T6SS2-deficient cells within these initially colonized crypts have been eliminated, resulting in crypts colonized exclusively by the T6SS2+ genotype.

Because this work focused on T6SS2, the functional role of T6SS1 remains unclear. Although we did not observe vasA_1-dependent killing of ES114 (Fig. 4A), T6SS1 may be active under different conditions compared with T6SS2. Indeed, other *Vibrio* species show T6SS activity that is conditionally regulated (13, 33, 84), and future work will need to explore how free-living versus host-like conditions modulate *V. fischeri* T6SS gene expression and function. Alternatively, T6SS1 may be used to interact with eukaryotic cells (15, 28, 72). In a previous transposon sequencing study during squid colonization, the genes in the predicted T6SS1 locus contained abundant transposon insertions (i.e., are not essential); however, a role for T6SS1 in host colonization was not apparent from that work (85).

Our observations also raise many questions from an evolutionary perspective. The distribution of killing activity and the T6SS2-encoding genomic island, and if its presence influences survival in both the host and greater marine environments.

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One model for the evolutionary dynamics of the T6SS2-containing island is that it is frequently transferred among strains by horizontal gene transfer, and the noted hallmarks of flanking repeats and tRNA locus support such a model. Given the presence of the \textit{V. fischeri} T6SS2 homologs in closely related \textit{Vibrio} spp. (SI Appendix, Fig. S4), an alternate model may be that the locus was present in a common ancestor of \textit{V. fischeri} and closely related Vibrios, and has since been lost among \textit{V. fischeri} strains in environments where interbacterial killing is no longer advantageous. Additional work is required to evaluate these options, and pursuing these ideas may provide evidence as to how selective pressure on the T6SS influences \textit{V. fischeri} evolution.

If \textit{V. fischeri} strains use T6SS2 to eliminate competitor populations when the symbiosis is first established, it is possible that once a crypt becomes clonally colonized, the T6SS2 no longer provides the population with an advantage. Moreover, maintaining a 50-kb genomic island could be a fitness cost to the cell, in addition to the energetic costs associated with synthesizing and using the T6SS apparatus (86). Given the presence of direct repeat sequences flanking the genomic island, it is possible that the island could be lost by excision through homologous recombination, leaving behind a single copy of this sequence as observed in strain ES114 (Fig. 3B). Because a functional T6SS is not needed for immunity to T6SS-exported effectors of isogenic strains (SI Appendix, Fig. S1E), which are encoded in auxiliary gene clusters elsewhere on the chromosome (SI Appendix, Table S3), loss of T6SS2 would not be detrimental to the cell’s survival. Such a scenario could explain the consistent mix of lethal and nonlethal strains isolated among naturally cooccurring light-organ symbionts, as well as the presence of auxiliary gene clusters in nonlethal strains lacking the T6SS2-encoding genomic island (ES114, MB13B1, SR5) (SI Appendix, Fig. S6). Further
Materials and Methods

The laboratory practices were carried out under the general principles described in the Guide for the Care and Use of Laboratory Animals (89) in communication with the Institutional Animal Care and Use Committee Office at Pennsylvania State University. See SI Appendix for additional experimental details, including media and growth conditions, isolation of symbiotic V. fischeri, strain and plasmid construction, contact-dependence assays, fluorescence microscopy, and phylogenetic analysis. Bacterial strains, plasmids, and oligonucleotides used in this study are in SI Appendix, Tables S6 and S7.

Coincubation Assays. V. fischeri strains were grown overnight on LBS agar plates supplemented with the appropriate antibiotic at 24 °C. Cells were scraped from agar surfaces, resuspended in LBS medium, and diluted to an OD₆₀₀ of 1.0. For each coincubation, strains were mixed in a 1:1 ratio and 10 μL of the mixture was spotted on LBS agar plates and incubated at 24 °C. After 5–24 h, each coincubation was resuspended in 1 mL LBS medium. Strains were quantified by plating serial dilutions onto LBS plates supplemented with antibiotics selective for each strain.

Fluorescence Metrscopy. For coincubation assays, fluorescence microscopy images were taken with a stereo microscope equipped with a Nipkow fluorescence adapter kit for green and red fluorescence detection. Images were taken using an OMAX 14MP camera with ViewPoint software and color changes were made by adjusting the HLS color mode. No brightness or contrast adjustments were made. For high-magnification images of cocultivated bacteria (64), spots were imaged using an Olympus BX51 microscope outfitted with a Hamamatsu ORCA RC camera and either a 4×0.13 UPlanFLN or 10×0.3 UPlanFLN objective lens. Images were captured using Improvision’s Velocity software. To visualize GFP-tagged T6SS sheath formation, strains carrying the IPTG-inducible vipA_2-gfp fusion expression vector were spotted onto a thin pad of 2% LBS agar with 0.5 mM IPTG and imaged using an Olympus BX51 microscope outfitted with a Hamamatsu C8484-03G01 camera and a 100×1.30 Oil SD apochromatic objective. Images were captured using MetaMorph software. Contrast on images was adjusted uniformly across images by subtracting background using imageJ software.

Phylogenetic Analysis. A multilocus phylogenetic analysis was performed using partial sequences of four loci: recA, mdh, katA, and pyrC. Published sequence data and newly amplified sequences of 35 total Vibrio isolates were collected, combined into a single concatenated sequence (ordered recA mdh katA pyrC − → − → 2,880 nucleotides), and aligned with ClustalX 2.1 (90). The concatenated sequence alignment was analyzed by ModelTest 2.1 v20160303 (53) via three information criteria methods (Akaike, Bayesian, and Decision Theory). Construction of the majority-rule consensus tree and statistical analysis of clade membership/presence was assessed by sampling an “appropriately stationary” posterior probability distribution (91). Sequences associated with this analysis were submitted to the GenBank database and their accession numbers are listed in SI Appendix, Table S6.

Squid Colonization Assays. For each treatment, 24–30 freshly hatched juvenile squid were exposed to the inoculum containing an even mix of YFP- and CFP-labeled strains at a final concentration ranging from 1,200–4,760 CFU/mL. Squid were exposed to this mixed inoculum for 20 h and then washed in fresh filter-sterilized seawater. After 44 h, animals were fixed, washed, and prepared for fluorescence microscopy by dissecting the ventral side of the mantle to reveal the light organ. YFP and CFP images were taken using a Zeiss 780 confocal microscope equipped with a 10× or 40× water lens. Each crypt space was scored separately for CFP and YFP fluorescence.

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**Figure S1.** Competitive *V. fischeri* strains eliminate ES114 in a contact-dependent manner. (A) Growth curves of ES114 and lethal squid 1 (EBS) and squid 2 (FQ1 and FQ2) isolates; growth rates calculated with data from 2 to 4 h for all strains. (B) Flow cytometry cell count data of GFP-tagged ES114 after 5 h co-incubation with lethal squid isolates. Dashed line indicates average ES114 CFU at 0 h. White circles indicate cell counts at or under the limit of detection (7140 cells) and asterisks indicate statistical difference for ES114 co-incubated with other strains compared to with itself at 5 h using a student’s t-test (p<0.01). (C) CFU counts for each co-incubation spot for co-incubations of ES114 (blue) with lethal squid isolates (green) where strains were physically separated by a 0.22 µm filter preventing direct cell-cell contact but still allowing diffusion of molecules. (D) CFU counts for each co-incubation spot for co-incubations of ES114 at 0 h (light gray) and after 24 h (dark gray) when incubated with water or kanamycin separated by a 0.22 µm filter. Fluorescence microscopy images were taken at 24 h; scale bar = 2 mm. Asterisk indicates statistical difference for ES114 incubated in water compared to kanamycin using a students t-test (p=0.0004). The dashed line indicates the limit of detection (200 CFUs) for the assay. (E) CFU counts for each co-incubation spot for co-incubations of lethal wild-type strains (blue) with the vasA-2 mutant derivative strain. Error bars indicate the standard error of the mean for biological replicates. Each experiment was performed at least three times and either combined data are shown (B and D, n=12) or a representative experiment is shown (A, n=1; C and E, n=4).
**Figure S2.** Co-incubations between *V. fischeri* isolates and ES114. Fluorescent microscopy images of co-incubations of GFP-tagged ES114 (blue) with 32 RFP-tagged *V. fischeri* isolates (green) taken at 24 h. Strains were mixed in a 1:5 ratio outnumbering ES114. Scale bar = 2 mm. RFP-tagged co-incubated strains are listed below the image pair. If ES114 (blue) is observed and not inhibited (ex. ES213), then the co-incubated strain is designated as non-lethal. If ES114 is not observed (ex. with ES12 or MB11B1), or if ES114 is only observed as outgrowth of survivors around the colony edge (ex: with MB13B3) then the co-incubated strain is designated as lethal.
**Figure S3.** Comparison of chromosome II for ES114 and MJ11. Geneious R8 software was used to align the genomes of ES114 and MJ11 and visualized using Mauve. (A) Alignment of chromosome II sequences for ES114 (NC_006841.2) and MJ11 (NC_011186.1). Regions with high identity (yellow) indicate conserved genes and low identity (white) are strain specific. (B) Enlargement of T6SS2-encoding genomic island that is absent in ES114 but present in MJ11 shows it is near a tRNA gene (asterisk).
Figure S4. T6SS2 is broadly conserved among Vibrio species. Hsp60 percent identity tree for fully-sequenced representative genomes of Vibrio species. Filled boxes indicate presence of T6SS1 or T6SS2 IcmF homolog based on >60% identity (Table S2). Genomes were also examined for conserved genetic structure of T6SS. Asterisks indicate species found associated with an animal host. Note this figure only identifies homologs of T6SS1 and T6SS2 from V. fischeri and does not include other more distantly related T6SSs that may be in these representative genomes.
Figure S5

**Figure S5. Distribution of the T6SS1 and T6SS2 among V. fischeri isolates.** Strains were screened for the presence/absence of structural T6SS2 genes (*icmF_2* and *vasA_2*), the genomic island’s left junction using primers specific to the left flanking gene and the first gene encoded in the genomic island, a structural T6SS1 gene *vasA_1*, and housekeeping gene *recA*, which is present in all strains. Asterisks indicate lethal strains.
Auxiliary gene cluster 1:

- Allele A genes share >93% ID and allele B genes share >99.9% seq identity. Allele A and Allele B sequences share ~60% sequence identity. The first 791 bp share >93% sequence identity and after 791 the two alleles share ~48% sequence identity. A Pfam search returned no significant predicted functional domains for either allele.

Auxiliary gene cluster 2:

- Allele B genes share >93% seq identity. Allele A and Allele B sequences share ~60% sequence identity. The first 155 bp share >94% sequence identity and after 155 bp the two alleles share ~58% sequence identity. A Pfam search returned a conserved LysM domain at the N terminus of both alleles (yellow). Homologs of auxiliary gene cluster 2 were not detected in ES114 or SR5.

Auxiliary gene cluster 3:

- Allele A sequences share >99.9% sequence identity. Allele B sequences are 100% identical and B* allele for ES114 is 93% identical to the other B alleles, however it is distinct and has a stop codon. Allele C sequences share 97% sequence identity. All three alleles have two DUF2235 domains that are uncharacterized alpha/beta hydrolase domains (boxes). The three alleles share a highly conserved N-terminus sequence (>94% ID), while the DUF2236 domains are least similar (78-85% ID) and a more conserved C-terminus domain (88-94% ID). Homologs of auxiliary gene cluster 3 were not detected in SR5.

Primary T6SS2 operon:

- All sequences share >96% identity. No significant predicted functional domains were identified from a Pfam search. The T6SS2 operon was not detected in ES114, SR5, or MB13B1.

**Figure S6**

### Predicted Compatibility Table

| Strain | Group | A1 | A2 | A3 | P | killer |
|--------|-------|----|----|----|---|--------|
| ES213  | 1     | B  | B  | B  | B | A      | No      |
| MB11B1 | 1     | B  | B  | B  | A | Yes    |
| MB13B2 | 1     | B  | B  | B  | A | No      |
| MB13B3 | 1     | B  | B  | B  | A | Yes    |
| MB13B1 | 2     | A  | B  | A  | NP| No      |
| KB4B5  | 3     | A  | B  | C  | A | Yes    |
| MB15A4 | 3     | A  | B  | C  | A | Yes    |
| ES114  | 4     | A  | NP | B* | NP| No      |
| MJ11   | 5     | A  | A  | A  | A | Yes    |
| SR5    | 6     | A  | NP | NP | NP| No      |

NP indicates the gene cluster was not detected either bioinformatically or with PCR.
Figure S6. T6SS toxin genotypes of 10 *V. fischeri* isolates. (A) Putative T6SS toxin alleles of 10 *V. fischeri* strain draft genomes for auxiliary gene clusters 1 (A1), 2 (A2), 3 (A3) and the primary T6SS2 operon (P). Protein sequences from the *V. fischeri* MJ11 genome were used to perform a tblastn search for homologs which were aligned using clustal omega, and an average distance tree was built using jalview. (B) Predicted compatibility table showing 6 compatibility groups based on toxin alleles; NP (not present) indicates absence of a gene cluster. (C) Fluorescence microscopy images for pairwise co-incubations of Group 1 and Group 3 strains taken at 24 h; “-” indicates *vasA*₂ mutants; scale bar is 2 mm.
Figure S7. **Inoculum size correlates with co-colonized animals.** (A) Scatter plot showing calculated frequency of co-colonized squid light organs for animals exposed to ES114 differentially expressing CFP or YFP (ESvES) and animals exposed to FQA001 differentially expressing CFP or YFP (FQAvFQA) at various inoculum sizes. The frequencies of co-colonized animals were determined by dividing the number of animals with both YFP- and CFP-positive infections by the total number of animals in the group. (B) Scatter plot showing the calculated frequency of co-colonized crypts. Proportions of co-colonized crypts were determined by dividing the number of crypts that were positive for both CFP and YFP by the total number of crypts that were CFP positive. (C) Number of crypts that were scored as CFP only, YFP only, or CFP+YFP for competitions using CFP- and YFP-tagged ES114 (n = 29 animals) or CFP- and YFP-tagged FQ-A001 (n = 22 animals) at indicated inoculation sizes. The proportion of CFP+ YFP+ crypts between the different competitions were compared using a two-proportion z-test and an asterisk indicates $p < 0.001$. 
# Table S1. Type VI Secretion System 2 Genes in MJ11 Genome

| VFMJ11# | Vas name | Other name | Predicted Function | % AA identity between T6SS1 and T6SS2 gene clusters |
|---------|----------|------------|--------------------|-----------------------------------------------|
| A0803   | Flanking Gene |           |                    |                                               |
| A0804   |           |           |                    |                                               |
| A0805   | VasC     | Impl, TagH*, | core component     | 24% to VFMJ11_1079                           |
| A0806   | VasD     | SciN*, TssJ*, EvpL* | (core component) | 28% to VFMJ11_1078                           |
| A0807   | VasE     | ImpV, SciO*, TssK*, EvpM* | (core component) | 34% to VFMJ11_1077                           |
| A0808   | VasF     | ImpK, TssL‡, IcmH‡, DotU‡, SciP*, EvpN* | (core component) | 30% to VFMJ11_1076                           |
| A0809   |           |           | Conserved hypothetical protein |                                               |
| A0810   |           |           | Hypothetical protein |                                               |
| A0811   |           |           | Conserved hypothetical protein |                                               |
| A0812   |           |           | Conserved hypothetical protein |                                               |
| A0813   |           |           | Hypothetical protein |                                               |
| A0814   |           |           | Putative lipoprotein |                                               |
| A0815   |           |           | M23 peptidase domain protein |                                               |
| A0816   | VasL     | ImpA, SciA*, EvpK* (((TssA) | ImpA-related N-terminal family protein (core component) | Absent in T6SS1 |
| A0817   | VasK/IcmF | ImpL, TssM‡, SciS*, EvpO* | Membrane transport protein … (core component) | 24% to VFMJ11_1075 |
| A0818   |           |           | Putative transcriptional regulator |                                               |
| A0819   | VasB     | ImpH, TssG*, AciB*, EmpG* | Baseplate (core component) | 33% to VFMJ11_1083 |
| A0820   | VasA     | ImpG, TssF‡, SciC*, EvpF* | Baseplate (core component) | 34% to VFMJ11_1084 |
| A0821   | VasS     |           | Lysozome-related protein (core component) | 37% to VFMJ11_1085 |
| A0822   | VipB     | ImpC, TssB‡ | Outer sheath (core component) | 67% to VFMJ11_1086 |
| A0823   | VipA     | ImpB, TssC‡ | Outer sheath (core component) | 57% to VFMJ11_1087 |
| A0824   | VasJ     | ImpA, DapB, SciA, EvpK | (core component) | 25% to VFMJ11_1088 |
| A0825   |           |           | Serine-threonine protein kinase |                                               |
| A0826   |           |           | Conserved hypothetical protein |                                               |
| A0827   |           |           | Conserved hypothetical protein |                                               |
| A0828   |           |           | Conserved hypothetical protein |                                               |
| A0829   |           |           | Conserved hypothetical protein |                                               |
| A0830   | VgrG     | TssI‡, VgrS* | Spiked tip (core component) | Absent in T6SS1 |
| A0831   | Hcp      | TssD‡, SciK*, SciM*, EvpC* | Inner tube (core component) | Absent in T6SS1 |
| A0832   | VasG     | ClpV, SciG*, TssH*, EvpH* | (core component) | 53% to VFMJ11_1082 |
| A0833   |           |           | Nitric Oxide reductase regulator |                                               |
| A0834 | Putative lipoprotein |
|-------|---------------------|
| A0835 | OmpA family protein |
| A0836 | Putative lipoprotein |
| A0837 | Conservative hypothetical protein |
| A0838 | Flanking Gene Chitodextrinase |
| A0839 | tRNA-Gly Aminoacyl-tRNA biosynthesis |

‡ Cianfanelli, F.R., Monlezun, L. and Coulthurst, S.J., 2016. Aim, load, fire: the type VI secretion system, a bacterial nanoweapon. *Trends in microbiology*, 24(1), pp.51-62.

*Cascales, E., 2008. The type VI secretion toolkit. *EMBO reports*, 9(8), pp.735-741.
### Supplemental Table S2. Distribution of \textit{V. fischeri} T6SS2-encoded proteins among \textit{Vibrio} spp.

| \textit{Vibrio} species | MJ11_A0817 homolog | % ID\textsuperscript{a} | MJ11_A0818 homolog | % ID\textsuperscript{a} | Ref.\textsuperscript{b} |
|-------------------------|---------------------|-----------------|---------------------|-----------------|----------------|
| \textit{V. fischeri}\textsuperscript{*} | ACH63505.1 | 100% | ACH63887.1 | 100% | (1) |
| \textit{V. logei}\textsuperscript{*} | WP_017020851.1 | 80% | WP_065611695.1 | 89% | (2) |
| \textit{V. wodanis}\textsuperscript{*} | WP_061004504.1 | 80% | WP_061004503.1 | 87% | (3) |
| \textit{V. proteolyticus}\textsuperscript{*} | WP_040902488.1 | 66% | GAD67006.1 | 78% | (4) |
| \textit{V. jasicida}\textsuperscript{*} | WP_038878560.1 | 64% | WP_045410168.1 | 56% | (5) |
| \textit{V. crassostreae}\textsuperscript{*} | WP_017064525.1 | 63% | OEE92195.1 | 76% | (6) |
| \textit{V. tasmaniensis}\textsuperscript{*} | WP_065104332.1 | 63% | WP_012600854.1 | 77% | (7) |
| \textit{V. splendidus}\textsuperscript{*} | WP_054543291.1 | 63% | KPL96954.1 | 76% | (8) |
| \textit{V. campbellii}\textsuperscript{*} | WP_005532464.1 | 63% | WP_005427005.1 | 74% | (9) |
| \textit{V. harveyi}\textsuperscript{*} | WP_045491053.1 | 63% | AIY08715.1 | 74% | (10) |
| \textit{V. alginolyticus}\textsuperscript{†} | WP_065645837.1 | 63% | EAS76637.1 | 73% | (8) |
| \textit{V. parahaemolyticus}\textsuperscript{†} | WP_053807876.1 | 63% | WP_005480668.1 | 72% | (11) |
| \textit{V. antiquarius} | WP_006741088.1 | 63% | EDN58559.1 | 73% | (12) |
| \textit{V. owensii}\textsuperscript{*} | WP_041052795.1 | 62% | KIF49344.1 | 74% | (13) |
| \textit{V. tubiashii}\textsuperscript{*} | WP_004749068.1 | 62% | WP_004743867.1 | 71% | (14) |
| \textit{V. azureus} | WP_021710015.1 | 61% | WP_021710014.1 | 71% | (15) |
| \textit{V. caribbeanicus}\textsuperscript{*} | WP_009603313.1 | 60% | WP_009603314.1 | 72% | (16) |
| \textit{V. coralliilyticus}\textsuperscript{*} | WP_045985165.1 | 58% | EEX32044.1 | 75% | (17) |
| \textit{V. neptunius}\textsuperscript{*} | WP_045976861.1 | 58% | WP_045976860.1 | 75% | (18) |

\textsuperscript{a}Percent identity based on BlastP results using VFMJ11_A0817 (IcmF_2) and VFMJ11_A0818 as sequence query.

\textsuperscript{b}Reference for host association or isolation

\textsuperscript{*} Indicates species is associated with a marine host

\textsuperscript{†} Indicates species is associated with human host

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Table S3. Type VI Secretion System Auxiliary Gene Clusters in MJ11 Genome

| VFMJ11# | Vas name       | Other name          | Auxiliary gene cluster          | % AA identity between MJ11 and ES114 |
|---------|----------------|---------------------|----------------------------------|-------------------------------------|
| 1309    | VgrG           | TssI‡, VgrS*        | Auxiliary cluster I              | 93% to VF_1229                      |
| 1310    | DUF4123 domain protein | Auxiliary cluster I        | 93% to VF_1230                      |
| 1311    | Conserved hypothetical | Auxiliary cluster I        | 94% to VF_1231                      |
| 1312    | Conserved hypothetical | Auxiliary cluster I        | 79% to VF_1232                      |
| 1313    | Conserved hypothetical | Auxiliary cluster I        | 77% to VF_1233                      |
| 1314    | Conserved hypothetical | Auxiliary cluster I        | 82% to VF_1232                      |
| 1315    | Conserved hypothetical | Auxiliary cluster I        | 87% to VF_1232                      |
| 1316    | Conserved hypothetical | Auxiliary cluster I        | 78% to VF_1232                      |
| 1495    | Hcp            | TssD‡, SciK*, SciM*, EvpC* | Auxiliary cluster II              | Absent in ES114                     |
| 1494    | VgrG           | TssI‡, VgrS*        | Auxiliary cluster II              | Absent in ES114                     |
| 1493    | DUF4123 domain protein | Auxiliary cluster II        | Absent in ES114                     |
| 1492    | LysM domain protein | Auxiliary cluster II        | Absent in ES114                     |
| A1070   | Hcp            | TssD‡, SciK*, SciM*, EvpC* | Auxiliary cluster III             | 100% to VF_A0954                    |
| A1069   | VgrG           | TssI‡, VgrS*        | Auxiliary cluster III              | 99% to VF_A0953                      |
| A1068   | DUF4123 domain protein | Auxiliary cluster III        | 97% to VF_A0952                      |
| A1067   | DUF2235 domain protein | Auxiliary cluster III        | 86% to VF_A0950                      |
| A1066   | DUF2931 domain protein | Auxiliary cluster III        | 72% to VF_A0945                      |
Table S4. Impact of T6SS2 on proportion of co-colonized animals

|                                | ES v FQ1 Trial 1 | ES v FQ1-Trial 1 | ES v FQ1-Trial 2 | ES v FQ1-Trial2 |
|--------------------------------|------------------|------------------|------------------|----------------|
| Co-colonized animals           | 24               | 20               | 24               | 22             |
| Total Squid                    | 30               | 24               | 26               | 27             |
| Proportion                     | 0.80             | 0.92             | 0.83             | 0.81           |
| P value by z test              | 0.107            |                  | 0.424            |                |
Table S5. Statistical analysis of co-colonized crypts

| Strains in inoculum | Trial 1 | Trial 2 |  | Strains in inoculum | Trial 1 | Trial 2 |
|---------------------|---------|---------|  | Strains in inoculum | Trial 1 | Trial 2 |
| CFP+ crypts         | 370     | 68      |  | CFP+ crypts         | 470     | 87      |
| CFP+YFP+ crypts     | 10      | 12      |  | CFP+YFP+ crypts     | 10      | 17      |
| Proportion          | 0.027   | 0.176   |  | Proportion          | 0.021   | 0.195   |
| p-value by z-test   | <0.001  | <0.001  |  | p-value by z-test   | <0.001  | <0.001  |
| Proportion needed for power = 0.8 | 0.105 | 0.099 |
| Effect size         | 0.369   | 0.4106  |  | Effect size         | 0.369   | 0.4106  |
| Strain | Collection Description | Geography          | Ecology                  | Source or Reference | NCBI GenBank Accession Numbers |
|--------|------------------------|--------------------|--------------------------|---------------------|-------------------------------|
| ABM004 | Oahu, HI, USA (Maunalua Bay) | Euprymna scolopes (squid light organ) | This study              | recA: MF076795, mdh: MF076808, katA: MF076821, pyrC: MF076834 |
| AGC005 | State College, PA, USA | Aquarium seawater containing *E. scolopes* collected from Maunalua Bay | "" | recA: MF076800, mdh: MF076813, katA: MF076826, pyrC: MF076839 |
| ANM004 | Oahu, HI, USA (Maunalua Bay) | *E. scolopes* (squid light organ) | "" | recA: MF076798, mdh: MF076811, katA: MF076824, pyrC: MF076837 |
| CG101 | Australia | Cleidopus gloriamaris (fish light organ) | (Lee 1994) | recA: HQ595306, mdh: EU907966, katA: EU907990, pyrC: JF509856 |
| CG103 | "" | "" | "" | recA: HQ595307, mdh: HQ595322, katA: HQ595331, pyrC: JF509855 |
| CHS319 | Oahu, HI, USA (Maunalua Bay) | *E. scolopes* | This study | recA: MF076801, mdh: MF076814, katA: MF076827, pyrC: MF076840 |
| EBS004 | "" | "" | "" | recA: MF076797, mdh: MF076810, katA: MF076823, pyrC: MF076836 |
| ECT001 | "" | "" | "" | recA: MF076804, mdh: MF076817, katA: MF076830, pyrC: MF076843 |
| EMG003 | "" | "" | "" | recA: MF076796, mdh: MF076809, katA: MF076822, pyrC: MF076835 |
| ES12 | Oahu, HI, USA (Kaneohe Bay) | "" | (Boettcher and Ruby 1994) | recA: HQ595309, mdh: HQ595323, katA: HQ595332, pyrC: JF509862 |
| ES114; ATCC 700601 | "" | "" | (Boettcher and Ruby 1990) | recA: VF_0535, mdh: VF_0276, katA: VF_A0009, pyrC: VF_A0412 |
| ES213 | Oahu, HI, USA (Maunalua Bay) | "" | (Boettcher and Ruby 1994) | recA: HQ595310, mdh: EU907971, katA: EU907995, pyrC: JF509863 |
| ES401 | "" | "" | (Lee 1994) | recA: HQ595311, mdh: HQ595324, katA: HQ595333, pyrC: JF509864 |
| ET101 | Victoria, Australia (Crib Point) Townsville, Australia | *Euprymna tasmanica* (squid light organ) | (Nishiguchi 2002) | recA: HQ595312, mdh: HQ595325, katA: HQ595334, pyrC: JF509865 |
| ET401 | "" | "" | (Nishiguchi 2002) | recA: HQ595313, mdh: HQ595326, katA: HQ595335, pyrC: JF509866 |
| Sample Code | Location | Organism | Organ Type | Sample Code | Accession Numbers |
|-------------|----------|----------|------------|-------------|-------------------|
| FQ-A001     | Oahu, HI, USA (Kaneohe Bay) | E. scolopes | squid light organ | This Study | KU756584, KU756585, KU756586, KU756587 |
| H905        |          | Planktonic |            |             | HQ595314, EU907972, EU907996, JF509867 |
| IRR001      |          | E. scolopes | squid light organ | This Study | MF076799, MF076812, MF076825, MF076838 |
| KB4B5       |          |          |            |             | JF509762, JF509787, JF509845, JF509873 |
| LFI1238     | Hammerfest, Norway | Gadus morhua | cod head kidney | (Hjerde et al 2008) | VSAL_I0634, VSAL_I0359, VSAL_II0215, VSAL_II0468 |
| MB11B1      | Oahu, HI, USA (Maunalua Bay) | E. scolopes | squid light organ | (Wollenberg and Ruby 2009) | JF509765, JF509789, JF509847, JF509876 |
| MB13B1      |          |          |            |             | JF509766, JF509790, JF509848, JF509877 |
| MB13B2      |          |          |            |             | JF509767, JF509791, JF509849, JF509878 |
| MB13B3      |          |          |            |             | JF509768, JF509792, JF509850, JF509879 |
| MB15A4      |          |          |            |             | JF509771, JF509793, JF509851, JF509882 |
| MJ1S        | Japan    | Monocentris japonicus | fish light organ | (Bose Stabb 2011?) | MF076792, MF076805, MF076818, MF076831 |
| MJ11        | Japan    | Monocentris japonicus | fish light organ | (Ruby and Nealson 1976) | VFMJ11_0538, VFMJ11_0264, VFMJ11_A0023, VFMJ11_A0452 |
| NAD004      | Oahu, HI, USA | E. scolopes |            | This Study | MF076802, MF076815, MF076828, MF076841 |
| ID   | Location                | Organism                  | Reference                  | Accession Numbers | Additional Notes |
|------|-------------------------|---------------------------|----------------------------|-------------------|------------------|
| PP3  | (Maunalua Bay) Oahu, HI, USA | (squid light organ) Planktonic | (Lee and Ruby 1992) | HQ595317          |                  |
|      | (Kaneohe Bay)           |                           |                            | HQ595329          |                  |
|      |                         |                           |                            | HQ595338          |                  |
|      |                         |                           |                            | JF509893          |                  |
| SA1  | Banyuls sur Mer, France | Sepiola affinis (squid light organ) | (Fidopiastis et al 1998) | HQ595318          |                  |
|      |                         |                           |                            | EU907986          |                  |
|      |                         |                           |                            | EU908010          |                  |
|      |                         |                           |                            | JF509894          |                  |
| SA6  | (V. logei)              | ""                       | (Fidopiastis et al 1998) | JF509782          |                  |
|      |                         |                           |                            | JF509796          |                  |
|      |                         |                           |                            | JF509854          |                  |
|      |                         |                           |                            | JF509895          |                  |
| SR5  | ""                     | Sepiola robusta (squid light organ) | (Fidopiastis et al 1998) | HQ595319          |                  |
|      |                         |                           |                            | EU907987          |                  |
|      |                         |                           |                            | EU908011          |                  |
|      |                         |                           |                            | JF509896          |                  |
| ZJH004² | Oahu, HI, USA (Maunalua Bay) | E. scolopes (squid light organ) | This Study             | MF076803          |                  |
|      |                         |                           |                            | MF076816          |                  |
|      |                         |                           |                            | MF076829          |                  |
|      |                         |                           |                            | MF076842          |                  |

¹ All strains are *V. fischeri* unless otherwise noted.
² Sequences collected in this study by PCR and Sanger sequencing.
³ Sequences collected in this study by next-generation sequencing via the Illumina platform.
### Supplemental Table S7. Strains, Plasmids, Oligo table

| Strains or Plasmids | Relevant characteristics | Source or Ref. |
|---------------------|--------------------------|---------------|
| **E. coli**          |                          |               |
| DH5α                | F′/endA1 hsdR17 glnV44 thi-1 recA1 gyrA relA1 Δ(lacIZYA-argF)U169deoR1806lacIΔ(lacZ)M15 | (Hanahan, 1983) |
| DH5αλpir            | pir derivative of DH5α    | (Dunn et al., 2005) |
| CC118Δpir           | Δ(ara-leu) araD Δlac74 galE galK phoA20 thi-1 rpsE rpsB argE(Am) recA λpir | (Herrero et al., 1990) |
| **V. fischeri**     |                          |               |
| ANS2098             | FQ-A001 with vasA_2 disruption (ErmR) | This study |
| ANS2099             | FQ-A002 with vasA_2 disruption (ErmR) | This study |
| LAS003              | EBS004 with vasA_2 disruption (ErmR) | This study |
| LAS005              | FQ-A001 with vasA_1 disruption (ErmR) | This study |
| LAS006              | MB11B1 with vasA_2 disruption (ErmR) | This study |
| LAS007              | MB13B3 with vasA_2 disruption (ErmR) | This study |
| **Plasmids**        |                          |               |
| pAS2038             | vasA_2 disruption vector; oriV_bsk, oriT, ErmR | This study |
| pLS04               | vasA_1 disruption vector; oriV_bsk, oriT, ErmR | This study |
| pSNS116             | vasAB_2 complementation vector; oriV_bsk, oriV_pES213, oriT, KnR | This study |
| pSNS119             | vipA_2-gfp fusion vector; oriV_bsk, oriV_pES213, oriT, KnR | This study |
| pSCV38              | P_stu-gfp, P_data-mCherry, oriV_bsk, oriV_pES213, oriT, CmR | Sun et al., 2016 |
| pYS112              | P_pro-cfp, P_data-mCherry, oriV_bsk, oriV_pES213, oriT, CmR | Sun et al., 2016 |
| pEV104              | conjugative helper, oriV_bsk, oriT, KnR | Stabb & Ruby, 2002 |
| pAKD601             | lacIq and IPTG-inducible promoter with optional GFP fusion, oriV_bsk, oriV_pES213, oriT, KnR | Dunn and Stabb, 2008 |
| pEV122              | oriV_bsk, oriT, ErmR | Dunn et al., 2005 |
| pVSV102             | gfp+, oriV_bsk, oriV_pES213, oriT, KnR | Dunn et al., 2006 |
| pVSV208             | dsRed+, oriV_bsk, oriV_pES213, oriT, CmR | Dunn et al., 2006 |

### Oligonucleotides

| AS1146          | TAGGTACCCTTGATGGAGCGCTTTATTAG | This study |
| AS1147          | ATCATGCAAGAATCTTGTCTGTTTTG | This study |
| AS1064          | ATGTACCCTCACAGCACTAGTTATTTTAG | This study |
| AS1066          | ATGTACCTTGAACAAATACCTGGAATATCAATG | This study |
| AS1140-R        | TATTAACTCCTACTACACATATAACTG | This study |
| AS1141-R        | ATGATTCATATATGTTTAAATACC | This study |
| AS1158          | TGCGCTCGATATAAATACCG | This study |
| AS1159          | TCACTTCTAGAATCAGG | This study |
| AS1204          | GCGAATTCCGACGCTCGGATACCACAGCTTATGACC | This study |
| AS1205          | GACCTTAGAGGATCCCGGGGATAGATACGTATCAAAGTGGG | This study |
| LS004           | TCGAAAGGCCTCGTTTTTTTAG | This study |
| SNS56           | GGTGAATCCGAGCTCGGATACCAAGGATATATGTCACGTGAT | This study |
| SNS57           | GCTCTTCTTATTAG | This study |
| SNS41           | GGGATTCCGAGCTCGGATACCAAGGATATATGTCACGTGAT | This study |
| SNS42           | TCCTTCTTGTGTCTCTATGATTAGTTACTGCGCCATTATAATTTCG | This study |
| recA outer-F    | GACGATAACAAGAAAAAGCACCTGG | Wollenberg 2012 |
| recA outer-R    | CGGTTCCTTAAATTTCCGGAAGAC | Wollenberg 2012 |
| recA inner-F    | TGARAARCARTTYGGAAGGAC | Wollenberg 2012 |
| recA inner-R    | GGGAGRCATACGTCTCCTG | Wollenberg 2012 |
| mdh outer-F     | AAGTAGCTGGTTATTG | Wollenberg 2012 |
| Gene       | Primer Type | Primers                                      | Reference       |
|------------|-------------|----------------------------------------------|-----------------|
| mdh        | outer-R     | CTTCGCCAATTTTGATATCG                         | Wollenberg 2012 |
| mdh        | inner-F     | GCCATTGGACAAGCGTTAGC                         | Wollenberg 2012 |
| mdh        | inner-R     | CGCCTCTTAGCTATCTAGC                         | Wollenberg 2012 |
| katA       | outer-F     | TGTCCTGTTGCACATAACC                         | Wollenberg 2012 |
| katA       | outer-R     | CGCTTACATCAATATCAAG                         | Wollenberg 2012 |
| katA       | inner-F     | CGTGGTATTCTCTGAACATAC                       | Wollenberg 2012 |
| katA       | inner-R     | CCGATACCTTCACATAAGC                        | Wollenberg 2012 |
| pyrC       | outer-F     | CTGATGATTGGCATTTACAC                        | Wollenberg 2012 |
| pyrC       | outer-R     | GCCACTCAACGCTTCACC                         | Wollenberg 2012 |
| pyrC       | inner-F     | CACTTACGTGATGATGATGATG                       | Wollenberg 2012 |
| pyrC       | inner-R     | GCCACTCAACGCTTCACC                         | Wollenberg 2012 |

*For complete list of *V. fischeri* strains used in this study see supplemental Table S6.

**Restriction sites are underlined.**
Supplemental Methods for Speare et al.

**Media and growth conditions.** *V. fischeri* strains were grown in LBS medium (1) at 24°C and *E. coli* strains were grown in either LB medium (2) or Brain Heart Infusion (Difco) at 37°C. Antibiotic selection for *V. fischeri* and *E. coli* strains were as described previously (3). Plasmids with the R6Kγ origin of replication were maintained in *E. coli* strain DH5αλpir (3) and plasmid pEVS104 (4) was maintained in strain CC118λpir (5). All other plasmids were maintained in *E. coli* strain DH5α (6).

**Isolation of symbiotic *V. fischeri.*** New *V. fischeri* isolates described in this study (Table S3) were isolated from *Euprymna scolopes* light organs. Briefly, adult *E. scolopes* squid were caught by dip-net in Kaneohe or Maunalua Bay, Oahu. After capture, animals were transported to a holding tank supplied with natural seawater. Adults were transported to Penn State where they were kept in an aquarium before anesthetizing, dissection, and plating of dilution series of light organ homogenate. Individual colonies were picked and re-streaked for purification.

**Strain and plasmid construction.** Bacterial strains, plasmids, and oligonucleotides used in this study are presented in Table S4. For mutant construction in *V. fischeri*, mutant alleles were mobilized on plasmids into recipients by triparental mating using CC118λpir pEVS104 as a conjugative helper. Potential mutants were screened for appropriate antibiotic resistance markers and verified using PCR. All primer design was based on the MJ11 genome sequence. To construct the *vasA_1* disruption mutant, approximately 1 kb of the *vasA_1* gene was PCR amplified using primers AS1204 and AS1205 from FQ-A001 gDNA. The resulting PCR product was cloned into the KpnI and SphI sites of plasmid pEVS122, resulting in the *vasA_1* disruption construct, pLS04.
The \textit{vasA\textsubscript{1}} disruption construct on pLS04 was moved into strain FQ-A001, resulting in strains LAS005. To construct the \textit{vasA\textsubscript{2}} disruption mutants, approximately 1 kb of the \textit{vasA\textsubscript{2}} gene was PCR amplified using primers AS1146 and AS1147 from FQ-A001 gDNA. The resulting PCR product was cloned into the KpnI and SphI sites of plasmid pEVS122, resulting in the \textit{vasA\textsubscript{2}} disruption construct, pAS2038. The \textit{vasA\textsubscript{2}} disruption construct on pAS2038 was moved into strains FQ-A001, FQ-A002, EBS004, MB11B1, and MB13B3 resulting in strains ANS2098, ANS2099, LAS003, LAS006, and LAS007, respectively.

To construct the VipA-GFP fusion expression vector, \textit{vipA\textsubscript{2}} was PCR-amplified from strain ES401 gDNA using primers SNS56 and SNS57. The forward primer includes 11 bp upstream of the \textit{vipA\textsubscript{2}} start codon to include the native ribosome binding site (RBS). The reverse primer excluded the native stop codon for \textit{vipA\textsubscript{2}} and a linker sequence was added (5’ GCAGCAGCAGGAGGAGGA 3’) for translational fusion of \textit{vipA\textsubscript{2}} to the \textit{gfp} gene encoded in pAKD601 (7). The \textit{vipA\textsubscript{2}} PCR product was cloned into KpnI and NheI digested pAKD601 using the standard sequence-and ligation-independent cloning (SLIC) technique (8). The \textit{vipA\textsubscript{2}}-\textit{gfp} fusion in the resulting plasmid (pSNS119) is located downstream of an IPTG-inducible promoter.

To construct a complementation vector for the \textit{vasA\textsubscript{2}} mutation, \textit{vasAB\textsubscript{2}} was PCR-amplified from strain FQ-A001 gDNA using primers SNS41 and SNS42. The forward primer includes 11 bp upstream of the start codon to include the native RBS. The reverse primer included the native stop codon to prevent a translational fusion to, or expression of, the downstream \textit{gfp} gene on pAKD601. The resulting \textit{vasAB\textsubscript{2}} PCR product was cloned downstream of an IPTG-inducible promoter in plasmid pAKD601 (cut with KpnI and NheI) using the standard SLIC cloning technique (8), resulting in plasmid pSNS116.
**Single-cell Fluorescence Microscopy.** To visualize GFP-tagged T6SS2 sheath formation in *V. fischeri* cells, we used a single-cell fluorescence microscopy approach adapted from Basler et al., 2012 (9). Overnight cultures of *V. fischeri* wild-type FQ-A001, the *vasA_1* mutant (LAS05), or the *vasA_2* mutant (ANS2098) strains carrying the IPTG-inducible *vipA_2-gfp* fusion expression vector (pSNS119) were diluted 1:100 into fresh LBS medium supplemented with 0.5 mM isopropyl-β-D-1-thiogalactopyranoside (IPTG) and cultivated at 24°C with shaking for 2.5-3 hours to an OD600 of approximately 1.5. Cells from 5 µL of these cultures were spotted onto a thin pad of LBS with 2% agar and 0.5 mM IPTG, covered with a glass cover slip and imaged after two hours at room temperature. Fluorescence images were captured using an Olympus BX51 microscope outfitted with a Hammatsu C8484-03G01 camera and a 100X/1.30 Oil Ph3 objective lens. Images were captured using MetaMorph software. Contrast on images was adjusted uniformly across images by subtracting background using ImageJ software.

**Contact-dependent Co-incubation Assay.** To test for contact-dependent interactions, strains were prepared as described in the methods section, except strains were separated using a 0.22 µm nitrocellulose membrane. Specifically, 5 µl of each strain was spotted onto a membrane and allowed to dry. These membranes were placed directly on top of one another (alternating which strain was on the top and bottom membranes) and placed onto LBS agar plates and incubated at 24°C for 5 h. After 5 h, both membranes were removed from the plate and suspended in 3 mL LBS medium. Strain were quantified by plating serial dilutions for T0 and T5 onto selective LBS agar plates. For each experiment four independent cultures of each strain were assayed and each experiment was repeated three times.
Squid Colonization Assays. Overnight cultures of the indicated strains were diluted 1/100 into LBS supplemented with 2.5 μg/ml chloramphenicol and grown to OD$_{600}$ ~ 1.0. For each inoculum, cultures were diluted into filter-sterilized seawater (FSSW) and sampled for CFU. For each treatment, 24-30 freshly hatched juvenile squid were exposed to the inoculum containing an even mix of YFP- and CFP- labeled strains (using pSCV38 and pYS112, respectively) at a final concentration ranging from 1600 to 8240 CFU/ml. Squid were exposed to this mixed inoculum for 20 h and then washed in fresh FSSW. After 44 h, animals were fixed in 4% paraformaldehyde/marine phosphate buffered saline (mPBS) for 24 h at 4°C, then washed exhaustively in mPBS. Animals were prepared for fluorescence microscopy by dissecting the ventral side of the mantle and removing the siphon to reveal the light organ. YFP, CFP, and differential interference contrast (DIC) images were taken using a Zeiss 780 confocal microscope (Carl Zeiss AG, Jena, Germany) equipped with a 10x or 40x water lens. Each crypt space was scored separately for CFP and YFP fluorescence.

Phylogenetic Analysis Details. A multi-locus phylogenetic analysis was performed using partial sequences of four loci: recA, mdh, katA, and pyrC. Published sequence data and newly amplified sequences of 35 total Vibrio isolates were collected, aligned with ClustalX 2.1 (10), analyzed via three independent runs of 2,000 samples each in ClonalFrame 1.2 (11), and visualized with a consensus network in Splitstree 4.12.2 (12) as described previously (13, 14). The resulting consensus network showed little evidence of phylogenetic incongruence (so-called “splits” represented by parallelograms visualized among nodes in the network) among these four partial loci. Because the ClonalFrame/Splitstree analysis revealed little evidence of phylogenetic incongruence among these four partial loci, for each isolate the four partial sequences were
combined into a single concatenated sequence (ordered *recA mdh katA pyrC* – approximately 2880 nucleotides). Concatenated sequences were analyzed by jModelTest 2.1 v20160303 (15) via three information criteria methods (Akaike, Bayesian, and Decision Theory). The latter two methods calculated the lowest likelihood score for a transitional model with a gamma shape parameter and a proportion of invariable sites (TIM3+$\Gamma$+$I$) while a general time-reversible model with a gamma shape parameter and a proportion of invariable sites (GTR+$\Gamma$+$I$) was given the lowest likelihood score with the Akaike method.

TIM3+$\Gamma$+$I$ evolutionary model parameter estimates calculated by jModelTest were used by the software program PAUP*4.0b10 (16) to infer phylogenetic trees and bootstrap those trees via two methods: Maximum Parsimony (MP) and Maximum Likelihood (ML). ML phylogenetic inference and bootstrapping was performed by searching heuristically using simple addition and subtree pruning and regrafting for swaps, treating gaps as missing, and swapping on “best only” with 1000 replicates and 1000 bootstrap pseudoreplicates. MP phylogenetic inference and bootstrapping was performed by searching heuristically using simple addition and tree bisection reconnection for swaps, treating gaps as missing, and swapping on “best only” with 1000 replicates and 1000 bootstrap pseudoreplicates.

A Bayesian approach (Ba) to phylogenetic inference was also completed with the program MrBayes 3.1.2 (17) by setting the “nst” variable to “6” and the “rates” variable to “invgamma” (this approximates a GTR+$\Gamma$+$I$ model); three heated chains were set using the “temp” variable to a value of 0.05 (to ensure appropriate chain swapping). Construction of the majority-rule consensus tree and statistical analysis of clade membership/presence was assessed by sampling an “appropriately stationary” posterior probability distribution. For the purposes of this study, an “appropriately stationary” distribution was defined, as recommended by Ronquist
and colleagues (18), as an average standard deviation of split frequencies of less than 0.01 for 70% to 90% of samples between two, independent Metropolis-coupled Markov Chain Monte Carlo runs. Approximately 3,000,000 total generations were sampled every 100 generations for a total of 30,000 samples – 10,000 of these samples were discarded via the “burnin” variable in MrBayes. Majority-rule consensus trees drawn from the resulting 20,000-sample, stationary distribution were used for the assessment of the posterior probabilities of all clades. The above methods were independently repeated twice; all three separate Ba “replicates” showed nearly identical phylogenetic patterns of clades and posterior probabilities. Sequences associated with this analysis were submitted to the GenBank database and their accession numbers are listed in Table S3.

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