Gene-Swapping Mediates Host Specificity among Symbiotic Bacteria in a Beneficial Symbiosis

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Abstract

Environmentally acquired beneficial associations are comprised of a wide variety of symbiotic species that vary both genetically and phenotypically, and therefore have differential colonization abilities, even when symbionts are of the same species. Strain variation is common among conspecific hosts, where subtle differences can lead to competitive exclusion between closely related strains. One example where symbiont specificity is observed is in the sepiolid squid-Vibrio mutualism, where competitive dominance exists among *V. fischeri* isolates due to subtle genetic differences between strains. Although key symbiotic loci are responsible for the establishment of this association, the genetic mechanisms that dictate strain specificity are not fully understood. We examined several symbiotic loci (*lux*–bioluminescence, *pil* = pili, and *msh*-mannose sensitive hemagglutinin) from mutualistic *V. fischeri* strains isolated from two geographically distinct squid host species (*Euprymna tasmanica*-Australia and *E. scolopes*-Hawaii) to determine whether slight genetic differences regulated host specificity. Through colonization studies performed in naïve squid hatchlings from both hosts, we found that all loci examined are important for specificity and host recognition. Complementation of null mutations in non-native *V. fischeri* with loci from the native *V. fischeri* caused a gain in fitness, resulting in competitive dominance in the non-native host. The competitive ability of these symbiotic loci depended upon the locus tested and the specific squid species in which colonization was measured. Our results demonstrate that multiple bacterial genetic elements can determine *V. fischeri* strain specificity between two closely related squid hosts, indicating how important genetic variation is for regulating conspecific beneficial interactions that are acquired from the environment.

Introduction

Environmentally transmitted symbioses occur through the acquisition of bacteria from the environment into a naïve, uncolonized juvenile host [1]. This type of transmission strategy can be complex, since bacteria are obtained anew for each generation of hosts, and is dependent upon the population and type of symbionts present when transmission occurs [2,3]. Both host and environment have strong influences upon symbiont fitness, and it is the interplay between these two forces that determine whether specific symbiotic strains are able to colonize and persist generation after generation [4,5]. One example where both host and environment exert notable selection pressures upon symbiotic bacteria is in the sepiolid squid-*Vibrio* mutualism [5].

Complex molecular dialogs (including genetic interdependence) exist between sepiolid squid hosts and their *Vibrio* bacteria, leading to a highly specific association and subsequent coevolution [5–8]. The complex processes by which hosts and symbionts find each other (among the tremendous marine bacterial community) in order to initiate a successful mutualism include a myriad of well-defined molecular signaling events that dictate a certain “conversation” between partners [9]. Additionally, it has been reported that both bacterial specificity (where Vibrios preferentially colonize particular species of hosts as well as environment [6,10,11]) dictates which symbionts are successful in squid light organ symbioses. Studies competing native strains with non-native strains in both allopatric Australian *Euprymna tasmanica* and Hawaiian *Euprymna scolopes* indicate the existence of competitive dominance and intraspecific recognition of environmentally transferred symbionts [6,8,11]. Along with host specificity, environmental temperature is also an important factor for colonization and dominance of specific *Vibrio* strains when colonizing different squid host species living in sympathy [4]. Thus, bacterial specificity is dictated both by host mechanisms of selection (particular *Vibrio* spp. are host specialists) or the environment (*vibrio* bacteria as a group are host generalists), but the exact means of this specificity have not been determined [5].

Recent studies have been devoted to defining bacterial mechanisms (for example gene activation, horizontally transmitted elements, mutations, duplications, etc.) for host specificity. Recently, a study demonstrated that, under laboratory conditions, host specificity between sepiolid squids and one species of monocentric fish was determined by the presence of a single gene in *V. fischeri* (*mshD*), which regulates luminescence and synthesis of the symbiosis
polysaccharide locus (gp) that is important for host colonization and biofilm formation [7]. What subtle genetic factors are responsible for the dramatic competitive fitness differences between various isolates of V. fischeri among all cephalopod squids, rather than between squids and a completely different vertebrate host, is the focus of this study.

Two closely related V. fischeri isolates (ETJB1H from the Australian host Euprymna tasmaniae, and ES114 from the Hawaiian host E. scolopes) were examined in order to determine whether differences in symbiotic loci were important for strain specificity and host recognition. These two V. fischeri strains were selected because they can each colonize aposymbiotic hatchlings from both species of squid equally well in 48 hour colonization assays so long as they are the only strain of V. fischeri present; and they nevertheless also demonstrate competitive dominance when they are presented together to their native host. That is to say Australian E. tasmaniae hosts are preferentially colonized by V. fischeri ETJB1H when V. fischeri ETJB1H and V. fischeri ES114 bacteria are both present, and E. scolopes are preferentially colonized by V. fischeri ES114 when V. fischeri ETJB1H bacteria are also present [6,8,11]. It is likely that subtle differences in specific symbiotic loci are responsible for this complex phenotype. The two particular bacterial strains were also selected because they have full or partial sequenced genomes, allowing easy genetic comparisons between both strains [10].

Materials and Methods

This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocol was approved by the Institutional Biosafety Committee of New Mexico State University (Permit Number: 1306NMD20103) and under the guidelines of the NMSU’s Institutional Animal Care and Use Committee (IS-R-009 and OLAW A4022-01 and IACUC license 2013-029). Animals were appropriately handled with care and under appropriate conditions to minimize any suffering [5]. Adult Euprymna tasmaniae were collected from Botany Bay, New South Wales, Australia with permits from the Australian Government, Department of Sustainability, Environment, Water, Population, and Communities (Export permit WT2013-10343), the New South Wales Government Industry and Investment (Collection permit P04/0014-6.0), and the Australian Government Department of Agriculture, Fisheries, and Forestry Biosecurity (AQIS invoice EL50015507329). Euprymna scolopes (Kane‘ohe Bay, Honolulu, O‘ahu, GPS coordinates- N 21° 26’ W 157° 47’) were not required to have any collection permits at this site. Both species of Euprymna are not endangered or are protected in either location.

Bacterial strains and growth conditions

Two V. fischeri strains were chosen for this study: V. fischeri ETJB1H isolated from the light organ of Euprymna tasmaniae from Jervis Bay, Australia and V. fischeri ES114 isolated from the light organ of Euprymna scolopes from Kane‘ohe Bay, Hawaii. Both strains were grown in Luria Bertani high Salt (LBS; per liter composition: 10 g tryptone, 3 g yeast extract, 20 g NaCl, 50 mL 1 M Tris pH 7.5, 3.75 mL 80% glycerol and 950 mL dH2O) media and shaken at 225 rpm at 28°C overnight.

Mutation construction

Campbell mutations. Luciferase (lux) mutants of both strains (V. fischeri ETJB1H and ES114) were constructed by insertion of the plasmid pEV122 as described previously (11) and constructs are listed in Table S2. Briefly, the luxA gene was partially amplified with specific primers designed from the sequenced strain V. fischeri ES114 (NCBI accession: NC_006040.2). PCR products were purified and cloned (after double digestion of PCR products and plasmid with XhoI and XmnI, with posterior ligation in a 1:5 plasmid-insert ratio) into the suicide vector pEV122, and wild type V. fischeri strains were transformed by tri-parental mating via conjugation through a helper strain [12]. Strains that had undergone single homologous recombination events with the native gene were selected on LBS plates enriched with erythromycin (25 μg/mL). Strains constructed were defined as ES114::pACH101 (for the lux mutant of the Hawaiian V. fischeri strain ES114) and ETJB1H::pACH102 (for the lux mutant of the Australian V. fischeri strain ETJB1H). Constructs were verified by Southern blotting.

Allelic exchange. msh and pil mutants (mshLNO and pilABCD) were constructed by allelic replacement of the chromosomal loci as described previously [13]. 500 bp of neighbor genes were amplified and cloned (the first insert was cloned after digestion of plasmid and PCR product with SamI and BamHI, with posterior ligation in a 1:5 plasmid-insert ratio; the second insert was cloned after digestion of plasmid and PCR product with XhoI and XbaI, with posterior ligation in a 1:10 plasmid-insert ratio) into the suicide vector pSW7848 containing the PBAD promoter and a chloramphenicol resistant cassette (Tables S1, S2). After transformation of ultracompotent cells (NEB 10-beta competent E. coli, New England Biolabs, MA, USA), selection was achieved through antibiotic enrichment (5 μg/mL chloramphenicol). Recipient cells (V. fischeri) were transformed by tri-parental mating as described above. Transformed strains (with respective deletions) were selected through colony patching after inoculation in LBS media enriched with 2% arabinose to allow dismissal of inserted constructs through expression of the toxic gene ccdB by activation of the PBAD promoter after incubation in LBS media with arabinose. To easily discriminate transformants, colonies from the original tri-parental mating that were initially resistant to chloramphenicol and eventually insensitive to CcdB toxicity were selected. Constructs were named by their respective deletion (Table S2) and were verified by PCR.

Complement construction

GAPtore or TAR cloning. Complementation of the lux operon with the opposite strain’s loci (lux in ETJB1H for ES114::pACH101 and lux in ES114 for ETJB1H::pACH102) was achieved through operon mobilization (“TAR cloning” or “GAPtore”) as previously described [14]. TAR cloning technique was achieved by using the yeast homologous recombination pathway. 700 bp of neighboring genes (upstream and downstream) from the luxCDABEG operon (with 40 nucleotides of the 5’ end that were homologous with vectors pCRG13 and pCRG25) were PCR amplified and purified. Vector pCRG25 was digested with SfiI and pCRG13 was digested with EcoRV. In Saccharomyces cerevisiae transformation, the yeast strain CRY1-2 (containing the genotype ura3-1, leu2-3,112, cyh2, that confers sensitivity to cyclohexamid and cannot grow in media without uracil) was co-transformed with the two digested plasmids and the amplified upstream and downstream genes using lithium acetate transformation. After transformation, yeast colonies were plated on synthetic URA medium (per liter composition: adenine hemisulfate 0.18 g, arginine HCl 0.12 g, glutamic acid 0.6 g, histidine HCl 0.12 g, myo-inositol 0.2 g, isoleucine 0.18 g, leucine 0.18 g, lysine HCl 0.18 g, methionine 0.12 g, p-aminobenzoic acid 0.02 g, phenylalanine 0.3 g, homoserine 0.6 g, tryptophan 0.24 g, tyrosine 0.18 g, valine 0.9 g, Difco yeast nitrogen base
without amino acids 6.67 g, glucose 20 g) and incubated at 30°C for 4 days. After incubation, colonies were suspended in fresh medium with 10 mL of TE and transferred into a 15 mL falcon tube, spun, and the construct (two plasmids + upstream/downstream genes) was extracted with glass beads and 200 μL of glass beads solution (per liter composition: 5 mL 20% sodium dodecyl sulfate, 10 mL 1 M NaCl, 1 mL 1 M Tris-HCl pH 8.0, 1 mL 0.1 M EDTA, 2 mL, Triton X100) and purified with Phenol/Chloroform.

Plasmid DNA was transformed with ultracompetent E. coli cells (NEB 10-beta competent E. coli, New England BioLabs, MA, USA) and incubated for 24 hours at 37°C. Transformed cells (~10 colonies) were re-isolated and plasmid DNA was extracted using the Qiagen plasmid Maxi kit (QIAGEN Inc., CA, USA). For lux operon cloning step, CRY 1–2 yeast cells were transformed (CaCl₂ spheroplast transformation procedure) with 1 μg of the plasmid extract and 5 μg of genomic DNA (from either V. fischeri ES114 or ETJB1H). Yeast cells were plated onto TYG1/ Cycloheximide plates (per liter composition: D-sorbitol 182.2 g, Difco yeast nitrogen base without amino acids 6.75 g, dextrose 0.98 g, adenine 0.2 g, arginine 0.2 g, aspartic acid 1 g, histidine 0.2 g, leucine 0.59 g, lysine 0.53 g, methionine 0.2 g, phenylalanine 0.4 g, threonine 2 g, tryptophan 0.2 g, tyrosine 0.3 g and 3 mg/mL cycloheximide) and incubated at 30°C for 7 days. Plasmids (containing the lux operon) were extracted from yeast spheroplasts using the Stratagene stastrapt smplid miniprep kit (Fisher Scientific, PA, USA). Constructs were verified by PCR and Southern blotting. E. coli ultracompetent cells were transformed and triparental mating was achieved as described previously. V. fischeri ES114::pACH101 was complemented with the lux operon of V. fischeri ETJB1H and strain V. fischeri ETJB1H::pACH102 was complemented with the lux operon of V. fischeri ES114.

Cloning using the conjugal vector pVSV105. Complete copies of the loci for msh and pil operons were amplified with specific primers for the entire locus (Table S1). PCR products and the vector pVSV105 [15] were double digested with XbaI and XmnI digests were ligated in a 1:3 plasmid-insert ratio and transformed into ultracompetent E. coli cells (NEB 10-beta competent E. coli, New England BioLabs, MA, USA). Cells were selected with chloramphenicol enrichment (25 μg/mL) and V. fischeri recipient cells were transformed by tri-parental mating and selected after chloramphenicol enrichment (5 μg/mL). Lastly, we constructed complements that contain each strain’s native loci by cloning the respective locus into vector pVSV105. Transformation was then performed as described previously (see Table S2 for complete list of complements). Complemented strains were verified by Southern blot.

Colonization assays

Colonization assays were performed as described previously [16]. Overnight cultures of V. fischeri wild-type strains (ES114 and ETJB1H), mutants and complements were regrown in 5 mL of fresh LBS media until they reached an OD₆₀₀ of 0.3. For single and competition infection experiments, cultures were then diluted to approximately 1×10⁶ CFU/mL in 5 mL of sterile artificial seawater and added to glass scintillation vials where newly hatched juvenile squid were placed (one individual/vial). Seawater was changed with fresh uninoculated artificial seawater every 12 hours over a period of 48 hours. Animals were maintained on a light/dark cycle of 12/12. After 48 hours, animals were sacrificed and homogenized, and the diluted homogenate was plated onto LBS agar plates for the wild-type V. fischeri, LBS with erythromycin (25 μg/mL) for the V. fischeri mutants, and LBS with chloramphenicol (5 μg/mL) for the V. fischeri complements. A second set of animals were selected for competition studies where juvenile squids were co-infected with the native strain and a respective complement (Table S2), sacrificed after 48 hours, homogenized, and plated onto the various media as reported above. Colony forming units (CFUs) were counted the next day to determine colonization efficiency of each strain. A total of 8 animals/strain were used for each competition assay, and 10 non-infected (aposymbiotic) juveniles were used as negative controls.

Statistical analysis

To compare bacterial populations (wild-type, mutant, and complement constructs), one way ANOVA followed by the Tukey comparison was performed on calculated CFU numbers. Three technical replicates and 10 biological replicates (representing 3 treatments with 10 animals/strain and one set of 10 non-infected or aposymbiotic animals for the negative control). Results of similar experiments completed in E. tasmanica E. scolopes against non-native Australian V. fischeri ES114 against non-native Australian lux- V. fischeri ETJB1H strain complemented with either the native ES114 lux or the ETJB1H lux genes (Figs. 1, S1). The lux- native V. fischeri ETJB1H had equal competitive ability against native V. fischeri ES114 exclusively when its lux mutation had been complemented with native ES114 lux (Fig. 1). That is, lux- non-native ETJB1H behaved like ES114 exclusively when the lux mutation was complemented with the ES114 lux operon. Furthermore, when the lux- native Hawaiian strain (ES114) was complemented with the non-native lux operon from the Australian strain (ETJB1H) and competed against native V. fischeri ES114, the wild type dominated the complemented strain (Fig. 1). That is to say the lux- ES114 strain complemented with the lux operon from non-native ETJB1H behaved like the non-cognate ETJB1H strain.

Results of similar experiments completed in E. tasmanica juveniles (where V. fischeri ETJB1H is native, and V. fischeri ES114 is non-native) produced the expected reciprocal results. For example, when lux from non-native Hawaiian V. fischeri (ES114) was replaced with native lux from Australian V. fischeri ETJB1H strain –luxES114::luxETJB1H, there was an increase in colonization efficiency of the competitor as if using the native wild type
though it may not be the only gene responsible for symbiont recognition and specificity.

The lux operon is responsible for biosynthesis of luciferase, which has a crucial role in V. fischeri bioluminescence and fitness. Light production is used by the squid to avoid predation via silhouette reduction in a behavior known as counterillumination [16,17]. The lux operon is present in V. fischeri as a conserved, contiguous, and coordinately expressed set of genes that have thought to have been acquired through horizontal gene transfer (HGT) among closely related bacterial clones and through vertical inheritance between bacterial families (e.g., Vibrionaceae and Enterobacteriaceae). Results from this part of our study indicate that although both lux operons produce bioluminescence and their structural proteins are similar in primary sequence, host specificity can be obtained through artificial HGT of the lux operon alone [18]. Our cloning method intentionally included 5’ and 3’ noncoding sequences flanking the operons; perhaps noncoding sequences contribute to the observed host preference. The lux operon might therefore drive evolutionary strain speciation through non-reproductive transmission of lux genes, when lux DNA is available in the environment and there are no other constraints on integration of operons into the recipient cell (e.g., the action of restriction endonucleases). Additionally, natural competence has been previously observed in V. fischeri after expression of the transcriptional regulators fokA and fokV (chitin-sensing regulators); this earlier study highlights a conserved mechanism of genetic exchange in the presence of chitin [19].

Multiple genes comprise the entire msh operon (including mshABCDGJMNPQ), which is responsible for the synthesis of type IV pseudopili, important for biofilm formation [20] and attachment (or adherence) to abiotic surfaces [21]. The msh operon has also been reported to be crucial for attachment to certain host tissues, which is an important step for successful colonization and persistence [11]. We specifically targeted mshI, mshQ, and mshN since these proteins exhibit high variability in their primary sequence among multiple strains of V. fischeri, including Hawaiian ES114 and Australian ETJB1H (C. Lostroh, unpublished data). Loci from the msh operon were mutated by means of insertional inactivation and complemented by extrachromosomal maintenance [13,15,22]. Similar to the lux operon experiments, ES114 mutant strains were complemented with the ETJB1H msh gene, and vice versa. Colonization tests using all mutant strains were then completed in both E. tasmanica and E. scolopes juvenile squids. Results of mshI, mshN and mshQ loci after colonization are illustrated in Figure 2. Due to the difficulty of obtaining a large number of animals from one clutch to complete all infection experiments with msh strains, we used three different clutches from E. scolopes and two from E. tasmanica. Inter-clutch colonization variability was observed between groups, and reflected in low numbers in competition experiments; however, animals from the same clutch were used to replicate the same competition experiment to avoid variation in colonization efficiency.

Mutation of the msh genes in Hawaiian V. fischeri ES114 caused a significant reduction in colonization efficiency in E. scolopes (data not shown), demonstrating that the msh operon is important for symbiotic competence. When Hawaiian E. scolopes were infected with non-native ETJB1H complemented with native mshI (mshIETJB1H::mshIES114), colonization efficiency was equal to the native wild-type (ES114). Results were similar for mshL and mshQ. These observations indicate that mshI, mshL, or mshQ are all important in conferring host specificity between E. scolopes and E. tasmanica squids. Interestingly, colonization in E. tasmanica juveniles did not mirror these results as they did for the lux operon. Overall levels of colonization in this case were very low (Fig. S1). The mshI
and mshN ES114 strains complemented with their ETJB1H msh counterparts were out-competed by wild type ETJB1H, while the mshQ ES114 strain complemented with its ETJB1H msh counterpart competed slightly better for colonization than wild type ETJB1H. Alternatively, ES114 complemented with native ETJB1H mshQ locus (-mshQ:ES114::mshQ ETJB1H) outcompeted the wild-type strain. Thus, results for mshQ closely resemble those for lax, whereas mshI and mshN favored one V. fischeri strain (ETJB1H) but not the other (ES114). In our mixed competitions using mshI or mshN mutants, E. tasmanica hosts select against all complemented bacteria, keeping total levels of each symbiont low (Figs. 2, S2). In addition to clutch variability, E. tasmanica hosts may exert stronger sanctions against non-native V. fischeri more than its congeners due to the presence of a genetically diverse group of V. fischeri symbionts available for colonization in the E. tasmanica habitat, whereas V. fischeri symbionts from E. scolopes are more homogeneous and host squids sample from only a small set of V. fischeri genotypes [5]. Having the ability to discern amongst a large, genetically diverse pool of V. fischeri may give squids an advantage to also differentiate cheaters to allow for a more successful beneficial symbiosis [23]. Recent work has demonstrated changes in particular symbiotic traits (luminescence, biofilm production, motility, carbon source utilization, growth) of Hawaiian V. fischeri strain ES114 when evolved in E. tasmanica hosts [24]. These traits differ quite dramatically, with the evolved strain gaining traits similar to the native strain over time. Thus, our results indicate that the msh operon is not only important for successful colonization of cephalopod squids, but also determines host range and accommodation from a large pool of available Vibrio symbionts.

We also created null pil mutants, and complemented them in trans to examine host selection. Colonization experiments in Hawaiian E. scolopes hatchlings indicate that pilA, pilB and pilD have important roles in host specificity (Figs. 3, S3). When constructs containing the native complemented gene were competed with either native or non-native wild-type strains, colonization efficiency of the constructed strains was equal or greater than the wild-type strain (Fig. 3). Similar results were observed in the case of E. tasmanica infection studies; however, pilD was the only locus that demonstrated host specificity in E. tasmanica (and not pilA or pilB). Genes from the pil operon (pilABCD) encode for assembly of type IV pili, and are essential for bacterial attachment to both abiotic surfaces and to host cells [16,19]. In V. fischeri, pilus subunits are synthesized by the pilABCD operon, where pilA contributes to colonization efficiency and encodes a protein similar to type IV-A pilins (where mshA is a close relative [25,26]. Phylogenetic and molecular differences have also been observed in pilB and pilD loci among multiple V. fischeri strains isolated from different squid hosts [27]. Our study demonstrates that V. fischeri ETJB1C pilC complemented with non-native Hawaiian V. fischeri ES114 pilC (-pilC:ETJB1H:pilC ES114) is dominant in E. scolopes, but loses in E. tasmanica, since there is a competitive dominance for the native pilC locus (Fig. S3). Similar results are observed with V. fischeri ES114 pilC complemented with non-native Australian V. fischeri ETJB1H from E. tasmanica. This may be due to PilC being a phase variable protein [with minor differences in 3–5 aminoacids [25], which besides being implicated in type IV pilus biogenesis, mediates cell adherence [26]. Also, the heterogeneity of pili morphology means that multiple minor proteins composed of PilC subunits have evolved to be variable in order to compete for pilus receptors in host cells [25]. The intriguing question of how Pil-dependent binding is modulated and controlled between closely related host species may explain how host-switching can be accomplished through slight variations at this locus.

Additionally, we constructed complements containing the native genes and performed single colonization experiments as well as competition studies. Single colonization assays indicated that the complement was able to regain the colonization efficiency observed in the wild-type strain (data not shown), and the competition experiments (where the wild-type strain is used to co-infect the host with the native complement) indicated equivalent colonization efficiency between the two strains (Fig. S4).

To determine whether deletion of the various genetic elements had polar effects, we performed additional experiments to observe...
whether our mutants can affect phenotypes related to the function of downstream genes in the operon. For example, msh and pil influence adhesion and biofilm formation [21], and lux is responsible for light production [9]. We quantified biofilm and light production in both mutants and mutants complemented with the native gene, or in the case of lux, genes. Biofilm production decreased in mutant msh and pil strains, and light production was also impaired in lux mutants. Each phenotype was recovered in the respective native complements (data not shown), indicating that polarity effects may not be present; however, to be certain of this assumption additional studies are planned to determine if there is an influence in additional phenotypes. Future studies include transcriptional profiling and genetic analyses of metabolic pathways that might be affected in the various mutant strains.

Previous work examining experimental evolution of *V. fischeri* demonstrated polymorphic changes in phenotype (e.g., bioluminescence, biofilm, motility, growth) when strains are evolved in a novel host, allowing greater colonization efficiency of evolved strains when competed against their non-evolved ancestor [3,24]. These results are consistent with our directionally mutated strains reported here, where non-native strains complemented with native loci (in both Hawaiian *E. scolopes* and Australian *E. tasmanica*) out-competed non-native strains and competed favorably over native parental strains during colonization in both host species examined. Our results indicate that the operons examined here are critical host-specificity factors and sufficient to dictate host recognition among closely related strains of *V. fischeri* from different geographical origins. Thus, strain specificity between two closely related *V. fischeri* symbionts from similar hosts is not mediated by a single or few loci, but rather multiple bacterial genetic elements that determine host range in allopatric Indo-west Pacific *Euprymna-Vibrio* associations.

Colonization of the squid host is multifactorial, and different studies from our laboratory have demonstrated that genes that are responsible for phenotypes associated to colonization are important for successful infection; additionally, experimental evolution of closely related strains does lead to competitive dominance of non-native strains [5,24]. Although these studies suggest that these loci are important for host preference, there is the possibility that compensatory mechanisms could overtake the effect of a mutation and regulatory mechanisms (along with genetic factors) are responsible for colonization efficiency.

This study provides additional support of how bacterial diversity can be maintained through host selection, and key symbiotic loci are just one factor in determining host specificity. Determining whether these loci are acting in concert with one another to further push the selective advantage of beneficial vibrios is crucial for our understanding the evolution of symbiotic associations. How these subtle differences arise in wild populations, and whether they confer a greater selective advantage in bacterial fitness, will give insight into the processes of ecological adaptation in *Vibrio* bacteria.

Supporting Information

Figure S1 A) Colonization assays 48-hour post-infection of juvenile *Euprymna scolopes* by wild-type, mutant, and complement strains of the *lux* operon for *Vibrio fischeri*. Single strain infection experiments are represented when only a single bar is shown (□). Competition experiments are represented with two bars (□ is the first strain and ■ is the second strain), and each strain used is indicated below each competition. Wild-type ES114 significantly colonized the host better than non-native ETJB1H. Apo = apo-symbiotic or non-infected juvenile squids. Data are plotted as the mean of Colony Forming Units (CFUs) counted for each strain. Multiple comparisons were calculated between groups using the Tukey PostHoc comparison. Different letters indicate significant differences (p<0.05) between groups or infection sets. See Table S1 for a complete description of strains and Table S2 for a complete description of colonization experiments. B) Colonization assays 48-hour post-infection of juvenile *Euprymna tasmanica* by wild-type, mutant, and complement strains of the *lux* operon for *Vibrio fischeri*. Single strain infection experiments are represented with a single bar (□). Competition experiments are represented with two bars (□ is the first strain and ■ is the second strain), and each strain used is indicated below each competition. Wild-type

Figure 3. pil operon data. Colonization assays 48-hour post-infection of juvenile (A) *Euprymna scolopes* and (B) *Euprymna tasmanica* by their respective wild-type (ES114 or ETJB1H), mutant, and complement strains of pil genes for *Vibrio fischeri*. Infection efficiency data is plotted as the log values of the relative competitiveness index (RCIs), calculated by dividing the ratio of mutant to wild-type by the starting ratio [28]. If the RCI is <1 the mutant strain was outcompeted by the wild-type, the wild-type strain was outcompeted by the mutant if the value is >1, and a RCI equal to 1 indicates no competitive difference. Data points represent individual animals and the position of the figures on the y axis is merely for spacing. Vertical line represents the median value of each data plot. doi:10.1371/journal.pone.0101691.g003

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1. **Euprymna scolopes**
   - ETJB1H vs ES114
   - pilAETJB1H::pilAES114 vs ES114
   - pilBETJB1H::pilBES114 vs ES114
   - Competition: competitor dominates

2. **Euprymna tasmanica**
   - ES114 vs ETJB1H
   - pilDES114::pilDETJB1H vs ETJB1H
   - Competition: competitor dominates

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**Figure 3. pil operon data.** Colonization assays 48-hour post-infection of juvenile (A) *Euprymna scolopes* and (B) *Euprymna tasmanica* by their respective wild-type (ES114 or ETJB1H), mutant, and complement strains of pil genes for *Vibrio fischeri*. Infection efficiency data is plotted as the log values of the relative competitiveness index (RCIs), calculated by dividing the ratio of mutant to wild-type by the starting ratio [28]. If the RCI is <1 the mutant strain was outcompeted by the wild-type, the wild-type strain was outcompeted by the mutant if the value is >1, and a RCI equal to 1 indicates no competitive difference. Data points represent individual animals and the position of the figures on the y axis is merely for spacing. Vertical line represents the median value of each data plot. doi:10.1371/journal.pone.0101691.g003
Australian *V. fischeri* ETJB1H significantly colonized the host better than the non-native Hawaiian *V. fischeri* ES114. Aposymbiotic or non-infected juvenile squids. Data are plotted as the mean of Colony Forming Units (CFUs) counted for each strain. Multiple comparisons were calculated between groups using the Tukey PostHoc comparison. Different letters indicate significant differences (p<0.05) between groups or infection sets.

**Figure S2 A** Colonization assays 48-hour post-infection of juvenile *Euprymna scolopes* by wild-type, mutant, and complement strains of *msh* genes for *Vibrio fischeri*. Single strain infection experiments are represented when only a single bar is shown ( []). Competition experiments are represented with two bars ( □ is the first strain and ■ is the second strain), and each strain used is indicated below each competition. Wild-type ES114 significantly colonized its native host better than non-native ETJB1H. Aposymbiotic or non-infected juvenile squids. Data are plotted as the mean of Colony Forming Units (CFUs) counted for each strain. Multiple comparisons were calculated between groups using the Tukey PostHoc comparison. Different letters indicate significant differences (p<0.05) between groups or infection sets.

**Figure S3 A** Colonization assays 48-hour post-infection of juvenile *Euprymna scolopes* by wild-type, mutant, and complement strains of the *pil* genes for *Vibrio fischeri*. Single strain infection experiments are represented when only a single bar is shown ( []). Competition experiments are represented with two bars ( □ is the first strain and ■ is the second strain), and each strain used is indicated below each competition. Wild-type ETJB1H significantly colonized the host better than the non-native ES114. Aposymbiotic or non-infected juvenile squids. Data are plotted as the mean of Colony Forming Units (CFUs) counted for each strain. Multiple comparisons were calculated between groups using the Tukey PostHoc comparison. Different letters indicate significant differences (p<0.05) between groups or infection sets.

**Figure S4 A** Colonization assays 48-hour post-infection of juvenile *Euprymna tasmania* by wild-type, mutant, and complement strains of native genes for *Vibrio fischeri*. Single strain infection experiments are represented when only a single bar is shown ( ■). Competition experiments are represented with two bars ( □ is the first strain and ■ is the second strain), and each strain used is indicated below each competition. Wild-type ETJB1H significantly colonized the host better than the non-native ES114. Aposymbiotic or non-infected juvenile squids. Data are plotted as the mean of Colony Forming Units (CFUs) counted for each strain. Multiple comparisons were calculated between groups using the one-way ANOVA test and Tukey PostHoc comparison. There was no significant difference between strains (as indicated by the *P* value of each column factor).

**Table S1** Plasmids used and constructed in this study.

**Table S2** Strains used and constructed in this study.

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**Author Contributions**

Conceived and designed the experiments: AC CPL MKN. Performed the experiments: AC CG. Analyzed the data: AC CG CPL MKN. Contributed reagents/materials/analysis tools: MKN. Wrote the paper: AC CPL MKN.

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