Population Dynamics of *Vibrio fischeri* during Infection of *Euprymna scolopes*

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The luminous bacterium *Vibrio fischeri* colonizes a specialized light-emitting organ within its squid host, *Euprymna scolopes*. Newly hatched juvenile squid must acquire their symbiont from ambient seawater, where the bacteria are present at low concentrations. To understand the population dynamics of *V. fischeri* during colonization more fully, we used mini-Tn7 transposons to mark bacteria with antibiotic resistance so that the growth of their progeny could be monitored. When grown in culture, there was no detectable metabolic burden on *V. fischeri* cells carrying the transposon, which inserts in single copy in a specific intergenic region of the *V. fischeri* genome. Strains marked with mini-Tn7 also appeared to be equivalent to the wild type in their ability to infect and multiply within the host during coinoculation experiments. Studies of the early stages of colonization suggested that only a few bacteria became associated with symbiotic tissue when animals were exposed for a discrete period (3 h) to an inoculum of *V. fischeri* cells equivalent to natural population levels; nevertheless, all these hosts became infected. When three differentially marked strains of *V. fischeri* were coincubated with juvenile squid, the number of strains recovered from an individual symbiotic organ was directly dependent on the size of the inoculum. Further, these results indicated that, when exposed to low numbers of *V. fischeri*, the host may become colonized by only one or a few bacterial cells, suggesting that symbiotic infection is highly efficient.

The luminous bacterium *Vibrio fischeri* lives in a cooperative association with the Hawaiian bobtail squid, *Euprymna scolopes*. The squid has developed a specialized light-emitting organ to house the bacteria within the animal’s mantle cavity. When juvenile *E. scolopes* squid hatch, they are symbiont free and must acquire *V. fischeri* from the surrounding seawater (27). The bacteria enter the nascent organ through six pores which connect to ducts that, in turn, lead to specialized, epithelium-lined crypts (18, 26, 30). Once inside these crypts, the bacteria divide rapidly and fill the light organ with approximately $5 \times 10^6$ cells (34). During this colonization process, after a critical cell density is reached, the bacteria induce the lux operon and begin to luminesce. The animal can control the amount of light emitted to the environment and uses this luminescence during its nocturnal activities, probably as a means to disguise itself from predators (42). At dawn, the host expels approximately 95% of the bacteria; the remaining symbionts multiply over the next 12 h, thereby providing the squid again with a source of luminescence by nightfall (34).

The relationship between *V. fischeri* and its squid host is complex and highly specific. For example, although surrounded by seawater containing hundreds of different bacterial species, juvenile *E. scolopes* squid become colonized only by *V. fischeri* (5, 35). Lee and Ruby (19) used a lux gene probe to determine that there were between 5 and 200 CFU of *V. fischeri* per 100 ml of seawater in *E. scolopes* habitats. However, subsequent work suggested that the actual number of *V. fischeri* cells was between 100 and 1,500/ml of seawater, although most of these cells, while symbiotically infective, were in a cryptic state that did not form colonies on typical media (20). Thus, it appeared that juvenile squid are normally colonized in seawater containing a concentration of only a few hundred *V. fischeri* cells per ml (34), although higher population densities might occur in sediments (20). That *V. fischeri* cells constitute less than 0.1% of the total bacterial community makes a strong argument for both the specificity and the efficiency of this benign infection process. In addition, successful colonization of the animal by *V. fischeri* has been shown to trigger a complex developmental program in the light-organ morphology of the juvenile, and this program does not occur if the squid remains uncolonized (11, 18, 30). This close linkage between colonization and juvenile development further emphasizes the importance of the initiation of the relationship between host and symbiont.

To understand the basis for both the efficiency and the specificity of colonization more fully, we developed derivatives of the site-specific transposon mini-Tn7 (22) to create differentially marked strains of *V. fischeri*. These strains were used to track the progeny of individual cells as they infected the light organ of juvenile *E. scolopes*. Because in *Escherichia coli* and other bacterial species Tn7 inserts itself in single copy, and with high fidelity, into a specific and apparently unimportant intergenic region (8, 13, 22, 33, 43), we adapted it as a useful tool to chromosomally mark *V. fischeri* strains without affecting their biological fitness.

In this study, we determined the efficiency of *V. fischeri* infection of juvenile squid and used this information, as well as two different mini-Tn7-tagged strains, to track symbiont population dynamics during the early stages of infection. Specifically, we determined (i) the number of cells associated with a
juvenile light organ in the first few hours following exposure to *V. fischeri*, (ii) the influence of inoculum size on the onset and ultimate level of animal luminescence within the first 48 h after inoculation, and (iii) the role that inoculum size plays in determining whether a symbiotic population is derived from a single bacterial cell.

**MATERIALS AND METHODS**

**Bacterial strains and media.** Unless mentioned otherwise, *V. fischeri* cells were grown with shaking at 28°C in either seawater-trypotone medium (SWT), which contained 5 g of Bacto Tryptone-Peptone (Difco, Inc., Sparks, Md.), 3 g of yeast extract, and 3 ml of glycerol per liter of 70% seawater (5), or LBS, which contained 10 g of Bacto Tryptone-Peptone, 5 g of yeast extract, 50 ml of 1 M Tris base (Sigma Chemical Co., St. Louis, Mo.) at pH 7.5, and 20 g of NaCl per liter (10). Strains of *E. coli* were grown in either Luria-Bertani medium (7) or brain heart infusion medium (Difco). All strains were stored at −80°C.

**Sample collection.** Squid used in this study were obtained from Waikiki Aquarium, Honolulu, Hawaii) for at least 30 min prior to inoculation of hatching from eggs laid in the laboratory and were held in natural seawater (31). To determine the 50% infective dose (ID50), determinations were carried out by inoculating 5-mm juvenile squid (approximately 1 mm in mantle length) were collected using a plastic eyedropper within minutes of hatching from eggs laid in the laboratory and were held in natural seawater (5929) required for colonization of 50% of the animals, juvenile squid were placed in vials containing 4 ml of seawater inoculated with serial dilutions of a culture of *V. fischeri* that previously been digested with EcoRV and allowed to self-ligate, resulting in the loss of its Em’ gene. After ligation, the resulting mini-Tn7-Cmr plasmid, pJRM3, was transfected into *E. coli* strain BW23474 and mobilized by conjugation into *V. fischeri* ES114 as described above, producing the Cmr-marked strain JRM200. The number and location of six independently arising mini-Tn7-Cmr insertions were determined by Southern blotting and/or PCR amplification. Briefly, 3 μg of genomic DNA was subjected to restriction enzyme digestion, separated by electrophoresis, transferred to a nitrocellulose membrane, and analyzed by Southern hybridization (38). The PCR amplification product from primers JRM3 and JRM4 (Table 1) was used as a probe. The specific insertion sites of the transposon into the chromosome of *V. fischeri* were determined by sequencing from either end of the transposon by using primers J1 and J2 (Table 1).

**Competition assays of Tn7-tagged strains in culture.** The ability of the transposon-tagged strains to grow as well as wild type in liquid medium was determined as follows. *V. fischeri* strains ES114, JRM100, and JRM200 were grown separately with shaking for 18 h in SWT at 28°C. Approximately equal numbers of cells of each strain were then combined, the mixture was diluted 1:100 in fresh SWT, and the culture was split into three flasks. After approximately 20 h of shaking, 1 ml of each culture was transferred into a new flask containing 15 ml of fresh SWT. To determine the relative ratio of strains in the culture at various times during growth, three 100-μl samples of each flask were removed, serially diluted, and spread onto SWT agar plates. Representative CFU were screened for antibiotic resistance and identified as either strain JRM100, strain JRM200, or strain ES114.

**Animal colonization experiments.** Juvenile *E. scolopes* squid (approximately 1 mm in mantle length) were collected using a plastic eyedropper within minutes of hatching from eggs laid in the laboratory and were held in natural seawater (Waikiki Aquarium, Honolulu, Hawaii) for at least 30 min prior to inoculation with strains of *V. fischeri*. This water contained fewer than 10² CFU/ml and did not harbor *V. fischeri* cells in sufficient numbers to colonize the squid (31). To determine the 50% infective dose (ID50), defined as the number of bacteria required for colonization of 50% of the animals, juvenile squid were placed in vials containing 4 ml of seawater inoculated with serial dilutions of a culture of
V. fischeri ES114. Samples of the inoculum were spread on agar medium, and the number of CFU that arose at 28°C was counted at 24 h. After a 3-h incubation period at 22°C, the animals were rinsed and placed in vials of natural seawater. The seawater was changed every 12 to 16 h. At 48 h postinoculation the luminescence of each juvenile E. scolopes was measured with a TD20/20 luminometer (Turner Designs, Sunnyvale, Calif.). Because uncolonized squid do not produce light (27), all animals with a luminescence reading significantly above background (i.e., >1.3 × 10^{4} quanta/s) were considered colonized. In other experiments the development of colonization was determined over a 48-h period by using a liquid scintillation counter (TriCarb 2100TR; Packard Inc., Meriden, Conn.) modified to measure the level of light emission, which is an indirect measure of the number of V. fischeri cells in the light organ (27, 29).

Competition of marked V. fischeri strains for host colonization. To determine whether the transposon-tagged strains have a competitive colonization defect in symbiosis, equal numbers of V. fischeri ES114 and either JRM100 or JRM200 organisms were coinoculated into seawater containing 16 juvenile E. scolopes squid and incubated as described above. The inoculum was spread onto SWT agar, and the resulting CFU were screened for antibiotic resistance to determine the ratio of ES114 and the mini-Tn7-marked strains in the inoculum. Approximately 24 h postinoculation, each animal was homogenized in 700 μl of sterile seawater in a 1.5-ml Eppendorf tube, and 100 μl of this homogenate (or serial dilutions of it) was spread on SWT agar to determine the number of V. fischeri CFU per light organ (34). When exposed for as little as 3 h to about 250 V. fischeri cells per squid, an average of 50% of juvenile E. scolopes squid became colonized (Fig. 1), as indicated by the appearance of host luminescence. Colonization was rare at inoculum levels below 100 bacteria per animal, but colonization efficiency rapidly improved as the inoculum size increased, reaching 100% at levels above 1,000 bacteria. Among juveniles that became infected, those exposed to higher doses of V. fischeri cells became luminescent earlier, and reached maximal luminescence sooner, than those animals that were exposed to a lower dose (Fig. 2). However, this enhancement reached a limit, and exposing animals to more than 36,000 cells did not lead to a more rapid colonization. Importantly, all animals became equally luminescent by the end of the experimental period, indicating that, while any of a wide range of doses is sufficient to initiate colonization, the length of time required for onset of host luminescence is inversely correlated with the size of the dose of V. fischeri administered.

Effect of mini-Tn7 carriage on V. fischeri ES114. We created genetically marked, but otherwise apparently normal, strains of V. fischeri so that we could describe the nature of the population of bacteria involved in the initiation of squid light-organ colonization. Based on the biology of Tn7 in other systems, we hypothesized that the mini-Tn7-Em^{R} constructs in V. fischeri ES114 would have a single chromosomal insertion site and that the insertion would not affect the ability of bacteria to grow or colonize. To test this hypothesis, the genomic fragment carrying the transposon was cloned and sequenced from each of five independently isolated clones (designated JRM100 to JRM104), revealing that the point of insertion was in essentially the same noncoding, intergenic region (Fig. 3). This region was downstream of an open reading frame encoding a protein with 76% amino acid identity to the glmS product in E.
coli and with 88% amino acid identity to the putative glmS product in Vibrio cholerae (2, 15). Similarly, PCR analysis of the insertion in JRM200 confirmed that the Cm\(^R\)-containing form of the transposon was inserted at the same genomic site (data not shown). Southern analysis confirmed that each strain received a single insertion.

To determine whether carriage of the insertion created a significant growth burden, we examined the effect of the transposon on cells growing in culture. Two Tn7-containing strains (JRM100 and JRM200) and their wild-type parent (ES114) were coinoculated into SWT, and the ratio of the strains to each other was determined during several rounds of subculturing. The starting ratio of strains was maintained throughout the 44 h of batch culture (data not shown), indicating that over this period neither the relative growth rate nor survival in stationary phase was significantly affected in the transposon-bearing strains.

**Light-organ colonization by Tn7 strains of V. fischeri.** To examine whether insertion of the mini-Tn7 affected symbiotic competence, wild-type V. fischeri was coinoculated with either strain JRM100 or strain JRM200 into seawater containing juvenile E. scolopes. After 24 h, the mini-Tn7 insertion mutants had colonized the light organ to the same degree as had the wild type, as indicated by a relative competitive index value of close to 1.0 (Table 2). We also examined the colonization ability of V. fischeri ESR1 (14), a spontaneously occurring R\(^R\) mutant of V. fischeri ES114, to determine whether it could also be used as a marked strain in multistrain colonization experiments. However, unlike the mini-Tn7-marked strains, ESR1 comprised a significantly smaller portion of the squid light-organ population than it did of the initial inoculum (Table 2), demonstrating that ESR1 could not compete effectively with its wild-type parent during colonization. For this reason V. fischeri ESR1 was not used in the multistrain infection experiments described below.

![FIG. 3. Site of mini-Tn7 insertions in the V. fischeri genome.](image)

**TABLE 2. Relative colonization abilities\(^a\) of V. fischeri strains carrying antibiotic resistance**

<table>
<thead>
<tr>
<th>Strain</th>
<th>24 h RCI(^a)</th>
<th>48 h RCI(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>JRM100</td>
<td>0.91 ± 0.29</td>
<td>ND(^d)</td>
</tr>
<tr>
<td>JRM200</td>
<td>0.76 ± 0.28</td>
<td>ND(^d)</td>
</tr>
<tr>
<td>ESR1</td>
<td>0.05 ± 0.82(^c)</td>
<td>0.03 ± 0.16(^c)</td>
</tr>
</tbody>
</table>

\(^{a}\) Colonization ability was measured during competition with the parent wild-type strain ES114.

\(^{b}\) RCI, percentage of a given strain in the light organ divided by percentage of that strain in the inoculum at 24 or 48 h postinoculation.

\(^{c}\) ND, not determined.

\(^{d}\) The ability to colonize the light organ was significantly less than that of wild-type V. fischeri (\(P < 0.05\)).
with the mucus or light-organ tissue, a value that compared well with CFU determinations (data not shown). However, animals of the same cohort that had been transferred into natural seawater after exposure to the *V. fischeri* inoculum for either 3 or 12 h became fully colonized. Therefore, at the lower doses (below 1,000 CFU/ml) described in this study, it appears that bacterial cells are infecting as individuals or a few cells and not aggregating into large groups prior to entering the light organ.

**Simultaneous colonization with three *V. fischeri* strains.** An objective of this study was to describe the numbers and composition of the founding population of bacteria that can colonize the light organ. The experimental results described above suggested (i) that only a small number of bacterial cells were necessary for a successful colonization and (ii) that the number of cells that participated in the colonization was dependent on inoculum density, at least at low inoculum concentrations. To determine whether a single cell, or small group of cells, could indeed be the sole symbiotic founder in a juvenile light organ, we designed the following experiment. Groups of *E. scolopes* squid were exposed to a mixed inoculum of *V. fischeri* containing equal cell numbers of wild type (ES114) and two differentiable Tn7-carrying strains (JRM100 and JRM200), but at different total inoculum concentrations. We reasoned that, if the light organ could be populated by a small initial population, or even a single colonizing bacterium, it would be possible to find light organs inhabited exclusively by only one of the three marked strains, even though all three were in the inoculum. We found that, when animals were exposed to large numbers of *V. fischeri* cells in the mixed inoculum, light-organ populations would consist of two or more strains. However, when exposed to a small inoculum, the animals were most often colonized by only one of the three strains (Fig. 5), which suggested colonization by a single cell or a small number of cells.

**DISCUSSION**

All horizontally transmitted microbial associations, whether beneficial or pathogenic, must develop mechanisms that ensure or enhance infection of the host. Often there is an attribution of the inoculum by host defenses, such as that encountered by *V. cholerae* cells during their passage through the stomach (6). Unfortunately, even for well-studied pathogeneses, the number of cells that survive and initiate the infection has been difficult to determine, and it is usually impossible to visualize and count the founding bacterial population in host tissue. In this study we investigated the number and composition of *V. fischeri* cells taking part in the initiation of light-organ colonization in *E. scolopes*. Using genetic and microscopic approaches, we determined that the efficiency of colonization is dependent on the number of *V. fischeri* bacteria present in the inoculum and that the symbiont population can be founded by a few cells of *V. fischeri*.

**FIG. 4.** Evidence of the association of *V. fischeri* cells with host tissue 3 h after inoculation. Three sets of 24 animals were exposed to three different doses of *V. fischeri* cells: 340, 1,800, and 3,400 per animal. The number of *V. fischeri* cells (as determined by CFU) associated with light-organ tissue at 3 h postinoculation was determined (left panel) and compared to the percentage of animals that eventually became colonized at these inoculation doses (right panel).

**FIG. 5.** Effect of the total dose on the frequency of colonization by one, two, or three strains of *V. fischeri* from a mixed inoculum. Animals were exposed to the following approximate doses of *V. fischeri* cells per animal: 500 (open bars; n = 8), 5,000 (stippled bars; n = 15), 16,000 (striped bars; n = 20), and 27,000 (solid bars; n = 14). All doses contained a mixed inoculum of approximately equal proportions of *V. fischeri* strains ES114, JRM100, and JRM200. The number of different strains detected in each light organ was determined 24 h postinoculation.
Mini-Tn7-marked \textit{V. fischeri} strains. We used a mini-Tn7 to differentially mark \textit{V. fischeri} strains with two antibiotic resistance cassettes and tracked their fate during the colonization process. We demonstrated that both the growth rate and the colonization efficiency (Table 2) of the \textit{V. fischeri} mini-Tn7 insertion strains that we created were comparable to those of the wild-type parent. We also showed that, as in \textit{E. coli}, the site of mini-Tn7 insertion in \textit{V. fischeri} is an intergenic region (Fig. 3), located downstream of a glmS-like gene (8, 13, 22). Because glmS encodes glucosamine–fructose-6-phosphate aminotransferase, an important enzyme in cell wall synthesis, it has been conserved among many gram-negative bacteria (22). In previous work the transposon Tn7 has been used to insert reporter constructs such as the gene for lux (luminescence) (36) or GFP (17) into the genomes of several species of gram-negative bacteria. The ability of Tn7 to integrate into the genome of \textit{V. fischeri} strain ESR1 was demonstrated previously (43), and it has been shown to integrate into \textit{Vibrio anguillarum} as well (21). The transposon has also recently been used for genetic complementation of chromosomal mutations (9), and similar derivatives can now be used to replace the carriage of other markers (e.g., GFP [32]) on plasmids, whose maintenance requires continuous selection with antibiotics that can lower the rate of normal colonization (data not shown).

**Determination of the ID$_{50}$** \textit{V. fischeri} is effective at colonizing the light organ of \textit{E. scolopes}, requiring a dose of only about 250 cells for 3 h to infect 50% of its squid hosts (Fig. 1). Such a low ID$_{50}$ had been suggested from previous work (25, 34, 40) and indicates that colonization is an efficient process. This ID$_{50}$ is especially remarkable because the dose is not directly introduced into host tissue and is present in the surrounding seawater for only a limited time. In comparison, the analogous 50% lethal dose, which is the bacterial dose that causes death in 50% of infected hosts, is typically much higher in other \textit{Vibrio} species. For instance, \textit{Vibrio parahaemolyticus} and \textit{Vibrio alginolyticus}, two pathogens of shellfish, have reported 50% lethal doses on the order of 10$^7$ bacteria per g of host tissue (23, 24). This dose is 100-fold higher than what is required for \textit{V. fischeri} colonization and suggests an unusually efficient mechanism of symbiotic infection by these bacteria. Such a mechanism is not unexpected, considering that the natural concentration of \textit{V. fischeri} cells in the seawater inhabited by \textit{E. scolopes} is usually well below a thousand cells per ml of seawater (20). Similarly, while potential hosts resist infection by pathogenic vibrios, \textit{E. scolopes} has evolved a morphology and behavior that encourage \textit{V. fischeri} colonization (42).

\textit{V. fischeri} colonization dynamics. Previous studies of the initiation of colonization (31, 32) have revealed that the juvenile squid light organ secretes a mucus layer that preferentially entraps \textit{V. fischeri} cells present in the ambient seawater by a process that is surprisingly like that occurring between rhizobia and their leguminous hosts (12). In the squid, the action of surface cilia removes approximately 1% of the \textit{V. fischeri} bacteria from the water (S. Nyholm, personal communication), forming bacterial aggregates in the mucus that, depending on the dose, contain a few to hundreds of bacteria. These aggregates eventually migrate to pores on the light-organ surface and through ducts that lead to symbiotic crypts of the light organ. Because the ducts are sites of a potential gaulette of host-produced oxidative stress (37), the question arose as to how many of the inoculating bacteria survive the initial colonization events and become the progenitors of the resulting population of symbionts.

In the work described here mini-Tn7-tagged \textit{V. fischeri} cells were used to examine the early colonization events during the initiation of symbiosis. We focused on low inoculum levels of a few hundred or thousand \textit{V. fischeri} cells, levels at which the rate of colonization appeared to be limited by dose (Fig. 1 and 2). Consistent with previous findings (34), the small number of CFU (one to four) that arose from homogenates of juvenile squid suggested that only a few cells had become associated with the light organs under these inoculation conditions. Nevertheless, such animals typically became subsequently colonized, suggesting that only a few \textit{V. fischeri} cells needed to be present to initiate a successful colonization. Direct visualization of \textit{V. fischeri} cells in contact with host tissue correlated with the number of CFU detected by plating, arguing against the possibilities that either (i) the bacteria might be present in larger numbers but are in aggregates that are not easily separated when spread on agar or (ii) many of the cells have entered a viable but nonculturable state (20).

When animals are exposed to a dose of over 10,000 \textit{V. fischeri} cells per ml, aggregates containing hundreds of these bacteria are seen (32). At such large doses symbiotic luminescence is detected 3 to 4 h sooner than in animals infected at or around the ID$_{50}$ (Fig. 2). Based on a doubling time in the light organ of about 30 min (34), the number of cells initiating colonization at these large doses is likely to be in the hundreds. That is, it would take an inoculum of at least 256 cells to be eight generations (i.e., 4 h) ahead of an inoculum as small as one cell. The ability to infect the squid with hundreds of cells is experimentally useful because this dose is sufficiently high to make feasible in vivo expression technology approaches for identifying host-induced bacterial genes (3). Traditional genetic studies have already revealed a number of activities, e.g., flagellar motility (14, 29) and gene regulation (9), as well as gene products, e.g., an outer membrane protein (1) and a pilus (40), that are required for normal initiation of symbiosis. Future microarray-based studies are likely to identify other bacterial activities that may underlie the high symbiotic infectivity of \textit{V. fischeri}.

Taken together, the data presented here indicate that \textit{V. fischeri} cells, even at the low concentrations that may be found in nature, are highly efficient at colonization, due to a combination of intrinsic properties of the bacterium and selective properties of the host (42). As a result only a few cells need to be in immediate proximity to the light-organ tissue in order for a full colonization to occur. To understand the mechanisms responsible for this efficiency, future investigations will focus on bacterial traits that underlie the species specificity that is characteristic of this association, uncovering those genes whose functions are important in the earliest stages of contact between bacterium and squid host tissue.

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