Impact and influence of the natural Vibrio-squid symbiosis

in understanding bacterial-animal interactions

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September 2016

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Abstract

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8 Animals are colonized by bacteria, and in many cases partners have co-evolved to perform mutually

beneficial functions. An exciting and ongoing legacy of the past decade has been an expansion of

technology to enable study of natural associations in situ/in vivo. As a result, more symbioses are

being examined, and additional details are being revealed for well-studied systems with a focus on the

interactions between partners in the native context. With this framing, we review recent literature from

the Vibrio fischeri-Euprymna scolopes symbiosis and focus on key studies that have had an impact

on understanding bacteria-animal interactions broadly. This is not intended to be a comprehensive

15 review of the system, but rather to focus on particular studies that have excelled at moving from pat-

6 tern to process in facilitating an understanding of the molecular basis to intriguing observations in the

field of host-microbe interactions. In this review we discuss the following topics: processes regulating strain and species specificity; bacterial signaling to host morphogenesis; multiple roles for nitric
oxide; flagellar motility and chemotaxis; and efforts to understand unannotated and poorly annotated
genes. Overall these studies demonstrate how functional approaches *in vivo* in a tractable system have
provided valuable insight into general principles of microbe-host interactions.

Studies of human, animal, and plant microbiomes have been advanced by novel culture-independent

Introduction

approaches and technological advancements in DNA sequencing. In recent years a prominent role for microbial communities of the gut, skin, and other organs has emerged as modulators of human health (Human Microbiome Project Consortium, 2012). These studies followed from influential animal studies in systems that are yielding critical insight into microbiome assembly, stability, communication, and evolution (McFall-Ngai et al., 2013; Ruby, 2008). The focus of this review is to examine one model system, the Vibrio fischeri-Euprymna scolopes symbiosis, and how key findings in that system have enabled an increasingly higher resolution of the processes and principles that underlie microbe-host communication. When Hawaiian bobtail squid hatch from their eggs, they are exposed to a million bacteria in each milliliter of seawater. Although V. fischeri make up less than 1 in 5,000 of these planktonic, environ-33 mental bacteria, the "light organ" of the hatchling squid becomes colonized exclusively with V. fischeri (Mandel, 2010; Ruby and Lee, 1998). The microbe-host specificity relies on a series of reciprocal communications between the partners, many of which are detailed in the sections below. Over the course of 48 hours the bacteria establish a mature colonization in epithelium-lined crypts of the squid light organ, and, at high cell density, produce light as a result of quorum-sensing. The bacterial bioluminescence is reflected by host tissue to camouflage the shadow or silhouette that the nocturnal-foraging squid would cast in the moonlight, thus protecting the host in a process termed counter-illumination (Jones and Nishiguchi, 2004; Ruby and McFall-Ngai, 1992). Initiation of colonization occurs in newly-hatched

squid, seeding an individual host's crypts for its lifetime. The bacteria produce light at night, then at dawn approximately 90-95% of the symbiotic population is expelled into the seawater (Boettcher et al., 1996; Lee and Ruby, 1994; Nyholm and McFall-Ngai, 1998). The remaining cells grow up during the day, produce light at night, and a diel cycle of growth, light production, and expulsion proceeds for the lifetime of the animal (Wier et al., 2010). Host cellular changes accompany this cycle, e.g. a daily reshaping of the epithelial brush border against which the bacteria reside during the final two hours prior to the daily expulsion (Wier et al., 2010).

As an environmentally-transmitted symbiosis, the *Vibrio*-squid model has a number of valuable characteristics that have served it well as a study system for identifying molecular mechanisms. First, the
binary system (two partners) is naturally reduced. Second, both partners can be raised separately and
then introduced for experimentation. Third, *V. fischeri* is genetically tractable, and unbiased mutagenesis as well as precise genetic alterations can be introduced with relative ease. Fourth, the bacteria colonize the host light organ directly under the semi-transparent mantle and funnel; this permits imaging
of the site of infection and direct analysis of bacterial behaviors and host responses. Fifth, synchronous
colonization of hatchlings has permitted developmental staging of the colonization process. For most
of the processes described below, many of these benefits were important in the advances described.

From pattern to process in the *Vibrio*-squid symbiosis

In each section below, we highlight key discoveries in the *Vibrio*-squid symbiosis with a specific focus on how this model system has revealed molecular processes that underlie mutually beneficial phenotypes.

Just the two of us

E. scolopes squid light organs are colonized only by *V. fischeri*, and this exclusivity has guided substantial inquiry and discovery in the system. This pattern was first explored by McFall-Ngai and Ruby

(McFall-Ngai and Ruby, 1991) and extended in subsequent works (Mandel et al., 2009; Ruby and Lee, 1998). The ability to image the live animal during colonization enabled the discovery of *V. fischeri* aggregating in close proximity to the ciliated epithelial fields of the light organ (Nyholm et al., 2000). Nyholm discovered that a narrow distance between the green fluorescent protein-expressing bacteria and the squid epithelial tissue was the result of host-produced mucus, which included N-acetylneuraminic acid and N-acetylgalactosamine. Recent work has demonstrated that V. fischeri bind to cilia within this mucus field (Altura et al., 2013). Whereas many bacteria can bind in host mucus, only specific 71 strains and species exhibit a competitive dominance over non-colonizing isolates, and only (some) V. fischeri strains proceed to fully initiate colonization (Mandel et al., 2009; Nyholm and McFall-Ngai, 2003; Nyholm et al., 2000). Around this same time, the genetic basis for bacterial aggregation was being discovered and characterized in the laboratory of Karen Visick. A forward genetic screen for colonization factors first identified an orphan histidine kinase, RscS (regulator of symbiotic colonization-sensor), but without a phenotype or target it was difficult to know how this factor connected to the colonization process (Visick and Skoufos, 2001). The same screen identified an eighteen gene locus that encoded regulatory proteins, glycosyltransferases, and other factors involved in exopolysaccharide production and export. Mutations in this region, the syp locus (symbiosis polysaccharide), conferred dramatic colonization defects in the animal as well as defects in biofilm formation in culture (Yip et al., 2005). A connection between these earlier studies was discovered when it was shown that RscS regulates expression of the syp locus (Yip et al., 2006). Overexpression of RscS provided a valuable tool in which bacterial colony formation took on a wrinkled or rugose colony morphology that is typical of biofilm formation (Yip et al., 2006). Phenotypes of rscS and syp alleles in colony-based biofilm assays map closely to their phenotypes during squid colonization, providing a valuable experimental tool for discovery and characterization of biofilm regulation. Further work has identified multiple layers of regulation, including a negative regulatory pathway that includes SypE and SypA, putative matrix proteins that integrate with the polysaccharide matrix, and a unique phosphorelay pathway (Morris and Visick, 2013; Norsworthy and Visick, 2015; Ray et al., 2015; Visick, 2009).

The genetic approaches described above (and in most studies in this review) were conducted in strain ES114, a squid isolate from Kaneohe Bay, Hawaii, that is used widely as a canonical squid symbiont. In addition to the biofilm regulatory pathway, a number of approaches including forward and reverse genetics studies had identified factors in strain ES114 that were important for squid colonization (Stabb and Visick, 2013). However, only some V. fischeri strains can colonize squid. Therefore, to examine the genetic basis for this host colonization specificity, Mandel and colleagues conducted a comparative 97 genomic analysis of strains ES114 and MJ11, the latter being a fish symbiont that does not colonize squid robustly (Mandel et al., 2009). The study determined that 91 % of ES114 genes were almost identical between the squid and fish symbiont, but that approximately 400 genes in each strain were 100 unique. Analysis of these factors revealed that the squid biofilm regulator, RscS, was encoded in the 101 squid symbiont but not in the fish symbiont. The known RscS target genes, sypA through sypR were en-102 coded in both genomes and fairly conserved (>85 % amino acid identity). It was known previously that 103 ES114 mutants that lacked RscS were unable to productively colonize the squid (Visick and Skoufos, 104 2001). Therefore, the study asked whether the absence of the regulator could explain the differential 105 colonization phenotype. Introduction of RscS into strain MJ11 was sufficient to allow it to colonize 106 the squid host. Phylogenetic analyses supported a model in which MJ11 was part of an ancestral group 107 of V. fischeri that lacked rscS, and that this gene was acquired coincident with colonization of squid in the North Pacific Ocean (i.e., Japan and Hawaii).

The idea that a single gene was sufficient to shift the animal hosts available to a bacterium was extreme 110 but consistent with emerging literature that individual loci could impact microbe-host specificity. Work 111 in entomopathogenic nematodes showed that symbiotic *Xenorhabdus nematophila* requires the three-112 gene nilABC locus for colonization, and that expression of these factors in a heterologous symbiont 113 is sufficient to enable colonization of Steinernema carpocapsae, the worm host that otherwise is spe-114 cific for X. nematophila (Cowles and Goodrich-Blair, 2008). Small genetic changes in Yersinia pestis 115 have been key to its ability to colonize new niches, including single gene acquisitions and even in-116 activation of a gene already present (Sun et al., 2008, 2014; Zimbler et al., 2015). In the human gut 117 microbiome there are examples in which single gene changes have been critical; e.g., in Bacteroides 118

fragilis, polysaccharide A (PSA) confers a key immunomodulatory benefit that cannot be obtained from the other seven capsular polysaccharides produced (Mazmanian et al., 2008).

Studies on host colonization specificity in general, and biofilm formation in particular, have highlighted many of the strengths of the squid model. Imaging *in situ* was key to the initial discovery
of the aggregates, forward genetics identified core exopolysaccharide synthetic and regulatory components, comparative genomics revealed the role of this pathway in the evolution and specificity of
the association, and high-throughput genetic approaches are identifying additional levels of regulation.
Additionally, this work highlights the value of model systems of beneficial bacteria, including *Vibrio*and *Xenorhabdus* models, to identify mechanistic details that resonate in beneficial and pathogenic
colonization models.

9 The Code Word is TCT

E. scolopes squid provide a particularly dramatic example of a role for bacteria influencing a spe-130 cific host developmental process. Development of the host tissue proceeds on different trajectories 131 depending on whether the specific symbiont V. fischeri is present. Only once the symbiont has colonized, the ciliated appendages of the host light organ undergo apoptosis, hemocyte infiltration, and 133 tissue regression during the subsequent five days (Koropatnick et al., 2004; McFall-Ngai and Ruby, 134 1991; Montgomery and McFall-Ngai, 1994). The host morphogenesis is striking, with appendages that begin as outstretched mucus factories to recruit colonizing bacteria being reduced to small stumps 136 (Montgomery and McFall-Ngai, 1994). As a result, it seems that initiation of the symbiosis is restricted 137 to the first few days of the animal's life while the appendages are present and secreting mucus. 138

How does the host know that the bacteria are inside to appropriately time the regression? It turns out that *V. fischeri* sheds envelope components that are received by receptors on the host. In particular, the bacterial peptidoglycan fragment, tracheal cytotoxin (TCT)–previously shown to induce a damaging apoptosis in ciliated epithelia upon release from *Bordetella pertussis*—was identified to perform a similar function in *V. fischeri*, but this time with a resulting beneficial outcome (Koropatnick et al., 2004).

To recapitulate the apoptosis phenotype observed when intact *V. fischeri* are presented to the host, in the absence of the bacteria both the Lipid A portion of lipopolysaccharide (LPS) and TCT are required. The cell death from these compounds, in conjunction with hemocyte trafficking that is also induced from TCT, results in the regression phenotype. Previously these compounds had only pathogenic associations, but this work underscored a remarkable conservation to the cell biology of microbial-host interactions, emphasizing the context of the interaction to understand the fitness effects on the partners involved (Koropatnick et al., 2004).

Once the bacteria announce their arrival, how does the host speak back? In addition to regression of
the appendages that recruit the bacteria, there are additional mechanisms by which the host receives
and likely modulates the bacterial signal. Host nitric oxide production, described in more detail below, is diminished as a result of bacterial signaling (synergistically with LPS) (Altura et al., 2011).
The host produces a peptidoglycan recognition protein, EsPGRP2, which is secreted into the bacterialcontaining crypts and has the ability to degrade TCT (Troll et al., 2010). Additionally, there are data
to suggest that host alkaline phosphatase, EsAP, modifies Lipid A after the initial signaling (Rader et
al., 2012). In each case the host response is to diminish the potency of the bacterial products, but only
after they have exerted their influence on host development.

This work in *V. fischeri* was influenced by studies in invertebrate systems that demonstrated host devel-160 opment in response to symbiont colonization and in vertebrates that showed general responses to con-161 sortia (reviewed in (Montgomery and McFall-Ngai, 1994) and more recently in (McFall-Ngai, 2014)), 162 and itself has influenced a field in which bacterial products play important roles in animal development. 163 An early mammalian example by Hooper and Gordon demonstrated that in response to colonization by 164 gut Bacteroidetes such as Bacteroides thetaiotaomicron, terminal tissue differentiation (e.g., fucosy-165 lation) is dependent on the presence of the symbiotic bacteria (Hooper and Gordon, 2001). There now 166 exist many examples of bacteria directing specific host development. Recent exciting examples include Algoriphagus machipongonensis sulfonolipid signaling for multicellular rosette development in the choanoflagellate Salpingoeca rosetta, and Pseudoalteromonas luteoviolacea phage tail-like structures that stimulate tubeworm metamorphosis (Alegado et al., 2012; Shikuma et al., 2014).

71 NO way in

There is a long history of the study of nitric oxide (NO) in eukaryotes, and this small diffusible molecule 172 has been implicated in many different cellular processes including signaling and innate immunity (Fang, 173 2004). Although the roles for NO in eukaryotic physiology and defense against pathogens were dis-174 covered many years ago, the study of this compound in the Vibrio-squid system and other symbioses 175 (Damiani et al., 2016) has revealed that NO also influences the establishment and maintenance of mu-176 tualistic microbe-host relationships as both a signal and a specificity determinant (Wang and Ruby, 177 2011). 178 Davidson, et al. (Davidson et al., 2004) first demonstrated that NO is produced in squid host tissue 179 through the activity of nitric oxide synthase (NOS), and this activity was attenuated after successful colonization by V. fischeri. Using staining and immunocytochemistry, NOS and NO were found located in the epithelium of the light organ, as well as in vesicles within mucus shed from these cells. 182 It is within this mucus that the bacterial cells aggregate prior to entering the light organ. Normally, 183 V. fischeri aggregate in the mucus, colonize the host, and after successful colonization NOS activity 184 and NO production are attenuated. Treatment of the animals with an NO-scavenging compound to di-185 minish NO levels allowed large aggregates of non-symbiotic vibrios to form, but these bacteria did not 186 successfully initiate colonization. (Davidson et al., 2004) The results suggested that NO acts as a speci-187 ficity determinant, helping to limit aggregation of non-symbiotic vibrios and select for symbiotically 188 competent *V. fischeri* from the mixed microbial population found in seawater. 189 If NO plays a role in specificity, then how do colonizing V. fischeri sense and respond to the host-190 produced NO to successfully establish the partnership? Using genetic approaches it was demonstrated 191 that a strain lacking the NO-detoxifying enzyme flavohemoglobin (Hmp) displayed a colonization deficiency (Poole and Hughes, 2000; Wang et al., 2010b). Expression of hmp is regulated by the NOresponsive negative regulator NsrR (Rodionov et al., 2005; Tucker et al., 2010). However, NsrR is not the only important NO-sensing regulator in *V. fischeri*. H-NOX, a heme NO/oxygen-binding protein,

also plays a role in symbiotically relevant NO-responsive regulation of genes in V. fischeri (Wang et

al., 2010a). Although H-NOX-like proteins are widely distributed in bacteria, this was the first report describing bacterial H-NOX function. Interestingly, it appears that one role for H-NOX in V. fischeri 198 is to sense NO and correspondingly suppress bacterial hemin uptake during the early stages of host 199 colonization. The authors predicted that early repression of iron uptake would protect the cells from 200 the potentially harmful effects of Fenton chemistry when they are exposed to host-generated oxidants 201 (Davidson et al., 2004; Graf and Ruby, 2000; Wang et al., 2010a). Consistent with this model, hemin 202 uptake genes in *V. fischeri* were shown to be induced during the later stages of symbiotic colonization, 203 and deletion of these genes negatively impacted colonization (Septer et al., 2011). Together, these 204 studies support a model whereby host NO stimulates repression of hemin uptake genes; once bacterial 205 colonization leads to an attenuation of host oxidant production, then hemin uptake genes are dere-206 pressed to support growth in the iron-limited light organ environment. Therefore, the ability to sense 207 and detoxify NO is important for symbiotic specificity, and NO acts as a temporal signal to modulate 208 bacterial gene expression and promote successful colonization. 209

Although these studies have led to a better understanding of the role of a few key proteins and regu-210 lators in the response of V. fischeri to NO and the initial stages of the symbiosis, there is much yet to be learned about the global effects of NO on V. fischeri gene expression and metabolism, how this molecule acts as a specificity determinant, and whether there is a role for NO in the mature symbiosis. 213 For example, the work of Wier et al. has suggested that NO may play a role in the daily symbiotic 214 rhythm in the adult animal (Wier et al., 2010). Their data predicted that nitrate/nitrite respiration is 215 used by the bacterial symbionts throughout the daylight hours. Similarly to Escherichia coli (Vine 216 and Cole, 2011), it is predicted that NO is produced by V. fischeri during respiration of nitrate/nitrite 217 in laboratory culture. Endogenously-produced NO could induce alternative respiratory pathways that 218 likely influence the physiology and metabolism of the bacterium (Dunn et al., 2010). Together these 219 separate lines of evidence suggest that NO may play a role beyond signaling and selection in the initi-220 ation of the symbiotic relationship. In the future it will be exciting to combine studies of NO and the 221 bacterial NO response with the more recently developed ability to rear squid to adulthood ((Koch et 222 al., 2013); see section below on light production).

The value of further studies of NO in the *Vibrio*-squid system lie not only in providing important information about the role of this molecule in beneficial host-microbe interactions, but also for comparative studies to host-pathogen responses. Our current understanding supports a view that NO is being produced by the host and sensed by the bacteria in similar ways in many of the studied host-microbe interactions, whether the outcome of the relationship is beneficial or detrimental (Fang, 2004; Wang and Ruby, 2011). The prevalence of NO in host tissues colonized by bacteria suggests that a better understanding of the role of NO in symbiosis may have wide-reaching consequences for microbes at the interface of health and disease.

232 Swimming against the flow

In the mucus field that serves as the entry point for bacteria heading into the host, colonizing bacteria 233 enter at one of three pores on either side of the bilaterally symmetrical light organ. Mucus is shed from 234 the pores of the host at the same time that V. fischeri aggregates in that mucus. The bacteria proceed to 235 migrate toward the pores, and each aggregate swims into a pore to colonize the ducts and crypts of the 236 host. How do colonizing bacteria travel against this powerful flow? A key role for flagellar motility 237 was identified over twenty years ago (Ruby and Asato, 1993). In that work Ruby and Asato confirmed 238 that planktonic *V. fischeri* were motile due to a polar tuft of sheathed flagella. However, by 24 hours-239 post-inoculation most cells in the light organ crypts were non-flagellated. Upon expulsion of bacteria 240 from the host, the bacteria regrow their flagella in 45-60 min even in nutrient-deplete seawater (Ruby and Asato, 1993). Therefore, the bacterial life cycle alternates between a motile planktonic lifestyle and a non-flagellated crypt-colonized state.

Significant details have since been elucidated about the molecular mechanisms that control flagellar development in *V. fischeri*, which in turn has solidified the importance of swimming motility for squid colonization. Random transposon mutagenesis provided evidence that nonmotile mutants could not colonize (Graf et al., 1994), and reverse genetics revealed that mutants defective for flagellar motility or chemotaxis did not establish productive colonization with the squid host (DeLoney-Marino and Visick,

²⁴⁹ 2012; Millikan and Ruby, 2003, 2004). Together these studies established a model of a hierarchy of flagellar gene expression in *V. fischeri* controlled by the σ54-dependent regulator FlrA. There is evidence for regulation by quorum sensing and magnesium, and other sensory inputs are likely (Cao et al., 2012; O'Shea et al., 2005).

Bacterial flagellar motility often occurs in a directed fashion in which rotation of the flagellar bundle 253 results in net movement toward preferred nutrient sources. Given the above information that chemo-254 taxis was required for colonization, it seemed likely that the bacteria were swimming toward a host 255 compound. The first evidence for chitin oligosaccharides as the specific attractant was obtained when 256 addition of exogenous chitobiose, the N-acetylglucosamine dimer, blocked colonization, whereas the 257 monomer did not have such an effect (Mandel et al., 2012). Given that N-acetylglucosamine is abun-258 dant on eukaryotic cell surfaces, yet chitin and its breakdown oligosaccharides are more specialized 259 in their localization, it seemed possible that oligosaccharides may be a specific cue to direct entry into the host crypts. Mutants defective for chemotaxis remained at the outer face of the light organ pore, the same stage at which wild-type V. fischeri arrested their symbiotic development in the presence of added chitin oligosaccharides (Mandel et al., 2012). These results strongly suggested that host chitin served as a signal for the bacteria to enter the pore. Direct imaging revealed the presence of insoluble chitin bound to hemocytes within the host (Heath-Heckman and McFall-Ngai, 2011; Mandel et al., 265 2012), which may be released through the action of a host endochitinase (Kremer et al., 2013). To-266 gether, this illustrates a specific colonization checkpoint that is regulated by both host and symbiont 267 factors. 268

Work on bacterial motility at the host interface has provided a valuable toolset to probe mechanisms of symbiosis and reveal novel signaling pathways. Many bacterial strains have dozens of genes that encode chemotactic sensory proteins, the methyl-accepting chemotaxis proteins (MCPs). The set of 43 MCPs in *V. fischeri* is typical in this regard, and despite difficulties in studying a large protein family, functions have now been assigned to three of these proteins. VfcA is the major amino acid chemoreceptor, and VfcB and VfcB2 are fatty acid chemoreceptors (Brennan et al., 2013; Nikolakakis et al., 2016). In addition to providing information directly about colonization, these tools provided

insight into the role of LPS during colonization and for the evolution and the generation of torque at
the flagellar motor (Beeby et al., 2016; Post et al., 2012). Furthermore, recent work suggests that the
rotation of the flagella—which is enclosed in an LPS sheath—stimulates outer membrane vesicle release
and triggers the host immune response by promoting LPS release (Aschtgen et al., 2016; Brennan et
al., 2014).

Satisfying answers to some of these questions are beginning to be addressed, including a role for cilia in modulating adhesion, as well as chemotaxis toward host-produced and host-cleaved chitin modulating a key developmental checkpoint. Still, important questions remain that suggest novel and interesting biology to be revealed through the symbiosis. Open questions include how bacteria transit through the mucus in a flagellar-independent manner; the molecular basis of chitin oligosaccharide sensing in the symbiont; and the processes that regulate the developmental switch between the aflagellate state in the host versus the swimming state in seawater.

88 Light up my life

An important aspect to mutualistic symbioses is the selection of appropriate and cooperative partners.

In both the rhizobium-leguminous plant (Kiers et al., 2003) and *Vibrio*-squid symbioses the microbial

partners provide costly services to their hosts (nitrogen fixation and light production, respectively). In

theory, these relationships could be exploited by symbionts that are less cooperative (i.e. "cheaters")

(Ghoul et al., 2014). However, it is rare to find bacterial symbionts associated with the hosts that do

not provide these services. Therefore, the *Vibrio*-squid mutualism provides an excellent model system

for studying cooperative partner stability, and studies to date indicate that bacterial light production is

required for bacterial cells to persist in the light organ.

V. fischeri is known to produce light in the squid host, and a key study demonstrated a role for luciferase, the enzyme that produces light, in bacterial symbiotic persistence (Visick et al., 2000). Mutants with defective luminescence structural genes or luminescence regulatory genes colonized juvenile squid to the same levels as wild type in the first 24 hours. However, by 48 hours there was a three- to four-fold reduction in colonization by the dark mutants relative to wild-type controls. In squid co-colonized with
both a luminescence mutant and wild type, levels of the mutant strains similarly decreased, indicating
that light-producing wild-type cells in the light organ could not complement the colonization defect of
the light-deficient cells. These results suggested that the ability of individual bacteria to produce light
was important for persistence in the light organ, and that somehow non-luminescent cells are selected
against during development of the symbiosis.

Interestingly, the light-deficient strains have a specific effect on host development. Although colonization by a luminescence mutant still triggered apoptosis-related developmental changes in the ciliated surface of the light organ, colonization of the tissue by these strains no longer increased cell swelling of the epithelial cells lining the light organ crypt spaces. Therefore, light production appeared to play a specific role in host developmental pathways. Notably, this was the first report of *V. fischeri* genes required for induction of bacterial-triggered differentiation of host tissue (Visick et al., 2000).

It was later discovered that the antibiotic markers and method for constructing the early luminescence mutants (Visick et al., 2000) resulted in colonization attenuation and pleiotropic effects. In a later study, newly developed genetic tools were used to construct luminescence mutants that were not negatively affected in growth and colonization (Bose et al., 2008). Using these strains, the early results were confirmed demonstrating that the strain lacking the luminescence structural genes displayed a four-fold reduction in colonization as compared to wild type at 48 hours-post-inoculation.

Previous studies suggested that maintenance of the symbiosis over the life of the animal requires a maturation process of several weeks (Montgomery and McFall-Ngai, 1998), leaving the question of how production of light influences symbiosis maturation beyond 72 hours. A major breakthrough for the field came with the development of protocols for simplified rearing of newly-hatched juvenile squid through and beyond the maturation process. These methods allowed investigation of how bacterial-produced light affects the development of the symbiosis over four weeks (Koch et al., 2013). In these studies, the levels of the luminescence-deficient mutant associated with the squid light organ continued to diminish over time, to the minimum level of detection after 28 days. Similar results were observed

in squid colonized with mixed inocula containing both wild type and the luminescence mutant, where
after 15 days the mutant was barely detected. Therefore the persistence defect observed during early
colonization becomes more pronounced as the symbiosis matures, with eventual loss (or near loss) of
non-luminescent strains in a matter of weeks.

Luminescence regulation is one of the hallmarks of the V. fischeri-squid symbiosis and has been studied 331 intensively, yet there are still exciting open questions. First, how are the dark mutants removed from 332 the population even in the midst of neighboring bright populations? A clue comes from studies testing 333 the influence of a previous colonization event on recolonization (Koch et al., 2013). Juvenile animals 334 were colonized with either wild type or a luminescence mutant. After 1-5 days, the animals were 335 treated with antibiotics to clear bacteria from the light organ and then exposed again to wild-type V. 336 fischeri to test whether light production is a "signal" to the host that influences symbiotic maturation. 337 Animals treated with antibiotics after one day were readily recolonized, regardless of the strain that initially colonized. However, after five days, wild-type V. fischeri induced a refractory state in the animal that prevented recolonization. In contrast, in animals initially colonized by a luminescence mutant, greater than 80% of the animals were recolonized by wild type. These results support the idea that the host is detecting light production by bacterial cells and/or is altering physiological conditions to sanction the non-luminescent strains. In addition, the host apparently is able to "eject" an inappropriate light deficient strain-directly or indirectly-while allowing future recolonization by a symbiotically 344 appropriate light-producing strain. The exact mechanisms by which the detection, sanctioning, and/or 345 ejection occurs remain to be described. The host does have the capacity to detect light but it is unknown 346 whether this capacity is connected to symbiont selection (Tong et al., 2009). 347

A second interesting question relates to how bacterial light production is matched to the moonlight in such an exquisite fashion. The squid contains elaborate tissues to physically reflect and modulate bacterial light production (Crookes et al., 2004). This physical response could be triggered through the activity of products of host cryptochrome and eye-specification genes; the expression of these genes appears to be influenced by the light produced by *V. fischeri* (Heath-Heckman et al., 2013; Peyer et al., 2014). The physical reflection and modulation of bacterial luminescence is also coordinated with

a molecular signaling response. For example, host epithelial cells swell in response to light-producing strains but not dark mutants (Visick et al., 2000). This swelling could release chemical cues into the 355 light organ environment. Recent evidence indicates that bacterial luminescence in the light organ is 356 controlled not only through quorum sensing, but also through response to environmental signaling 357 (Septer and Stabb, 2012). These results suggest there is complex chemical and physical control of 358 light production in the symbiosis. Bacterial luminescence is a particularly intriguing and engaging 359 aspect of the Vibrio-squid symbiosis, and it is clear that there are abundant questions remaining to 360 be addressed as to how the interaction with the host and the environment lead to specific phenotypic 361 output in the host. 362

Nice to meet you... now what is it you do?

The *Vibrio*-squid symbiosis has provided a useful framework for identifying the function of bacterial genes and studying novel genes *in vivo*. Due to the wealth of genetic tools that have been developed for *V. fischeri* and the ability to access the host interface with direct imaging, it is possible to test the effects of gene loss in the real-world environment of the host. Two examples discussed below are using the *Vibrio*-squid system to broaden understanding of gene function for alternative oxidase (AOX) and for discovering the role of the biofilm inhibitor BinK.

AOX is a terminal respiratory oxidase that is ubiquitous in plants, and is unusual because its activity is not directly linked to generation of the proton motive force (Vanlerberghe and McIntosh, 1997). 371 The study of the function of AOX in plants is an active area of research, and AOX function has been 372 linked to both abiotic and biotic stress responses (Vanlerberghe, 2013). Only with the explosion of 373 genome and metagenome sequencing was it discovered that certain bacterial genomes also encode 374 this protein (Stenmark and Nordlund, 2003), and that aox-like genes are abundant in metagenomic 375 sequences from ocean surface waters (McDonald and Vanlerberghe, 2005). However, early progress 376 towards understanding the physiological benefit of AOX function in bacteria was limited by the lack of 377 genetic tools for many of the AOX-encoding organisms. A path to revealing a functional role for AOX 378

came with the discovery that the genome of *V. fischeri* strain ES114 encoded AOX (Ruby et al., 2005). A transcriptomic analysis of the *V. fischeri* response to NO revealed that nitric oxide induces expression of aox (Wang et al., 2010a). The connection to NO was further clarified through characterization of the 381 role of the NO-responsive negative regulator NsrR in regulation of aox expression, and identification 382 of the ability of *V. fischeri* AOX to function as an NO-resistant oxidase (Dunn et al., 2010). Despite the 383 known connections between aox and NO, and between NO and the early stages of host colonization, 384 no discernible phenotypic difference between the aox mutant and wild type in early colonization of 385 the squid host has been observed. Although there is the possibility that AOX does not play a role in 386 bacterial physiology during host colonization, an alternative explanation is that the benefit of AOX 387 expression during colonization does not result in a phenotype dramatic enough to be detected in the 388 short time frame of the experiments (1-3 days). Experiments to test this possibility are in progress 380 and would be consistent with studies above described for luminescence mutants in which colonization 390 phenotypes change over the course of symbiosis and effects are magnified over a multi-week time 391 course.

Studying AOX regulation and function in *V. fischeri* as a model organism will provide a framework for understanding how bacteria in ocean surface waters utilize this respiratory pathway in growth and survival. Work is underway to clarify the physiological benefit of AOX function in *V. fischeri* and other *aox*-containing bacteria, with the ultimate goal of better understanding how bacteria cope with changing conditions in the environment. Studying AOX in the context of the symbiosis has provided insight into the expression and function of this interesting protein, and provides a framework for broad studies of how AOX function influences bacterial physiology in the environment.

Study of AOX followed a reverse-genetic approach, starting with identification of an interesting gene through genome sequencing, and through directed experimental approaches leading to a better understanding of gene function. However, in many cases forward genetic approaches have identified genes whose products are relevant for a specific colonization process. An excellent example is *binK*, which encodes a histidine kinase. Above we described a key role for biofilm formation in the colonization process as regulated by RscS and Syp. In a recent global genetic screen for mutants with an advantage

in squid colonization, binK was identified as a locus that when disrupted resulted in substantially better colonization of the *V. fischeri* strain (Brooks and Mandel, 2016). Typical means to predict protein 407 function (e.g., homology, neighboring genes) were not helpful, so phenotypes of cells lacking binK 408 were examined in culture and in the host and revealed a substantial increase in symbiotic biofilm for-400 mation. BinK (biofilm inhibitor kinase) is therefore a negative regulator of biofilm formation and an 410 additional membrane-bound histidine kinase that is critical for proper regulation of the Syp biofilm. 411 In the case of both AOX and BinK, the depth of the V. fischeri-squid system has provided a means 412 to assign function to novel and poorly-understood proteins. A striking number of genes are poorly 413 understood in bacterial genomes, exemplified by the 149 (32 %) of the minimal 473 genes in the JCVI-414 syn3.0 genome with functions that remain to be discovered (Hutchison et al., 2016). The ability to study 415 biological function in the context of the host thus provides a useful lens through which to identify and 416 characterize genes and their products. 417

418 Conclusions

The Vibrio-squid system has proven to be a valuable study system for identifying principles of microbehost interactions, continues to serve as a fertile field for discovery, and provides a useful road map for 420 moving from patterns of intriguing phenotypes to discerning the molecular communication between 421 microbe and host that is responsible for those patterns. By integrating approaches in genetics, genomics, 422 molecular biology, imaging, physiology, evolutionary biology, and cell biology, each of the topic areas 423 highlights an integrated and mechanistic view of how symbiotic partners functionally communicate 424 in a model microbiome. In this manner, the Vibrio-squid system provides a durable example for how 425 to move from fascinating observations to molecular understanding of the processes by which very 426 different organisms communicate and establish a productive partnership. 427

Figure Legend

Figure 1. (A) Juvenile Euprymna scolopes hatchling, ventral view. White box highlights the ink sac 420 and the light organ. (B) Confocal micrograph of the bilaterally symmetric light organ. Host tissue is 430 counterstained in red and the colonizing bacteria are visible in green. Arrowheads point to the three 431 pores on each side of the organ, into which V. fischeri swim into the internal anatomy (ducts, antecham-432 ber, bottleneck, and crypts) of the organ. White box highlights one half of the organ, which is shown 433 in cartoon view in the next panel. (C) Current state of knowledge about the temporal and spatial action 434 of key processes discussed in this review, including Syp biofilm formation and aggregation (red), host 435 nitric oxide production (yellow), bacterial motility and chemotaxis toward host chitin oligosaccharides 436 (orange), symbiont TCT release (green), and luminescence (blue). In general the location of the colo-437 nizing bacteria are highlighted; e.g., for TCT release the bacteria colonize the crypts and release TCT 438 (indicated), though the effect of this release on the host is apoptosis and regression of the ciliated ep-439 ithelial appendages (not indicated in this representation). Panels A and B are adapted from (Mandel et 440 al., 2012).

442 Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

445 Author Contributions

446 MJM and AKD wrote the manuscript.

447 Funding

Research in the authors' laboratories in supported by National Science Foundation awards IOS-1456963 (MJM) and MCB-1050687 (AKD), and National Institutes of Health Awards R35GM119627

- (MJM) and R21AI117262 (MJM). The content is solely the responsibility of the authors and does not
- necessarily represent the official views of the funding agencies.

Acknowledgments

We thank Ella Rotman and Denise Tarnowski for comments on the manuscript.

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