The haem-uptake gene cluster in *Vibrio fischeri* is regulated by Fur and contributes to symbiotic colonization

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Summary

Although it is accepted that bacteria-colonizing host tissues are commonly faced with iron-limiting conditions and that pathogenic bacteria often utilize iron from host-derived haem-based compounds, the mechanisms of iron acquisition by beneficial symbiotic bacteria are less clear. The bacterium *Vibrio fischeri* mutualistically colonizes the light organ of the squid *Euprymna scolopes*. Genome sequence analysis of *V. fischeri* revealed a putative haem-uptake gene cluster, and through mutant analysis we confirmed this cluster is important for haemin use by *V. fischeri* in culture. LacZ reporter assays demonstrated Fur-dependent transcriptional regulation of cluster promoter activity in culture. GFP-based reporter assays revealed that gene cluster promoter activity is induced in symbiotic *V. fischeri* as early as 14 h post inoculation, although colonization assays with the haem uptake mutant suggested an inability to uptake haem does not begin to limit colonization until later stages of the symbiosis. Our data indicate that the squid light organ is a low iron environment and that haem-based sources of iron are used by symbiotic *V. fischeri* cells. These findings provide important additional information on the availability of iron during symbiotic colonization of *E. scolopes* by *V. fischeri*, as well as the role of haem uptake in non-pathogenic host-microbe interactions.

Introduction

Pathogenic bacteria associated with animal or plant host tissues often face an iron-limited environment, and must use specific acquisition systems to obtain this essential element. Such iron scavenging is especially well studied in pathogenic bacteria that infect vertebrates. Although these bacteria can possess uptake systems for ferrous iron, transferrin, lactoferrin and/or ferritin (Wandersman and Delepelaire, 2004), one of the most abundant iron sources in vertebrate tissue is haem-based (Anzaldi and Skaar, 2010). As a result, many pathogenic bacteria have specialized systems to facilitate haem uptake and utilization (Anzaldi and Skaar, 2010). For example, haem uptake systems have been described in fish pathogens among the *Vibrionaceae*, and in certain cases these systems may be expressed and play a role in iron uptake during infection (Lemos and Osorio, 2007). Regulation of the haem uptake system genes has been described in *Vibrio anguillarum*, where expression is controlled in response to iron concentration through the activity of the ferric uptake regulation (Fur) protein (Mourino et al., 2006).

In contrast to pathogenic host-microbe interactions, relatively little is known about the role of haem uptake in beneficial relationships of bacteria with their hosts. Animals and plants often support the growth and persistence of mutualistic bacteria in specific specialized tissues, and for the most part it remains unclear whether the bacteria supported by the host in these environments face the same iron limitations and sources as pathogens invading these or other tissues. With the exception of the interactions between *Bradyrhizobium japonicum* and soybean plants (Nienaber et al., 2001), and *Sodalis glossinidius* and the tsetse fly (Runyen-Janecky et al., 2010), the role of microbial haem uptake in mutualistic symbioses has not been explored. In the case of *B. japonicum*, disrupting bacterial haem uptake does not affect symbiotic nitrogen fixation, suggesting that haem is not a required iron source for development or maintenance of the symbiotic relationship between the bacterium and plant (Nienaber et al., 2001). In contrast, *in vivo* expression analysis demonstrated that portions of the haem uptake system of *S. glossinidius* are expressed during
insect colonization, suggesting that a haem-based iron source could be important for the interactions between the bacterium and invertebrate host (Runyen-Janecky et al., 2010).

_Vibrio fischeri_ is a bioluminescent marine bacterium that forms a beneficial symbiotic relationship with the Hawaiian bobtail squid _Euprymna scolopes_, during which the bacterium colonizes a specific organ in the animal called the light organ (Visick and Ruby, 2006). _Vibrio fischeri_ grows rapidly during the first day of light-organ infection, doubling as rapidly as every 30 min (Ruby and Asato, 1993); however, it has also been reported that the symbionts may experience iron-limited growth in the light organ later in infection (Graf and Ruby, 2000). The squid vent a subpopulation of symbionts each day, with the residual bacteria regrowing (Boettcher et al., 1996), and it is possible that _V. fischeri_ must actively scavenge some form of iron to persistently colonize the host through these cycles of regrowth. Based on genome sequence analysis, _V. fischeri_ appears to possess multiple types of iron acquisition systems similar to those in pathogenic bacteria, including a predicted haem-uptake system (Ruby et al., 2005). Although a detailed analysis of iron source availability in the host has not been performed, it has been suggested that free ferrous or ferric iron is not the main form of iron present for _V. fischeri_ in the light organ environment (Wang et al., 2010), and that iron may be acquired by the bacterium from a haem-based source during host association (Wang et al., 2010; Wier et al., 2010).

To investigate the role of haem uptake in _V. fischeri_, we used mutant analysis to test and confirm that the genome-predicted haem-uptake gene cluster is important for haem-in uptake and/or utilization in _V. fischeri_, and to explore the role of haem uptake during symbiotic colonization. These studies were combined with reporter-based analyses to characterize transcriptional regulation of the genes of the haem uptake cluster during growth both in culture media and in the host light organ. The results of this study not only advance our understanding of the role of iron acquisition in _V. fischeri_ during host colonization, but provide further insight into how haem acquisition may contribute to beneficial host-microbe interactions in general.

**Results**

**Identification of a haem-uptake gene cluster in** _V. fischeri_ **ES114**

Analysis of the _V. fischeri_ ES114 genome sequence (Ruby et al., 2005) indicated the presence of a gene cluster (ORFs VF_1220-VF_1228) with similar gene content and arrangement as compared with haem-uptake cluster genes in other members of the _Vibrionaceae_ (Lemos and Osorio, 2007) (Fig. 1, Table 1). Based on this similarity, we predicted that these _V. fischeri_ genes encode proteins involved in providing energy for the transport process (TonB, ExbB, ExbD), a periplasmic haem-binding protein (HutB), an inner membrane permease (HutC), and an ABC-transporter ATPase (HutD) (Table 1). This gene cluster also encodes the proteins HutW, HutX and HutZ. HutW is a member of the radical SAM superfamily (Sofia et al., 2001), and is often annotated as a coproporphyrinogen oxidase (as in _V. fischeri_ ES114) due to its homology to HemN. However, the _hutW_ from _Vibrio cholerae_ was unable to complement a _Salmonella enterica_ serovar Typhimurium hemN mutant (Wyckoff et al., 2004), suggesting that HutW is not a coproporphyrinogen oxidase. As a result, the specific function of HutW remains undefined. The _V. fischeri_ _hutX_ and _hutZ_ genes are annotated as proteins of unknown function; however, it has been shown that HutZ in _V. cholerae_ and the homologous HuvZ in _V. anguillarum_ are required for optimal haem utilization (Mourino et al., 2004; Wyckoff et al., 2004). Similar to many of the _Vibrionaceae_, a gene encoding a putative TonB-dependent haem receptor protein (VF_1234, VF_A0332) is not directly linked to the _V. fischeri_ VF_1220-VF_1228 cluster (Lemos and Osorio,
2007), and therefore the contribution of haem receptor proteins was not explored in this study. To determine whether proteins encoded by the VF_1220-VF_1228 cluster were necessary for use of haemin as a sole iron source by *V. fischeri*, we constructed mutant strain AKD910, which contains an in-frame deletion of VF_1220-VF_1228.

AKD910 is unable to use haemin as a sole iron source

Growth of AKD910 (lacking VF_1220–1228) was compared with that of wild type in iron-limited mineral-salts medium either in the absence or presence of haemin. Growth density assays demonstrated that AKD910 was unable to use haemin as a sole iron source (Fig. 2), indicating that this gene cluster encodes proteins necessary for haem uptake and/or utilization as an iron source. However, when ferrous sulfate was provided as an iron source, both AKD910 and wild-type cultures grew to similar final optical densities of 1.29 ± 0.01 and 1.28 ± 0.02 respectively.

Promoters for the gene cluster are repressed by Fur and induced in response to low iron

Plasmid-based promoter–*lacZ* fusions were constructed to determine the influence of iron levels on haem-uptake gene promoter activity. The organization of this gene cluster in *V. fischeri* (Fig. 1) suggested that the genes are expressed in two transcriptional units. Two promoter–*lacZ* fusion plasmids were constructed by directionally cloning DNA fragments corresponding to the sequence upstream of VF_1225 and VF_1226. These plasmids were then moved via conjugation into the appropriate *V. fischeri* strains.

Comparative analysis of expression patterns between the two reporters in various strains and culture conditions indicated congruent promoter activities (Fig. 3A and B). Promoter activity in wild-type cells was relatively low in LBS, an iron-replete rich medium; however, promoter activity was significantly increased in iron-limited conditions.

Table 1. Comparison of the *Vibrio fischeri* haemin uptake gene cluster to known proteins.

<table>
<thead>
<tr>
<th><em>V. fischeri</em> ORF</th>
<th>Assignment</th>
<th>Database comparison</th>
<th>% identity</th>
<th>% similarity</th>
</tr>
</thead>
<tbody>
<tr>
<td>VF_1220</td>
<td>HutD (259 aa)</td>
<td><em>Vibrio anguillarum</em>, HuvD, ABC transporter ATPase (199 aa), CAF25489 (Mourino et al., 2004)</td>
<td>61% (118/195 aa)</td>
<td>77% (145/195 aa)</td>
</tr>
<tr>
<td>VF_1221</td>
<td>HutC (345 aa)</td>
<td><em>V. anguillarum</em>, HuvC, inner membrane permease (314 aa), CAF25488 (Mourino et al., 2004)</td>
<td>74% (230/314 aa)</td>
<td>89% (277/314 aa)</td>
</tr>
<tr>
<td>VF_1222</td>
<td>HutB (282 aa)</td>
<td><em>V. anguillarum</em>, HuvB, periplasmic haemin-binding protein (282 aa), CAF25487 (Mourino et al., 2004)</td>
<td>61% (180/282 aa)</td>
<td>75% (209/282 aa)</td>
</tr>
<tr>
<td>VF_1223</td>
<td>ExbD (139 aa)</td>
<td><em>V. anguillarum</em>, ExbD, energy transducer for active transport (136 aa), CAD43043 (Mourino et al., 2004)</td>
<td>59% (80/136 aa)</td>
<td>76% (103/136 aa)</td>
</tr>
<tr>
<td>VF_1224</td>
<td>ExbB (227 aa)</td>
<td><em>V. anguillarum</em>, ExbB, energy transducer for active transport (235 aa), CAD43042 (Mourino et al., 2004)</td>
<td>59% (120/221 aa)</td>
<td>64% (162/221 aa)</td>
</tr>
<tr>
<td>VF_1225</td>
<td>TonB (254 aa)</td>
<td><em>V. anguillarum</em>, TonB, energy transducer for active transport (240 aa), CAD43041 (Mourino et al., 2004)</td>
<td>47% (124/264 aa)</td>
<td>63% (164/264 aa)</td>
</tr>
<tr>
<td>VF_1226</td>
<td>HutW (458 aa)</td>
<td><em>Vibrio cholerae</em>, HuvW (455 aa), VCA0909 (Wyckoff et al., 2004)</td>
<td>59% (261/443 aa)</td>
<td>77% (338/443 aa)</td>
</tr>
<tr>
<td>VF_1227</td>
<td>HutX (178 aa)</td>
<td><em>V. anguillarum</em>, HutX (171 aa), CAD43040 (Mourino et al., 2004)</td>
<td>64% (102/161 aa)</td>
<td>82% (131/161 aa)</td>
</tr>
<tr>
<td>VF_1228</td>
<td>HutZ (175 aa)</td>
<td><em>V. anguillarum</em>, HutZ (176 aa), CAD43039 (Mourino et al., 2004)</td>
<td>70% (119/172 aa)</td>
<td>84% (143/172 aa)</td>
</tr>
</tbody>
</table>

a. *V. anguillarum* serovar 01, accession or ORF number and reference provided.
b. HutW is not present in *V. anguillarum*, but is present in *V. cholerae* O1 biovar El Tor strain N16961.

Fig. 2. The putative haem uptake genes are necessary for growth of *V. fischeri* with haemin as the sole iron source. Cultures of wild type and AKD910 (ΔVF_1220–1228) were grown in mineral salts medium containing iron chelator, with and without haemin added. Optical density was measured over time for wild type grown without haemin (open squares), wild type grown with haemin (filled squares), AKD910 grown without haemin (open triangles), and AKD910 grown with haemin (filled triangles). Error bars represent standard error of the mean.

activity was approximately 30-fold higher in LBS medium when iron chelator was added (Fig. 3A). This influence of iron levels on promoter activity, and a putative Fur box in the VF_1225 upstream sequence (Wang et al., 2010), suggested that the V. fischeri homologue of the ferric uptake regulator repressor protein (Fur, VF_0810) could influence VF_1225 promoter activity. To test this possibility, we introduced the reporter plasmids into a V. fischeri strain lacking fur (YLW111), and as described above, assayed for promoter activity both in the presence and absence of iron chelator (Fig. 3B). These data demonstrate that V. fischeri Fur mediates repression of promoter activity in response to iron availability.

Although promoter activity increases under low-iron conditions, there was the possibility that activity could also be influenced by the addition of haemin to the assays. Our results suggest that under the conditions tested, promoter activity does not increase in the presence of haemin (Fig. 3A). However, a trend of a decrease in promoter activity in wild-type cells grown in medium amended with haemin and an iron chelator as compared with iron chelator alone was observed (Fig. 3A), although it was not always statistically significant. We hypothesized that this trend was due to the processing of haemin by the proteins encoded in the gene cluster, effectively increasing intracellular iron levels, resulting in moderate Fur-mediated repression of promoter activity. When promoter activity levels were measured under identical growth conditions (LBS medium amended with iron chelator, with or without haemin) in the mutant strain AKD910, which lacks the ability to utilize haemin as an iron source, no significant decrease in promoter activity was observed in the presence of haemin (data not shown). These data support the hypothesis that products of this gene cluster are involved in processing of haemin as an iron source, and the corresponding increase in intracellular iron is sensed by Fur in wild-type cells.

Haem uptake gene expression is induced during symbiotic colonization

To determine whether gene cluster expression is induced during colonization of the host, P_{VF_1225}−_gfp and P_{VF_1226}−_gfp reporter plasmids, which also encode a constitutively expressed red fluorescent protein gene (rfp), were constructed and introduced into wild-type V. fischeri. Juvenile aposymbiotic squid colonized by these strains were monitored using epifluorescence microscopy. Similar results were obtained with the two reporters, therefore only P_{VF_1225}−_gfp (pAKD911) data are shown. Results of these assays demonstrated that V. fischeri colonizing the juvenile squid light organ are expressing the gene cluster at 28 h post inoculation (Fig. 4C and D), with similar results obtained at earlier (14 h) and later (48 h, 72 h) time points (data not shown). Similar to the β-galactosidase assays, measureable GFP expression is only observed in culture when iron is limiting (Fig. 4E–H), suggesting that the genes are expressed in response to low iron conditions present in the squid light organ.

The haem-uptake genes influence V. fischeri symbiotic competency

The expression of the gene cluster during host colonization indicated that V. fischeri could be using host-derived
haem compounds as a source of iron. To determine whether the ability to utilize haem was important for host colonization, we assayed the ability of AKD910 (ΔVF_1220–1228) to colonize aposymbiotic juvenile E. scolopes squid. In experiments where E. scolopes squid were exposed to clonal inocula containing either AKD910 or the wild type, colonization levels were similar in the two treatments at time points 48 and 72 h post inoculation, suggesting that AKD910 does not have a detectable symbiotic defect compared with the wild type (data not shown). However, it has been documented that symbiotic defects in certain mutants are only apparent when they are competed with the wild type in mixed inocula (Stabb and Ruby, 2003, Visick and Ruby, 1998). Competitive colonization assays revealed AKD910 was less competitive for colonization (average relative competitive index or RCI value of less than one, see materials and methods), but this deficiency was not observed until several days post inoculation, and became increasingly robust at later time points (Fig. 5). These results indicate that although promoter activity is induced in symbiotic V. fischeri as early as 14 h post inoculation, an inability to transport and use haem does not detectably limit colonization competitiveness until 48 h post inoculation, with the most significant effects at later stages of the symbiosis (~96 h).

Discussion

Bacterial pathogens often encounter iron-limiting conditions in hosts, and must employ iron-scavenging strategies for successful infection (Weinberg, 1978). Although there are numerous forms of iron that these bacteria could acquire during host colonization, haem in particular has been shown to influence bacterial virulence, and strains defective in haem uptake demonstrate reduced virulence (reviewed in Lee, 1995).

In contrast, the role of haem as an iron source in beneficial host-microbe interactions is less clear. Bacterial growth in mutualistic symbioses is often supported by the host, making it uncertain whether these bacteria also face similar iron-limited host environments, and, if so, the role of iron limitation in the relationship. Previous studies of the V. fischeri – E. scolopes mutualistic symbiosis suggested that the squid light organ may be an iron-limited environment after the first day of infection (Graf and Ruby, 2000), and that haem-based compounds could be important sources of iron for symbiotic V. fischeri (Wang et al., 2010; Wier et al., 2010). Based on analysis of the V. fischeri genome sequence (Ruby et al., 2005) and identification of a putative haem-uptake gene cluster (Fig. 1) similar to those characterized in other Vibrionaceae (Lemos and Osorio, 2007), we hypothesized that this gene cluster would be expressed in response to iron limitation and that an ability to uptake and utilize haem-based iron would be important for V. fischeri during the symbiotic lifestyle.

Our experimental results confirm that the identified gene cluster is involved in haemin uptake and utilization, and that regulation of gene expression under culture conditions is similar to other members of the Vibrionaceae that form pathogenic relationships with multicellular hosts (Mourino et al., 2006; Lemos and Osorio, 2007). The idea that pathogenic and mutualistic members of the Vibrionaceae have nearly identical uptake systems that are regulated similarly suggest that, although the outcome of the host-bacterial relationships are very different, there are fundamental similarities in how these bacteria obtain host-derived iron sources. Further understanding of these similarities and differences will provide useful insight into what defines pathogenesis versus mutualism.

Beyond providing a comparison of culture-based haem-uptake gene regulation in pathogenic and mutualistic Vibrionaceae, our results provide important insight into how iron influences the symbiotic relationship between V. fischeri and E. scolopes. Fluorescence-based reporter assays during symbiotic colonization provide strong support for the idea that the squid light organ is an iron-limited environment (Fig. 4). Results from culture-based reporter assays indicate that gene cluster expression is induced in response to low iron conditions through the regulatory activity of Fur, and squid colonization assays demonstrate that the same reporter constructs are induced during colonization (Figs 3 and 4). In combination, these experiments provide data to support the model of the light organ being an iron-limited environment, as

![Graphical representation of colonization competitiveness in the combined 94 and 100 h assays. Each symbol represents a RCI value from one animal. The dashed line indicates the average RCI value (0.33) for all animals in the experiment. The animals represented by open symbols were clonally colonized by wild-type V. fischeri.](image)
Haem uptake in Vibrio fischeri 2861

previously suggested (Graf and Ruby, 2000; Wang et al., 2010), although there may be other currently undescribed conditions and regulators that influence gene cluster expression during host colonization.

In addition, our results indicate that haem uptake and/or utilization does play a role in the establishment of a successful symbiotic relationship (Fig. 5), providing further information concerning the role of this physiological process in beneficial host-microbe interactions. One interesting observation from these analyses is that although the promoters are active at early time points post inoculation, colonization is not measurably limited in a strain lacking the gene cluster (AKD910) until later stages of the symbiosis (48–100 h) (Figs 4 and 5). This apparent discrepancy could, in part, be explained by what is currently known about host physiology during symbiotic colonization.

The levels of haem-based iron available to host-associated V. fischeri could be related to morphological changes that occur in the host epithelium lining the light organ crypts during the day-night cycle of the symbiosis. The animals use the light produced by bacteria during nocturnal behaviours, and at dawn expel approximately 90% of the bacterial population, the remainder of which will divide and recolonize the light organ (Nyholm and McFall-Ngai, 2004). Corresponding with these daily ventings, the epithelial cells bleb into the interior of the light organ (Wier et al., 2010). These blebs could be a source of haem-based iron, and this is reflected in the transcriptional response of the bacterial population to this stage of the symbiosis, which indicates an increase in transcript levels for members of the haem-uptake genes (Wier et al., 2010). These changes were documented in adult animals with a ‘mature’ symbiosis, and it is likely that during the onset of colonization (prior to 94 h) the morphological changes to the host epithelium are less pronounced during the day-night cycle, resulting in less haem-based iron available to the bacteria. In support of this hypothesis, squid colonization data using haem biosynthesis mutants suggest haem levels are too low to support the growth of V. fischeri 2861 at lower cell densities (Haygood and Nealson, 1985; Dunlap, 1992). Experiments designed to identify the molecular mechanism of iron influence on bioluminescence were performed in transgenic E. coli carrying the lux genes (Dunlap, 1992), yet it remains unclear how iron is physiologically linked to these phenotypes in V. fischeri. However, since this earlier publication, genetic tools have been constructed for V. fischeri that will facilitate future studies to characterize the link between iron levels, luminescence, and successful colonization and maintenance of the symbiosis.

This study represents the first genetics-based investigation of haem-based iron acquisition in V. fischeri, and this information in combination with previous and future studies can be used to further define the parameters important for successful establishment and maintenance of the mutualistic symbiosis between V. fischeri and E. scolopes. As this system is an important model for beneficial host-microbe interactions, further understanding of this relationship provides an important comparison to pathogenic host-bacterial interactions and contributes to the ultimate goal of defining the boundaries between pathogenesis and mutualism.

Experimental procedures

Growth media

Vibrio fischeri strains were grown at 28°C (unless noted) in either LBS (Stabb et al., 2001), SWT (Boettcher and Ruby, 1990) wherein seawater was replaced with Instant Ocean (Aquarium Systems; medium referred to as ASWT), or mineral salts medium (per litre: 378 µl 1 M NaPO₄ (pH 7.5), 50 ml 1 M Tris (pH 8.0), 0.59 g NH₄Cl, 3 mg FeSO₄·7H₂O, 13.6 g MgSO₄·7H₂O, 0.83 g KCl, 19.5 g NaCl, 1.62 g CaCl₂·2H₂O] containing 40 mM glycerol and 1 g l⁻¹ vitamin-free casamino acids (Difco) as carbon and nitrogen sources. When indicated, 100 µg ml⁻¹ kanamycin or 5 µg ml⁻¹ erythromycin was added to the culture medium. Escherichia coli strains were grown at 37°C in LB medium (Miller, 1992) with 40 µg ml⁻¹ kanamycin or 22.5 g l⁻¹ HiVeg special infusion (HIMEDIA) containing 150 µg ml⁻¹ erythromycin.

Strains and plasmids

Strains and plasmids used in this study are listed in Table 2. General cloning procedures were used in E. coli strains DH5α (Hanahan, 1983) or DH5α::pir (Dunn et al., 2005), and constructs containing the RP4 origin of transfer were intro-
duded into *V. fischeri* using triparental mating (Stabb and Ruby, 2002). All *Vibrio* plasmids used in this study contain the pES213 origin of replication, which is maintained at approximately 10 copies per chromosome (Dunn et al., 2005). Primer sequences are available upon request.

**Construction of V. fischeri mutant strains.** A strain lacking the putative haemin uptake and processing genes (VF_1220-VF_1228), which are divergently arranged at one locus (Fig. 1), was constructed using allelic exchange (Bose et al., 2008), resulting in an in-frame deletion mutant (AKD910). Briefly, approximately 1.6 kb of DNA downstream of VF_1220 was PCR-amplified and fused to an approximately 1.6 kb DNA fragment downstream of VF_1228 using an engineered Nhel 6 bp restriction site. Additional codons were included in order to design specific primers with reasonable G + C content, resulting in inclusion of the last six codons of VF_1220 and the last four codons of VF_1228. Due to the large size of the gene cluster (~10 kb) plasmid-based complementation methods were not used with AKD910; however, given the gene arrangement (Fig. 1) and the complete deletion of the cluster, polar effects of the deletion are not expected.

A strain lacking *fur* (VF_0810; strain name YLW111) was constructed using an identical approach, with inclusion of the first seven codons of VF_0810.

**Plasmid-based PVF_1225 and PVF_1226 transcriptional reporters.** The plasmid-based reporters pAKD911 (PVF_1225-\textit{gfp}) and pAKD912 (PVF_1225-\textit{lacZ}) were constructed by PCR amplifying a 243 bp fragment that included 221 bp upstream of the start codon of VF_1225 (\textit{lonB}, Fig. 1), and directionally cloning this fragment into pSVS209 or pAKD701 respectively, whereas pAKD913 (PVF_1226-\textit{gfp}) and pAKD914 (PVF_1226-\textit{lacZ}) were constructed by PCR amplifying a 576 bp fragment that included 499 bp upstream of the start codon of VF_1226 (\textit{furW}; Fig. 1) and directionally cloning this fragment into pVSV209 or pAKD701 respectively.

**Growth of wild-type V. fischeri and AKD910 using haemin as an iron source.**

To assay growth of *V. fischeri* using haemin as an iron source, cultures of wild type and AKD910 were grown overnight in LBS medium and diluted 1000-fold into mineral salts medium lacking FeSO₄ and supplemented with 50 μM of the iron chelator 2,2'-bipyridyl (Sigma, St. Louis, MO, USA), either with or without 77 μM (50 μg ml⁻¹) bovine haemin (Sigma) or 10.8 μM ferrous sulfate. The diluted cultures were grown in 200 μl aliquots in a Falcon polystyrene 96-well plate (Becton Dickinson, Franklin Lakes, NJ, USA) for 3 h at 28°C. Growth was measured by absorbance at 595 nm using a BioTek Synergy 2 plate reader, and absorbance readings (optical density) were adjusted to a 1 cm path length. Each experiment included two independent cultures of each strain grown in duplicate, and the experiment was repeated three times. One representative experiment is shown.

**β-Galactosidase assays of promoter activity.**

Overnight cultures of wild-type or YLW111 (\textit{Δfur}) containing either the control plasmid pAKD701 or the haem uptake reporter plasmids pAKD912 or pAKD914 were grown in LBS medium containing kanamycin. Cultures were diluted 1000-fold into LBS medium containing kanamycin, either with or without 100 μM of the iron chelator 2,2'-bipyridyl and/or 31 μM haemin, and grown to an OD₅₆₂ of between 0.4 and 0.5. Cells were harvested by pelleting 2 ml of culture at 15 000 g for 5 min. Supernatant was discarded and pellets
frozen at −20°C for no longer than 24 h. Cell pellets were resuspended in one ml of Z buffer and β-galactosidase assays performed using a modified Miller assay as previously described (Bose et al., 2008). The average Miller unit values for the strains containing pAKD701 were subtracted from the corresponding pAKD912 or pAKD914 individual sample Miller unit values prior to calculating averages to account for effects of medium amendments on absorbance readings. For each experiment three independent cultures of each strain were assayed, and each experiment was repeated at least three times. Data shown is from one representative experiment.

**Squid colonization and reporter assays**

For all squid colonization assays, *V. fischeri* strains were grown in ASWT medium at 28°C without shaking to an OD595 of between 0.3 and 0.6 and diluted in Instant Ocean to between 1360 and 2600 cfu ml⁻¹ inocula. Aposymbiotic juvenile squid were exposed to inocula for 15 h before being transferred to fresh filter-sterilized Instant Ocean.

For RFP- and GFP-based reporter assays in the squid, juvenile aposymbiotic animals were inoculated as described above with wild-type *V. fischeri* containing either the control plasmid pVSV209, or the haem uptake cluster gene reporter plasmids pAKD911 or pAKD913. At various time points post inoculation, animals were anesthetized using 0.12 M MgCl₂ and the light organ tissue imaged using a Nikon (Melville, NY, USA) Eclipse E600 epifluorescence microscope equipped with a Nikon 96157 red filter cube, a Nikon 41017 green filter cube, and a Nikon Coolpix 5000 camera for image capture. Similar results were observed with *V. fischeri* harbouring pAKD911 and pAKD913, therefore images are only shown for pAKD911-containing *V. fischeri*. At least five animals were analysed at each time point, with images from one representative animal shown.

In colonization competition experiments, juvenile squid were exposed to mixed inocula with roughly equal numbers of each cell type for 15 h before being rinsed in inoculum-free Instant Ocean. The unmarked deletion mutant AKD910 was competed against AKD100, an erythromycin-resistant mini Tn-7-ermR marked derivative of *V. fischeri* wild-type strain ES114. AKD100 has no apparent colonization defects when competed against wild-type *V. fischeri* ES114 (data not shown), and therefore was used as a marked strain with wild-type competitiveness in these assays. Strain ratios were determined after dilution plating by patching to LBS medium containing erythromycin. Competitiveness is presented as a RCI, which was calculated by dividing the AKD910 to AKD100 ratio in the squid by the AKD910 to AKD100 ratio of the inoculum. A RCI value of 1 indicates the strains competed equally during host colonization, whereas a RCI value of < 1 indicates AKD100 (wild-type) out-competed the mutant. Data shown for the −24 and −96 h time points are combined from two independent experiments, whereas the data shown for the −48 and −72 h time points are each from a single experiment.

**GFP-based reporter assays in culture**

*Vibrio fischeri* ES114 harbouring pAKD911 or pAKD913 was grown overnight in LBS medium containing kanamycin and either with or without 100 μM of the iron chelator 2,2′-bipyridyl at 28°C with shaking. Cultures were diluted 100-fold into 25 ml of the same medium type in 125 ml Erlenmeyer flasks and incubated at 24°C with shaking until the cultures reached an OD₅₉₅ of between 0.93 and 0.99. Ten microlitres of culture was transferred onto a spot of LBS with 1% agarose on a glass slide and images were taken using 100x objective using the epifluorescence microscope described above. All images used an aperture setting of F4.6, and the exposure time was 0.03 s for bright field and 1.0 s for GFP images. Identical results were obtained in experiments with cultures at varying optical densities (OD₅₉₅ of 0.5 to 1.5), and cultures grown at 28°C (data not shown).

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**References**


