LuxU connects quorum sensing to biofilm formation in Vibrio fischeri

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Summary

Biofilm formation by Vibrio fischeri is a complex process involving multiple regulators, including the sensor kinase (SK) RscS and the response regulator (RR) SypG, which control the symbiosis polysaccharide (syp) locus. To identify other regulators of biofilm formation in V. fischeri, we screened a transposon library for mutants defective in wrinkled colony formation. We identified LuxQ as a positive regulator of syp-dependent biofilm formation. LuxQ is a member of the Lux phosphorelay and is predicted to control bioluminescence in concert with the SK AinR, the phosphotransferase LuxU and the RR LuxO. Of these, LuxU was the only other regulator that exerted a substantial impact on biofilm formation. We propose a model in which the Lux pathway branches at LuxU to control both bioluminescence and biofilm formation. Furthermore, our evidence suggests that LuxU functions to regulate syp transcription, likely by controlling SypG activity. Finally, we found that, in contrast to its predicted function, the SK AinR has little impact on bioluminescence under our conditions. Thus, this study reveals a novel connection between the Lux and Syp pathways in V. fischeri, and furthers our understanding of how the Lux pathway regulates bioluminescence in this organism.

Introduction

Bacteria readily adapt to changing environmental conditions by sensing and integrating different cues present in their surroundings to produce a response appropriate for survival or growth. One mechanism by which bacteria co-ordinate such responses is through the use of twocomponent systems, with the sensor kinase (SK) component involved in detecting and relaying an environmental cue, and the response regulator (RR) component promoting the response, often a change in gene expression

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(Stock *et al.*, 2000; West and Stock, 2001). A classic example of signal integration via two-component regulators is the Lux pathway in the marine bioluminescent bacterium *Vibrio harveyi* (Ng and Bassler, 2009). This organism integrates distinct signals (small molecules termed autoinducers or Als) using specific SKs that funnel their activities into a common phosphorelay pathway to control the production of cellular bioluminescence.

In V. harveyi, the Lux pathway (Fig. 1A) is composed of the SKs LuxQ (in association with the periplasmic protein LuxP), LuxN and CqsS (not depicted), the histidine phosphotransferase LuxU and the RR LuxO (reviewed in Ng and Bassler, 2009). Under low cell densities (low AI concentrations), the SKs exhibit net kinase activity and serve as phosphoryl-donors to LuxU, which serves as a phosphoryl-donor to the RR LuxO. Phosphorylated LuxO (LuxO~P) then promotes the transcription of five sRNAs (grr1-5), which bind to and destabilize (in conjunction with Hfq) the transcript for the master transcriptional regulator LuxR_{VH}. This regulator promotes the transcription of the lux operon, which encodes the proteins necessary for light production. Without LuxR_{VH}, light is not produced. However, as the cell density increases, the AI concentrations also increase, causing a shift in the equilibrium of SK activity (i.e. from net kinase to net phosphatase activity). This switch to net phosphatase activity promotes the removal of phosphoryl groups from LuxO through LuxU. Without LuxO~P, LuxR_{VH} is produced and promotes the expression of the lux operon, ultimately leading to light production.

Similar pathways exist in a variety of other *Vibrio* species, including *V. parahaemolyticus*, *V. cholerae*, *V. anguillarum*, *V. vulnificus* and *V. fischeri* (Milton, 2006; Zhang *et al.*, 2012). Like *V. harveyi*, the *V. fischeri* Lux pathway also controls bioluminescence (reviewed in Stabb *et al.*, 2008; Miyashiro and Ruby, 2012). The *V. fischeri* Lux components appear similar to those in *V. harveyi* (Fig. 1). For example, *V. fischeri* encodes (or is predicted to encode) homologues of LuxP, LuxQ, LuxU and LuxO, which are proposed to function in a manner similar to their counterparts in *V. harveyi* (Miyamoto *et al.*, 2000; Visick, 2005; Stabb *et al.*, 2008; Miyashiro *et al.*, 2010); however, differences between the Lux pathways in *V. fischeri* and *V. harveyi* exist. For example, *V. fischeri* does not encode a homologue of CqsS or LuxN, but encodes another SK,





A. In *V. harveyi*, bioluminescence is regulated by the Lux phosphorelay, composed of the sensor kinases (SKs) LuxQ (which interacts with the periplasmic protein LuxP), LuxN and CqsS (not depicted), the phosphotransferase LuxU and the response regulator (RR) LuxO. Phosphoryl-transfer (dashed, double-sided arrows) occurs between the SKs, LuxU and LuxO. Under low cell density conditions, LuxO is phosphorylated by the kinase activity of the SKs and activates transcription of the *qrr* sRNAs, which bind to the transcript of *luxR*_{VH} and prevent its translation. Without LuxR_{VH}, the *lux* operon (not depicted) is not expressed and light (bioluminescence) is not produced. At high cell densities, LuxO is dephosphorylated by the phosphatase activity of the SKs, leading to subsequent production of LuxR_{VH} and bioluminescence. Autoinducers (Als) or the Al synthases (LuxM, LuxS or CqsA) are not depicted for simplicity. B. In *V. fischeri*, the Lux phosphorelay functions in largely the same manner to regulate bioluminescence, with homologues of LuxP, LuxQ,

LuxU and LuxO, but not CqsS or LuxN; *V. fischeri* encodes another putative SK, AinR. Additionally, *V. fischeri* uses the LuxR_{VH} homologue, LitR, to activate transcription of LuxR_{VF} (not similar to LuxR_{VH}), which promotes transcription of the *lux* operon (when bound to the Al produced by LuxI) (not depicted), leading to subsequent light production (bioluminescence). Regulators shaded in grey indicate those found in this study to be involved in biofilm formation. RscS is a SK known to control biofilm formation. Phosphorylation of the RR SypG is predicted to activate transcription of the *syp* locus, which encodes proteins thought to regulate, produce and transport a polysaccharide necessary for biofilm formation. The specific activity of LuxU in activating biofilm formation is unknown, but it appears from the current study to work at or above the level of *syp* transcription, likely at the level of SypG activation (indicated by a question mark). This figure is adapted from Visick (2005).

AinR, predicted to function at the same level as LuxQ (Gilson *et al.*, 1995; Stabb *et al.*, 2008; Miyashiro and Ruby, 2012). Furthermore, LuxO~P controls the transcription of a single *qrr* sRNA, which likely binds to and destabilizes the transcript of the transcriptional regulator LitR (a LuxR_{VH} homologue) (Miyamoto *et al.*, 2003; Miyashiro *et al.*, 2010). Finally, LitR controls the transcription of an additional downstream regulator, LuxR_{VF} (not similar to LuxR_{VH}) (Fidopiastis *et al.*, 2002), which activates the *lux* operon when bound by its AI (Stevens *et al.*, 1994; Sitnikov *et al.*, 1995). Thus, *V. fischeri* integrates AI cues not only to regulate the phosphorylation state of the RR LuxO, but also to regulate the transcription of the *lux* operon, adding another level of control to the production of cellular bioluminescence in this organism.

In addition to controlling bioluminescence, the Lux pathway in *V. fischeri* impacts other processes such as acetate metabolism (Studer *et al.*, 2008) and motility (Lupp and Ruby, 2005; Hussa *et al.*, 2007; Cao *et al.*, 2012). Lux

regulates a variety of processes in other vibrios as well. For example, *V. cholerae, V. anguillarum, V. vulnificus* and *V. parahaemolyticus* utilize the Lux pathway to control biofilm formation via regulators downstream of LuxO (Croxatto *et al.*, 2002; Hammer and Bassler, 2003; Enos-Berlage *et al.*, 2005; Lee *et al.*, 2007; Zhang *et al.*, 2012). For *V. fischeri*, no clear connection between the Lux pathway and biofilm formation has been observed, with the exception that a *litR* mutant exhibits a change in colony morphology (from translucent to opaque) consistent with a possible alteration in biofilm formation (Fidopiastis *et al.*, 2002). Thus, it seems likely that *V. fischeri* could utilize the Lux pathway to regulate biofilm formation, but the exact role, if any, has yet to be determined.

Vibrio fischeri is known to promote biofilm formation through the <u>sy</u>mbiotic <u>polysaccharide</u> (*syp*) locus (Fig. 1B). The *syp* locus is set of 18 genes thought to be involved in the regulation, production and transport of a polysaccharide involved in biofilm formation (Yip *et al.*, 2005; 2006;

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Shibata and Visick, 2012). This locus is transcriptionally controlled by the RR SypG, encoded within the syp locus, and σ^{54} (Yip *et al.*, 2005). Another regulator of the *svp* locus is the SK RscS, which functions upstream of SypG to promote syp transcription (Yip et al., 2006; Hussa et al., 2008). Under standard laboratory conditions, V. fischeri does not form robust biofilms (i.e. wrinkled colony formation and pellicle production). However, robust biofilms that are dependent on the syp locus can be induced by overexpression of either rscS (Yip et al., 2006) or svpG; for SypG to induce biofilm formation, the biofilm inhibitor protein SypE must be absent or inactivated (Hussa et al., 2008; Morris et al., 2011). This enhanced biofilm production correlates with a competitive advantage for colonization by V. fischeri of its symbiotic host, the squid Euprymna scolopes (Yip et al., 2006; Morris et al., 2011).

In the current study, we sought to identify other components involved in biofilm formation by V. fischeri and found that the Lux pathway plays a role in this phenotype. Regulation of biofilm formation by the Lux pathway in V. fischeri is novel, as LuxU plays an important role but LuxO involvement is minimal. Our data indicate that the Lux pathway in *V. fischeri* branches at LuxU to regulate both bioluminescence and syp-dependent biofilm formation. Additionally, we have evidence to suggest that the SK AinR plays a minimal role in regulating light production and no role in regulating biofilm formation, under our conditions. Thus, this work provides new insight into the mechanism by which the Lux pathway functions in V. fischeri, and also helps further our understanding of the complex regulatory network involved in controlling biofilm formation in this organism.

Results

Transposon mutagenesis reveals a regulatory connection between syp and lux

To better understand the requirements for biofilm formation in V. fischeri, we performed a random transposon mutagenesis of KV3299, a strain that lacks the syp biofilm inhibitor protein SypE. We then induced biofilm formation by introducing the *sypG* overexpression plasmid pEAH73; under these biofilm-inducing conditions, V. fischeri forms wrinkled colonies instead of smooth colonies (Hussa et al., 2008). We screened approximately 5000 mutants for those exhibiting a defect in wrinkled colony formation and found 27 that appeared to form smooth colonies. To verify the phenotypes of these mutants, we cured them of their sypG overexpression plasmid and then reintroduced it. All of the mutants remained defective in wrinkled colony formation and fell into two classes. Class 1 mutants (24 total) exhibited smooth colony morphology (compare Fig. 2A and B), while class 2 mutants (three total) exhib-



Fig. 2. Transposon mutagenesis reveals other regulators of biofilm formation in *V. fischeri*. A–C. Time-course assays of wrinkled colony formation induced by *sypG* overexpression using plasmid pEAH73. Cultures were constant and the Bernetic as a constant of the sector of the se

spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 45.5 h post spotting for the following strains: $\Delta sypE$ control (pEAH73/KV3299) (A), a representative class 1 mutant (pEAH73/KV5872; *syp::*Tn5 $\Delta sypE$) (B) and a representative class 2 mutant (pEAH73/KV4431; *luxQ::*Tn5 $\Delta sypE$) (C). An asterisk indicates the time at which wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments. D. A graphical depiction of the predicted *luxPQ* genes (block arrows) and Tn insertion sites (black triangles). There are 2 bp between the predicted translational stop site of *luxP* and the predicted translational start site of *luxQ*.

ited a substantial delay (approximately 6 h) in the start of wrinkled colony formation, but appeared similar to that of the parent strain at later times (compare Fig. 2A and C; asterisks indicate the time at which wrinkled colony formation is apparent, typically identified by ridge formation around the outer edge of the spot). To further evaluate these mutants, we performed Southern blot analysis on each mutant to determine whether the transposon had inserted within the syp locus, a location predicted to disrupt wrinkling. We found that the class 1 (smooth) mutants mapped within this locus, while the class 2 (delayed) mutants were unaltered in their syp regions (data not shown). These results confirm the assumption that wrinkled colony formation induced by sypG overexpression depends on the syp locus. Because our goal was to identify novel (non-syp) factors involved in biofilm formation, we pursued characterization of the class 2 mutants. Upon cloning and sequencing the DNA flanking the site of the Tn insertion in each class 2 mutant, we found that one insertion mapped near the end of *VF_0707*, while the other two were within *VF_0708* (Fig. 2D). These genes are predicted to encode LuxP and LuxQ, respectively, two proteins proposed to be involved in controlling bioluminescence in *V. fischeri* (Fig. 1B).

Loss of LuxQ affects bioluminescence and biofilm formation

In V. fischeri, LuxP and LuxQ are predicted to regulate bioluminescence due to their sequence similarity to the well-characterized proteins of V. harveyi (58% and 44% identical, 72% and 67% similar respectively) and to the functional conservation of other members of the lux regulatory pathway between V. harveyi and V. fischeri (reviewed in Stabb et al., 2008); however, the functions (bioluminescence or otherwise) of these two proteins in V. fischeri have not yet been assessed through mutagenesis studies. Thus, to understand the functions of these putative regulators, we asked whether these genes were involved in controlling bioluminescence, in addition to probing their role in controlling biofilm formation. As the Tn insertions were in a $\Delta sypE$ background, it was first necessary to ask whether loss of SypE impacted luminescence; we found that it did not substantially impact luminescence (Fig. S1A), regardless of whether we used OD₆₀₀ to estimate cell number (Fig. S1B) or determined the number of colony-forming units (Fig. S1C) to calculate the specific luminescence. Next, we assessed the impact of the Tn mutations on luminescence. The model (Fig. 1B), generated from work in V. harveyi (Bassler et al., 1994; Neiditch et al., 2005), predicts that a luxP mutant should fail to transmit the AI signal to LuxQ, causing LuxQ to remain a kinase; as a result, the levels of LuxO~P should be higher and luminescence should be lower. The model also predicts that the *luxQ* mutant should exhibit a decrease in LuxO~P levels, leading to increased luminescence. We found that all three mutants exhibited an increase in luminescence relative to their parent ($\Delta sypE$) (Fig. 3A).

Since the *luxP* Tn mutant did not exhibit the predicted luminescence phenotype, we hypothesized that the Tn insertion, which was located at the end of *luxP*, was polar on *luxQ*. To test this prediction, we constructed in-frame deletions of both *luxP* ($\Delta luxP$) and *luxQ* ($\Delta luxQ$) in both the $\triangle sypE$ and wild-type backgrounds. Neither mutation impacted growth of V. fischeri (data not shown). Regardless of the background, loss of LuxP decreased bioluminescence, while loss of LuxQ increased bioluminescence as predicted (Fig. 3B and data not shown). The luminescence of the $\Delta luxQ$ mutant could be restored to the level of the $luxQ^+$ control by expression of an epitope-tagged version of *luxQ* (*luxQ-FLAG*) in single copy from the chromosome (Fig. 3C). Together, these data indicate that: (i) LuxP and LuxQ are involved in controlling bioluminescence, as predicted, (ii) the Tn insertion within *luxP* was polar on *luxQ*, and (iii) *luxP* and *luxQ* likely comprise an operon (Fig. 2D).

As the Tn insertion in *luxP* was polar on *luxQ*, we predicted that the 6 h delay in biofilm formation initially observed from the Tn mutants was likely due to loss or disruption of *luxQ*. To test this prediction, we examined wrinkled colony formation by the $\Delta luxQ$ ($\Delta sypE$) mutant that overexpressed sypG. Like the Tn mutants, this mutant also exhibited a delay in wrinkled colony formation relative to the control (compare Fig. 4A and B). This delay in biofilm formation could be complemented by expression of *luxQ-FLAG* in single copy from the chromosome of the $\Delta luxQ$ mutant (Fig. 4A–C). Thus, LuxQ appears to control both bioluminescence and biofilm formation in V. fischeri. In contrast, the luxP mutation exerted relatively little effect on biofilm formation (Fig. S2); thus, we focused our subsequent studies on LuxQ and other Lux regulators.

LuxU exerts a more substantial impact on biofilm formation than LuxO

In V. harvevi, LuxQ functions through the phosphotransferase LuxU, to control the phosphorylation state of the RR LuxO (Fig. 1A) (Freeman and Bassler, 1999a,b). As LuxQ is involved in controlling biofilm formation in V. fischeri, we asked whether LuxU and LuxO were also involved. Thus, we generated deletions of both luxU ($\Delta luxU$) and luxO $(\Delta luxO)$ in the $\Delta sypE$ background. However, it was necessary to first confirm that our mutants exhibited the predicted pattern of luminescence [i.e. increased bioluminescence; for *luxO* mutants, this has been previously reported (Lupp et al., 2003; Hussa et al., 2007)] (Fig. 1B). As expected, both mutants exhibited an increase in bioluminescence relative to their parent (Figs 3D and S1). Neither mutant exhibited a growth defect (data not shown). Finally, a $\Delta luxU$ $\Delta luxO$ ($\Delta sypE$) mutant exhibited a luminescence phenotype similar to that of the individual mutants (Fig. 3D). Overall, these data confirm that LuxU functions to control bioluminescence in V. fischeri, as predicted.

Next, we introduced the *sypG* plasmid into the $\Delta luxU$ and $\Delta luxO$ mutants and assessed wrinkled colony formation. Similar to the loss of LuxQ, loss of LuxU resulted in a delay (about 7 h) in wrinkled colony formation (compare Fig. 5A and B). However, loss of LuxO resulted in only a slight, but reproducible delay (1.5 h) in wrinkled colony formation (compare Fig. 5A and C); we also observed the same slight delay for a *luxO*::*kan* mutant (data not shown), confirming the results of the $\Delta luxO$ mutant. These data suggest that, under our conditions, LuxU plays a more critical role than LuxO in controlling biofilm formation. These results also suggest that LuxU may function independently of LuxO to control biofilm formation. To investigate this possibility further, we evaluated biofilm formation



Fig. 3. Luminescence of *lux* mutants in culture. Cultures were grown in SWTO and incubated at 24°C with vigorous shaking. Luminescence and OD₆₀₀ were measured over time until maximum luminescence was achieved (between OD₆₀₀ 1.5 and 2). All data are plotted as specific luminescence (Sp. lum.; relative luminescence divided by OD₆₀₀) versus OD₆₀₀ and are representative of at least three independent experiments.

A. $\Delta sypE$ control (black squares; KV3299), *luxP*::Tn5 $\Delta sypE$ (white circles; KV4430), *luxQ*::Tn5 $\Delta sypE$ (grey diamonds; KV4431), *luxQ*::Tn5 $\Delta sypE$ (black triangles; KV4432).

B. $\Delta sypE$ control (black squares; KV3299), $\Delta luxP \Delta sypE$ (white circles; KV5347), $\Delta luxQ \Delta sypE$ (black circles; KV5394), $\Delta luxU \Delta sypE$ (grey triangles; KV4830), $\Delta luxQ \Delta luxU \Delta sypE$ (black diamonds; KV6008).

C. ΔsypE attTn7::erm control (black squares; KV4390), ΔluxQ ΔsypE attTn7::erm (black circles; KV5973), ΔluxQ ΔsypE attTn7::luxQ-FLAG (white circles; KV5902), ΔluxQ ΔsypE attTn7::luxQ-A216P-FLAG (grey triangles; KV5904), ΔluxQ ΔsypE attTn7::luxQ-H378A-FLAG (black diamonds; KV5903).

D. Δ sypE control (black squares; KV3299), Δ luxO Δ sypE (white circles; KV5468), Δ luxU Δ sypE (black triangles; KV4830), Δ luxO Δ luxU Δ sypE (grey diamonds; KV5472).

E. ΔsypE attTn7::erm control (black squares; KV4390), ΔluxU ΔsypE attTn7::erm (black triangles; KV5974), ΔluxU ΔsypE attTn7::luxU-FLAG (white triangles; KV5905), ΔluxU ΔsypE attTn7::luxU-H61A-FLAG (white circles; KV5906).

F. Δ sypE control (black squares; KV3299), Δ ainR Δ sypE (grey circles; KV6196), Δ luxU Δ sypE (black triangles; KV4830), Δ ainR Δ luxU Δ sypE (black diamonds; KV6259).

The data for KV3299 in (B) are the same as that shown in (D). The data for KV4390 in (C) are the same as that shown in (E).

by the $\Delta luxO \Delta luxU$ ($\Delta sypE$) mutant. We predicted that if LuxU functions through LuxO to regulate biofilm formation, then loss of both LuxU and LuxO would result in a phenotype similar to loss of LuxO alone (i.e. a 1.5 h delay). This was not the case: loss of both regulators resulted in an 8 h delay in wrinkled colony formation (compare Fig. 5A and D). This delay supports the hypothesis that the two regulators function independently to impact biofilm formation. Furthermore, this result contrasts with the luminescence results, in which the phenotypes of the *luxU*, *luxO* and *luxU luxO* mutants were similar (Fig. 3D), and which suggest that LuxU likely functions through LuxO to control bioluminescence. Together, these data suggest that the Lux pathway branches at LuxU to control both bioluminescence and biofilm formation (Fig. 1B). Since loss of LuxU resulted in a more severe biofilm phenotype than loss of LuxO, we chose to pursue the role of LuxU (and its inputs) in the current study.



Fig. 4. The effect of *luxQ* point mutations on biofilm formation. Time-course assays of wrinkled colony formation induced by sypG overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 45 h post spotting for the following strains: △sypE attTn7::erm control (pEAH73/KV4390) (A), $\Delta luxQ \Delta sypE attTn7::erm$ (pEAH73/KV5973) (B), ∆luxQ ∆sypE attTn7::luxQ-FLAG (pEAH73/KV5902) (C), ∆luxQ ∆sypE attTn7::luxQ-A216P-FLAG (pEAH73/KV5904) (D) and $\Delta luxQ \Delta sypE$ attTn7::luxQ-H378A-FLAG (pEAH73/KV5903) (E). An asterisk indicates the time at which wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments.

LuxQ kinase activity promotes biofilm formation

Our current data suggest that LuxQ functions as a positive regulator of biofilm formation under our conditions. Because this SK is predicted to function as both a kinase and a phosphatase (Freeman and Bassler, 1999a; Neiditch et al., 2006), we asked whether the ability of LuxQ to positively regulate biofilm formation depended upon its kinase and/or phosphatase activity. Previous work from V. harveyi had demonstrated that certain point mutations cause the loss of one activity but not the other (i.e. kinase activity is lost, while phosphatase activity is retained and vice versa) (Neiditch et al., 2006). Thus, we generated point mutations in the V. fischeri luxQ gene that are predicted to cause either loss of phosphatase activity (luxQ-A216P; kin+/phos-) or loss of kinase activity (luxQ-H378A; kin-/phos+), while retaining the other activity respectively. We then expressed these *luxQ* alleles in single copy from the chromosome of the $\Delta luxQ$ mutant. To confirm that these LuxQ derivatives were functional, we examined their ability to control light production. According to the model (Fig. 1B) and work from V. harveyi (Neiditch et al., 2006), a phosphatase mutant (LuxQ-A216P, kin+/phos-) should exhibit a decrease in bioluminescence (due to an increase in LuxO~P), while a kinase mutant (LuxQ-H378A, kin-/phos+) should exhibit an increase in bioluminescence (due to a decrease in LuxO~P). Indeed, each mutant exhibited the expected pattern of luminescence (Fig. 3C), indicating that the proteins produced were functional and behaved as predicted.

We next assessed the ability of these alleles to complement the $\Delta luxQ$ mutant with respect to the timing of wrinkled colony formation. We found that the phosphatase mutant, LuxQ-A216P (kin+/phos-), could complement the *luxQ* mutant, restoring the timing of wrinkled colony formation to approximately that of the control strain (*luxQ*⁺) and the wild-type-complemented $\Delta luxQ$ mutant (compare Fig. 4A, C and D). In contrast, the kinase mutant, LuxQ-H378A (kin-/phos+), failed to complement the *luxQ* mutant; this strain exhibited wrinkled colony formation that was indistinguishable from the $\Delta luxQ$ parent (compare Fig. 4A, B and E). These data suggest that the kinase activity of LuxQ, but not its phosphatase activity, is necessary to regulate biofilm formation.

The impact of LuxQ on biofilm formation depends on LuxU

Our data indicate that LuxQ (specifically its kinase activity) and LuxU are necessary to regulate biofilm formation. According to the model (Fig. 1B), LuxQ is predicted to function through LuxU. To test this hypothesis, we first



Fig. 5. Wrinkled colony formation by *luxU* and *luxO* mutants. Time-course assays of wrinkled colony formation induced by *sypG* overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 45 h post spotting for the following strains: $\Delta sypE$ control (pEAH73/KV3299) (A), $\Delta luxU \Delta sypE$ (pEAH73/KV4830) (B), $\Delta luxO \Delta sypE$ (pEAH73/KV5468) (C) and $\Delta luxO \Delta luxU \Delta sypE$ (pEAH73/KV5472) (D). An asterisk indicates the time at which wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments.

asked whether LuxQ functioned through LuxU to regulate bioluminescence. If this were the case, we would expect that a *luxQ luxU* mutant would phenocopy a *luxU* mutant, and indeed it did (Fig. 3B). To further evaluate this regulatory connection, we expressed the *luxQ-A216P* (kin+/ phos-) allele in the *luxQ luxU* mutant. Whereas, in the context of the *luxQ (luxU*⁺) background this allele decreased luminescence, it failed to do so when *luxU* was also disrupted: the levels of luminescence produced by the *luxQ luxU* mutant expressing *luxQ-A216P* (kin+/phos-) were indistinguishable from that of the *luxU* mutant (Fig. 6A). These data suggest that LuxQ functions through LuxU to regulate bioluminescence.

Next, we asked whether LuxQ functioned through LuxU to control biofilm formation. We first evaluated biofilm formation by the *luxQ luxU* (*sypE*) mutant. We found that the double mutant exhibited a delay in wrinkled colony formation similar to that seen with the individual *luxQ* and *luxU* mutants (Fig. S3), rather than an additive delay. Thus, these results suggest that LuxQ and LuxU function in the

same pathway to regulate biofilm formation. To probe this relationship further, we utilized the *luxQ-A216P* allele, which permits complementation of the *luxQ* mutation (Fig. 4). We hypothesized that if LuxU were necessary for LuxQ to regulate biofilm formation, then disruption of *luxU* in the *luxQ* mutant expressing *luxQ-A216P* should delay biofilm formation relative to the *luxU*⁺ strain. Indeed, this was the case (Fig. 6B). These data suggest that the ability of LuxQ to positively regulate biofilm formation depends upon LuxU.

Biofilm formation depends on the conserved site of phosphorylation in LuxU

In *V. harveyi*, LuxU serves as a phosphotransferase, shuttling phosphoryl groups between the SKs and the RR LuxO (Fig. 1A). This role depends upon the conserved site of phosphorylation, His58 (Freeman and Bassler, 1999b). To determine whether the *V. fischeri* homologue functions in a similar manner, we first constructed an epitope-tagged version of *luxU* (*luxU-FLAG*). Expression of this allele in



Fig. 6. Luminescence and wrinkled colony formation by *luxQ* and *luxU* mutants. A. Luminescence of lux mutants in culture. Cultures were grown in SWTO and incubated at 24°C with vigorous shaking. Luminescence and OD₆₀₀ were measured over time until maximum luminescence was achieved (between OD₆₀₀ 1.5 and 2) for the following strains: ∆luxQ attTn7::erm control (black triangles; KV5973), ∆luxU ∆sypE attTn7::erm (black diamonds; KV5974), *\(\Delta\luxQ\) \(\Delta\symbol{sypE\)* attTn7::luxQ-A216P-FLAG (white circles; KV5904) and $\Delta luxQ \Delta luxU \Delta svpE$ attTn7::luxQ-A216P-FLAG (black circles; KV6054). All data are plotted as specific luminescence (Sp. lum.; relative luminescence divided by OD₆₀₀) versus OD₆₀₀ and are representative of at least three independent experiments.

B. Time-course assays of wrinkled colony formation induced by sypG overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 42 h post spotting for the following strains: ∆*sypE att*Tn7::*erm* control (pEAH73/KV4390), *\(\Delta\) luxQ \(\Delta\) sypE* attTn7::luxQ-A216P-FLAG (pEAH73/KV5904) and $\Delta luxQ \Delta luxU \Delta sypE$ attTn7::luxQ-A216P-FLAG (pEAH73/KV6054). An asterisk indicates the time at which wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments.

single copy from the chromosome of the $\Delta luxU$ mutant restored luminescence to that of the control (Fig. 3E), as well as the normal timing of wrinkled colony formation (compare Fig. 7A-C). Next, we substituted the predicted, conserved histidine for an alanine (H61A) in the luxU-FLAG construct and introduced this allele into the chromosome of the $\Delta luxU$ mutant. The $\Delta luxU$ mutant expressing the *luxU-H61A* allele failed to restore luminescence to the level of the parent (Fig. 3E) and exhibited the same 6 h delay in wrinkled colony formation as the uncomplemented $\Delta luxU$ mutant (compare Fig. 7B and D). To ensure that the lack of complementation was not due to a reduction or loss of the protein, we performed Western blot analysis and found that protein was expressed from both alleles (Fig. S4). Together, these data suggest that the conserved site of phosphorylation in LuxU is necessary to regulate biofilm formation. Thus, it appears that key residues predicted to be involved in phosphotransfer are required for

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AinR does not function as predicted to regulate bioluminescence and exerts no impact on biofilm formation

It has been predicted that, like the V. harveyi Lux pathway, multiple SKs feed into LuxU to control bioluminescence in V. fischeri (Visick, 2005; Stabb et al., 2008; Miyashiro and Ruby, 2012). In particular, the SK AinR is proposed to function at the same level as LuxQ to control bioluminescence in V. fischeri (Fig. 1B) (Gilson et al., 1995; Stabb et al., 2008). To determine whether AinR is involved in controlling bioluminescence and biofilm formation, we generated a $\Delta ainR$ ($\Delta sypE$) mutant and first assessed its luminescence phenotype; no study of AinR has assessed its role in controlling bioluminescence in liquid culture. The model (Fig. 1B) predicts that, similar to loss of LuxQ, loss of AinR would result in an increase in luminescence. However, this was not the case: loss of AinR resulted in a consistent but very slight decrease in luminescence as



Fig. 7. Wrinkled colony formation by complemented $\Delta luxU$ mutants. Time-course assays of wrinkled colony formation induced by *sypG* overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 45 h post spotting for the following strains: $\Delta sypE$ attTn7::erm control (pEAH73/KV4390) (A), $\Delta luxU \Delta sypE$ attTn7::luxU-FLAG (pEAH73/KV5905) (C) and $\Delta luxU \Delta sypE$ attTn7::luxU-H61A-FLAG (pEAH73/KV5906) (D). An asterisk indicates the time at which wrinkled colony formation was apparent,

wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments.

compared to the control (Figs 3F and S5). To determine whether AinR functioned through the known phosphorelay pathway (i.e. through LuxU), we generated a $\Delta ainR \Delta luxU$ ($\Delta sypE$) mutant and assessed its luminescence phenotype. We expected that the double mutant would exhibit the luminescence phenotype of the *luxU* single mutant. Surprisingly, this mutant consistently exhibited in an intermediate luminescence phenotype: the $\Delta ainR \Delta luxU$ mutant was brighter than the $\Delta ainR$ mutant, but not as bright as the $\Delta luxU$ mutant (Fig. 3F). These data suggested that AinR may play only a minor role in controlling bioluminescence under these conditions.

As a putative SK, AinR is predicted to recognize and respond to the AI N-octanoyl-homoserine lactone (C8-HSL) (Gilson et al., 1995). Thus, the diminished luminescence phenotypes of the $\Delta ainR$ and $\Delta ainR \Delta luxU$ mutants could result from a failure of this mutant to respond to C8-HSL. Alternatively, deletion of ainR could impact expression of the upstream gene ainS, which encodes the C8-HSL synthase protein. An impact on AinS synthesis could lead to decreased amounts of C8-HSL and decreased light production, potentially via direct control of the lux operon, as previously demonstrated (Kuo et al., 1996; Egland and Greenberg, 2000). To distinguish between these possibilities, we added exogenous C8-HSL to the ainR mutants and controls. We reasoned that, if the ainR mutants were defective in their response to C8-HSL, then they would still exhibit diminished luminescence relative to their controls. This appeared not to be the case, however, as addition of C8-HSL to the $\Delta ainR$ and $\Delta ainR$ $\Delta luxU$ mutants increased their luminescence levels to those of the control strain and the $\Delta luxU$ mutant respectively (Fig. S5B). These data suggest that the decrease in luminescence by both the $\Delta ainR$ and $\Delta ainR \Delta luxU$ mutants is likely due to decreased levels of C8-HSL, whose activity in promoting luminescence is largely or fully independent of the function of AinR, at least under our conditions. Thus, the role of AinR in controlling luminescence remains unclear.

Although AinR did not function as predicted in controlling bioluminescence, we wondered whether loss of AinR would impact biofilm formation. This was not the case: the $\Delta ainR$ mutant exhibited no defect in biofilm formation, while the $\Delta ainR \Delta luxU$ mutant exhibited the defect of the luxU mutant (Fig. S6) and could be complemented when the wild-type allele of luxU-FLAG was expressed in single copy from the chromosome (data not shown). Thus, AinR has no impact on biofilm formation, and its role in controlling bioluminescence remains unclear. Further work will be necessary to determine what role AinR plays, if any, in controlling luminescence in *V. fischeri*.

LuxU, but not RscS, is necessary to regulate syp-dependent biofilm formation under SypG-inducing conditions

Since the only known role of LuxU is to serve as a phosphoryl-donor (Freeman and Bassler, 1999b; Shikuma *et al.*, 2009) and LuxU impacts *syp*-dependent biofilm formation, we hypothesized that it could function







upstream of SypG, a RR known to be required for transcription of the syp locus (Yip et al., 2005). As previous studies had demonstrated that the SK RscS functions upstream of the RR SypG to control syp-dependent biofilm formation (Fig. 1B) (Yip et al., 2006; Hussa et al., 2008), we questioned the relative importance of these two potential inputs, RscS and LuxU, on SypG-induced biofilm formation. We thus generated svpE mutants with deletions in *luxU*, *rscS* or both and evaluated SypGinduced biofilm formation. Surprisingly, only loss of LuxU exerted an impact: whereas the luxU mutant exhibited a delay in wrinkled colony formation (compare Fig. 8A and C), the rscS mutant showed no significant defect in biofilm formation under these conditions (compare Fig. 8A and B). Even when the *rscS* and *luxU* mutations were combined, this mutant exhibited the same delay as the luxU mutant alone and could be complemented when the wild-type allele of *luxU-FLAG* was expressed in single copy from the chromosome (compare Fig. 8C and D and data not shown). Overall, these data indicate that LuxU plays a more important role than RscS in controlling biofilm formation when sypG is overexpressed.

LuxU functions at or above SypG to impact syp transcription

As LuxU is necessary to promote *syp*-dependent biofilm formation, we sought to determine whether LuxU functioned upstream of SypG to control its activation (phosphorylation). If so, then we would expect that a phosphorylation-independent allele of SypG would be 'blind' to the presence or absence of LuxU. We thus overexpressed a version of SypG in which the conserved site of phosphorylation, D53, was substituted for a glutamate (D53E). This substitution in RRs has previously been shown to promote the active state of the RR (Sanders et al., 1989; 1992; Freeman and Bassler, 1999a). Indeed, this substitution in SypG caused an increase in SypG activity, as measured by syp transcription (Hussa et al., 2008). Consistent with this increased activity, when overexpressed in the $\triangle svpE$ mutant, svpG-D53E induced wrinkling sooner than when the wild-type allele of sypG was overexpressed (9-10 h vs. 13-15 h respectively). When sypG-D53E was overexpressed in the $\Delta luxU \Delta sypE$ mutant, the timing of wrinkled colony formation was not delayed, but rather was similar to that of the $luxU^+$ control (Fig. 9A). These data are consistent with a model in which LuxU functions at or above the level of SypG.

It remains formally possible that the accelerated wrinkling effects of the *sypG-D53E* allele, combined with the delayed wrinkling caused by the loss of LuxU, result in a strain with a net timing of biofilm formation similar to the wild-type strain. Thus, to further probe the level at which LuxU exerts its impact on biofilm formation, we asked whether loss of LuxU affected transcription of the SypGcontrolled *sypA* gene using a *lacZ* reporter fusion. We assayed β -galactosidase activity from the reporter expressed from the chromosomes of the $\Delta luxU$ and the



Fig. 9. The role of LuxU in syp activation.

A. Time-course assays of wrinkled colony formation induced by *sypG-D53E* overexpression using plasmid pKV276. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 24 h post spotting for the following strains: $\Delta sypE$ control (pKV276/KV3299) and $\Delta luxU \Delta sypE$ (pKV276/KV4830). Data are representative of at least three independent experiments.

B. SypG-induced *syp* transcription from P_{sypA} -*lacZ* reporter strains. Cultures of *sypG* overexpressing strains $\Delta sypE$ attTn7:: P_{sypA} -*lacZ* (white bars; pEAH73/KV4926) and $\Delta luxU \Delta sypE$ attTn7:: P_{sypA} -*lacZ* (grey bars; pEAH73/KV5516) were inoculated in LBS containing Tet and grown at 28°C with shaking. Samples were collected at 12 and 24 h and assessed for β -galactosidase activity (in Miller units) as a measure of promoter activity. All experiments were performed in triplicate. Data are a combination of two independent experiments with error bars representing the standard error. The *P*-value refers to the variation between the two samples as indicated by the brackets.

luxU⁺ strains that overexpressed the wild-type allele of *sypG*. Loss of LuxU resulted in a decrease in *syp* transcription at two time points tested (12 and 24 h) (Fig. 9B). Thus, these data suggest that LuxU functions at or above the level of *syp* transcription, potentially due to an impact on SypG activation (Fig. 1B).

Discussion

In this study, we identified a novel connection between the Lux pathway and biofilm formation in *V. fischeri*. Specifically, we found that disruption of either the gene encoding the SK LuxQ or the gene encoding the phosphotransferase

LuxU caused a delay in SypG-induced biofilm formation. Surprisingly, this effect was independent of LuxO, which exerted only a minor impact on biofilm formation. However, LuxU does seem to function through LuxO to regulate bioluminescence. Thus, the Lux pathway appears to branch at LuxU to regulate bioluminescence through LuxO and biofilm formation via a SypG-dependent pathway.

From the data presented in this work, we propose a model in which LuxQ functions through LuxU to regulate svp-dependent biofilm formation through activation of the RR SypG (Fig. 1B). Support for the idea that LuxQ and LuxU serve as phosphoryl-donors to a downstream regulator of biofilm formation is as follows: (i) the kinase activity of LuxQ is necessary to promote biofilm formation, (ii) the predicted, conserved site of phosphorylation in LuxU is necessary to regulate biofilm formation, (iii) the only known role of LuxU in the literature is as a histidine phosphotransferase (HPt) (Freeman and Bassler, 1999b; Shikuma et al., 2009), (iv) LuxQ depends on LuxU to regulate biofilm formation, and (v) the downstream RR of the Lux pathway, LuxO, is not required for the effect of LuxU on biofilm formation. Together, these data suggest that phosphotransfer is necessary for LuxQ and LuxU to regulate biofilm formation via a regulator distinct from LuxO. In support of the idea that LuxU serves as an input to regulate the activity of the RR SypG, we found that: (i) a 'constitutively active' allele of SypG overcomes the requirement for LuxU, and (ii) LuxU functions at or above the level of syp transcription. Overall, these data suggest that LuxU functions at or above the level of SypG, potentially at the level of SypG phosphorylation. This possibility is further supported by the fact that SypG and LuxO have similar domain structures (both are σ^{54} -dependent RRs) and exhibit 50% identity to each other. However, proof of such a possibility awaits additional biochemical experimentation; to date, attempts to examine the phosphorylation state of SypG have been unsuccessful. Thus, while our data support the hypothesis that LuxU could serve as a phosphoryl-donor to SypG, the regulation is clearly complex and may include currently unknown regulators.

It has been previously proposed, in two other *Vibrio* species, that LuxU can function independently of LuxO to control the activity of downstream targets of the Lux pathway. The first example is from the fish pathogen *V. anguillarum*. In this organism, VanU functions through VanO to regulate the expression of the LuxR_{VH} homologue, VanT, by activating expression of qrr1-4 (Croxatto *et al.*, 2004). VanU also appears to act through a VanO-independent mechanism to inhibit the expression of qrr1-4 (Croxatto *et al.*, 2004; Weber *et al.*, 2011). Weber *et al.* (2011) thus hypothesize that VanU functions through another RR to repress expression of qrr1-4. Similarly, in *Vibrio alginolyticus*, Liu *et al.* (2011) propose that LuxU functions, at least in part, independently of LuxO to control expression of a

downstream regulator, LuxT, likely through another RR. However, in neither case has a downstream RR been identified.

There are at least a couple of examples in the literature in which a single domain HPt protein interacts with more than one target RR. *Caulobacter crescentus* ChpT is one such example. ChpT phosphorylates the RRs CtrA and CpdR with equal affinity *in vitro* (Biondi *et al.*, 2006); these phosphorylation events are critical during cell cycle progression. Phosphorylation of CtrA activates this protein, permitting it to bind DNA and control, among other things, DNA replication (Domian *et al.*, 1997; Quon *et al.*, 1998; Jacobs *et al.*, 2003). In contrast, it is the unphosphorylated form of CpdR that is active; in this state, CpdR indirectly promotes degradation of (unphosphorylated) CtrA (Iniesta *et al.*, 2006), permitting the cell to replicate its DNA. Thus, the same phosphorelay controls two separate RRs to exert opposite effects on protein activity.

Another well-studied example of an HPt protein interacting with two RRs occurs in the yeast Saccharomyces cerevisiae. In this organism, the HPt protein YPD1 serves as a phosphoryl-donor to the RRs SSK1 and SKN7 under hypo-osmotic conditions (Li et al., 1998). However, YPD1 interacts differently with each RR. For example, YPD1 stabilizes the phosphorylated state of the RR SSK1 via protein-protein interactions, but does not form stable complexes with the RR SNK1 (Janiak-Spens et al., 2000). Phosphorylation of SSK1 inactivates this regulator until the cell experiences hyperosmotic conditions, in which case SSK1 is rapidly dephosphorylated and activates a downstream pathway involved in controlling osmotic stress genes (Posas et al., 1996; Posas and Saito, 1998). In contrast, phosphorylation of SNK1 promotes activation of a downstream pathway involved in controlling genes for the cell wall and cell cycle (Morgan et al., 1995; Li et al., 1998; Bouquin et al., 1999). These activities of YPD1 allow for the co-ordinated regulation of multiple pathways in S. cerevisiae. It is possible that LuxU similarly provides a mechanism for co-ordination of two distinct pathways in V. fischeri.

One question that remains is why only the kinase activity of LuxQ, but not its phosphatase activity, is important for biofilm formation. Furthermore, making LuxQ a 'constitutive' kinase through three predicted routes (LuxQ-A216P mutation, deletion of *luxP* or deletion of *luxS*) did not (reproducibly, in the case of LuxQ-A216P) promote accelerated biofilm formation by *V. fischeri* (Figs 4D and S2, and V. A. Ray and K. L. Visick, unpubl. data). Potentially, similar to the yeast system described above, LuxU could interact differently with LuxO and SypG, serving as a phosphoryldonor to both, but only removing the phosphoryl groups from LuxO. Additional work is necessary to better understand the role of the Lux pathway in influencing biofilm formation.

Our work provides insight into the control of biofilm formation by V. fischeri, but also challenges the current model of how the Lux pathway regulates bioluminescence in V. fischeri. While LuxP, LuxQ, LuxU and LuxO appeared to function to regulate bioluminescence as predicted [or as previously shown for LuxO (Miyamoto et al., 2000)] (Figs 1B and 3B and D), the SK AinR did not. Loss of AinR led to a slight decrease in luminescence compared to the control strain (Figs 3F and S5), while loss of both AinR and LuxU resulted in an intermediate level of luminescence (Fig. 3F). However, the decreased luminescence of the ainR and ainR luxU mutants could be overcome by the addition of exogenous C8-HSL, which is normally produced by the AI synthase AinS. From these data, we conclude that deletion of ainR impacts expression of the gene encoding ainS (located directly upstream of ainR), and that loss of AinR itself has little impact on bioluminescence. Furthermore, it seems possible that AinR may not function through the known phosphorelay, or at least not through LuxU, to control bioluminescence. These data are not inconsistent with those reported by Lyell et al., who showed that, on solid media, loss of AinS resulted in an increased luminescence phenotype that did not depend on AinR function (Lyell et al., 2010). Thus, additional work is necessary to understand what role AinR may have in controlling this process in V. fischeri.

If AinR does not function through the known phosphorelay to regulate bioluminescence, does LuxQ serve as the only input? Our data suggest that this is not the case: loss of LuxU resulted in a greater increase in luminescence than did the loss of LuxQ (Fig. 3B). In both *V. cholerae* and *V. harveyi*, three SKs feed into LuxU (LuxQ, CqsS and VpsS, and LuxQ, LuxN and CqsS respectively) (Ng and Bassler, 2009). In *V. fischeri*, no gene for CqsS exists, but one for VpsS is present (Shikuma *et al.*, 2009). Thus, it is possible that VpsS may also feed into LuxU and serve as another input to regulate light production and possibly biofilm formation. Identifying a missing SK(s) would be an interesting future direction.

This is the first study to examine the role of RscS in biofilm formation under SypG-inducing conditions (overexpression of *sypG* in a $\Delta sypE$ background). Previous studies have already demonstrated that RscS functions upstream of SypG to induce *syp* transcription in a manner that depends on *sypG* (Hussa *et al.*, 2008) and that RscS is critical in symbiotic biofilm (aggregate) formation and colonization (Visick and Skoufos, 2001; Yip *et al.*, 2006); however, these previous studies have only explored how overexpression of *rscS* impacts biofilm formation or how its loss impacts colonization. Our findings indicate that LuxU is more important than RscS for biofilm formation under the conditions we used here. In contrast to these results, our preliminary data for the impact of a *luxU* mutation on the ability of *V. fischeri* to colonize squid

Table 1. V. fischeri strains used in this study.

Strains	Relevant genotype	Reference
CL42	luxO::kan	Lupp <i>et al.</i> (2003)
ES114	Wild-type	Boettcher and Ruby (1990)
KV3299	∆sypE	Hussa et al. (2008)
KV4390	<i>∆sypE att</i> Tn <i>7</i> :: <i>erm</i>	Morris et al. (2011)
KV4430	<i>luxP</i> ::Tn5 ∆ <i>sypE</i>	This study
KV4431	luxQ::Tn5 ∆sypE	This study
KV4432	<i>luxQ</i> ::Tn5 ∆ <i>sypE</i>	This study
KV4828	luxO::kan ∆sypE	This study
KV4830	$\Delta luxU \Delta sypE$	This study
KV4926	<i>∆sypE att</i> Tn <i>7</i> ::P _{sypA} -lacZ (Erm ^r)	This study
KV5347	$\Delta luxP \Delta sypE$	This study
KV5394	$\Delta luxQ \Delta sypE$	This study
KV5468	$\Delta luxO \Delta sypE$	This study
KV5472	$\Delta luxO \Delta luxU \Delta sypE$	This study
KV5516	<i>∆luxU ∆sypE att</i> Tn <i>7</i> ::P _{sypA} -lacZ (Erm')	This study
KV5872	<i>syp</i> ::Tn <i>5</i> ∆ <i>sypE</i>	This study
KV5902	∆luxQ ∆sypE atfTn7::luxQ-FLAG	This study
KV5903	∆luxQ ∆sypE atfTn7::luxQ-H378A-FLAG	This study
KV5904	∆luxQ ∆sypE attTn7::luxQ-A216P-FLAG	This study
KV5905	∆luxU ∆sypE atfTn7::luxU-FLAG	This study
KV5906	∆luxU ∆sypE attTn7::luxU-H61A-FLAG	This study
KV5973	∆luxQ ∆sypE atfTn7::erm	This study
KV5974	∆luxU ∆sypE attTn7::erm	This study
KV6008	$\Delta luxQ \Delta luxU \Delta sypE$	This study
KV6054	∆luxQ ∆luxU ∆sypE atfTn7::luxQ-A216P-FLAG	This study
KV6196	$\Delta ainR \Delta sypE$	This study
KV6259	$\Delta ainR \Delta luxU \Delta sypE$	This study
KV6268	$\Delta rscS \Delta sypE$	This study
KV6269	ΔrscS ΔluxU ΔsypE	This study

revealed, at most, a mild defect due to loss of LuxU (V. A. Ray and K. L. Visick, unpubl. data). These data suggest, perhaps not surprisingly, that our biofilm-inducing (*sypG* overexpression) conditions do not fully reflect the dynamics in nature (during colonization). It is of interest to note, however, that not all symbiosis-competent strains of *V. fischeri* encode a functional RscS protein (Mandel *et al.*, 2009; Gyllborg *et al.*, 2012). Therefore, it is not unreasonable to imagine that another pathway such as Lux could contribute to *syp* induction and biofilm formation during colonization. This work thus provides an important framework for deepening our understanding of the complex regulatory control over processes critical to colonization by *V. fischeri*.

Experimental procedures

Bacterial strains and media

Vibrio fischeri strains utilized in this study are shown in Table 1. Strains used in this study were derived from strain ES114, a bacterial isolate from *E. scolopes* (Boettcher and Ruby, 1990). For routine culturing, *V. fischeri* strains were grown in LBS (Luria–Bertani salt) medium (Graf *et al.*, 1994). For luminescence studies, *V. fischeri* strains were grown in Sea Water Tryptone (SWT) (Yip *et al.*, 2005) and SWTO (Bose *et al.*, 2007). All derivatives of *V. fischeri* were gener-

ated via conjugation, as previously described (DeLoney et al., 2002). Escherichia coli strains GT115 (InvivoGen, San Diego, CA, USA), Tam1 λ pir (Active Motif, Carlsbad, CA, USA), β3914 (Le Roux et al., 2007), π3813 (Le Roux et al., 2007) and CC118 (Herrero et al., 1990) were used for cloning and conjugation. All E. coli strains were grown in Luria-Bertani medium (Davis et al., 1980). Solid media were made using agar to a final concentration of 1.5%. Antibiotics were added to cultures when appropriate to the following final concentrations: ampicillin (Ap) at 100 µg ml⁻¹ (E. coli), tetracycline (Tet) at 15 µg ml⁻¹ (E. coli) or 5 µg ml⁻¹ (V. fischeri), chloramphenicol (Cm) at 20 or 25 µg ml⁻¹ (E. coli) or 5 µg ml⁻¹ (V. fischeri), kanamycin (Kan) at 50 µg ml⁻¹ (E. coli) or 100 µg ml⁻¹ (V. fischeri) and erythromycin (Erm) at 5 µg ml⁻¹ (*V. fischeri*). Along with any necessary antibiotics, diaminopimelate (DAP) was added to a final concentration of 0.3 mM for *E. coli* strain β3914 and thymidine was added to a final concentration of 0.3 mM for *E. coli* strain π 3813.

Transposon mutagenesis and identification of mutants with wrinkling defects

Transposon mutants were generated as described previously (Lyell *et al.*, 2008). Briefly, plasmid pEVS170, containing the mini-Tn5 transposon, was introduced into *V. fischeri* strain KV3299 via conjugation. Ex-conjugates were then pooled and the *sypG* overexpression plasmid pEAH73 was introduced via conjugation. The resultant ex-conjugates were then screened for their ability to form wrinkled colonies. Any mutants found to

Table 2. Plasmids used in this study.

Discussion	Description	0
Plasmids		Source or reference
pEAH73	pKV69 carrying wild-type sypG; Cm ^r Tet ^r	Hussa <i>et al</i> . (2008)
pEAH90	pEVS107 the P _{sypA} promoter region (generated with primers 714 and 782) upstream of promoterless <i>lacZ</i>	This study
pEVS104	Conjugal helper plasmid (<i>tra trb</i>); Kan ^r	Stabb and Ruby (2002)
pEVS107	Mini-Tn7 delivery plasmid; oriR6K, mob; Kan', Erm'	McCann <i>et al.</i> (2003)
pJET1.2	Commercial cloning vector; Ap ^r	Fermentas
pKV69	Mobilizable vector; Cm' Tet'	Visick and Skoufos (2001)
pKV276	pEAH73 with D53E mutation in <i>sypG</i> ; Cm' Tet'	Hussa <i>et al</i> . (2008)
pKV363	Mobilizable suicide vector; Cm ^r	Shibata and Visick (2012)
pKV456	pKV363 containing 1.1 kb sequence flanking <i>rscS</i> using primers 1494-97	This study
pSW7848	Mobilizable suicide vector; Cm ^r	Marie-Eve Val
pVAR17	pSW7848 containing 2 kb sequence flanking <i>sypE</i> derived from pCLD19 (Hussa <i>et al.</i> , 2008)	This study
pVAR18	pSW7848 containing 3.3 kb sequence flanking <i>luxU</i> using primers 995, 996, 1017 and 1018	This study
pVAR29	pKV363 containing 850 bp sequencing flanking <i>luxQ</i> using primers 1286, 1287, 1288 and 1304	This study
pVAR30	pKV363 containing 1.1 kb sequence flanking <i>luxP</i> using primers 1282, 1283, 1284 and 1303	This study
pVAR36	pKV363 containing 1.1 kb sequence flanking <i>luxO</i> using primers 1319, 1320, 1344 and 1345	This study
pVAR37	pKV363 containing 1.1 kb sequence flanking <i>luxO</i> and <i>luxU</i> using primers 1319, 1321, 1344 and 1346	This study
pVAR52	pEVS107 with P _{lacz} containing 2.3 kb <i>luxQ</i> -FLAG allele using primers 1314 and 1437	This study
pVAR53	pEVS107 with P _{lacz} containing 2.3 kb <i>luxQ-A216P</i> -FLAG allele using primers 849 and 1425	This study
pVAR54	pEVS107 with P _{lacz} containing 2.3 kb <i>luxQ-H378A</i> -FLAG allele using primers 849 and 1426	This study
pVAR55	pEVS107 with P _{lacz} containing 400 bp <i>luxU</i> -FLAG allele using primers 1312 and 1422	This study
pVAR56	pEVS107 with P _{lacz} containing 400 bp <i>luxU-H61A</i> -FLAG allele using primers 849 and 1427	This study
pVAR62	pKV363 containing 1.1 kb sequence flanking <i>ainR</i> using primers 1323 and pr_NL35 (Lyell <i>et al.</i> , 2010)	This study

be defective for wrinkled colony formation after 2 days were then cured of their sypG overexpression plasmid and the plasmid was reintroduced. Any mutant that still exhibited a defect was subject to further analysis as described below.

Southern blot analysis

Southern blot analysis was performed as described previously (Visick and Skoufos, 2001; Yip *et al.*, 2005), except that chromosomal DNA was digested with KpnI and probed for the *syp* locus or with PstI and probed for Tn sequences. All Tn mutants exhibited a pattern consistent with only one Tn insertion.

Molecular techniques

All plasmids were constructed using standard molecular biological techniques, with restriction and modification enzymes obtained from New England Biolabs (Beverly, MA, USA) or Fermentas (Glen Burnie, MD, USA). Plasmids utilized in this study are shown in Table 2. To identify the site of insertion of the three non-syp Tn mutants, we cloned the Tn, with flanking DNA, as previously described (Lyell et al., 2008). Unmarked deletions in V. fischeri were generated as previously described (Le Roux et al., 2007; Shibata and Visick, 2012). V. fischeri ES114 was used as the template in PCR amplifications to obtain the DNA containing or flanking the genes of interest using primers listed in Table S1. PCR products were cloned into the pJET1.2 cloning vector (Fermentas, Glen Burnie, MD, USA) or pCR1.2-TOPO (Life Technologies, Grand Island, NY, USA), then subcloned into appropriate final vectors using standard molecular techniques. Site-directed For complementation in single copy from the chromosome, luxQ and luxU alleles were cloned downstream of a P_{lacZ} promoter in the mini-Tn7 delivery vector pEVS107. Insertion at the Tn7 site of the chromosome was performed via tetraparental mating, as previously described (McCann *et al.*, 2003). All plasmids constructed in this study were sequenced at the Genomics Core Facility at the Center for Genetic Medicine at Northwestern University (Chicago, IL, USA) or ACGT (Wheeling, IL, USA) to ensure that the insertion contained the desired sequence or mutation.

mutagenesis was performed using the QuikChange Site-

Directed Mutagenesis Kit (Stratagene, La Jolla, CA, USA).

Luminescence assays

Vibrio fischeri cultures were grown in SWT overnight at 24°C with shaking, then diluted to an optical density at 600 nm (OD_{600}) of ~0.01 in 30 ml of SWTO and incubated at 24°C with vigorous shaking. Samples were taken every 30–60 min. At each time point, bioluminescence (using a Turner Designs TD-20/20 luminometer at the factory settings and a large, clear scintillation vial) and OD₆₀₀ (using a cuvette) were measured for each sample. Maximum luminescence was observed at OD₆₀₀ measurements between 1.5 and 2 for all strains. Specific luminescence was calculated as relative luminescence (the relative light units of 1 ml of culture integrated over a 6 s count) divided by the OD₆₀₀.

Wrinkled colony assays

Vibrio fischeri strains were cultured overnight at 28°C with shaking in LBS containing Tet, then subcultured 1:100 into

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fresh LBS containing Tet and grown under the same conditions for 3 to 4 h the next day. Subcultures were standardized to an OD_{600} of 0.2 and 10 μ l aliquots were spotted onto LBS agar plates containing Tet and incubated at 28°C. Spotted cultures were then monitored from the time the start of wrinkled colony formation became apparent to the point at which wrinkled colony development ceased or the appropriate data set was collected. Each set of strains for a particular experiment was spotted onto the same plate to account for any minor plate-to-plate variations. Each assay was performed at least two to three times, and most were done much more than three times. To ensure that cultures spotted at an OD₆₀₀ of 0.2 resulted in the same number of cells inoculated per spot, we evaluated the correlation between cell number and OD. Specifically, we determined the cell number of the pEAH73containing strains $\triangle sypE$, $\triangle sypE \triangle luxU$, $\triangle sypE \triangle luxQ$ and $\Delta sypE \Delta ainR$ using cultures normalized to an OD₆₀₀ of 0.2, and found no significant difference in the number of colonyforming units obtained from dilutions of the normalized cultures of these strains.

β-Galactosidase assay

Cultures of the reporter strains KV4926 and KV5516 carrying the *sypG* overexpression plasmid pEAH73 were grown in LBS containing Tet at 28°C with shaking. Samples (50 μ l) were collected at 12 and 24 h and 50 μ l of Pierce β -galactosidase Assay Reagent (Pierce Biotechnology, Rockford, IL, USA) were added to each sample. Measurements were taken in a microtitre dish using an ELx800 Absorbance Microplate Reader (BioTek, Winooski, VT, USA) with the appropriate settings. β -Galactosidase activity was determined as previously described (Miller, 1972). *P*-values were calculated using the student's *t*-test.

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Supporting information

Additional supporting information may be found in the online version of this article.

Supporting Information

LuxU Connects Quorum Sensing to Biofilm Formation in Vibrio fischeri

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Fig. S1. Luminescence of ES114, $\Delta sypE$, and $\Delta luxU \Delta sypE$ in culture. Cultures were grown in SWTO and incubated at 24°C with vigorous shaking. Luminescence and OD₆₀₀ were measured over time until maximum luminescence was achieved for the following strains: ES114 (black circles), $\Delta sypE$ (black squares; KV3299), and $\Delta luxU \Delta sypE$ (black triangles; KV4830). (A) Data are plotted as specific luminescence (Sp. lum.; relative luminescence divided by OD₆₀₀) versus OD₆₀₀. The inset depicts a close-up of the luminescence levels of ES114 and $\Delta sypE$ to show the error bars. (B) Data are plotted as luminescence divided by OD at one time point around maximal luminescence (OD ~1.5). (C) Data are plotted as luminescence divided by CFU at one time point around maximal luminescence (OD ~1.5), in which samples were taken and the CFUs were calculated. Samples were taken in triplicate for each strain, with the average and standard deviation (error bars) represented. These data are representative of 2 independent experiments.



Fig. S2. Wrinkled colony formation by $\Delta luxP$. Time-course assays of wrinkled colony formation induced by *sypG* overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 40 h post-spotting for the following strains: $\Delta sypE$ control (pEAH73/KV3299) and $\Delta luxP \Delta sypE$ (pEAH73/KV5347). An * indicates the time at which wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments.





Fig. S3. Wrinkled colony formation of *luxQ* and *luxU* mutants. Time-course assays of wrinkled colony formation induced by *sypG* overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 45.5 h post-spotting for the following strains: $\Delta sypE$ control (pEAH73/KV3299) (A), $\Delta luxQ \Delta sypE$ (pEAH73/KV5394) (B), $\Delta luxU \Delta sypE$ (pEAH73/KV4830) (C), and $\Delta luxQ \Delta luxU \Delta sypE$ (pEAH73/KV6008) (D). An * indicates the time at which wrinkled colony formation was apparent, typically identified by the presence of ridges around the outer edge of the spot. Data are representative of at least three independent experiments.



Fig. S4. Western blot analysis of LuxU-FLAG and LuxU-H61A-FLAG. Cells extracts from $\Delta luxU$ (KV4830) expressing either *luxU-FLAG* from pVAR44 (lane 1) or *luxU-H61A-FLAG* from pVAR49 (lane 2) were probed with an anti-FLAG antibody. LuxU is predicted to be 13 kD. This blot is representative of three independent experiments.



Fig. S5. Luminescence of *ainR* mutants in the presence and absence of C8-HSL. Cultures were grown in SWTO and incubated at 24°C with vigorous shaking. Luminescence and OD_{600} were measured over time until maximum luminescence was achieved for the following strains: $\Delta sypE$ control (squares; KV3299), $\Delta ainR \Delta sypE$ (circles; KV6169), $\Delta luxU \Delta sypE$ (triangles; KV4830), $\Delta ainR \Delta luxU \Delta sypE$ (diamonds; KV6259). (A) Luminescence in the absence of C8-HSL (black symbols). (B) Luminescence in the

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presence of 120 nM C8-HSL (white symbols). The inset for panel A depicts a close-up of the luminescence levels between the $\Delta sypE$ control and $\Delta ainR \Delta sypE$ mutant. The data for $\Delta sypE$ in panel A (black squares) is the same as those in panel B (black squares). Data are plotted as specific luminescence (Sp. Lum.; relative luminescence divided by OD₆₀₀) versus OD₆₀₀ and are representative of at least 3 independent experiments.

Fig. S6



Fig. S6. Wrinkled colony formation by *ainR* mutants. Time-course assay of wrinkled colony formation induced by *sypG* overexpression using plasmid pEAH73. Cultures were spotted onto LBS medium containing Tet and incubated at 28°C. Wrinkled colony formation was monitored up to 49 h post-spotting for the following strains: $\Delta sypE$ control (pEAH73/KV3299), $\Delta ainR \quad \Delta sypE$ (pEAH73/KV6196), $\Delta luxU \quad \Delta sypE$ (pEAH73/KV4830), and $\Delta ainR \quad \Delta luxU \quad \Delta sypE$ (pEAH73/KV6259). Data are representative of at least three independent experiments.

Experimental Procedures

Western blot analysis

To evaluate the levels of protein produced by the *luxU*-H61A mutant, we expressed epitope (FLAG)-tagged versions of this allele and the wild-type control from plasmids (pVAR44 and pVAR49, respectively) in the $\Delta luxU$ mutant (KV4830). Strains were grown overnight at 28°C with shaking in LBS containing Tet. Samples were collected (1 ml) and pelleted, then resuspended in 500µl 2X SDS loading buffer (4% SDS, 10% 2-mercaptoethanol, 0.005% bromophenol blue, 20% glycerol, 0.1M Tris pH 7), boiled for 5 min, and then loaded onto a 15% SDS polyacrylamide gel. After electrophoresis, proteins were transferred to a polyvinylidene fluoride membrane and probed with anti-FLAG antibody (Sigma-Aldrich, St. Louis, MO). Protein bands were visualized using a horseradish peroxidase-conjugated secondary antibody and ECL reagents (SuperSignal West Pico Chemiluminescent Substrate, Pierce Biotechnology, Rockford, IL).

Luminescence assays supplemented with N-octanoyl-homoserine lactone (C8-HSL)

V. fischeri cultures were grown in SWT overnight at 24°C with shaking, then diluted to an optical density at 600 nm (OD₆₀₀) of ~0.01 in 30 ml of SWTO and incubated at 24°C with vigorous shaking. Samples were taken every 30-60 minutes. At each time point, bioluminescence (using a Turner Designs TD-20/20 luminometer at the factory settings and a large, clear scintillation vial) and OD₆₀₀ (using a cuvette) were measured for each sample. Maximum luminescence was observed at OD₆₀₀ measurements between 1.5 and 2 for all strains. Specific luminescence was calculated as relative luminescence (the relative light units of 1 ml of culture integrated over a 6-second count) divided by the OD₆₀₀. For cultures supplemented with exogenous C8-HSL, we used 120 nM purified C8-HSL (Sigma Aldrich, St. Louis, MO) dissolved in a carrier (ethyl acetate and 0.01% (v/v) glacial acetic acid), which was spotted onto the bottom of the flask and allowed to dry before SWTO and cells were added. Addition of the carrier alone had no effect on growth or luminescence. **Table S1.** Primers used in this study¹

- # Primer
- 460 GCCTTGATAGGAGCATTATAATG
- 714 AAACTAGTCATTAACCGATGGCGTCCATATCACCTTGAA
- 782 AAGGGCCCCTCTTAAGTCGATTAATATTCTGCAAACTGCA
- 849 CCTGTGTGAAATTGTTATCCG
- 995 AGACATGCCTGAGGTTTCAT
- 996 CACTAATACTGACTTGTCCTG
- 998 aaaggatccTGCCATTGTTGCAAGCTTATCT
- 1017 aaaggatccTACTGACTCACGATAGTCCC
- 1018 aaaggatccAGATAAGCTTGCAACAATGGC
- 1123 GATAGTAAGAGTGGTGTAAAT
- 1160 taggcggccgcacttagtatgGATGCACTGAATAATTGAGATACC
- 1282 taggcggccgcacttagtatgTAAAACAGGATCAGCTAAAACCAG
- 1283 catactaagtgcggccgcctaGAACACTAGTAATGAATAAAGGT
- 1284 ATGCAACAATTCTACCTGAATC
- 1286 taggcggccgcacttagtatgGTAGGTATGTATGGTTATACCTAC
- 1287 catactaagtgcggccgcctaGACTCGTAATTAGCTTCGCTTAAC
- 1288 GCAATAGGATAGCTATCACCTTC
- 1303 CGTGGCATTTATTAGCTACTGGG
- 1304 GATATTGCTTTAGGTGCTATTGATG
- 1312 GAAAACTACAGTCTTGGAATG
- 1314 CTGGTGTAGAACACTAGTAATG
- 1319 taggcggccgcacttagtatgGGAAGCAGTATCTTCTACCAT
- 1320 catactaagtgcggccgcctaTGGAATGAAAGATAAGGGGAC
- 1321 catactaagtgcggccgcctaTAAATCAATAATCGCGTCTTTTAAC
- 1323 GATCCGTCGCTTAGATGAC
- 1344 CGTAAAGTTGTTGCACCTAAG
- 1345 GCAGGTAAGATGGATCATAGG
- 1346 GGTGAGCTGCACGGGCAC
- 1422 aaaaaggtacettatttatcatcatcattttataatcCTTAAGTAAGAGTTCTCGATAAG¹
- 1425 GATAATAAGATATTAcCATCAAGTGTAAGTTCC
- 1426 TTAGCAAGAATGAGTgcTGAAATAAGAACGCCT
- 1427 GTAAGAGAGATTAGCgcTTGTCTGAAGAGTAGT
- $1437 \quad a a a a a a g g t a c c t t a t t a t c$
- 1494 TACTGACGTATCCGTGTTGC
- 1495 GGCCGATGCTAAAGATTCAG
- $1496 \quad taggcggccgcacttagtatgAATGATTGTGATAAGGCTATAACG$
- 1497 catactaagtgcggccgcctaAAGTATGAAACACAATAAACTTCG
- ¹Non-native sequences are shown in lower case