

Report

Regulating Alternative Lifestyles in Entomopathogenic Bacteria

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Summary

Bacteria belonging to the genera *Photorhabdus* and *Xenorhabdus* participate in a trilateral symbiosis in which they enable their nematode hosts to parasitize insect larvae [1]. The bacteria switch from persisting peacefully in a nematode's digestive tract to a lifestyle in which pathways to produce insecticidal toxins, degrading enzymes to digest the insect for consumption, and antibiotics to ward off bacterial and fungal competitors are activated. This study addresses three questions: (1) What molecular signal triggers antibiotic production in the bacteria? (2) What small molecules are regulated by the signal? And (3), how do the bacteria recognize the signal? Differential metabolomic profiling in *Photorhabdus luminescens* TT01 and *Xenorhabdus nematophila* revealed that L-proline in the insect's hemolymph initiates a metabolic shift. Small molecules known to be crucial for virulence and antibiotic production in addition to previously unknown metabolites are dramatically upregulated by L-proline, linking the recognition of host environment to bacterial metabolic regulation. To identify the L-proline-induced signaling pathway, we deleted the proline transporters *putP* and *proU* in *P. luminescens* TT01. Studies of these strains support a model in which acquisition of L-proline both regulates the metabolic shift and maintains the bacterial proton motive force that ultimately regulates the downstream bacterial pathways affecting virulence and antibiotic production.

Results and Discussion

During their infective juvenile stage, insect-killing (entomopathogenic) nematodes sequester symbiotic bacteria in their digestive tracts and hunt insect larvae in the soil [2, 3]. When a nematode successfully invades a larva's circulatory system, it releases the bacteria. In this new food-rich environment, the bacteria switch from quiescent nematode residents to highly effective insect pathogens while the infective juvenile nematodes develop into adults and begin reproducing. Whereas nematode and bacteria have a specific association—*Photorhabdus* spp. bacteria with *Heterorhabditis* spp. nematodes and *Xenorhabdus* spp. bacteria with *Steinernema* spp. nematodes—a given nematode-bacteria pair can parasitize a wide variety of insect larvae. Identifying the signal (or signals) driving the switch from one bacterial phenotype to another would provide a more complete description of this trilateral symbiosis and of bacterial virulence regulation. In addition, the signal (or signals) could provide access to a potentially

important suite of antibacterial and antifungal small molecules that have so far eluded characterization (Figure 1) [4].

Photorhabdus luminescens TT01, the best studied of these bacterial symbionts, has a sequenced genome containing a wide array of antibiotic synthesizing genes with at least 33 genes in 20 loci similar to those used to make nonribosomal peptides and polyketides—two large families of biologically active small molecules [5]. The genome of *Xenorhabdus nematophila* ATCC19061 has recently been sequenced, and it appears to have a similar size and comparable number of small-molecule biosynthetic genes (www.genoscope.cns.fr/agc/mage/). The majority of these encoded molecules are cryptic, meaning that their existence can be inferred from the genome but not yet confirmed in the laboratory, and analyses of sequenced bacterial genomes indicate that cryptic metabolites greatly outnumber known metabolites—arguably by as much as a factor of ten [6]. Most cryptic metabolite biosyntheses are tightly regulated, and discovering the regulatory key for a given metabolite has proved challenging. The *Photorhabdus/Xenorhabdus* system provides an opportunity to discover the regulatory signal for wholesale production of an entire suite of bioactive small molecules that control a broad swath of microbial competitors.

Insect Hemolymph Regulates Bacterial Secondary Metabolism

Crude small-molecule extracts of *Photorhabdus*- and *Xenorhabdus*-infected insect larvae have enhanced titers of antibacterial, antifungal, and insecticidal activities versus the levels extracted when the bacteria are grown on standard microbiological media [7, 8]. The host factor (or factors) responsible for turning on production of these activities is likely a shared feature of insect larvae, and we used small-molecule production, as judged by chromatographic analysis, to identify it.

Aqueous and organic extracts from a common insect host, *Galleria mellonella* (greater wax moth) larvae, were assessed for their ability to stimulate metabolite production by both *P. luminescens* TT01 and *X. nematophila* ATCC19061. Addition of aqueous insect extracts or hemolymph dramatically induced metabolite production in both bacterial strains as judged by high-pressure liquid chromatography (HPLC) of culture extracts (Figure 2, top panels). Individual metabolites were upregulated by as much as an order of magnitude, and some previously cryptic metabolites became evident in the differential profiles. Larval *G. mellonella* hemolymph was subjected to HPLC-based bioassay-guided fractionation to identify fractions capable of stimulating secondary metabolite production in either strain, and over successive chromatographic steps, identical fractions were shown to induce activity in both. Nuclear magnetic resonance (NMR) spectroscopy and mass spectrometry (MS) of the purified induction factor led to its identification as proline (see Figure S1 available online). Supplementing bacterial cultures with L-proline, but not D-proline, increased secondary metabolite production in both strains (Figure S1). The differential profiles induced by L-proline were very similar to the hemolymph-induced chromatograms in both *Photorhabdus* and *Xenorhabdus* species

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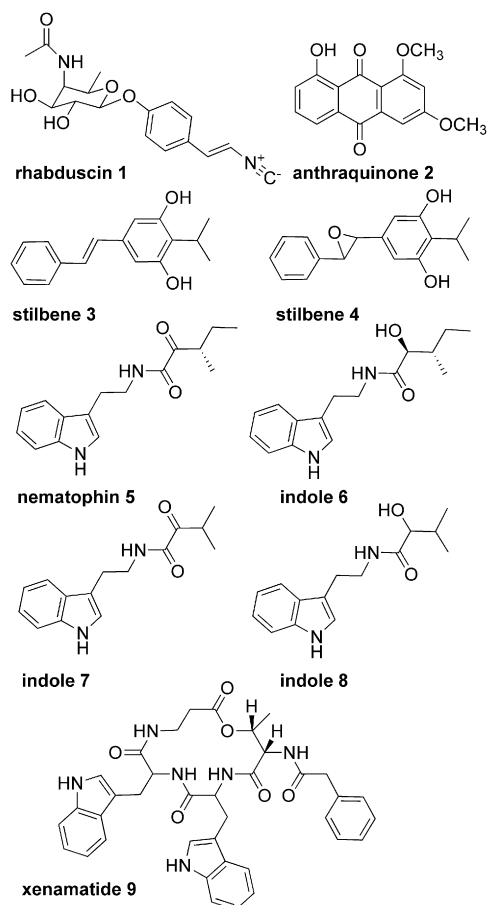


Figure 1. Selected *P. luminescens* and *X. nematophila* Metabolites
Structures 1–4 are *P. luminescens* metabolites; structures 1 and 5–9 are *X. nematophila* metabolites. See also Figure S2 and Tables S2–S5.

(Figure 2, bottom panels). Because a variety of insect larvae contain high levels of L-proline [9], it could be a generic activation signal.

L-Proline Regulates Known Virulence Factors

Structural characterization by NMR and MS established that known virulence factors and antibiotics were regulated in both bacteria by exogenously supplied L-proline. In *P. luminescens*, L-proline substantially upregulated an anthraquinone (structure 2 in Figure 1) and two stilbenes (structures 3 and 4) previously identified in this bacterium (Figure 1 and Figure 2) [10, 11]. In *X. nematophila*, L-proline significantly altered three indole-containing metabolites: nematophin (structure 5), its reduction product (structure 6), and xenamatide (structure 9) [12, 13]. Production of nematophin decreased with a concomitant increase in its reduction product, which illustrates that L-proline (or hemolymph) regulates a metabolic shift rather than a general upregulation.

To evaluate whether proline-mediated metabolite regulation occurs at ecologically relevant L-proline concentrations, we generated dose-response curves for select metabolites in the two model bacterial strains. Marked metabolite regulation was induced by L-proline concentrations greater than 6.4 and 4.8 mM for *P. luminescens* and *X. nematophila*, respectively (Figure 3). Hemolymph, which carries nutrients as insects respire by diffusion, has high free amino acid, carbohydrate,

and inorganic salt concentrations [9]. Complete amino acid analysis conducted on hemolymph samples from four representative host insect larvae revealed individual L-proline concentrations of 73, 3, 31, and 59 mM (Table S1). Although hemolymph L-proline concentrations have a substantial range (~20-fold), these findings show that the low-mM L-proline concentrations needed to regulate metabolite production are physiologically relevant.

The previously described molecules exhibit a range of biological activities, and their regulation by L-proline underscores the connection between recognition of a specific host environment and subsequent bacterial metabolic adjustment. Stilbene 3, for example, possesses antibiotic activity [10], is critical for maintaining symbiosis with nematodes [14], and is a potent inhibitor of phenoloxidase, a component of the insect innate immune system [15]. Nematophin (structure 5) also exhibits antibacterial activity, whereas other *X. nematophila* compounds (structures 6 and 9) have shown little to no activity in various biological assays to date [12, 13], but their regulation by L-proline implies important biological functions. Differential metabolomic profiling with biologically relevant triggers provides a system-wide approach to coupling small-molecule metabolites to the bacteria's physiological state.

L-Proline Upregulates Cryptic Metabolite Production

In order to evaluate the ability of L-proline addition to reveal previously cryptic metabolites, we characterized several compounds from *X. nematophila* that became evident with L-proline addition. After purification of the metabolites, high-resolution MS (Table S2) and a combination of one- and two-dimensional NMR experiments (¹H, COSY, HSQC, HMBC, and ROESY) were performed (Tables S3–S5), identifying two of them as the indole-containing metabolites 7 and 8 (Figure 1). Compound 7 is a new natural product, but it has been prepared by synthesis and shown to be a potent antibiotic against multidrug-resistant *Staphylococcus aureus* [12]. Compound 8 is its previously unreported reduction product. The third selected metabolite from *X. nematophila*, which was upregulated by over an order of magnitude in L-proline-induced cultures, was also characterized. NMR analyses revealed a new isocyanide- and aminoglycosyl-functionalized tyrosine derivative, which we have named rhabduscin (structure 1) (Table S3).

The biosynthetic genes *isnA* and *isnB* were recently discovered from environmental DNA cosmid libraries and found to introduce isocyanide functional groups into their encoded products [16]. The *X. nematophila* ATCC19061 genome contains a candidate biosynthetic gene cluster for rhabduscin (1): *isnA*, *isnB*, and a glycosyltransferase gene (Figure S2). Heterologous expression of the cluster in *Escherichia coli* reconstituted rhabduscin production, confirming assignment of the previously cryptic biosynthetic gene cluster (Figure S2). By using these three genes as bioinformatics probes, we also discovered homologs in the genome sequence for *P. luminescens* TT01. Rhabduscin was, not surprisingly, then identified in *P. luminescens* culture extracts, and its occurrence in both genera suggests an important role in both systems.

A structurally related isocyanide metabolite with an altered sugar moiety was previously isolated from an *Enterobacter* sp., and it was found to possess potent antiphenoloxidase activity [17]. As noted above, phenoloxidases are critical components of the insect innate immune system. Because of the structural similarity of rhabduscin to the known inhibitor,

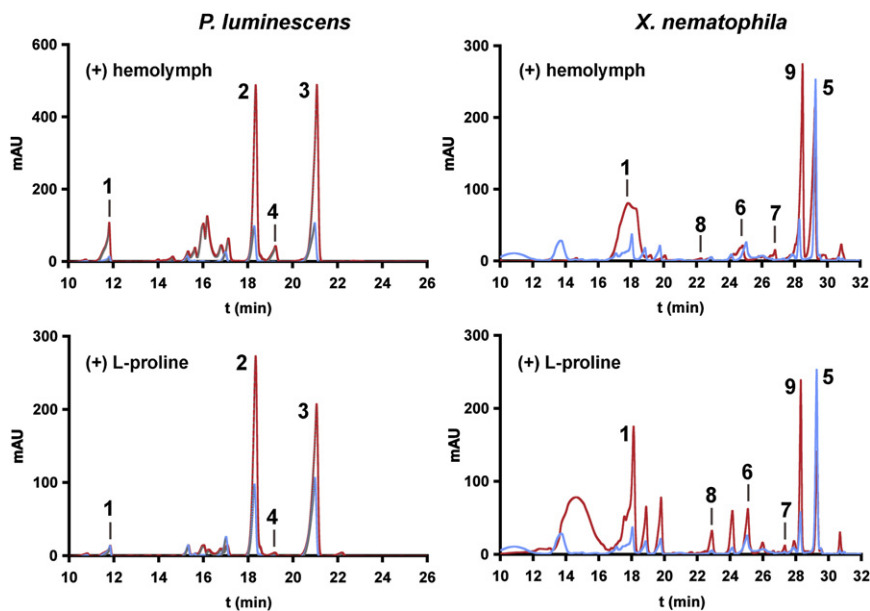


Figure 2. Differential Metabolomic Profiling

Bacterial cultures with or without sterile-filtered insect hemolymph or L-proline were extracted after 72 hr, and organic extracts were assessed by high-pressure liquid chromatography (HPLC) (280 nm). Hemolymph supplementation (red traces, top panels) to *P. luminescens* (left panels) and *X. nematophila* (right panels) alters metabolite production compared to control cultures (blue traces). Proline supplementation (red traces, bottom panels) similarly regulates metabolite production in both bacteria compared to controls (blue traces). Numbers above peaks refer to metabolites in Figure 1. See also Figure S1.

it is possible that it could enhance the bacteria's immune-evading capabilities.

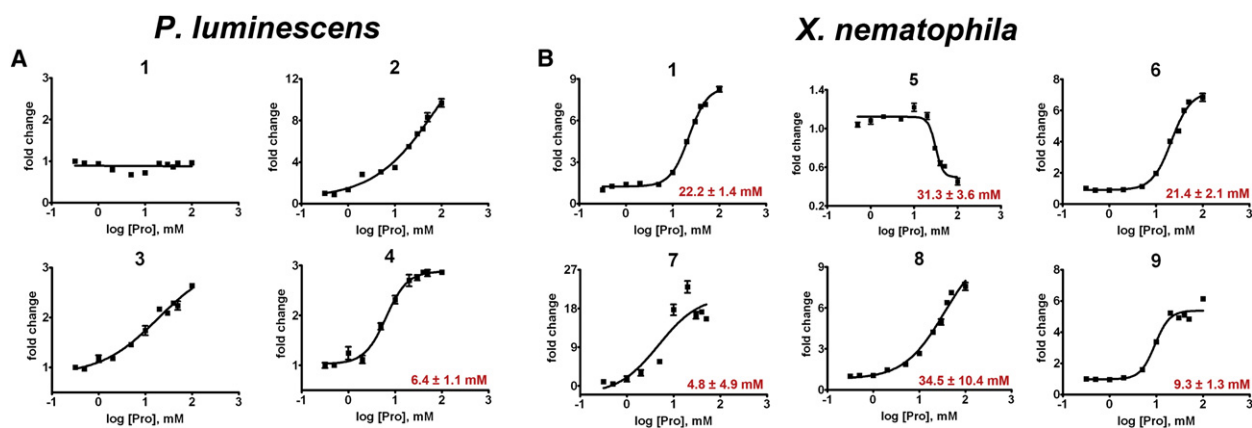
Genetic Analysis of Proline Transport and Response

Because osmolarity is typically much higher in the host than in other bacterial environments, it affects virulence in several bacterial pathogens [18]. To mimic hemolymph's high solute and inorganic salt concentrations [9], we challenged *P. luminescens* and *X. nematophila* cultures with osmotic stressors (sorbitol and NaCl) with and without proline. These osmotic challenges contributed to the regulation of virulence factors, and the introduction of L-proline provided an enhanced effect (Figure S3).

To further probe proline's role and decouple the connection between its transport and metabolite stimulation, we deleted candidate proline transporters in *P. luminescens*. The Gram-negative gammaproteobacteria *E. coli*, *Photorhabdus* spp., and *Xenorhabdus* spp. share many similarities in physiology

and metabolism. *E. coli* is known to acquire proline through three different proline transport systems: PutP, ProP, and ProU [18]. PutP is a high-affinity uptake system for proline metabolism, whereas ProP and ProU typically acquire the organic osmolytes proline and betaine for osmoregulation. Two *P. luminescens* TT01 homologs, *putP* and *proU* (which consists of *proV*, *proX*, and *proW*), were identified and individually excised from the genome by allelic exchange mutagenesis for further interrogation.

Production of stilbenes 3 and 4 was abolished in *P. luminescens* Δ *proU* compared to wild-type cultures, linking L-proline transport or accumulation to production of secondary metabolites (Figure 4). Conversely, anthraquinone 2 was dramatically upregulated in the Δ *proU* mutant, implying altered downstream regulatory control of the two natural product classes (Figure 4). *P. luminescens* Δ *putP* cultures showed similar trends, with virtually no production of stilbenes 3 and 4 and upregulated production of anthraquinone compared to wild-type. To validate the specificity of PutP for L-proline, we introduced *P. luminescens putP* driven by the lactose operon into the *E. coli* proline auxotroph WG389 [19], which fully restored growth under conditions lacking osmotic challenges compared to controls (Figure S4). Furthermore, the observed

Figure 3. Fold Change in Metabolite Production in *P. luminescens* and *X. nematophila* with Increasing Concentrations of Supplementary L-Proline

Numbers above curves refer to compounds in Figure 1, and half maximal effective concentration (EC₅₀) values are shown in red for metabolites with sigmoidal dose-response curves (error = standard deviation). Metabolites were extracted at 72 hr, after the bacteria had reached stationary phase (Figure S1). Rhabduscin (structure 1 in Figure 1) production remained constant in *P. luminescens* cultures upon addition of proline but was stimulated in the presence of elevated osmolarity. See also Figure S3.

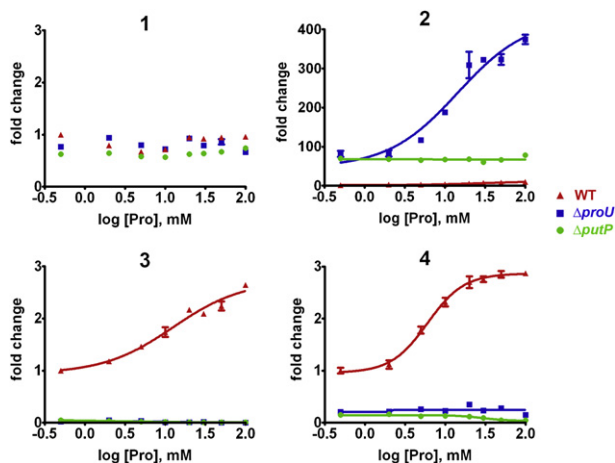


Figure 4. Metabolic Shift of Wild-Type versus $\Delta proU$ and $\Delta putP$ *P. luminescens* with Increasing Proline

Stilbene production (structures 3 and 4 in Figure 1) was abolished in both mutants (blue squares and green circles), but anthraquinone production (structure 2) was upregulated in the mutants compared to wild-type (WT, red triangles) (error bars = standard deviation). Whereas elevation of anthraquinone production by the $\Delta proU$ mutant (blue squares) was even more dramatic with addition of L-proline to the medium, levels were high but unchanged over the proline concentration window for the $\Delta putP$ mutant (green circles), indicating relative insensitivity of the ProU transporter to high levels of proline in a nonosmotically challenged medium. See also Figure S4.

preference for transporting and metabolizing L-proline led us to determine that both *P. luminescens* and *X. nematophila* could grow on a defined MOPS minimal medium with L-proline substituted as the sole carbon source (Figure S4). Taken together, the results argue that osmolarity alone is insufficient to account for the regulation of virulence factors contributing to pathogenesis and support the view that L-proline acquisition and its subsequent metabolism serve as indicators of environmental conditions to direct bacterial gene expression.

Proton Motive Force Manipulation Alters Metabolite Regulation

For reasons not understood, *Photorhabdus* and *Xenorhabdus* species undergo phenotypic variation in stationary phase denoted as primary and secondary forms [1, 20], which differ in proline consumption, respiratory enzyme activity, and proton motive force (PMF) depending on their growth conditions [21] (see Supplemental Discussion). The primary form maintains an ability to produce antibiotics in rich medium, whereas the secondary form does not. Because PMF is implicated in this phenotypic switch, and because it contains osmotic and electrical components, we reasoned that it might play a role in insect host recognition and metabolite stimulation as described above. Indeed, proline oxidation to pyrroline-5-carboxylate (and then to glutamate) by the trifunctional proline dehydrogenase PutA provides an electron source to generate a PMF in *E. coli* [22], and PMF has been implicated in many membrane-dependent roles such as sensory transduction and solute accumulation.

To evaluate the link between membrane energy levels and proline utilization, we first evaluated growth on solid rich medium with increasing osmotic stress (NaCl or sorbitol) with and without proline and/or the PMF diffusion agent carbonyl cyanide-m-chlorophenylhydrazone (CCCP) (Figure S5).

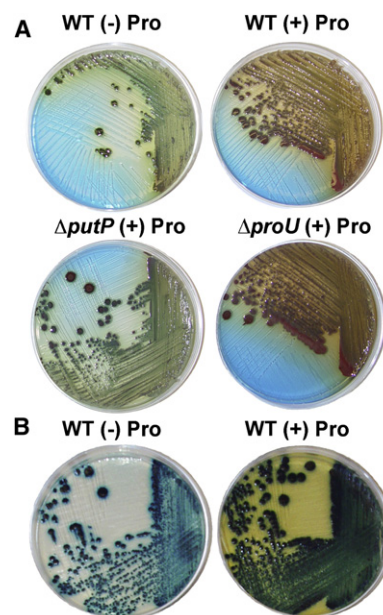


Figure 5. *P. luminescens* and *X. nematophila* L-Proline Growth Effects

P. luminescens (A) and *X. nematophila* (B) were grown on rich solid media (2 g tryptone and 5 g yeast extract per liter) containing the dye bromothymol blue and the redox indicator 2,3,5-triphenyl tetrazolium chloride (TTC) for 3–5 days, with and without 100 mM L-proline. Phenotypic growth effects on various media are illustrated in Figure S5. Staining patterns reveal a reddening of WT colonies and the $\Delta proU$ mutant upon proline supplementation, indicating increased reduction of TTC. A darkening of colonies is also observed with proline, indicating increased dye uptake. Given our differential metabolic observations, it is logical that the *P. luminescens* $\Delta putP$ culture with proline resembled the WT strain grown in the absence of proline. *P. luminescens* $\Delta proU$ was visually similar to WT in these dye uptake studies, in keeping with proline's proposed primary role.

P. luminescens was incapable of forming colonies on rich medium containing 10 μ M CCCP. Proline rescued growth of *P. luminescens* with or without osmotic stress, presumably by providing an electron source to outcompete CCCP-dependent membrane energy diffusion.

Another characteristic difference between primary and secondary forms is their differential staining patterns on agar plates containing the dye bromothymol blue and the redox indicator 2,3,5-triphenyl tetrazolium chloride (TTC) [20]. Both forms reduce TTC, giving a red color phenotype, but only the antibiotic-producing primary form takes up bromothymol blue. Proline addition to both *P. luminescens* and *X. nematophila* cultures showed a darkening of the colonies, indicating increased dye uptake (Figure 5). Additionally, intense reddening of the colonies was observed with proline, showing markedly increased reduction of TTC when proline was present, further supporting proline's role as an electron source.

To more directly test the link between membrane energization and metabolite stimulation, we added the PMF antagonists CCCP and valinomycin to proline-supplemented mid-exponential suspension cultures of *P. luminescens*. The cultures continued to grow until stationary phase before the metabolites were extracted. *P. luminescens* stilbene virulence factors 3 and 4 were substantially upregulated (2.8 ± 0.3 to 9.8 ± 1.0 fold change) in the presence of both CCCP and valinomycin compared to cultures supplemented with proline alone, indicating that PMF control is linked to metabolite production (Figure S3).

The connection between PMF and metabolite production may be applicable to other bacterial systems. For example, subinhibitory levels of a panel of translation-inhibiting antibiotics greatly alter global transcription patterns, and changes to as many as 10% of the gene transcripts by factors as large as 100-fold have been observed [23, 24]. The same families of protein translational inhibitors have also been shown to significantly affect the PMF in *Clostridium sporogenes* [25], supporting the link between antibiotic-regulated metabolite production and PMF regulation. For example, motility, another characteristic of the antibiotic-producing form, is directly controlled by the PMF because the electrical potential component of the PMF affords energy for flagellar motion [26]. Additionally, pathogenic *S. aureus* mutants defective in PutP exhibit significantly reduced mammalian infectivity [18], hinting that PMF control could be a general phenomenon for host recognition in bacterial pathogenesis.

Downstream Regulation

Investigations of *Photorhabdus temperata* (a close relative of *P. luminescens*) and *X. nematophila* have revealed that these species employ dissimilar downstream regulatory mechanisms for controlling general antibiotic activity as judged by zone of inhibition [27, 28]. *P. temperata* employs the LysR-type transcriptional regulator HexA to repress antibiotic activity, as evidenced by the $\Delta hexA$ mutant exhibiting a derepressed, larger zone of inhibition [27]. Deletion of the *X. nematophila hexA* homolog (*IrhA*), however, results in no observed differences in general antibiotic activity [29]. Instead, *X. nematophila* utilizes a leucine-responsive regulatory protein (Lrp) homolog [28], which governs many pathogen-associated activities, including positively influencing flagellar motion and antibiotic production (Δlrp exhibits no zone of inhibition) [3]. In *E. coli*, Lrp is known as a feast-or-famine transcriptional regulator, and nutrient sources such as amino acid ligands bind at the dimer interface, altering its activity [30]. Similarly, small-molecule coinducers modulate LysR-type transcriptional regulatory activity [31]. Given the drastic proline response observed, proline could either directly act as a transcriptional protein ligand or regulate the production of the cognate ligand. Although it is not yet clear how the initial signal translates to the disparate downstream regulatory machinery in *Photorhabdus* and *Xenorhabdus*, both genera evolved to recognize proline as a nutrient signal to induce phenotypic change, linking host physiology to bacterial pathogenesis.

Conclusions

Systematic examination of the insect host's small-molecule repertoire with differential metabolomic profiling provides a straightforward approach to bacterial metabolite discovery with an ecological rationale. The hemolymph nutrient availability and high osmolarity common in the wide spectrum of insect prey restricted bacterial evolution to only a handful of carbon/nitrogen sources—primarily free amino acids (concentrations of the 20 proteinogenic amino acids varied among insect species from undetectable to 100 mM; Table S1). The chemical and microbiological analyses reported here support a model in which both *Photorhabdus* and *Xenorhabdus* species evolved to utilize L-proline in the hemolymph as a preferred nutrient source, which can also protect against osmotic stress, to regulate the production of metabolites linked to pathogenesis through its ability to enhance the PMF. Because proline also functions as a preferred energy source for insects [32], its consumption by the bacteria

depletes the insect's resources in the conflict. These results define a link between metabolic status (PMF) and virulence factor regulation in *Photorhabdus* and *Xenorhabdus* species and provide an approach to the wholesale discovery of cryptic metabolites with biomedical potential involved in this trilateral symbiosis.

Experimental Procedures

Metabolite Stimulation

P. luminescens and *X. nematophila* were grown on Luria-Bertani (LB) agar for 2 days at 30°C. Single colonies were selected and grown for 2 additional days in LB broth. For metabolite stimulation assays, 50 μ l of the saturated suspension cultures was used to inoculate 5 ml of a rich tryptone-yeast extract-based medium (2 g tryptone, 5 g yeast extract, and 10 g NaCl per liter) with and without indicated supplements. Triplicate cultures were extracted at 72 hr with ethyl acetate, dried, and resuspended in methanol for HPLC analysis.

Induction Factor Identification

Hemolymph was collected from 500 *G. mellonella* larvae. Fresh hemolymph stored on ice was diluted with 100 mM potassium phosphate buffer (pH 7.4) and filtered through a 3000-molecular-weight cutoff membrane (Amicon Ultra, Millipore); the activity resided in the flowthrough. The flowthrough was concentrated and separated via reversed-phase HPLC. The activity eluted within the first isocratic step (2:98, acetonitrile:water). The polar fraction was iteratively collected, pooled, and concentrated for further purification with a Hypercarb column (Thermo Scientific), and again the compound was eluted in the initial isocratic step (100% water). NMR experiments in D₂O showed chemical shifts (¹H) matching that of a proline standard (Figure S1). LC-MS analysis and two-dimensional (gCOSY, gHSQC, and gHMBC) NMR correlations were also consistent with proline.

NMR Analysis

NMR experiments (¹H, gCOSY, gHSQC, gHMBC, and ROESY) were performed in deuterated methanol, water, chloroform, or dimethyl sulfoxide with a symmetrical NMR microtube susceptibility matched with the appropriate solvent (Shigemi, Inc.) on a Varian INOVA 600 MHz NMR spectrometer.

Allelic Exchange Mutagenesis

Deletion constructs were generated in pDS132 [33] and introduced by conjugation from the DAP auxotroph *E. coli* WM6026 lambda pir [34] via standard molecular and microbiological procedures. The *P. luminescens* $\Delta putP$ and $\Delta proU$ deletions were validated by polymerase chain reaction and sequence verified.

Phenotypic Growth Effects

P. luminescens and *X. nematophila* single colonies were selected and grown for 2 days in LB at 30°C. Thirty microliter aliquots of the suspensions were streaked onto agar plates (salt experiments: 2 g tryptone, 5 g yeast extract, and 0, 10, or 35 g NaCl per liter of agar; sorbitol experiment: 2 g tryptone and 5 g yeast extract per liter of agar, with 750 mM sorbitol) containing bromothymol blue and TTC and grown for 3–5 days [20]. These experiments were conducted with and without 100 mM L-proline and/or 10 μ M CCCP.

PMF and Metabolite Stimulation

Bacteria were grown as described in the metabolite stimulation experiments. Proline-supplemented (50 mM) cultures were grown to mid-log phase and then antagonized by 10 μ M CCCP, 10 μ M valinomycin and 150 mM KCl, or 20 μ M valinomycin and 150 mM KCl. The cultures continued to grow for a total time of 72 hr before extraction.

Supplemental Information

Supplemental Information includes Supplemental Discussion, five figures, five tables, and Supplemental Experimental Procedures and can be found with this article online at doi:10.1016/j.cub.2009.10.059.

Acknowledgments

This work was supported by National Institutes of Health grant CA24487 to J.C. J.M.C. is a Damon Runyon Fellow supported by the Damon Runyon

Cancer Research Foundation (DRG-2002-09). We are grateful to R. Kolter and C.T. Walsh (Harvard Medical School) for helpful suggestions and critical reading of a preliminary manuscript. We thank J. Wood (University of Guelph) and W. Metcalf (University of Illinois) for providing *E. coli* strains WG389 and WM6026, respectively.

Received: September 7, 2009

Revised: October 16, 2009

Accepted: October 20, 2009

Published online: December 17, 2009

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