COMPARISONS OF BACTERIA FROM THE GENUS PROVIDENCIA
ISOLATED FROM WILD DROSOPHILA MELANOGASTER

A Dissertation
Presented to the Faculty of the Graduate School
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Doctor of Philosophy

by
Madeline Rose Galac
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Comparisons of Bacteria from the Genus Providencia Isolated from Drosophila melanogaster

Madeline Rose Galac, Ph. D.
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Multiple strains representing four species of bacteria belonging to the genus Providencia have been isolated from wild caught Drosophila melanogaster: Providencia sneebia, Providencia burhodogranariea strain B, Providencia burhodogranariea strain D, Providencia rettgeri, and Providencia alcalifaciens. Using this laboratory-friendly and natural host, D. melanogaster, I determined how these bacteria differ in their ability to cause host mortality, replicate within the fly and trigger the fly’s immune response as measured by transcription of antimicrobial peptides. Although each bacterium has a unique profile of these phenotypes, in general the greater amount of mortality a given bacterium causes, the more proliferative it is and the greater antimicrobial peptide transcription they evoke in the host. An exception to this was P. sneebia which killed about 90% of infected flies and reached greater numbers within the fly than any of the other bacteria, but induced less antimicrobial peptide transcription than the less virulent Providencia. Coinfections in D. melanogaster with P. sneebia and P. rettgeri, which induces greater antimicrobial peptide expression and is less virulent than P. sneebia, allowed me to conclude that P. sneebia is actively avoiding recognition by the immune response. I sequenced and annotated draft genomes of these four species then compared them to each other. I found that about 50% of each genome belongs to
the *Providencia* core genome and about 15% of each genome consists of genes unique to that species. Four *Providencia* genomes of isolates originating from the human gut have also been sequenced, which include additional isolates of *P. rettgeri* and *P. alcalifaciens* as well as isolates of the species *Providencia stuartii* and *Providencia rustigianii*. When I compared orthologs of all eight sequenced *Providencia* genomes, I found that the percent of each genome that consists of the core genome and the unique genes stays largely the same. I found that each bacterium has type 3 secretion system, a known virulence factor. This means that presence of a type 3 secretion system does not correlate to those found to be virulent toward *D. melanogaster*. This work establishes *D. melanogaster-Providencia* as a model system for the study of host-pathogen interactions.
BIOGRAPHICAL SKETCH

Madeline R. Galac was born to Alan and Mary Galac in 1983 in Buffalo, NY. Her love of science was fostered early as a result of her mother’s interest in science. She remembers very fondly the large tank of *Xenopus* in the house for most of her young life. In junior high and high school, biology classes taught by Mr. Heinomen and Mr. Leavell helped Madeline to specify her scientific interest in biology. Madeline tried out different biological jobs by interning at both the Buffalo Zoological Society in Buffalo, NY and the Niagara Falls Aquarium in Niagara Falls, NY. She found that she loved the hands on interactions with the animals and was saddened by the difficulty of getting and then maintaining such a job.

Madeline continued her scientific education by pursuing a bachelor’s degree in biology at the State University of New York at Stony Brook in Stony Brook, NY. In 2002, the summer of after her freshmen year, she decided to do another internship to see if she liked primary scientific research. Dr. Joel Huberman at the Roswell Park Cancer Institute in Buffalo, NY agreed to let her help his grad student Sanjay Kumar for the summer. It was this summer that she not only determined that she enjoyed research but also learned about some of the quirks that go along with it, such as trying to explain to others why the very specific thing you are studying is important and the awkwardness of graduate students. Once back in Stony Brook, Madeline sought out a lab to do research in while completing her degree. She joined Dr. Jen-Chih Hsieh’s lab studying the interactions of proteins involved in Wnt signaling. In the summer of 2003, Madeline went to Woods Hole, MA to do research at The Woods Hole Oceanographic Institute with Dr. Sonya Dyhrman which included her only field work to date, an 11 day research cruise collecting toxic dinoflagellates in the Gulf of Maine. In her final
semester at SUNY Stony Brook, Madeline took a class entitled “Darwinian Medicine” which had a strong influence on her research interests as a graduate student. This class focused on host-pathogen interactions and gave the most in depth explanation of evolution that she had been exposed to during her education. The paper she wrote for that class about the Bubonic Plague pathogen, *Yersinia pestis*, cultivated her interest in the evolution of pathogenic bacteria.

In 2005 Madeline came to Cornell University in Ithaca, NY to pursue a Ph.D. in the field of Genetics and Development. She joined Dr. Brian Lazzaro’s lab to look at host-pathogen interactions using the well-studied model host *Drosophila melanogaster* infected with different strains of natural pathogens of the genus *Providencia*. During her research, she was tricked into doing whole genome sequencing of these *Providencia*, which lead to Madeline becoming a bioinformatician.

Madeline hopes to continue researching the way that bacteria interact with other organisms in their environment using both bioinformatics and wet lab approaches.
ACKNOWLEDGMENTS

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I am extremely sad to leave my lab behind, as I fear I will never find a group of people that are as much fun to work with as they were. I think we always did a good job of balancing science with fun, whether it was science related fun or not. They were all a great group of people that helped and supported me in my work when I needed it. It is amazing to have a group of people that you know will always lend an extra hand when you need it for an experiment. I will always think fondly on these days every Halloween. I hope that the tradition of dressing up that started while I was in the lab continues for many years to come.

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groups which include all of them: the G&D and MCB classes that started with me, the
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appreciated those parties not only for the dancing but for also reminding me that there are
things beyond this degree.

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a beer” while in grad school.
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CHAPTER 1

INTRODUCTION
1. Introduction

When closely related bacteria vary in a complicated phenotype, comparing them across a detailed dissection of the phenotype can give insight into how that phenotype is accomplished. For example, virulence is a complex and broad phenotype which can often be broken down into a more detailed aspects such as bacterial load during infection, bacterial location within the host, or host cell shape upon bacterial contact. By comparing the way closely related bacteria differ in the details of virulence, one can reveal how the more lethal ones are acting on the host. Differences in complex phenotypes among closely related bacteria are often the result of horizontal gene transfer as it can provide the bacteria with the multiple genes it may need for a complicated phenotype all at once [1]. Horizontal gene transfer notwithstanding, closely related bacteria have more similar gene content aiding the identification of those genes which do differ. Hence, when genomic information is available for such bacteria, this phenotypic information can lead to hypotheses for the genomic differences that are the cause.

In this thesis, both pathological and genomic comparisons are presented for five distinct strains of bacteria in the genus *Providencia*: *Providencia sneebia*, *Providencia rettgeri*, *Providencia alcalifaciens*, *Providencia burhodogranariea* strain B, and *Providencia burhodogranariea* strain D. These strains were all isolated from infected wild caught *Drosophila melanogaster*. The second chapter examines the infection biology of these bacteria in their laboratory-friendly natural host. In the third chapter, draft genomes of the four different species were sequenced and compared to examine both their general genomic differences and identify putative virulence factors.
2. *Drosophila melanogaster* as a model host

*D. melanogaster* has been used as a model host to study varied aspects of infection of many different bacteria. The innate immune response of *D. melanogaster*, particularly the humoral response and the protein pathways involved, have been well characterized, mostly using generic immune elicitors such as non-pathogenic bacteria or bacterially derived molecules that stimulate the innate immune response [2]. Recognition of these immune elicitors in the fly causes a signaling cascade that results in the transcription of numerous genes [2]. Among the genes that are transcribed as part of the humoral immune response are antimicrobial peptides, which directly kill the bacteria [2]. The triggering of the humoral immune response signaling pathways and the resulting response is the idealized fly immune response to the presence of bacteria as it should allow the fly to kill or remove the infecting bacteria and survive the infection. There are many different ways by which a bacteria can thwart this idealized fly response, such as suppression of the immune signaling, avoidance of recognition by the immune response, or being located somewhere in the host where the molecules like antimicrobial peptides cannot reach. Such tactics by the bacteria can allow it to replicate during infection and possibly kill the insect host.

The *D. melanogaster* signaling cascades of the humoral innate immune response have been found to be conserved in many other organisms, including humans [2]. This conservation of proteins and the laboratory tools available for *D. melanogaster* often leads to the fly being chosen as a model host to study the interactions of the innate immune response and human bacterial pathogens (i.e. [3–5]). While there are many things to be learned from the fly-bacteria interaction for human pathogens, the infections
may not accurately reflect the biology that occurs when the pathogen is in its natural host. Despite the similarity in innate immunity protein pathways, there are a lot of differences between flies and humans, such as vastly different body temperatures, circulatory systems, and internal organs, which may lead to different infection dynamics and, therefore, not accurately reflect the natural host-pathogen interaction. These types of problems with model host-pathogen pairs have been identified in other systems. The human bacterial pathogen *Burkholderia* has been studied in many model hosts including nematodes, wax moths, and mice [6]. It has been found that *Burkholderia* does not require the same virulence factors during infection across these different model hosts [6]. Similar levels of specificity in the genes involved in a host-pathogen interaction have been seen with mosquitoes and *Plasmodium*. Due to the specificity of which animals mosquitoes take their blood meal from, a particular mosquito species will only encounter a subset of *Plasmodium* species in nature. When mosquitoes are infected in the lab with either a *Plasmodium* species they encounter in nature or one which they do not, it has been found that different host factors are required for the establishment of infection by each *Plasmodium* species [7]. This has implications for studies that use a non-naturally occurring mosquito-*Plasmodium* pair as a model for what happens in nature. For these reasons, one should be cautious when using unnatural or model host-pathogen pairs as the information may not accurately reflect what the pathogen does during infection with its natural host or how the natural host responds to infection.
3. Natural D. melanogaster Pathogens

While the focus has mostly been on D. melanogaster as a model host, there has been some work on those bacteria that naturally infect them. These bacteria hold the potential to have co-evolved with D. melanogaster or to be generalist which infect other hosts as well. The bacteria Wolbachia spp. and Spiroplasma spp. have evolved to be heritable endosymbiont pathogens in D. melanogaster [8,9]. These bacteria live within the fly and are vertically transmitted. Therefore Wolbachia spp. and Spiroplasma spp. will not have the same infection dynamics as pathogenic infections a fly can have during its lifetime, which it either survives or succumbs to. Some such infectious bacterial pathogens of D. melanogaster have been identified and studied [10–14].

The isolates of Providencia studied here were isolated from the hemolymph of wild caught D. melanogaster [15,16]. The hemolymph of a healthy fly should be sterile so the presence of bacteria indicates that the bacteria are causing an infection in the fly. These bacteria were grouped into 5 distinct strains which represented four species, based on their 16S and housekeeping gene sequences and other identifiers [16]. Since these bacteria are closely related, their differences in virulence towards D. melanogaster provides a good system for comparisons to identify virulence mechanisms. Additionally, because these bacteria are natural pathogens of D. melanogaster, there is no concern over artifactual phenotypes during infection from unnatural host-pathogen pairs.

The infections in this study are established through pin-prick infections with a small needle coated in bacteria. This allows us to circumvent how the flies get infected to then focus on what happens once the bacteria are within the fly. Nevertheless, I think this method may actually mimic one way flies can get infections in nature. I hypothesize that
these bacteria may be making the transition from present in the environment to infectious agent through cuticle breaches caused by ectoparasitic mites [17]. Wild caught flies often have healed wounds as indicated by melanization independent of natural pigmentation patterns and frequently carry ectoparasitic mites that could be the cause of some of these wounds (P. Juneja, personal communication). Mite wounds in honey bees have been shown to be secondarily colonized by environmental bacteria [18]. Mites might not just be the source of cuticle breaches exposing the fly to infections, but could also be carrying bacteria that could lead to infection. One of the bacteria studied here, \( P. \) burhodogranariea, has additionally been isolated from a mite removed from a wild caught \( D. \) melanogaster (P. Juneja, personal communication).

4. Bacteria of the genus \textit{Providencia}

Bacteria of the genus \textit{Providencia} are a ubiquitous Gram-negative bacteria isolated from numerous locations and environments around the world. They belong to the tribe \textit{Proteeae} with the closely related genera \textit{Proteus} and \textit{Morganella} [19]. \textit{Providencia} isolated from soil samples from India and Los Alamos National Laboratory have been shown to breakdown toxic soil contamination [20,21]. \textit{Providencia} have been found to cause urinary tract infections and traveler’s diarrhea but are also known to be part of the normal human gut flora [22–24]. The ability of \textit{Providencia} isolated from such infections to invade human cells has been found to vary across strains and species [22,23,25,26]. They have also caused more serious bacterial infections in humans such as meningitis [27]. \textit{Providencia}’s role in human infections has led to many studies on the variation in the structure of their cell surface proteins and the genes which encode for them [28].
In addition to their associations with human infections, *Providencia* have been isolated from varied animal related locations such as penguin feces in German zoos, sea turtles in the Mediterranean, shark mouths in Brazil, entomopathogenic nematodes all over the world, and snakes from Vietnam [29–33]. This extends to also being found in association with different insects such as blowflies, stable flies, Mexican fruit flies, and house flies [34–37]. *Providencia* have been isolated on external surfaces or in the gut of *Drosophila melanogaster* both in the wild and in the laboratory [38,39]. *Providencia* have been found to be part of the normal gut flora of numerous species of *Drosophila* both from the lab and from the wild, though they make up a larger part of the gut microbiome in wild flies [39]. Although these associations say that these bacteria are present in and around these insects, it is unclear if they are capable of causing diseases. Two exceptions to this are an isolate of *Providencia rettgeri* isolated from entomopathogenic nematodes was found to be pathogenic to waxmoth larvae [32] and an unidentified species of *Providencia* which was found to be pathogenic to *Drosophila paulistorum* [40]. All this points to *Providencia* generally being around in the environment, which suggests that they were isolated as generalists and opportunistic pathogens.

5. Chapter 2: Comparative Pathology of *Providencia* in *D. melanogaster*

The second chapter of this thesis is about *D. melanogaster* infections with these five different strains of *Providencia*. I measured how much fly mortality results from bacterial infection, how much the bacteria replicate within the fly during infection, and how strong of an immune response the flies mount to the infection as determined by
antimicrobial peptide gene expression. Each strain has a unique profile of these phenotypes indicating that there are differences in virulence mechanisms ranging from being passively cleared to rapidly replicating and killing the fly. I identified a general trend that those bacteria that caused the most mortality were also able to replicate the most and lead to more antimicrobial peptide transcription. Identifying this trend allowed us to see that *P. sneebia* was an outlier as it causes about 90% fly mortality and replicates to significantly higher numbers during infection but induces a lower amount of antimicrobial peptide mRNA compared to the others examined. Through co-infections of *P. sneebia* and *P. rettgeri*, which induces greater antimicrobial peptide expression and is less virulent than *P. sneebia*, I concluded that *P. sneebia* is not causing a strong fly immune response because it is actively avoiding recognition by the immune response and is protected from antimicrobial peptides expressed due to the presence of *P. rettgeri*. I tested two possible mechanisms *P. sneebia* could be using to avoid recognition, biofilm formation and intracellular replication, but did not find evidence for either. This work establishes these *Providencia* strains as highly varied in their pathology to *D. melanogaster*.

6. Chapter 3: Genomic Comparisons of *Providencia*

To find genes that could potentially lead to the phenotypic differences seen in these strains as well as look at other information in the genomes, I sequenced the genomes of *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranaririae* strain B. While a single bacterial genome provides information about what genes are present in that bacterium, comparisons to other genomes can add more detail such as uniqueness or
conservation of a gene [41,42]. Comparing bacteria that are closely related but vary in specific phenotypes, such as virulence, can help narrow down what genetic differences are the cause of the phenotypic differences [41,42]. Next generation sequencing has aided whole genome sequencing and subsequent genomic comparisons of many organisms, particularly bacteria, by lowering the cost and increasing the ease of obtaining genomes. This increase in genomes and comparative studies allows for organisms at different phylogenetic distances to be compared which has broadened the types of questions that can be addressed [41].

I sequenced, assembled and annotated draft genomes of *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* strain B then compared their genomic content. I found these bacteria to have a large core genome and about 15% of the total genes of each genome are unique to that species. There have been additional *Providencia* genomes sequenced as part of the Human Microbiome Project [43], which include additional *Providencia* genomes sequenced as part of the Human Microbiome Project [43], which include additional isolates of *P. alcalifaciens* and *P. rettgeri* as well as the species *Providencia rustiganni* and *Providencia stuartii*. When all eight of the *Providencia* genomes are analyzed, the percentage of each genome that comprises of the core genome or unique genes stays relatively consistent. This suggests that the large amount of similarity will extend to the genomes of additional members of the genus. When looking for well known virulence factors I found that each genome encodes at least one type 3 secretion systems though all 4 *D. melanogaster* isolated strains did not have homologous type 3 secretion system [44]. The universal presence of a T3SS in all of the *Providencia* genomes indicates that bacterial virulence towards *D. melanogaster* or other hosts cannot be predicted solely based on the existence of this virulence factor for these strains.
7. Appendix: Attempted Verification of Putative Virulence Factors

The genomic comparisons revealed a putative virulence factor in the type 3 secretion systems (T3SS). These genes produce proteins which form a large needle-like apparatus which inject proteins, called effector proteins, into host cells [44]. I sought to create bacterial strains that are missing an essential T3SS gene to see how the virulence phenotypes change in both *P. sneebia* and *P. rettgeri* during infection in *D. melanogaster*. By creating the knockout in both species I would be able to compare the results to see if and how the T3SS is involved in virulence and then make hypotheses as to the function of the effector proteins during infection. I successfully made the constructs for genomic integration of an in-frame deletion of an essential T3SS protein for both species and integrated the whole plasmid into *P. rettgeri*’s genome but was unable to get the plasmid into the genome of *P. sneebia*. Without genomic integration of the plasmid, I was unable to excise the plasmid to create the in-frame deletion in *P. sneebia*. I was therefore unable to resolve the amount the type 3 secretion systems contribute to the virulence of *P. sneebia* or *P. rettgeri* towards *D. melanogaster*.

8. Summary

The *D. melanogaster-Providencia* system used here is a powerful and informative system to examine host-pathogen interactions and infection biology in naturally occurring pairs. The initial pathology of infections with *Providencia* isolated from wild infected *D. melanogaster* has been determined and the genomes of the four bacterial species have been sequenced. We found these isolates of *Providencia* to vary in their infection phenotypes in the host from which they were isolated but their gene content is highly
similar. Their genomes are also very similar to isolates of *Providencia* found in human feces suggesting that equally high amounts of homogeneity among gene content will be found in additional members of the genus. This, with the fact that *Providencia* have been isolated from a wild variety of locations and environments, suggests that they are generalist and opportunistic pathogens of varied hosts.

References


CHAPTER 2

COMPARTIVE PATHOLOGY OF BACTERIA
IN THE GENUS PROVIDENCIA TO A
NATURAL HOST, DROSOPHILA MELANOFSATER

This work has been published and reprinted with permission: Galac MR, Lazzaro BP (2011) Comparative pathology of bacteria in the genus Providencia to a natural host, Drosophila melanogaster. Microbes and Infection 13: 673–683.
Abstract

Bacteria in the genus *Providencia* are pathogens of many organisms, including humans and insects. We and colleagues have isolated five different strains belonging to four distinct *Providencia* species as natural infections of *Drosophila melanogaster* captured in the wild. We found that these isolates vary considerably in pathology to infected *D. melanogaster*, differing in the level of mortality they cause, their ability to replicate within the host and the level that the fly’s immune response is elicited. One interesting bacterium was *Providencia sneebia*, which causes nearly complete mortality and reaches large numbers in the fly but does not elicit a comparably strong immune response. Through coinfection experiments, we determined that *P. sneebia* avoids recognition by the immune system. We tested for biofilm formation and replication within *D. melanogaster* cells as possible mechanisms for *P. sneebia* escape from host immunity, but did not find evidence for either. *D. melanogaster* and *Providencia* provide a powerful system for studying general host-pathogen interactions, and for understanding how the well-studied immune model host *D. melanogaster* interacts with its natural bacterial pathogens.
1. Introduction

Closely related bacterial pathogens may utilize a wide range of mechanisms to infect hosts, in part because virulence mechanisms are genetically labile and are often horizontally transferred between reasonably distantly related microbes [1]. Understanding differences in pathology between closely related bacteria highlights recent shifts in virulence, and can ultimately lead to the identification of the underlying genetic basis. Several strains and species of the γ-Proteobacterial genus Providencia have recently been isolated from field infections of wild caught Drosophila melanogaster ([2,3], P. Juneja and S. M. Short unpublished), and in the present work we contrast the pathological interactions of these bacterial species within their Drosophila host. D. melanogaster is a well established model host for studying innate immunity [4] and the pathology of virulent bacteria (e.g. [5–8]), but few natural pathogens of D. melanogaster have been identified or extensively studied (but see [9–11]). We believe that Drosophila and Providencia comprise a powerful natural system for the study of variation in virulence and host-pathogen interactions. Because many microbial virulence strategies focus on conserved aspects of eukaryotic physiology and cell biology, inferences from this system can apply across broad host ranges, including from insects to humans.

Bacteria of the genus Providencia are Gram-negative opportunistic pathogens that have been isolated from a wide variety of environments and organisms ranging from humans to insects to sea turtles and shark mouths [12–15]. Providencia rettgeri, Providencia alcalifaciens, and Providencia stuartii have all been isolated from human stool samples both as part of the natural human gut flora and as the cause of gastric upset such as traveler’s diarrhea [16–18]. Some strains of P. alcalifaciens, but no strains of P. rettgeri or P. stuartii, have been found to be intracellularly invasive in human cell lines [16,17,19,20]. Providencia also cause urinary tract
and other nosocomial infections in humans [12,13]. Numerous studies surveying bacteria associated with insects such as blowflies, stable flies and Mexican fruit flies have isolated *Providencia* species either from the whole insect or specifically from the gut (e.g. [21–23]), although it is unclear whether these and other associations have meant the bacteria were acting as pathogens or were simply present in the insects’ environment. *Providencia* have been recurrently found in association with *D. melanogaster*, including in a survey for bacterial associates in a natural population [24], in the hemolymph of laboratory cultures of domino mutant larvae that are void of hemocytes and generally sick with bacterial infections [25], and as natural infections in wild caught *D. melanogaster* ([2,3], P. Juneja and S. M. Short unpublished).

The *D. melanogaster* innate immune system has been well described, primarily from experiments measuring the response to injection of avirulent bacteria or generic immune elicitors (reviewed in [4]). *D. melanogaster* is also an excellent model for studying the pathology of virulent bacteria, since many virulence mechanisms are effective across a broad range of hosts. As a result, Drosophila has successfully been used as an experimental host to model clinical pathogenesis in humans and animals, insect vectoring of human disease and microbiological control of insect agricultural and medical pests. For example, Drosophila has been used to study opportunistic human infectors such as *Serratia marcescens* and the pathologies of *Pseudomonas aeruginosa* communities found in cystic fibrosis patients [7,26]. *D. melanogaster* has also stood in as a model host for other arthropods such as ticks and mosquitoes that bear *Francisella tularensis* infections, ticks that host *Ehrlichia chaffeensis*, and caterpillars infected with *Photorhabdus luminescens* vectored by entomopathogenic nematodes [27–29].

Despite previous studies of bacterial pathogens of other animals using *D. melanogaster* as a model host, very little is known about the bacteria that infect *D. melanogaster* itself in its...
natural habitat. In some of the few efforts to identify bacterial pathogens of wild Drosophila, four different species belonging to the genus *Providencia* were recovered along with isolates of other bacteria from the hemolymph of wild caught *D. melanogaster* ([2,3], P. Juneja and S. M. Short unpublished). Since the hemolymph of a healthy fly should be sterile, the presence of bacteria can be considered to constitute an infection. Two of the recovered *Providencia* species are the previously described *P. rettgeri* and *P. alcalifaciens* [12]. The other two species were identified as novel species named *Providencia sneebia* and *Providencia burhodogranariea*, the latter of which has two distinct strains designated B and D [3].

In this paper, we determine the pathology of *Providencia* species and strains in *D. melanogaster*, where pathology is defined as the proportion of mortality caused by the bacteria, the bacterial ability to proliferate within the fly, and the levels of host immunity induced by infection as measured by the expression of antimicrobial peptide (AMP) genes. We find *Providencia* to be highly variable in all three phenotypes. The ability of the bacteria to proliferate within the fly, the amount of AMP expression, and the level of mortality the bacterial cause are often all positively correlated, with the most deadly bacteria reaching the highest amount within the fly and inducing the highest levels of AMP expression. A notable and interesting variation to this pattern is *P. sneebia*, which kills about 90% of infected flies and reaches very large numbers in these flies but induces less AMP expression than other *Providencia* species, even those that cause significantly lower mortality and do not proliferate as effectively within flies. Through coinfections with *P. sneebia* and *P. rettgeri*, we concluded that *P. sneebia* is able to actively avoid recognition by the fly’s immune system and is resistant to ectopic immune induction. Two possible hypotheses to explain these observations are that *P. sneebia* invades and replicates within insect cells or forms a biofilm during infection, but we do not find evidence supporting
either hypotheses in vitro, suggesting that P. sneebia virulence mechanisms are more complicated. The diversity of virulence profiles we observe among these Providencia isolates indicates they will be a rich substrate for future study of Providencia infection dynamics in a natural and experimentally tractable host.

2. Methods and materials

2.1. Fly stocks and bacteria strains

_D. melanogaster_ fly stocks that were used were either wild type OregonR (OreR), OR;imd^{10191};OR [30], Toll 1-RxA,ry,h,st,e/Tm3 Ser [31], or expressing green fluorescent protein (GFP) under the promoter of the AMP Dipterisin _A_ (DptA), _DptA_-GFP [32]. They were maintained on standard glucose medium (12 g agar, 100 g glucose and 100 g Brewer’s yeast per 1.2 L of water, plus a final concentration of 0.04% phosphoric acid and 0.4% propionic acid added to inhibit microbial growth in the food) and kept at room temperature (22-24°C). Table 1 provides a complete list of Providencia bacterial strains. All Providencia strains were grown in LB media at 37°C overnight with shaking, except for _P. burhodogranariea_ strains, which were grown at 25°C. _Listeria monocytogenes_ 10403S was grown at 37°C in BHI medium with shaking. _Escherichia coli_ Mach1-T1, a cloning strain (Invitrogen Corp), was grown at 37°C in LB medium with shaking.

2.2. Mortality

Overnight cultures used for infecting flies were grown to saturation and then diluted to an _A_{600nm} of 1.0. To deliver infections, a 0.15 mm minute pin (Fine Science Tools) mounted on a 200 µL pipet tip was dipped into the diluted overnight culture and poked into the thorax of a CO₂
anesthetized fly. This delivers about $10^3$ to $10^4$ bacteria to each fly. Sterilely wounded flies were pricked with a needle that was sterilized in 95% ethanol. Anesthetized control flies were handled in the same way as the others but were not wounded. Flies were maintained in vials with food at room temperature and surviving flies were counted once a day for 6 days after infection. Infection with each bacterium was performed on at least 2 days with controls done on each day. Product limit survival estimates and homogeneity by log-rank tests were conducted using proc lifetest in SAS version 9.1 (SAS Institute). \( P \)-values were corrected for multiple tests in some cases by a Bonferroni correction with a cut off value of \( p = 0.0025 \) for comparing all strains that are the focus of the paper, \( p = 0.00625 \) for comparing among \( P. \) sneebia isolates only, and \( p = 0.025 \) for comparing only among \( P. \) burhodogranariea strains. In contrasts of different strains of \( P. \) burhodogranariea, only those infections that were preformed on the same day were compared.

2.3. Bacterial load

To measure systemic bacterial load in infected flies, single OreR flies were infected by pinprick as described in Section 2.2, then homogenized in 500 \( \mu \)L LB and plated by robotic spiral platers (manufactured by Don Whitley Scientific and Spiral Biotech) on LB agar plates at 0, 2, 4, 6, 10, 18, 24, and 32 h post infection. Flies were kept in vials with food at room temperature between infection and homogenization. The LB agar plates were incubated overnight at 25°C for \( P. \) burhodogranariea or 37°C for \( P. \) rettgeri, \( P. \) alcalifaciens, \( P. \) sneebia and sterile wound. Gut commensal bacteria grow more slowly than \( Providencia \) under these conditions, so by limiting incubation to overnight we exclude any commensal bacteria from our assay. The number of colony forming units (CFU) on each plate was recorded using a counter.
associated with the spiral platers, allowing the concentration of viable bacteria in each homogenate to be calculated based on the number and position of colonies on the plates.

Bacterial loads for flies infected with *P. alcalifaciens* and *P. sneebia* were compared at each individual time point using proc glm in SAS version 9.1 with the model: ln(CFU+1) = bacterial treatment + sex. The boxplot was generated using the function boxplot in R. A small number of surviving flies from each treatment were also homogenized at 7-10 days post infection as described above.

2.4. Antimicrobial peptide expression

We first examined *DptA*-GFP flies to determine how much AMP expression occurred during infection. We infected flies on replicate days as described in Section 2.2 and kept them in vials with food until the time examined. Other AMP promoters examined which had undetectable levels of fluorescence were *Defensin*, *Drosocin*, *Attacin* and *Cecropin* [32]. At 6, 24, and 32 h post infection flies were anesthetized and examined under a dissecting scope and scored for the intensity of GFP fluorescence blind of the treatment. This assay was restricted to females because males were found to have too much background fluorescence.

For quantification of AMP expression by QPCR, OreR flies were either infected with a bacterium or sterilely wounded as described in Section 2.2 then were frozen at -80°C in pools of 8 flies at 2, 4, 6, 10, 18, 24, and 32 h post treatment. Flies were maintained in vials with food at room temperature between infection and freezing. Each treatment was performed on at least two different days. Total RNA was extracted with Trizol (Invitrogen Corp) using the manufacturer’s suggested protocol, then reverse transcribed to cDNA from poly-T primers using standard procedures. The abundances of the AMPs *Diptericin A* (*DptA*), *Drosomycin* (*Drs*) and
Defensin (Def) and the housekeeping gene rp49 were quantified by QPCR on an ABI 7000 Sequence Detection System (Applied Biosystems) using specific TaqMan primers and the manufacturer’s suggested protocol (primer and probe sequences available upon request). For statistical analysis, gene expression at each hour was examined separately in proc glm in SAS version 9.1 using the model: AMP Ct = Rp49 Ct + treatment + date infected. Correction for multiple tests was achieved using the Tukey-Kramer method. Least squares means were recovered at the mean Rp49 Ct. Fold induction was calculated as 2 to the power of the difference between the Ct of the sterile wound control and the Ct of the infection treatment for each time post infection.

2.5. Coinfection

For coinfections, overnight cultures of *P. rettgeri* and *P. sneebia* were grown to saturation and then diluted to an A$_{600nm}$ of 2.0. The bacteria were then mixed at proportions 1:1, 1:3 or 3:1 with either the alternate bacteria or LB. Flies were then infected in the thorax with a small needle dipped in the culture as described in Section 2.2, replicated on two different days. Although three different proportions of each bacterium were examined, we found that the results were the same for each infection class (singly infected *P. rettgeri*, singly infected *P. sneebia*, or coinfected) regardless of the mixing proportion, allowing us to pool all proportions in final analyses. We only examined male flies for AMP expression and bacterial load in the coinfection because we had found no difference between the sexes in our primary examination of mono-infections. At 6, 24, and 32 h post infection flies were frozen at -80°C. RNA extraction, QPCRs, and statistical analysis for AMP expression were performed as described in Section 2.4. Fisher’s combined probability was used to summarize the independent expression experiments.
For the examination of AMP expression in *DptA*-GFP flies, infected or control flies were placed in vials with standard fly food and examined blind of treatment at 6, 24, and 32 h post infection with a dissecting scope. Here, only female flies were examined due to male background fluorescence. The survival of these same flies was monitored up to six days post infection and statistically analyzed as in Section 2.2.

Determination of the bacterial load of coinfected flies and statistical analysis was carried out as described in Section 2.3. To distinguish between *P. sneebia* and *P. rettgeri*, we took advantage of *P. rettgeri*’s natural resistance to tetracycline. All samples were plated on LB plates without antibiotic and on plates with a tetracycline concentration of 10 µg/mL. The number of CFU on the tetracycline plates was inferred to be the count of *P. rettgeri* and the difference in CFU between the paired plates was assumed to be the *P. sneebia* count. PCR and restriction enzyme digestion of the 16S gene looking for species-specific digestion pattern was done to check that the proper species were growing on the correct plates. This experiment was carried out twice on different days.

2.6. Biofilm formation

Overnight bacteria cultures were diluted to an $A_{600\text{nm}}$ of 1.0, then gently centrifuged into a pellet and washed three times with 1 x PBS, and ultimately concentrated to 20×. 5 µl of bacteria or PBS, as a control, were added to 200 µL of Schneider’s media with 10% fetal calf serum in a 96-well plate. Bacteria that received the antibiotic treatment sat in media alone for approximately 1 h before the antibiotics were added to the well. The antibiotics ceftazidime and kanamycin were added to a final concentration of 1 mg/mL and 200 µg/mL, respectively. At 6 and 24 h after the bacteria or antibiotics were added to the media, the wells were washed three times with
sterile water before the addition of 0.1% crystal violet, and then incubated for 15 min. The wells were then washed twice with water before drying for 5 h. 30% acetic acid was added to the wells to solubilize the crystal violet. The A₅₄₀nm was read using Multiskan Spectrum plate reader (Thermo Scientific). The final absorbance was calculated as the difference from the PBS control well at that time.

2.7. Antibiotic protection assay

*D. melanogaster* S2 cells were maintained in Schneider’s media with 10% fetal calf serum at 25°C. For the antibiotic protection assay, cells were seeded in 6 well plates the day before the assay was carried out so that there would be approximately 10⁵ cells/mL the next day. Overnight cultures of bacteria were washed three times with PBS before addition to the wells containing S2 cells at a multiplicity of infection of 10. After 2 h the media was removed and the cells, which lightly adhere to the bottom of the wells, were washed while still in the wells three times with PBS. Schneider’s media with 10% fetal calf serum containing 1 mg/mL ceftazidime and 20 µg/mL kanamycin was then added to the wells. Neither ceftazidime nor kanamycin should penetrate eukaryotic cell membranes, so only extracellular bacteria should be killed. The cells were incubated with the antibiotics for 2 h to kill extracellular bacteria. At 0, 6 and 24 h following this 2 h incubation, the media only was removed from the wells and centrifuged. The pellet was then washed with water before being plated on BHI or LB plates, depending on the bacteria, to provide an estimate of the number of viable bacteria in suspension (this number should be near zero because of the presence of antibiotics). The S2 cells were then washed off with water and spun down and washed again with water. The pellet was then resuspended in BHI media before being plated on either BHI or LB. CFUs were manually counted to yield the
number of viable bacteria residing inside the S2 cells.

3. Results

3.1. Mortality

Given that closely related bacteria often vary in their virulence to a given host, we hypothesized that the different strains of Providencia isolated from wild caught Drosophila melanogaster might also vary in pathology (Table 2.1). There was minimal mortality (5-10%) among control flies either only anesthetized on CO₂ or wounded with a sterile needle. When flies are infected with P. burhodogranariea strain D, less than 10% of infected flies died by six days post infection (Figure 2.1). This is not significantly different from the amount of mortality from either control (Figure 2.1; P. burhodogranariea strain D contrasted to CO₂ control: p = 0.0612, P. burhodogranariea strain D-sterile wound: p = 0.0436, not significant after correcting for multiple tests). About 40% of flies infected with P. burhodogranariea strain B die from the infection, which is highly significantly different from P. burhodogranariea strain D (p < 0.0001), although for unknown reasons, P. burhodogranariea strain B infections displayed more day-to-day variation in mortality than infections with any other bacteria with mortality rates ranging from 20% to 60%. P. rettgeri strain Dmel likewise caused moderate mortality, with fewer than 50% of the flies dying. The amounts of mortality caused by P. rettgeri and P. burhodogranariea strain B are not significantly different from each other (p = 0.0303), although both infections caused significantly higher mortality than is observed in controls (all p < 0.0001). P. sneebia strain Type and P. alcalifaciens strain Dmel each caused much greater mortality than any of the other species. Within the first two days of infection, P. sneebia kills about 90% of infected flies and P. alcalifaciens causes mortality in 99% of infected flies. Mortality from
Figure 2.1. Mortality of *D. melanogaster* from *Providencia* Infection. Wild type *D. melanogaster* were infected through pinprick infections with different strains of *Providencia*. All treatments result in highly significant differences in mortality (all pairwise contrasts $p < 0.0001$), except the difference between sterile needle and CO$_2$ controls, between infection with *P. burhodogranariea* strain D and either control, and between *P. rettgeri* and *P. burhodogranariea* strain B (in all cases $p > 0.0025$, the Bonferroni-corrected cut off value).
infections with each *P. sneebia* and *P. alcalifaciens* is significantly different from all other treatments, including each other (all \( p < 0.0001 \)). Thus, there are three major classes of virulence among our isolated *Providencia* as defined by mortality: *P. burhodogranariea* strain D causes minimal mortality, *P. rettgeri* and *P. burhodogranariea* strain B cause moderate amounts of mortality, and *P. sneebia* and *P. alcalifaciens* are highly virulent.

Multiple isolates of *P. sneebia* and *P. burhodogranariea* strain B have been recovered from the hemolymph of wild caught *D. melanogaster* (Table 2.1) [2,3]. We infected flies with each of these to test whether there is heterogeneity among isolates in the mortality caused by these strains. Of the two other isolates of *P. burhodogranariea* strain B, only isolate B97 is significantly different than the Type strain B, with B97 causing less mortality (Figure 2.2; Supplementary Table 1; \( p = 0.0003 \)). Eight *P. sneebia* isolates were tested and all caused greater than 80% mortality, although some of them cause slightly but significantly different mortality than the Type strain (Table 2.2). This suggests that while there is some variation among isolates, *P. sneebia* can be considered to always be highly virulent while *P. burhodogranariea* is never highly virulent.

The *Drosophila* humoral immune response is activated by two major signaling pathways, the Toll pathway and the Imd pathway [4]. The Imd pathway tends to be more responsive to Gram-negative bacteria, whereas the Toll pathway preferentially activated by Gram-positive bacteria. We therefore hypothesized that the Imd pathway would be more important in fighting *Providencia*. We measured the mortality of flies that were mutationally deficient in either the Toll or Imd pathway after infection with *Providencia*. We found Toll pathway mutants showed no significant difference in mortality compared to wild type flies after infection with either strain of *P. burhodogranariea* or with *P. rettgeri* (\( p > 0.05 \), in all cases). *P. sneebia* and *P.*
Table 2.1. Bacterial Strains Used. (*) indicates the strains that are the main focus of this work.

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<th>Citation</th>
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<td>wild <em>D. melanogaster</em> hymolymph</td>
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Figure 2.2. Mortality of *D. melanogaster* from Infection with Other Isolates of *P. sneebia* and *P. burhodogranariea* strain B. Wild type *D. melanogaster* were infected through pin prick infections with different strains of *Providencia*. All isolates of *P. burhodogranariea* strain B cause low to moderate mortality, while infection with any isolate of *P. sneebia* results in high mortality.
Table 2.2. Statistics of Mortality of other isolates of *P. sneebia* and *P. burhodogranariea* strain B. *P*-value cut off was corrected for multiple testing using a Bonferroni correction resulting in a cut off value of *p* = 0.00625 for *P. sneebia* isolates and *p* = 0.025 for *P. burhodogranariea* strains. Values in bold are those that are significant.

<table>
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<th><em>P. sneebia</em> strain</th>
<th>Type</th>
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</thead>
<tbody>
<tr>
<td><em>P. sneebia</em> strain A101</td>
<td>0.099</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A102</td>
<td>0.0003</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A104</td>
<td>0.0002</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A16</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A36</td>
<td>0.0019</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A75</td>
<td>0.8125</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A83</td>
<td>0.6826</td>
</tr>
<tr>
<td><em>P. sneebia</em> strain A91</td>
<td>0.2144</td>
</tr>
<tr>
<td><em>P. burhodogranariea</em> strain B</td>
<td></td>
</tr>
<tr>
<td><em>P. burhodogranariea</em> strain B18</td>
<td>0.0326</td>
</tr>
<tr>
<td><em>P. burhodogranariea</em> strain B97</td>
<td>0.0003</td>
</tr>
</tbody>
</table>
*alcalifaciens* did cause significantly greater mortality in the Toll mutant flies compared to the wild type flies (*p* < 0.05, in both cases), but flies of both genotypes suffered severe mortality within 2 days of infection with these bacteria (Figure 2.3A). In contrast, Imd mutant flies infected with any strain of *Providencia* suffered very high mortality within 2 days post infection (Figure 2.3B). Notably, we observed high mortality in flies infected with the *P. rettgeri* and *P. burhodogranariea* strains, which cause only moderate to low mortality in wild type flies (Figure 2.3B). All *Providencia* infections in Imd mutant flies were significantly different than those seen in infected wild type flies (*p* < 0.05, in all cases). These data indicate that the Imd pathway is essential to fighting *Providencia* infection, and that *P. rettgeri* and *P. burhodogranariea* infections are controlled by the host immune system and not simply limited by inherent failure of the bacteria to be able to colonize the fly.

We were intrigued by the recurrent isolation of diverse *Providencia* species from *Drosophila*, so we examined the amount of mortality caused in *D. melanogaster* by the Type strains of 6 *Providencia* species isolated in other contexts, including *P. rettgeri* and *P. alcalifaciens* isolates not derived from *Drosophila* (Table 2.1; Figure 2.4). Except for *P. alcalifaciens* strain Type, all species caused less than 20% fly mortality in wild type flies. The Type strain of *P. alcalifaciens* caused less mortality than our Dmel strain (Figure 2.4; *p* < 0.0001), which suggests there are genetic differences between the strains. The Type strain of *P. rettgeri* also caused less mortality than our Dmel strain (Figure 2.4; *p* < 0.0001). These data indicate that the high amount of *Providencia*-induced *D. melanogaster* mortality is specific to those strains that were isolated from wild flies.
Figure 2.3. Mortality of Immune Mutant *D. melanogaster*. (A) Toll (B) Imd pathway deficient flies. When the Imd pathway is non-functional, all bacteria are highly virulent, while when the Toll pathway is mutated, the bacteria cause the same degree of mortality as seen in the wild type flies.
Figure 2.4. Mortality of *D. melanogaster* After Infection with Type strains of other *Providencia* species. Only those *Providencia* strains that were isolated from wild *Drosophila* are able to cause a high mortality. The Type strains of *P. rettgeri* and *P. alcalifaciens* are both significantly different from the Dmel strains (*P. rettgeri* strain Dmel contrasted to *P. rettgeri* strain Type: \( p < 0.0001 \). *P. alcalifaciens* strain Dmel contrasted with *P. alcalifaciens* strain Type: \( p < 0.0001 \)).
3.2. Bacterial load

For a given host and pathogen pair, bacterial proliferation and host mortality may or may not be correlated. To test our hypothesis that the *Providencia* species that cause the highest mortality are those that are best able to replicate in flies, we measured the number of bacteria present in *D. melanogaster* at multiple time points for the first 32 h after infection. Plates from control flies that were sham-infected with a sterile needle did not have any bacteria growth after the overnight incubation period (data not shown), indicating that the control flies did not have any *Providencia* within or on them. Commensal bacteria from the gut begin to appear on all plates after they have been incubated for at least 24 h. Infections with the five bacteria start to diverge in CFU counts around 10 h post infection (Figure 2.5). There are a few individual flies that are able to clear the infection during the first few hours. It is unclear why some flies are able to clear their infections and others are not, although we suspect it reflects minor heterogeneities in the infection process.

Flies infected with either strain of *P. burhodogranariea* cleared their infections or maintained stable bacterial loads around the level of the initial introduction over the first 32 h of infection (Figure 2.5). These bacteria are eventually cleared from all surviving flies, as survivors have no bacteria present 7-10 days post infection (data not shown). *P. rettgeri, P. sneebia* and *P. alcalifaciens* all show an increase in the number of CFU per fly after 6 h of infection. Among the flies infected with *P. rettgeri*, there is a large amount of variation in the number of bacteria present in individual flies at 24 and 32 h post infection, ranging from $10^3$ to $10^7$ CFU per fly (Figure 2.5). It seems likely that this variation reflects divergence in the infection trajectory among individual flies, where those with the highest bacterial loads probably succumb to the infection and the others survive. Flies that survived their infections for 7-10 days post infection
Figure 2.5. *Providencia* Bacterial Load in *D. melanogaster*. Boxplot of the number of CFU present in *D. melanogaster* during the first 32 hours post infection. Note that the y-axis is a log scale. Whiskers approximate two times the standard deviation. The table under the graph has the number of flies that had no CFU at each time point for each treatment, as well as the total infected flies per treatment at each time point. Flies with no CFU present were not included in the boxplot. Sterilely wounded control flies never had any CFU at any time point.
carried either no CFU or between $10^2$ and $3 \times 10^4$ CFU per fly (data not shown). Both *P. sneebia* and *P. alcalifaciens* are able to rapidly proliferate to very high numbers in the fly by 32 h post infection, which is shortly before flies die from these infections. The number of bacteria present in the infected flies is significantly different between *P. sneebia* and *P. alcalifaciens* at 18 h post infection ($p = 0.0115$), but not at 24 and 32 h (both $p > 0.05$). Approximately 10% of the total *P. sneebia* infected flies had no bacteria present at their time of sampling. These flies were most likely able to clear the bacteria within the first few hours of infection and probably represent the small percent of flies that survive in the mortality assays (Figure 2.1). This hypothesis is supported by the observation that flies infected with *P. sneebia* that survive 7-10 days post infection are free of *Providencia* (data not shown). In total, across all species, these data demonstrate that the *Providencia* species that are best able to proliferate within the fly are those that cause the highest mortality.

### 3.3. *D. melanogaster* immune response to infection

Insects respond to the presence of bacteria by activating their humoral immune system, which results in the production of antimicrobial peptides (AMPs). Induction of AMP gene expression varies among different microbes and immune elicitors [33], and we hypothesized that the *Providencia* bacteria that were most proliferative during infection would cause the highest induction of the immune response. To initially test this hypothesis, we infected transgenic flies that express GFP driven by AMP promoters [32] then examined individual flies by eye at 6, 24, and 32 h post infection. Although the expression patterns of several different AMPs were examined (see Section 2.4), only *DptA* produced a strong fluorescence after infection with most *Providencia*. Both *P. burhodogranariea* strains failed to drive detectable fluorescence signal.
even with \textit{DptA}. As expected, \textit{DptA}-GFP expression was localized to the immune responsive fat body. We repeatedly saw that flies that were infected with \textit{P. sneebia} showed lower fluorescence than flies infected with \textit{P. alcalifaciens} or \textit{P. rettgeri}, both in the intensity of the GFP expression as well as the proportion of a single fly expressing GFP (Figure 2.6A). This contrasted with our expectation based on the high levels of \textit{P. sneebia} proliferation within flies and host mortality caused by infection.

We used QPCR of AMP mRNAs to better quantify the immune response of infected flies for the first 32 h of infection relative to control flies that were wounded with a sterile needle. By calculating the fold induction over the sterile wound, we could determine the amount of AMP expression that was specifically attributable to the bacteria and not to the wound in delivering the infection (Figure 2.6B). Consistent with our observations of the \textit{DptA}-GFP flies, \textit{P. sneebia} infections consistently resulted in lower expression of \textit{DptA} than \textit{P. rettgeri} and \textit{P. alcalifaciens} did at later times in the infection progression. At 24 and 32 h after infection with \textit{P. sneebia}, \textit{DptA} expression was not significantly different from expression in response to the sterile wound alone (Figure 2.6B; both \(p > 0.05\)). In contrast, \textit{P. rettgeri} and \textit{P. alcalifaciens} induced significantly higher levels of expression than the sterile wound at 24 and 32 h after infection (Figure 2.6B; in all cases \(p < 0.05\)). None of the bacterial infections drove \textit{DptA} expression above the level seen from sterile wound alone prior to 24 h post infection, and flies infected with either strain of \textit{P. burhodogranariea} never showed \textit{DptA} expression above what is seen for the sterile wound treatment at any time point (Figure 2.6B).

\textit{Providencia} induction of \textit{Drs} over sterile wound was generally much smaller, and none of the infection treatments differ significantly from the sterile wound until 32 h post-infection (Figure 2.7A). The pattern of \textit{Def} expression was more complex, with strong induction in
Figure 2.6. *DptA* Expression in Flies Infected with *Providencia*. (A) *DptA*-GFP flies infected with (left to right) a sterile needle, *P. sneebia*, or *P. rettgeri* at 32 hours post infection. (B) Graph of *DptA* expression as measured by QPCR. The fold induction was calculated as the level of expression above that caused by a sterile wound alone. Error bars represent the standard error. At each time point, treatments labeled with “a” are not significantly different from the sterile wound alone while those with “b” are significantly different from the sterile wound (corrected for multiple tests by Tukey-Kramer method, cut off $p = 0.05$).
Figure 2.7. AMP Expression in Flies Infected with *Providencia* as measured by QPCR. (A) *Drs* (B) *Def*. The fold induction was calculated as the level of expression above that caused by a sterile wound alone. Error bars represent the standard error. (*) indicates those treatments that were significantly different from wounding with a sterile needle (corrected for multiple tests by Tukey-Kramer method, cut off $p = 0.05$).
response to \textit{P. rettgeri} and \textit{P. alcalifaciens} infections at 18 and 24 h post infection (Figure 2.7B). The induction of \textit{Def} in response to \textit{P. sneebia} is much delayed relative to \textit{P. alcalifaciens} and \textit{P. rettgeri} infection, with strong induction not appearing until 32 h post infection.

In summary, we observed that some of \textit{Providencia} species that proliferate the most within the fly and cause the greatest host mortality also drive higher AMP expression. An interesting departure from this trend is \textit{P. sneebia}, which is highly virulent and reaches the highest abundance within the fly, but expression of \textit{DptA} caused by \textit{P. sneebia} infection is never significantly higher than that caused by a sterile wound (Figure 2.6B).

3.4. Coinfections with \textit{P. rettgeri} and \textit{P. sneebia}

\textit{P. sneebia} could avoid inducing a strong immune response by actively evading detection by the host or by actively suppressing the immune response. To distinguish between these two possibilities, we took advantage of the differences in mortality and immune induction resulting from \textit{P. rettgeri} and \textit{P. sneebia} infections. We coinfected flies with both bacteria simultaneously and then measured AMP expression, host mortality, and bacterial load. We hypothesized that if \textit{P. sneebia} actively suppresses the immune response, we would see low levels of AMP expression even in the presence of \textit{P. rettgeri}. Alternatively, if \textit{P. sneebia} is not detected by the immune system, we would expect to see high levels of AMP expression induced by the presence of \textit{P. rettgeri} in the coinfection.

We measured \textit{DptA}, \textit{Drs} and \textit{Def} levels in groups of flies either coinfected or infected with an individual bacteria at 6, 24, and 32 h post infection by QPCR (Figure 2.8-2.9). Across all 3 AMPs, infection with \textit{P. sneebia} alone caused a lower immune response than infection with \textit{P. rettgeri} alone at 32 h post infection (Fisher’s combined probability, \(p = 0.00061\)), consistent with
Figure 2.8. *DptA* expression by *D. melanogaster* Infected with *P. sneebia, P. rettgeri* or Both Measured by QPCR. The fold induction was calculated as the level of expression over that caused by a sterile wound alone. Error bars represent the standard error.
Figure 2.9. AMP Expression in Flies Coinfected with *P. sneebia* and *P. rettgeri* as measured by QPCR. (A) *Def*(B) *Drs*. The fold induction was calculated as the level of expression above that caused by a sterile wound alone. Error bars represent standard error.
the results presented in the previous section. Expression of the 3 AMPs at 32 h post infection in coinfected flies was not significantly different from expression in flies singly infected with P. rettgeri (Fisher’s combined probability, p = 0.528), but coinfected flies had significantly higher expression than flies infected with P. sneebia (Fisher’s combined probability, p = 0.014). This result was further supported by visually examining the level of GFP expression in individual DptA-GFP flies, in which coinfected and P. rettgeri infected flies fluoresced more intensely than flies infected with P. sneebia. These data show that the lower expression of AMPs in flies infected with P. sneebia alone is not due to suppression of the immune response.

Consistent with previous mortality measurements (Figure 2.1), approximately 40% of the flies infected with P. rettgeri alone died from their infections, whereas about 95% of the flies infected with P. sneebia died within 72 h (Figure 2.10). Coinfected flies exhibited 85% mortality. While all three treatments are significantly different than each other (in all cases p < 0.01), the overall mortality of coinfected flies is clearly more similar to that of flies infected with P. sneebia alone (Figure 2.10). When bacterial load was measured at 32 h post infection, we observed that the abundance of each individual bacterium in the coinfected flies was not significantly different than their levels in flies that are singly infected (Figure 2.11; in all cases p > 0.05). Thus, it appears that the growth trajectories of the bacteria are completely independent of each other, and coinfected flies carry a bacterial load equivalent to the sum of each single infection. Considering all three coinfection phenotypes together, it is evident that P. sneebia is not able to suppress the host immune response, but is able to proliferate and cause host mortality even in the presence of an immune response triggered by P. rettgeri.
Figure 2.10. Survival of *D. melanogaster* from infections with *P. sneebia*, *P. rettgeri* or both.
Figure 2.11. Bacterial load of *D. melanogaster* infected with *P. sneebia*, *P. rettgeri* or both. The two bacteria are plotted separately in pale colors for the coinfected flies. Whiskers approximate two times the standard deviation. Coinfected flies show full induction of the immune system, but succumb to their infections and permit bacterial growth that is not different than what is observed in single infections.
3.5. Biofilm formation

One way that *P. sneebia* could protect itself from recognition and the microbicidal AMPs is by forming a biofilm within the fly [34]. The fly would only be able to detect bacteria at the perimeter of the biofilm, and thus the magnitude of the immune response would not be proportional to the total number of bacteria present. Additionally the bacteria within the biofilm would be able to freely multiply without being affected by expressed AMPs. We tested all of our *Providencia* isolates for their ability to form biofilms *in vitro* in 96-well plates. *E. coli* was used as a control that can form a biofilm [35]. None of our *Providencia* isolates formed a biofilm (Table 2.3). However, we cannot definitively rule out the possibility that *P. sneebia* might form a biofilm within the fly, since there could be host-specific molecules that act as signal to trigger *P. sneebia* biofilm formation *in vivo*.

3.6. Antibiotic protection assay

Another way that *P. sneebia* could evade detection and proliferate would be if it were able to invade cells and replicate within them. Strains of *P. alcalifaciens* that were isolated from human patients with diarrhea have been shown to invade human cells, demonstrating that some *Providencia* are able to do the first step in this process [16,17,19,20]. We used an antibiotic protection assay to test whether *P. sneebia* is able to divide within *D. melanogaster* cells. We also tested whether *P. alcalifaciens* and *P. rettgeri* are able to divide within *D. melanogaster* cells since they proliferate within the fly during infection. *E. coli* was used as a negative control bacteria that would be passively phagocytosed by the cells but is unable to replicate within. *L. monocytogenes* was used as a positive control that is able to replicate within insect cells [36]. Bacteria were exposed to a phagocytic *D. melanogaster* cell line for 2 h before antibiotics were
Table 2.3. *Providencia* isolated from *D. melanogaster* are not able to form biofilms. Measurements of absorbance of crystal violet at 540nm of replicate wells after biofilm formation *in vitro*. *E. coli*, which is capable of forming a biofilm, has much higher absorbance than any of the *Providencia* species.

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>Relative biomass (A&lt;sub&gt;540nm&lt;/sub&gt;)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>6 hours average</td>
</tr>
<tr>
<td><em>E. coli</em></td>
<td>0.0717 +/- 0.0277</td>
</tr>
<tr>
<td><em>P. sneebia</em></td>
<td>-0.0027 +/- 0.0047</td>
</tr>
<tr>
<td><em>P. alcalifaciens</em></td>
<td>0.0039 +/- 0.0108</td>
</tr>
<tr>
<td><em>P. rettgeri</em></td>
<td>0.0056 +/- 0.0098</td>
</tr>
<tr>
<td><em>P. burhodogranariea</em></td>
<td>0.0043 +/- 0.0002</td>
</tr>
<tr>
<td>strain B</td>
<td></td>
</tr>
<tr>
<td><em>P. burhodogranariea</em></td>
<td>0.0048 +/- 0.0083</td>
</tr>
<tr>
<td>strain D</td>
<td></td>
</tr>
</tbody>
</table>
added to the media to kill all extracellular bacteria. At 0, 6 and 24 h post antibiotic killing of extracellular bacteria, both the media and the cells were plated separately and CFU were counted. The CFU found in the media were minimal by comparison to those within the cells. Since *E. coli* will only be passively phagocytosed by the cells, it was used as a standard for determining if any of our strains are actively invading the cells. Among all replicates, there were consistently fewer *P. sneebia* and *P. rettgeri* than *E. coli* inside host cells at the 0 h time point (Table 2.4), suggesting that these bacteria have some resistance to phagocytosis by these *D. melanogaster* cells. By contrast, *P. alcalifaciens* had higher numbers of CFU than *E. coli* at the initial time point suggesting that our strain of *P. alcalifaciens* is invasive. The positive control, *L. monocytogenes*, was able to replicate to high numbers within the cells. *P. rettgeri*, *P. sneebia*, *P. alcalifaciens*, and the negative control, *E. coli*, all had fewer intracellular CFU 24 h after addition of antibiotic than at 0 h, indicating that none of them are able to replicate within the insect cells. These data suggest that *P. sneebia* does not avoid recognition by the immune response by invading and proliferating in *D. melanogaster* cells.

4. Discussion

We have established that closely related bacteria in the genus *Providencia* vary in their pathology in a natural host, *Drosophila melanogaster*, as measured by the amount of mortality they cause, their ability to replicate within the host and the magnitude of the host immune response to their presence. Those bacteria which are able to grow most effectively in the fly often also trigger the most robust immune response and result in the most host death. However, one of these bacteria, *P. sneebia*, causes nearly complete mortality and quickly replicates to high numbers within the fly but does not induce a strong immune response. Through coinfections with
Table 2.4. *Providencia* isolated from *D. melanogaster* are not intracellular pathogens. Number of CFU within *D. melanogaster* S2 cells at 0, 6, and 24 hours post antibiotic killing of extracellular bacteria. Two replicate wells were measured each day and the experiment was carried out on multiple days with similar results. *Listeria monocytogenes*, which is capable of intracellular invasion and proliferation, grows to high density within host cells, whereas *Providencia* and *E. coli*, which is not capable of intracellular invasion, are progressively eliminated.

<table>
<thead>
<tr>
<th></th>
<th>well</th>
<th>0 hour</th>
<th>6 hour</th>
<th>24 hour</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. sneebia</em></td>
<td>1</td>
<td>543</td>
<td>468</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>466</td>
<td>460</td>
<td>22</td>
</tr>
<tr>
<td><em>P. rettgeri</em></td>
<td>1</td>
<td>285</td>
<td>110</td>
<td>49</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>138</td>
<td>120</td>
<td>88</td>
</tr>
<tr>
<td><em>P. alcalifaciens</em></td>
<td>1</td>
<td>2541</td>
<td>2988</td>
<td>1620</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>2265</td>
<td>2061</td>
<td>1170</td>
</tr>
<tr>
<td><em>E. coli</em></td>
<td>1</td>
<td>1734</td>
<td>942</td>
<td>154</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>1500</td>
<td>465</td>
<td>105</td>
</tr>
<tr>
<td><em>L. monocytogenes</em></td>
<td>1</td>
<td>7570</td>
<td>8617</td>
<td>109680</td>
</tr>
<tr>
<td></td>
<td>2</td>
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<td>141200</td>
</tr>
<tr>
<td>PBS</td>
<td></td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
the less virulent *P. rettgeri*, we concluded that *P. sneebia* is able to actively avoid detection by the immune system as well as protect itself from the immune response. We did not find evidence that *P. sneebia* forms a biofilm or replicates intracellularly *in vitro*. Although we were unable to determine the exact virulence mechanisms used by *P. sneebia* during infection of *D. melanogaster*, our data imply that *P. sneebia* implements more complicated or multiple strategies to subvert the immune system.

We note that the proportion of flies that die from each bacterial infection in the mortality assays is approximately equivalent to the proportion of flies that sustain high numbers of bacteria in the load experiments, suggesting that the individual flies in which *Providencia* is able to replicate are those that succumb to the infection. The data we have for flies infected with *P. burhodogranariea* strain B does not conform to this hypothesis, as that bacterium causes a moderate amount of mortality despite not replicating within the fly as much as the similarly virulent *P. rettgeri*. This suggests that *P. burhodogranariea* strain B might do proportionally more damage to the fly, possibly by producing a harmful compound, at a lower density than the other bacteria. This also points at a distinction between the two *P. burhodogranariea* strains, as they both have similar levels of bacteria present during the first 32 h of infection but strain D causes significantly less mortality. The two strains are defined as distinct based on differences in metabolic profiles and in sequence of some housekeeping genes [3]. Our data suggest there are likely to be further genetic differences between the strains, including in genes involved in the phenotypes examined here.

Although we are primarily interested in *Providencia* species that are natural pathogens of *D. melanogaster*, we also examined mortality due to infection by other species in the genus, which have been isolated as clinical infections of humans and other animals, or in one case,
Providencia vermicola, as an associate of entomopathogenic nematodes [12,13,37]. The only bacteria we found to cause high mortality in infected Drosophila are those which were originally isolated from wild caught D. melanogaster. Two of these species were also previously described as clinical pathogens of humans, but in both of these species, P. alcalifaciens and P. rettgeri, the Dmel strain isolated from D. melanogaster caused greater mortality than the Type strain of the species. These results suggest that Providencia strains may become highly adapted to the host species they infect, and that the isolates recovered from D. melanogaster may be genetically suited to infect Drosophila and its close relatives. More detailed genomic and pathological examination of Providencia should reveal genes specifically involved in virulence to Drosophila.

Because we are specifically interested in the D. melanogaster-Providencia interaction after infection has occurred, we have relied on artificial infections to deliver the bacteria. Nevertheless, it is worth considering how the bacteria may establish infections in the wild. Our Providencia isolates do not cause mortality after being fed to flies in reasonable doses in the laboratory (data not shown), so it is not likely that they orally infect flies in the wild unless they are aided by coinfectors (e.g. [38,39]). There is good reason to believe, however, that our method of infecting through a pinprick wound may mimic infections that wild D. melanogaster can receive [40]. Wild caught flies often have melanization independent of natural pigmentation patterns, indicating healed wounds, and frequently carry ectoparasitic mites that could be the cause of some of these wounds (unpublished observation). Mite wounds in honey bees have been shown to be secondarily colonized by environmental bacteria [41]. P. burhodogranariea strain B has been isolated from a mite pulled from a wild caught D. melanogaster (P. Juneja, personal communication), suggesting that mites may also directly vector bacterial infections, although fly-to-fly transmission of Providencia via mites has not been experimentally demonstrated.
We anticipate that *D. melanogaster-Providencia* system will be an excellent one for continued examination of many aspects of host and pathogen interactions. There is ample phenotypic diversity in the host-pathogen interaction, with clear variation among *Providencia* species in pathological phenotypes. Both the bacteria and the insect host can be easily and inexpensively manipulated in the lab, providing a valuable setting to conduct research that will not only give insight into interactions specific to this host-pathogen pairing, but also into generic virulence mechanisms and their genetic basis. *D. melanogaster* has been extensively studied as a generic host for pathogenic bacteria and a model for innate immune system function, and these *Providencia* isolates now provide an opportunity to study how flies fight those bacteria that infect them in their natural environment.

**Acknowledgments**

We would like to thank Punita Juneja, Sarah Short, Jacob Crawford, Susan Rottschaefer, and Jennifer Comstock for helpful comments on the manuscript, discussion and help collecting the data. This work was supported by NSF grant DEB-0415851 and NIH grant AI083932.

**Reference**


CHAPTER 3

COMPARATIVE GENOMICS OF BACTERIA IN THE GENUS PROVIDENCIA ISOLATED FROM WILD DROSOPHILA MELANOGASTER
Abstract

Background

Comparative genomics can be an initial step in finding the genetic basis for phenotypic differences among bacterial strains and species. Bacteria belonging to the genus *Providencia* have been isolated from numerous and varied environments. We sequenced, annotated and compared draft genomes of *P. rettgeri*, *P. sneebia*, *P. alcalifaciens*, and *P. burhodogranariea*. These bacterial species that were all originally isolated as infections of wild *Drosophila melanogaster* and have been previously shown to vary in virulence to experimentally infected flies.

Results

We found that these *Providencia* species share a large core genome, but also possess distinct sets of genes that are unique to each isolate. We compared the genomes of these isolates to draft genomes of four *Providencia* isolated from the human gut and found that the core genome size does not substantially change upon inclusion of the human isolates. We found many adhesion related genes among those genes that were unique to each genome. We also found that each isolate has at least one type 3 secretion system (T3SS), a known virulence factor, though not all identified T3SS belong to the same family nor are in syntenic genomic locations.

Conclusions

The *Providencia* species examined here are characterized by high degree of genomic similarity which will likely extend to other species and isolates within this genus. The presence of T3SS islands in all of the genomes reveal that their presence is not sufficient to indicate virulence towards *D. melanogaster*, since some of the T3SS-bearing isolates are known to cause little mortality. The variation in adhesion genes and the presence of T3SSs indicates that host cell adhesion is likely an important aspect of *Providencia* virulence.
**1. Background**

*Providencia* are ubiquitous Gram-negative bacteria in the family Enterobacteriaceae that are often opportunistic pathogens. They are commonly found to cause traveler’s diarrhea and urinary tract infections in humans but have also been isolated from more severe human infections such as meningitis [1–4]. They have been identified as part of the normal human gut flora and the genomes of some strains have been sequenced as part of the Human Microbiome Project [5]. Additionally, *Providencia* have been associated with numerous animals worldwide, including isolation from penguin feces in German zoos [6], sea turtles in the Mediterranean [7], shark mouths in Brazil [8], entomopathogenic nematodes globally [9], and snakes from Vietnam [10]. *Providencia* have also been found in association with insects such as blowflies [11], stable flies [12], Mexican fruit flies [13], and house flies [14]. *Providencia* have also been found in the guts and external environment of *Drosophila melanogaster* [15,16].

*Providencia* strains have been additionally isolated as infectious agents of *D. melanogaster* and have been shown to have distinct phenotypes including varied virulence towards *D. melanogaster* [17,18]. Two species, *Providencia sneebia* and *Providencia alcalifaciens*, were found to be highly virulent, causing 90-100% mortality in infected flies. Infections with the other two, *Providencia rettgeri* and *Providencia burhodogranariea*, caused only moderate mortality, with 30-40% of infected flies succumbing to the infection [18]. The more lethal bacteria tended to proliferate to higher densities in the fly, and triggered greater expression of antibacterial immune genes, with the exception of *P. sneebia*, which did not induce substantial antimicrobial peptide gene expression despite rapid and lethal proliferation [18].

In the present work, we have sequenced, annotated, and compared the draft genomes of the four species isolated from infections in wild *D. melanogaster: P. sneebia, P. rettgeri, P. 
burhodogranariea and P. alcalifaciens. We compared our sequences to draft genomes of four sequenced species of Providencia isolated from the human gut [5]. We sought specifically to identify the core Providencia genome and accessory genes, to establish which genes may be evolving under positive selection, and to annotate differences in gene content that contribute to physiological differences among the isolates.

2. Methods

2.1 Bacteria Strains Sequenced

The four bacteria strains that were sequenced in this study were isolated from the hemolymph of wild caught Drosophila melanogaster [18]. They are the Providencia sneebia Type strain (DSM 19967) [GenBank:AKKN00000000], Providencia rettgeri strain Dmel1 [GenBank:AJSB00000000], Providencia alcalifaciens strain Dmel2 [GenBank:AKKM00000000], and Providencia burhodogranariea Type strain (DSM 19968) [Genbank:AKKL00000000].

2.2 Genome Sequencing and Assembly of P. rettgeri and P. sneebia

Bacterial DNA was extracted using Puregene DNA Purification Kit (Gentra Systems Inc., Minneapolis, MN) according to the manufacturer’s directions for Gram-negative bacteria. The DNA was then sequenced using FLX Roche/454 Sequencing Technology at Cornell University’s Life Science Core Laboratory Center in Ithaca, NY.

P. sneebia and P. rettgeri were sequenced with approximately 500,000 reads of an average length of 250 bp, providing about 30X coverage for each genome. The sequences for each species were obtained from separate full-plate sequencing runs and independently
assembled using the LaserGene SeqMan (version 8) software with the manufacturer’s recommended parameters (Figure 3.1). An additional 180,000 reads were obtained from a 454 sequencing plate on which the DNA from *P. sneebia* and *P. rettgeri* was separated by a rubber gasket. During the sequencing process, this gasket leaked allowing a very small amount of reciprocal contamination. We did not want to discard these sequences entirely, but we also wanted to avoid any contaminating reads fouling our assemblies. Therefore our second assembly step was to assemble the reads from the half-plate to those contigs initially assembled with the uncontaminated full-plate reads using SeqMan (Figure 3.1). From this second step of assembly, we retained: (1) contigs that contained half-plate reads assembled to full-plate contigs, increasing the depth of those contigs, (2) contigs in which half-plate reads bridged previously separate contigs from the full-plate assemblies, allowing them to be stitched together, and (3) novel contigs containing only half-plate reads but with a coverage depth of 30X or greater. Contaminating sequences in the half-plate reads would presumably fail to map to full-plate assemblies or would result in low-coverage contigs, so we infer that the small number of molecules that leaked through the gasket have been effectively discarded. After the second round of assembly, the *P. sneebia* genome was mapped into 72 contigs and the *P. rettgeri* genome was mapped into 71 contigs.

As we were annotating the *P. sneebia* and *P. rettgeri* genome sequences (see “2.5 Annotation Methods” section below), we noticed several instances of sequential open reading frames (ORFs) that were annotated with the same predicted function and whose combined length equaled the size of genes with the same functional annotation in other bacteria. Closer inspection revealed that these instances were generally due to a stop codon or frameshift mutation that interrupted the ORF, causing it to be annotated as two genes with identical function. These
Figure 3.1. Flow chart illustrating steps taken in the assembly of the *P. sneebia* and *P. rettgeri* genomes.
inferred mutations tended to happen after short homopolymer runs. Individual reads varied in the lengths of these homopolymer sequences, and the contig assembly often did not reflect the most common sequence length among the reads. It is a known problem that Roche/454 Sequencing often results in errors in homopolymer run lengths [19]. To improve the accuracy of inferred homopolymer lengths, we re-aligned all of the Roche/454 sequencing reads to our assembled reference sequences using the program BWA [20] (Figure 3.1). The consensus homopolymer length from the BWA alignment was used to fix the assembled contigs before any further analysis was performed. This correction improved our gene annotations by eliminating sequencing errors that interrupted ORFs.

We aimed to improve the assemblies of our *P. sneebia* and *P. rettgeri* genomes by joining contigs through PCR followed by direct Sanger sequencing. However, the order and orientation of the contigs was unknown. We hypothesized that there would be synteny among the genomes of *Providencia* species and isolates as well as species in the closely related genus *Proteus*, which we could use to predict the order and orientations of the contigs in our assemblies (Figure 3.1). We used MUMmer (version 3.22) [21] to compare *P. sneebia* and *P. rettgeri* to the draft genomes of *Providencia rettgeri* DSM 1131 (283 contigs) [GenBank:ACCI00000000], *Providencia alcalifaciens* DSM 30120 (79 contigs) [GenBank:ABXW00000000], *Providencia stuartii* ATCC 25827 (120 contigs) [GenBank:ABJD00000000], and *Providencia rustigianii* DSM 4541 (127 contigs) [GenBank:ABXV00000000] as well as the completed genome of *Proteus mirabilis* strain HI4320 [GenBank:NC_010554.1]. Where two of our *P. rettgeri* or *P. sneebia* contigs had similarity to a single contig of one of the other genome sequences, we designed PCR primers to amplify across the inferred gap. Successful amplifications were sequenced by primer walking and the resultant sequences were used to bridge contigs in the
assemblies. PCR and sequencing methods are described below in the “2.4 PCR and Sanger Sequencing Methods” section. We found that designing the primers inset about 900 bp from the contig breakpoints helped to ensure specificity in amplification, especially because repetitive sequences in the genome can be the cause of contig breaks in genome assemblies. Using this method, we reduced the number of contigs in the *P. sneebia* assembly from 72 to 67 and in the *P. rettgeri* assembly from 71 to 64.

To further connect the *P. sneebia* and *P. rettgeri* assemblies, we contracted the MapIt optical mapping service from OpGen, Inc. (Gaithersburg, MD) and analyzed the resulting data using their program MapSolver (version 2.1.1) (Figure 3.1). An *in silico* digestion of our contigs allowed them to be oriented onto an *in vitro* restriction digestion map of each bacterium’s physical genome. This ordered and oriented the contigs and allowed us to identify those contigs that comprised the majority of each genome. We used the optical map to identify physically consecutive contigs, then designed primers for PCR and Sanger sequencing to close most of the remaining gaps. The optical map also indicated a small number of computational misassemblies and allowed them to be fixed. After optical mapping and final gap closing, our draft genome sequences were assembled into 14 contigs for *P. sneebia* and 9 contigs for *P. rettgeri*.

2.3 Genome Sequencing and Assembly of *P. alcalifaciens* and *P. burhodogranariea*

*P. alcalifaciens* and *P. burhodogranariea* were sequenced by paired-end 454 sequencing. Libraries were constructed for each bacterium with an approximately 3 kb insert size, and roughly 1 million paired-end sequence reads of an average length of 250 bases were collected. The paired-end reads were assembled using Roche/454’s Newbler Assembler (version 2.5.3),
which resulted in 15 scaffolds for *P. alcalifaciens* and 8 scaffolds for *P. burhodogranariea*, sequenced to roughly 35X coverage.

2.4 PCR and Sanger Sequencing Methods

PCR primers for gap closing were designed either using Primer3 [22] or with a primer design function within SeqMan. PCRs were performed using a genomic DNA template with a final concentration of 1.2 ng/µl in each PCR reaction volume.

When the size of the expected product was unknown or was expected to be less than 5 kb, the PCR was done with Taq polymerase (New England Biolabs, Beverly, MA) with an annealing temperature gradient ranging from 2°C higher to 2°C lower than the melting temperature of the primers. PCR cycling parameters were as follows: (1) 2 minutes at 95°C, (2) 30 seconds at 95°C, (3) 30 seconds at annealing temperature gradient, (4) 1 minute at 72°C, (5) repeat steps 2-4 for 34 more cycles, (6) 5 minutes at 72°C. 3.5 µl of each PCR product was prepared for sequencing by treatment with 5 units of Exonuclease I (USB Corp., Cleveland, OH) and 0.5 units of shrimp alkaline phosphatase (USB Corp., Cleveland, OH) at 37°C for one hour before heat-killing the enzymes at 65°C for 15 minutes. PCR products were then directly sequenced using ABI BigDye Terminator (Applied Biosystems, Foster City, CA) according to the manufacturer’s directions.

PCR for products with an expected size greater than 5 kb was done using high fidelity iProof polymerase (Bio-Rad, Hercules, CA). Annealing temperatures and extension times were determined using manufacture’s suggested methods. The PCR cycling parameters were: (1) 30 seconds at 98°C, (2) 7 seconds at 98°C, (3) 20 seconds at appropriate annealing temperature, (4) appropriate extension time at 72°C, (5) repeat steps 2-4 for 29 more cycles, (6) 7 minutes at
72°C. Products were prepared for sequencing with PCR purification clean up columns (Invitrogen, Calsbad, CA) before being sequenced directly.

2.5 Annotation Methods

Genomic open reading frames were determined and annotated using the RAST Server (version 4) [23]. Gene ontology terms (GO terms) were assigned to the ORFs identified by RAST using Blast2GO (version 2.5) [24]. Fisher’s Exact Test for enriched GO categories was done within Blast2GO using a \( p \)-value cut off of 0.05 after adjusting for a false discovery rate (FDR) of 0.05 for multiple testing.

2.6 Plasmid Identifications and Analysis

The circular DNA structure of plasmids means that they will appear to be arbitrary broken when forming linear contigs during assembly of sequencing reads. We tested all potential plasmid contigs for a circular physical structure by designing PCR primers approximately 500 bases from the ends of the contig facing outward off each end of the contig. This primer design means that a product would be formed only if the ends of the contig were connected in the physical DNA. Any PCR product amplified from such primers was then sequenced with Sanger sequencing to confirm that the sequence supported a circular physical arrangement of the sequence. PCR reactions and Sanger sequencing was done as described above in “PCR and Sanger sequencing methods”.

Putative plasmids were identified in multiple ways. We speculated that one \( P. \) rettgeri contig might be a plasmid because it had substantially higher depth of coverage than other contigs, and when we compared the contig to itself using MUMmer, we found that it was
composed of the same sequence repeated multiple times. We hypothesized that the contig might actually represent a completely sequenced, high copy number plasmid, and that the circular shape of the physical DNA sequence was resulting in a tandem repeat of the sequence in the *in silico* assembly. PCR and Sanger sequencing confirmed that this contig is a plasmid.

To more systematically assess whether contigs from the assemblies were plasmids, we looked for contigs with identical sequence present at both ends. We hypothesized that the arbitrary break point of the physical circular structure to form a linear contig could result in identical sequence at each end of the contig. We used MUMmer to compare contigs to themselves. For *P. sneebia* and *P. rettgeri*, we examined all contigs which did not align to the optical map since we do not expect plasmids to map to the chromosomal genome. We identified three *P. sneebia* plasmids using this method, all of which were confirmed by PCR and Sanger sequencing. We identified no additional *P. rettgeri* contigs as putative plasmids using this approach. For *P. alcalifaciens* and *P. burhodogranariea*, we examined every scaffolds smaller than 6 kb in length, but none contained the same sequence at both ends of the contig.

When examining the synteny of the genomes (see “2.10 Synteny and Regional Comparisons” section below), we noticed that some of the contigs of *P. alcalifaciens* had no similarity to sequences in any of the other genomes. We hypothesized that these contigs could be plasmids that are unique to *P. alcalifaciens*. We tested four contigs by PCR and Sanger sequencing. One contig was confirmed to be a plasmid while the other three did not produce PCR products and therefore showed no evidence of being plasmids.

All putative plasmids were compared to each other using MUMmer. In order to determine whether our confirmed plasmids or previously sequenced *Providencia* plasmids [25–27] were present but undetected in the remaining sequences from this study, we constructed a
BLAST database of all of the reads from the Roche/454 sequencing of each individual species and searched for reads matching each *Providencia* plasmid using BLAST+ (version 2.2.25). Four previously identified *Providencia* plasmids were used as query sequences: pDIJ09-518a [GenBank:HQ834472.1], pGHS09-09a [GenBank:HQ834473.1], pMR0211 [GenBank:JN687470.1], and R7K [Genbank:NC_010643.1].

2.7 Identification of Orthologs

Orthologous genes were identified as shared among three different sets of bacteria: (1) the strains of *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* sequenced in this study; (2) the strains of *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* sequenced in this study plus *Proteus mirabilis* strain HI4320; (3) the strains of *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* sequenced in this study plus the strains of *P. rettgeri*, *P. alcalifaciens*, *P. stuartii*, and *P. rustigianii* sequenced as part of the Human Microbiome Project [5]. Orthologous gene clusters were identified using OrthoMCL (version 2.0.2) [28]. BLAST results used within OrthoMCL were performed with an e-value cut off of $10^{-10}$. The output from OrthoMCL was parsed using custom Python scripts.

2.8 Alignments of Orthologs

Orthologous gene clusters found among the strains sequenced in this study and *Proteus mirabilis* were aligned for phylogenetic analysis (see “2.9 Phylogenetic Analysis” section below). Those orthologs of the strains sequenced in this study only were aligned for use in the recombination and positive selection analyses (see “2.11 Recombination Analysis” and “2.12 Positive Selection Analysis” sections below). Only clusters of clear 1:1:1:1:1 or 1:1:1:1
orthology were retained, respectively. Alignments of the protein translation of the genes were done using ClustalW (version 2.1) [29] followed by back-translation to the nucleotide alignment using PAL2NAL (version 13) [30]. Alignments were visually inspected and poor alignments were removed as follows. We eliminated alignments where the difference in amino acid identity between the most-similar and least-similar pairs of species were greater than 40% out of concern that these might not be true orthologs. We also excluded alignments that had both an average protein identity that was less than 60% and a difference between the highest and lowest pair-wise protein identities greater than 20%. Those alignments that had an average protein identity of less than 70% were examined by hand to ensure proper alignment.

2.9 Phylogenetic Analysis

The alignments of all 1689 ortholog clusters that included the *Proteus mirabilis* outgroup were concatenated using FASconCAT (version 1) [31]. RAxML (version 7.2.8) [32] was used to construct the phylogenetic trees for the concatenation of all orthologous genes as well as for each individual orthologous cluster. *Proteus mirabilis* was set as an outgroup.

2.10 Synteny and Regional Comparisons

Synteny among genomes was examined using Mauve (version 2.3.1) [33], Artemis Comparison Tool (version 1) [34], and MUMmer. Comparisons of particular regions of the genomes were done using EasyFig (version 1.2) [35].
2.11 Recombination Analysis

Evidence for recombination was examined by executing the programs GENECONV which implements the Sawyer method [36] and PhiPack [37]. GENECONV was run using the default settings, which estimates p-values on 10,000 permutations of each alignment. PhiPack runs 3 separate tests: Pairwise Homoplasy Index, Maximum $\chi^2$, and Neighbor Similarity Score. The Pairwise Homoplasy Index was calculated on a window size of 50 while Maximum $\chi^2$ was calculated on a window that is $2/3$ the size of the polymorphic sites. We did 1000 permutations in PhiPack to calculate each p-value. The p-values of all tests were corrected for multiple testing using the program Q-value [38] with a FDR of 10%.

2.12 Positive Selection Analysis

Positive selection analysis was done using PAML (version 4.4) [39] on the 1937 orthologous clusters which are *Providencia*-specific. Site-model tests were implemented using codeml to compare model M8a (beta+\(\omega=1\)) to M8 (beta+\(\omega\)) [39,40]. The log-likelihoods from each test were compared in a likelihood ratio test assuming a $\chi^2$ distribution of the test statistic. We corrected for multiple testing using a q-value cut off which was calculated with the program Q-value [38] using a FDR of 20% [41,42].

2.13 Phage Identification

Phage genes were identified and classified using PHAST [43].
3. Results and Discussion

3.1 Basic Genome Information

The genomes of *Providencia sneebia*, *Providencia rettgeri*, *Providencia alcalifaciens* and *Providencia burhodogranariea* were sequenced in this study and assembled into 14, 9, 15 and 8 contigs or scaffolds, respectively (Table 3.1, Figures 3.2-3.5). The sequenced isolates were obtained from the hemolymph of wild *Drosophila melanogaster* and therefore will be referred to collectively as Dmel isolates in this paper. The summed contig or scaffold lengths of these assemblies vary from 4.5 Mb to 3.5 Mb, with *P. burhodogranariea* having the largest genome and *P. sneebia* having the smallest (Table 3.1). Because we know the sizes of the *P. sneebia* and *P. rettgeri* physical genomes from optical maps, we can discern that our *P. sneebia* assembly is missing approximately 300 kb of sequence and our *P. rettgeri* assembly is missing approximately 100 kb of sequences (Table 3.1). Based on the average gene size for each genome, we estimate that our annotated gene sets are missing roughly 275 out of 3750 for *P. sneebia* and 107 genes out of 4650 total for *P. rettgeri* (Table 3.1). Thus, we estimate that we have assembled approximately 93% of the *P. sneebia* coding genome and 98% of the *P. rettgeri* coding genome. Both *P. alcalifaciens* and *P. burhodogranariea* were sequenced with paired-end reads, resulting in scaffolds with sequence gaps of known sizes. These gaps can break up open reading frames, resulting in the absence of some genes from our inferred annotations. Since we did not optically map *P. alcalifaciens* or *P. burhodogranariea*, we do not know the precise sizes of their physical genomes. Because of the small assembly gaps, our annotations bear the caveat that a small number of genes may be inferred as absent when they are actually present in the physical genome.
Table 3.1. Basic genomic information of all four *D. melanogaster* isolated *Providencia*.

<table>
<thead>
<tr>
<th>Species</th>
<th>Strain</th>
<th>DSM¹</th>
<th>Sequenced size²</th>
<th>Physical size³</th>
<th># of contigs or scaffolds</th>
<th>Average GC %</th>
<th># of genes</th>
<th>Est. # of missing genes⁴</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Providencia sneebia</em></td>
<td>Type</td>
<td>19967</td>
<td>chromosome</td>
<td>3.5 Mb</td>
<td>3.8 Mb</td>
<td>14</td>
<td>38.08</td>
<td>3482</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>pPSN1</td>
<td>10787 bp</td>
<td>10787 bp</td>
<td>1</td>
<td>33.50</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>pPSN2</td>
<td>7592 bp</td>
<td>7592 bp</td>
<td>1</td>
<td>35.05</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>pPSN3</td>
<td>4321 bp</td>
<td>4321 bp</td>
<td>1</td>
<td>31.50</td>
<td>4</td>
</tr>
<tr>
<td><em>Providencia rettgeri</em></td>
<td>Dmel 1</td>
<td></td>
<td>chromosome</td>
<td>4.2 Mb</td>
<td>4.3 Mb</td>
<td>9</td>
<td>40.20</td>
<td>4532</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>pPRET1</td>
<td>5567 bp</td>
<td>5567 bp</td>
<td>1</td>
<td>40.81</td>
<td>6</td>
</tr>
<tr>
<td><em>Providencia alcalifaciens</em></td>
<td>Dmel 2</td>
<td></td>
<td>chromosome</td>
<td>4.2 Mb</td>
<td>unk</td>
<td>15</td>
<td>41.17</td>
<td>3900</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>pPALC1</td>
<td>14114 bp</td>
<td>14114 bp</td>
<td>1</td>
<td>37.57</td>
<td>17</td>
</tr>
<tr>
<td><em>Providencia burhodogranariea</em></td>
<td>Type</td>
<td>19968</td>
<td>chromosome</td>
<td>4.5 Mb</td>
<td>unk</td>
<td>8</td>
<td>38.23</td>
<td>3985</td>
</tr>
</tbody>
</table>

¹Strain number in DSM Culture Collection  
²Sum of the contigs or scaffolds lengths  
³Size of the physical chromosome as determined by optical maps. “unk” indicates unknown physical size.  
⁴Estimated from the known length of sequence missing from these assemblies divided by the average gene length. “unk” indicates that the number of missing genes cannot be calculated due to an unknown physical size.
Figure 3.2. Circular maps of the *Providencia sneebia* genome and plasmids. (A) *P. sneebia* genome. The contigs are ordered and oriented as they are in the physical genome based on an optical map, including the sizes of the gaps between contigs. Rings from the outermost to the center: 1) genes on the forward strand (blue), 2) genes on the reverse strand (blue), 3) tRNA and rRNA genes (black), 4) genes unique to *P. sneebia* when comparing all 8 Dmel and HMP *Providencia* genomes (orange), 5) individual assembly contigs (alternating shades of grey), 6) GC skew. (B) pPSN1. (C) pPSN2. (D) pPSN3. All plasmids have the genes on the forward strand on the outermost ring and genes on the reverse strand on the inner ring.
Figure 3.3. Circular maps of the *Providencia rettgeri* genome and plasmid. (A) *P. rettgeri* genome. The contigs are ordered and oriented as they are in the physical genome based on an optical map, including the sizes of the gaps between contigs. Rings from the outermost to the center: 1) genes on the forward strand (red), 2) genes on the reverse strand (red), 3) tRNA and rRNA genes (black), 4) genes unique to *P. rettgeri* when comparing all 8 Dmel and HMP *Providencia* genomes (orange), 5) individual assembly contigs (alternating shades of grey), 6) GC skew. (B) pPRET1. The plasmid has the genes on the forward strand on the outermost ring and genes on the reverse strand on the inner ring.
Figure 3.4. Maps of the *Providencia alcalifaciens* genome and plasmid. (A) *P. alcalifaciens* genome. The contigs are ordered and oriented for maximum synteny with *P. rettgeri*. The contigs are not positioned in a complete circle because the order and orientation of the contigs are not empirically known. The size of the gaps between contigs is unknown. Rings from the outermost to the center: 1) genes on the forward strand (purple), 2) genes on the reverse strand (purple), 3) tRNA and rRNA genes (black), 4) genes unique to *P. alcalifaciens* when comparing all 8 Dmel and HMP *Providencia* genomes (orange), 5) individual contigs (alternating shades of grey), 6) GC skew. (B) pPACL1. The plasmid has the genes on the forward strand on the outermost ring and genes on the reverse strand on the inner ring.
Figure 3.5. Map of the *Providencia burhodogranariea* genome. The contigs are ordered and oriented for maximum synteny with *P. rettgeri*. The contigs are not positioned in a complete circle because the order and orientation of the contigs are not empirically known. The size of the gaps between contigs is unknown. Rings from the outermost to the center: 1) genes on the forward strand (yellow), 2) genes on the reverse strand (yellow), 3) tRNA and rRNA genes (black), 4) genes unique to *P. burhodogranariea* when comparing all 8 Dmel and HMP *Providencia* genomes (orange), 5) individual contigs (alternating shades of grey), 6) GC skew.
3.2 Plasmids

We found 3 plasmids in *P. sneebia*, 1 in *P. rettgeri*, 1 in *P. alcalifaciens* and none in *P. burhodogranariea* (Table 3.1; Figures 3.2-3.5). We compared these plasmids to each other and found them to have no similarity in gene content. We also checked whether any of these plasmids were undetected in the sequencing reads of the Dmel isolates other than the one that each plasmid was assembled from. We found that while some plasmid genes were present on the chromosomes of other species, the complete plasmids were found only in the species from which they were initially recovered. Four plasmids have been isolated and sequenced by other groups studying other *Providencia* isolates [25–27], but we did not find any of these plasmids in the sequencing reads of our isolates. None of our novel *Providencia* plasmids contain genes with functional annotations that lead to a clear functional designation for the overall plasmid, such as “virulence” or “antibiotic resistance.” Our findings indicate that plasmids found in *Providencia* vary considerably in their identity, conservation, and probable functions.

3.3 Genomic Synteny

The optical maps of *P. sneebia* and *P. rettgeri* allowed us to order and orient the contigs as they are found in the physical chromosome for those two assemblies. The contigs of *P. burhodogranariea* and *P. alcalifaciens* were ordered and oriented so that they were as similar to the *P. rettgeri* and *P. sneebia* genomic orientations as possible, assuming the most parsimonious evolution of genome arrangements (Figure 3.6). It is in principle possible that any of the *P. burhodogranariea* and *P. alcalifaciens* contigs could be inverted or rearranged relative to their positions on our comparative syntenic plot, but only if the rearrangement breakpoints lie at contig breakpoints. While there are many small rearrangements found among the genomes, there
Figure 3.6. Alignments of the protein translations of the whole genomes of all four *Providencia* species isolated from *D. melanogaster*. The contigs of *P. sneebia* and *P. rettgeri* were ordered and oriented as they are in the physical genome based on optical maps made of each genome. *P. alcalificiens* and *P. burhodogranariea* contigs were ordered and oriented for maximum synteny with *P. rettgeri*, therefore assuming parsimony in the number of genomic arrangements. Similarity was calculated using promer function in MUMmer [21]. Red dots represent similar sequence in the same orientation in each genome pair while blue indicates that the similarity is in the opposite orientations in the genome pairs.
are only two large inversions apparent across the four species. Both inversions are in *P. sneebia* relative to the other genomes (Figure 3.6). The ends of the largest *P. sneebia* inversion, which is about 800 kb in length, fall within single contigs of both *P. burhodogranariea* and *P. alcalifaciens*, supporting the hypothesis that the inversion is derived in and unique to *P. sneebia*.

### 3.4 Phylogeny

We determined an overall phylogenetic relationship from a concatenated alignment of the single-copy orthologs shared by all four Dmel *Providencia* sequenced here, rooted with *Proteus mirabilis* (Figure 3.7). This tree is based on 1689 single copy orthologs shared among these five species. The phylogenetic tree indicates that *P. sneebia* and *P. burhodogranariea* share a common ancestor before either of them share a common ancestor with the *P. rettgeri* and *P. alcalifaciens* species pair (Figure 3.7). This is consistent with the phylogenetic relationships proposed previously based on the 16S rRNA gene and five housekeeping genes [17].

### 3.5 Orthologs and Unique Genes

We wanted to know how much gene content is shared among the Dmel *Providencia* isolates so we used a BLAST-based method to find all orthologous gene clusters. There are a total of 3644 orthologous clusters containing between 10 and 2 genes, as well as one exceptional cluster containing 20 fimbrial-related usher genes with at least 3 genes originating from each genome. Fimbrial-related usher proteins chaperone other proteins to the bacterial cell surface to form a proteinaceous extension involved in surface adhesion [44]. We found that 3293 genes are present as single copies in each genome, or 90% of the total ortholog clusters. The majority of these clusters, 1983, comprise the core genome of these *Providencia* isolates (Figure 3.8),
Figure 3.7. Phylogenetic relationships of all four *Providencia* species isolated from *D. melanogaster* and *Proteus mirabilis*. This phylogeny is constructed from a concatenation of 1689 orthologous genes shared by all five species and was inferred using maximum likelihood methods within the program RAxML [32]. The scale bar indicates number of substitutions per site. Each node of the tree is supported by a bootstrap value of 100. Above the line for each bacterial lineage is the number of inferred gene gains and losses on that lineage. Losses are preceded by (-) and gains by (+).
Figure 3.8. Shared and unique gene totals among all four *Providencia* species isolated from *D. melanogaster*. Numbers are the gene counts within each sector of the Venn diagram. Orthologous genes were determined with OrthoMCL [28].
meaning that the genes are shared as single-copy orthologs across all four sequenced species.

The core genome is 49-62% of the total genes in each genome, revealing that the species have a substantially homogeneous gene content. The next largest group of orthologous clusters is that which contains genes present in *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* but absent from *P. sneebia* (Figure 3.8). This is consistent with these three genomes each being almost one megabase bigger than that of *P. sneebia*. Given that *P. sneebia* shares a common ancestor with *P. burhodogranariea* before either shares a common ancestor with the more distantly related *P. rettgeri* and *P. alcalifaciens* (Figure 3.7), the absence of these genes in *P. sneebia* appears to reflect genome reduction in *P. sneebia*. To specifically examine putative gene loss, we determined the number of orthologous clusters that were missing a gene from only 1 of the 4 genomes (Figure 3.7 and 3.8). While *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* have between 74 and 89 genes that are apparently lost in their lineages, *P. sneebia* appears to have specifically lost 398 genes. As revealed in the analysis of genomic synteny above, differences in gene number and genome sizes generally are the result of small duplications, deletions, or insertions. In particular, the missing genes in *P. sneebia* do not result from large block deletions but instead arise from many small deletions eliminating individual genes distributed around the genome (Figure 3.6).

We found 137 orthologous clusters which contained paralogous duplications unique to a single genome. The majority of these clusters contain genes that are related to mobile elements or phages suggesting that there are families of transposons or phages that are specific to individual species. Most other clusters of species-specific paralog groups were annotated as hypothetical proteins.
We were able to determine genes unique to each genome by identifying those genes that were not assigned to any orthologous cluster (Figure 3.7 and 3.8). The genomes varied in the absolute number of unique genes, with the smallest genome, that of *P. sneebia*, having the fewest. Despite their variation in number among the genomes, unique single-copy genes represent 15-19% of their total genome content for each species.

The genes unique to each bacterium were tested for enrichment of gene ontology (GO) terms compared to those genes found in either the Dmel *Providencia* core genome or the individual whole genome from which the unique genes were drawn (Figure 3.9). GO terms enriched among the genes unique to each species were often related to interactions with phage or bacteria, including genes encoding phage lysozymes, bacteriocins, and restriction enzymes. This strongly suggests that these *Providencia* have acquired or developed different ways to deal with varied genome parasites and competitor organisms. There may also imply variation in the phage or bacteria to which these *Providencia* are most often exposed.

Based on GO annotation, the genes unique to *P. rettgeri* are enriched for those involved in rhamnose metabolic process compared to both the core and whole genome. While the particular strain of *P. rettgeri* we sequenced has not been tested for its ability to metabolize rhamnose, the type strain of *P. rettgeri* has been shown to metabolize this sugar while the same strains of *P. sneebia* and *P. burhodogranariea* sequenced in this study were unable to [17]. The type strain of *P. alcalifaciens* has also been tested for its ability to metabolize rhamnose, but the data were ambiguous [17]. The collective data are suggestive that rhamnose metabolism may be unique to *P. rettgeri* among these strains of *Providencia*, although previous work has shown that not all strains of *P. rettgeri* are able to metabolize rhamnose [45].
Figure 3.9. GO terms enriched in the unique genes of each species compared to the full set of Providencia isolated from *D. melanogaster*. GO terms were assigned and calculation of enrichment was done using a Fisher’s Exact Test using Blast2GO [24]. GO terms have been collapsed to only the most specific child term when multiple terms described the same group of genes.
All four species have genes with the GO term “pilus” significantly overrepresented in their unique genes compared to the core genome, although the absolute number of these genes varies for each genome (Figure 3.9). Pili are protein structures that extend from the surface of bacterial cells to allow the bacteria to adhere to substrates [44]. The majority of the unique genes given the pilus GO term are annotated as fimbrial proteins, which are the proteins that constitute the pilus structure [44]. The genic diversity indicates a high amount of variation in these fimbrial proteins. Pilus related proteins are often antigenic, so this variation in pilus protein could be a result of pressure to avoid host immune responses [46]. Alternatively, the distinction in fimbrial proteins could be due to variation among the species in adherence to specific surfaces [46]. As mentioned above, the largest orthologous gene cluster shared among the sequenced Providencia consisted of fimbrial-related usher proteins. These observations in combination show conservation of the genes which function to form the pilus but diversity of the genes encoding the proteins of the physical pilus structure itself.

3.6 Recombination and Positive Selection

Positive selection and recombination are two primary forces in bacterial evolution. Recombination rates have been found to vary widely in bacteria and it has been hypothesized that generalist bacteria or those in the process of adapting to new environments have higher rates of recombination [47]. These Providencia are closely related and were isolated from similar environments, so it is possible that there would have been opportunity for recombination among them. We examined 1937 orthologs in the core genome of the Dmel Providencia species for recombination, which is slightly reduced from the total 1983 core genome orthologs shown in Figure 3.8 due to the removal of comparatively poor sequence alignments (see Methods section
Among the 1937 orthologs examined, 781 orthologous clusters that showed evidence for recombination. The genes belonging to clusters exhibiting recombination are evenly distributed around the physical genomes of each bacteria, and we do not find evidence for hotspots of recombination.

We used the program PAML (Yang, 2007) to test for evidence of positive selection in the *Providencia* core genome, excluding genes that showed evidence of recombination since these violate the assumptions of the tests in PAML [48]. This left 1156 orthologous clusters in the core genome shared by the four Dmel *Providencia* species isolates. We used PAML to compare the likelihood of a model which does not allow for positive selection, termed model M8a, to a model that does allow for selection at various sites along the gene, model M8 [39,40]. We found 21 genes that yielded nominal \( p \)-values of less than 0.05, indicating that the model allowing for selection fit the data significantly better than the neutral model (Table 3.2). However, none of these 21 genes remained significant after application of a FDR of 20% [38]. Our selection test is extremely underpowered within *Providencia* given the small number of species examined and their close phylogenetic relationship to one another. We considered running the same site model tests on the four Dmel *Providencia* species plus *Proteus mirabilis*, but we there are too many synonymous changes on the lineage leading to *Proteus mirabilis* for the tests to be conducted appropriately. We present some genes that show nominal evidence for positive selection below, but stress that the selection results are provisional and further investigation into the biological function and adaptive significance of these genes is warranted.

One gene exhibiting evidence for positive selection is the *TolC* precursor \( (p = 0.003) \). The *TolC* gene has also been found to be under positive selection in *E. coli* [49]. This gene encodes an outer membrane protein that is part of a transporter system which transports toxins or
Table 3.2. Orthologous gene clusters with evidence for positive selection. *P*-values were calculated using codeml within PAML [39] by comparing models M8a and M8 and are uncorrected for multiple testing.

<table>
<thead>
<tr>
<th>Annotation</th>
<th>SEED Subsystem1</th>
<th>EC2</th>
<th><em>p</em>-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>homocysteine methyltransferase</td>
<td>Methionine biosynthesis</td>
<td>2.1.1.14</td>
<td>0.000496</td>
</tr>
<tr>
<td>Xanthine/uracil/thiamine/ascorbate permease family protein</td>
<td>Purine utilization</td>
<td>-</td>
<td>0.002595</td>
</tr>
<tr>
<td>ToLC precursor</td>
<td>Multidrug Efflux pumps</td>
<td>-</td>
<td>0.003340</td>
</tr>
<tr>
<td>Acetate permase ActP</td>
<td>acetogenesis from pyruvate</td>
<td>-</td>
<td>0.003703</td>
</tr>
<tr>
<td>hypothetical protein with DUF177</td>
<td>-</td>
<td>-</td>
<td>0.007188</td>
</tr>
<tr>
<td>2-octaprenyl-6-methoxyphenol hydroxylase</td>
<td>Ubiquinone Biosynthesis</td>
<td>1.14.13.-</td>
<td>0.009478</td>
</tr>
<tr>
<td>UDP-N-acetylglucosamine</td>
<td>-</td>
<td>2.4.2.227</td>
<td>0.010464</td>
</tr>
<tr>
<td>D-tyrosyl-tRNA(Tyr) deacylase</td>
<td>Stringent Response</td>
<td>-</td>
<td>0.014392</td>
</tr>
<tr>
<td>Sulfate transport system permease protein CysT</td>
<td>Cysteine Biosynthesis</td>
<td>-</td>
<td>0.014526</td>
</tr>
<tr>
<td>yihD</td>
<td>-</td>
<td>-</td>
<td>0.015702</td>
</tr>
<tr>
<td>antibiotic biosynthesis monooxygenonase</td>
<td>-</td>
<td>-</td>
<td>0.015830</td>
</tr>
<tr>
<td>4-diphosphocytidyl-2-C-methyl-Derythritol kinase</td>
<td>Isoprenoid Biosynthesis</td>
<td>2.7.1.148</td>
<td>0.018713</td>
</tr>
<tr>
<td>Acyl-phosphate:glycerol-3-phosphate O-acyltransferase P1sY</td>
<td>Glycerolipid and Glycerophospholipid Metabolism</td>
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</tr>
<tr>
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<td>Biotin biosynthesis</td>
<td>6.3.3.3</td>
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<tr>
<td>LSU ribosomal protein L9p</td>
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<td>0.029278</td>
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<tr>
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<td>Ribosomal biogenesis, RNA processing &amp; degradation</td>
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<tr>
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<td>protein Glycerolipid and Glycerophospholipid Metabolism</td>
<td>-</td>
<td>0.031606</td>
</tr>
<tr>
<td>Predicted Fe-S oxidoreductase</td>
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<td>-</td>
<td>0.041952</td>
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<tr>
<td>Flagellar transcription activator FlhC</td>
<td>Flagellum</td>
<td>-</td>
<td>0.044270</td>
</tr>
<tr>
<td>Magnesium transporter</td>
<td>-</td>
<td>-</td>
<td>0.048181</td>
</tr>
</tbody>
</table>

1The subsystem and protein function designation were determined through annotation with RAST [23] with the exception of the proteins annotated as antibiotic biosynthesis monooxygenonase which was designated by Blast2GO [24].

2Enzyme Commission number
antibiotics out of the cell [50,51]. It is possible that different proteins or varying amounts of proteins are transported by TolC in the various Providencia species, in which case selection may act on TolC to optimize interaction with different partners or vary secretion amounts.

Two different orthologous gene clusters showing weaker signal for positive selection were annotated as being involved in glycerolipid and glycerophospholipid metabolism ($p = 0.022$ and $p = 0.032$). This subsystem is involved in making lipids that are transported to the bacterial cell surface. The signatures of positive selection suggest possible adaptation in lipid and protein structures on the surface of these Providencia species.

We found the protein FlhC to also show weak evidence for positive selection ($p = 0.044$). This protein forms a heterodimer with FlhD to become the master regulatory complex of flagellar protein production, and these proteins have also been shown to regulate the expression of many other genes, including virulence genes [52–54].

### 3.7 Similarity of Dmel Isolates to Providencia Isolates from the Human Gut

Four Providencia isolated from human feces have been sequenced as part of the Human Microbiome Project [5]. These included isolates of P. rettgeri and P. alcalifaciens as well as isolates of the species Providencia stuartii and Providencia rustigianii. These isolates will be referred to as the HMP isolates, to distinguish them from the Dmel isolates sequenced in our study.

We hypothesized that the four HMP isolates might have specialized genes to facilitate living in the human gut while the Dmel isolates would have genes enabling infection of D. melanogaster and other insects. To test this hypothesis, we extracted all orthologs of the eight genomes, yielding 4926 total ortholog clusters. Only 177 orthologous clusters, 3.5% of the total,
contained no genes from any HMP isolates and were therefore specific to the Dmel isolates. None of these contained genes found in all four Dmel isolates, meaning there are no genes that are exclusive to and universal in these isolates. Similarly, only 354, or 7.2% of the total ortholog clusters contain no genes from any Dmel isolate and therefore were exclusive to the HMP isolates. The majority of the HMP-specific ortholog clusters, 235 clusters, only contain two orthologous genes. Eleven of the HMP-specific ortholog clusters are found in all four of the isolates. Eight of these are assigned annotations that relate to phage activity and are physically co-localized in their respective genome. It is therefore unlikely that they are required for colonization of the human gut per se, but instead they probably reflect the shared phage pressure in the common environment. These isolates did not all originate from the same human, suggesting that the relevant phage may be pervasive in human guts. These phage genes were identified as belonging to Fels-2 in \textit{P. alcalifaciens} HMP and the Myoviridae prophages PSP3 in \textit{P. stuartii} HMP, \textit{P. rettgeri} HMP, and \textit{P. rustigianii} HMP. These two prophages are closely related to each other [55]. Of the 354 HMP isolate specific orthologs and paralogs, 69 of them are found only in \textit{P. stuartii}. Most of these, 47, are annotated as hypothetical proteins. The total data indicate that there are no endogenous \textit{Providencia} genes that are specific to and ubiquitous in either isolation environment we examined, but we do find evidence that the bacteria are exposed to different phages in the respective environments.

The core genome derived from all eight \textit{Providencia} genome sequences contained 1925 genes, only very slightly fewer than the 1983 genes in the core genome of the Dmel \textit{Providencia} isolates. This gives us added certainty that the core genome of \textit{Providencia} as a whole is highly conserved.
The proportion of genes unique to any one of the HMP isolates is approximately the same as the unique gene complement in the Dmel strains. Unique genes represent 9% to 17% of the gene total for each of the isolates when all eight genomes are examined. Unsurprisingly, the HMP isolates of *P. alcalifaciens* and *P. rettgeri* share genes with only the Dmel isolates of these same species. Whereas 16% of the *P. alcalifaciens* Dmel genome consisted of unique genes when the Dmel isolates were considered alone, that value drops to 9% when the HMP isolates are also considered. The number of unique genes in *P. rettgeri* Dmel shows a similar decrease, from 19% of the total genes when Dmel isolates are considered in isolation to 10% after inclusion of the HMP isolates. However, these decreases in the count of unique genes are not only due to the addition of another isolate of the same species. Of the 800 genes initially considered unique to the Dmel isolate of *P. rettgeri*, 167 are found in the HMP isolates of *P. stuartii*, *P. alcalifaciens*, or *P. rustigianii*. Half of the decrease in apparently unique genes in *P. alcalifaciens* Dmel is due to orthologs in the HMP isolates of *P. stuartii*, *P. rettgeri* and *P. rustigianii*.

The unique, single copy genes for each of the eight genomes were assigned GO terms and then examined for GO term enrichment relative to either the whole genome or the core genome of all eight sequenced *Providencia* (Figure 3.10). Both the Dmel isolate of *P. burhodogranariea* and the HMP isolate of *P. alcalifaciens* had no GO terms enriched in their unique genes compared to either their respective whole genomes or the core genome. This suggests that the distinct genes acquired by these isolates have a wide variety of functions. There is also no shared enrichment for GO terms among all isolates originating from the human gut or among all those isolated from *D. melanogaster*, further emphasizing that there is no class of genes that tie the isolates together based on isolation environment. *P. rustigianii* HMP’s unique genes also are
Figure 3.10. GO terms enriched in the unique genes of each species when comparing the isolates collected from *D. melanogaster* (Dmel) to the isolates from the Human Microbiome Project (HMP). GO terms were assigned and calculation of enrichment was done using a Fisher’s Exact Test using Blast2GO [24]. GO terms have been collapsed to only the most specific child term when multiple terms described the same group of genes. No GO terms were enriched in *P. burhodogranaria* Dmel or *P. alcalifiaciens* HMP.
enriched in GO terms relating to the endoplasmic reticulum (ER) membrane compared to both
the core and *P. rustigianii* HMP whole genome (Figure 3.10). These proteins each have a
PGAP1-like domain, which is known to function in the ER, but since bacteria do not have an ER,
they are unlikely to be functioning the same way in *P. rustigianii*. Although this domain is found
in other bacteria, to our knowledge, no bacterial function has been determined for this domain.

GO categories related to transposons or transposition were enriched in several genomes,
suggesting that the genomes have unique transposable elements (Figure 3.10). Additionally, *P.
rustigianii* HMP unique genes have a number of GO terms related to viruses that are enriched
over the core and whole genome (Figure 3.10). These genes are present in multiple locations
throughout the genome and are either annotated as phage proteins or are surrounded by prophage
genes. This, again, emphasizes the variety of prophages and other mobile elements inserted in
these genomes.

As seen when considering the Dmel genomes alone, the HMP genomes are also enriched
for unique genes with the GO term “pilus” relative to the core genome (Figure 3.10). The
enrichment in the GO term “pilus” unexpectedly disappears from the Dmel isolates of *P.
burhodogranariea* and *P. alcalifaciens* after addition of the HMP isolates because some of the
genews with this GO term previously considered “unique” are additionally found in the HMP
isolates. *P. sneebia* and *P. rettgeri* retain enrichment of the pilus GO term as well as other GO
terms related to adhesion in their unique genes over the core genome.

### 3.8 Species Specific Genes

Two different isolates of both *P. rettgeri* and *P. alcalifaciens* have now been sequenced.
One isolate of each species was sequenced from infected wild *D. melanogaster* (Dmel) while the
other was isolated from human feces (HMP). There are 202 ortholog clusters that contain genes found only in the Dmel and HMP isolates of *P. rettgeri*, with six clusters containing more than two genes. The two isolates of *P. alcalifaciens* uniquely share 190 species-specific orthologous clusters. Unfortunately, most of these species-specific genes are annotated as hypothetical proteins so they do not lend any insight into biological distinction of these species from others in the *Providencia* genus. As described above for the individual isolates, genes annotated with the GO term “pilus” are enriched in genes specific to *P. alcalifaciens* relative to the *Providencia* core genome, and *P. rettgeri* is enriched for pilus genes and genes involved in rhamnose metabolic processes.

### 3.9 Type 3 Secretion Systems

A type 3 secretion system (T3SS) is a needle-like apparatus used by Gram-negative bacteria for injecting effector proteins into host cells [56]. The genes encoding the proteins of the needle machinery are physically clustered and may be acquired via horizontal gene transfer as a single “pathogenicity island”. Transcription of the genes encoding the T3SS machinery and secretion of the effector proteins is triggered by external signals indicating that the bacteria is in the infection environment [57]. The functions of the translocated effector proteins vary greatly among bacteria, and include toxins that kill the host cells and proteins that manipulate host cell cytoskeletal activity or other cell biology to the advantage of the bacterium. Effector proteins are not necessarily encoded for within the same pathogenicity island as the genes encoding the T3SS needle apparatus, and may be acquired and evolve independently of the machinery [58].

All four *Providencia* sequenced in this study have at least one T3SS island (Figure 3.11A). *P. sneebia* Dmel, *P. alcalifaciens* Dmel, and *P. rettgeri* Dmel all have a similar T3SS island, which
Figure 3.11. Similarity of Type 3 Secretion Systems (T3SS). (A) Alignment of the whole genomes of *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *P. burhodogranariea* with the locations of each T3SS marked by blue boxes. (B) Alignment of T3SS-1 and some surrounding genes in *P. sneebia*, *P. rettgeri*, *P. alcalifaciens*, and *Proteus mirabilis*. Since this island falls within the inversion in *P. sneebia’s* genome, the region is displayed as the reverse-complement for ease of viewing. (C) Alignment of T3SS-2 and some surrounding genes in *P. sneebia* and *P. burhodogranariea*. (D) Alignment of T3SS-1 and T3SS-2 and some surrounding genes in *P. sneebia*. Figures were made using EasyFig [35]. Lines connecting the sequence schematics indicate regions of similarity, with darker grey indicating greater similarity. Colored arrows indicate individual genes and their direction. The black boxes behind the genes indicate the approximate boundaries of the T3SS island based on gene annotation.
will be referred to as T3SS-1. These T3SS-1 islands are similar in sequence, gene content, gene orientation, and ATPase homology (Figure 3.11B). It is likely that this island was acquired prior to speciation of the sequenced Providencia as it is shared with Proteus mirabilis and the HMP Providencia isolates (Figure 3.11 A and B) and is found in the same syntenic region in all three Dmel genomes, although this position lies inside the P. sneebia inversion. The genomic region surrounding the T3SS-1 island is much less conserved than the genes of the island itself. Even though P. sneebia, P. alcalifaciens, and P. rettgeri all share this T3SS, these bacteria have been shown to vary in virulence towards D. melanogaster [18]. Some work has been done to characterize the T3SS-1 island of Proteus mirabilis during infection of the mouse ascending urinary tract, but disrupting the function of the secretion machinery had no effect on the bacteria’s ability to colonize the mouse [59].

There is a second T3SS island present in the P. sneebia genome that is also found in P. burhodogranariea. We refer to this as T3SS-2. The T3SS-2 island is not found in any of the Providencia HMP isolates. Although T3SS-2 is not syntenically conserved in its genomic location between P. sneebia and P. burhodogranariea, the sequences and gene contents of the islands from the two species are much similar to each other than they are to T3SS-1 of P. sneebia, P. alcalifaciens, and P. rettgeri, or even than the respective T3SS-1 islands are to each other (Figure 3.11). It would be equally parsimonious to conclude that the P. sneebia and P. burhodogranariea genomes have separately acquired T3SS-2 islands or to infer a single acquisition by their common ancestor followed by a subsequent relocation of the island in one of the genomes.

The ATPases of T3SS-1 and T3SS-2 belong to different families. The T3SS-1 ATPase belongs to the Inv-Mxi-Spa ATPase family, which generally functions in cell invasion [58]. The
ATPase of T3SS-2 belongs to the Ysc ATPase family, which is commonly found in extracellular pathogens [58]. Since *P. sneebia* carries both T3SS islands, it might be capable of functioning as both an extracellular pathogen and an intracellular one depending on infection context. However, the ATPase of T3SS-1 in our sequenced *P. sneebia* isolate contains a premature stop codon at codon 281 of the 420 codon gene, which likely abolishes the function of T3SS-1 in this isolate. We confirmed this stop codon by PCR and Sanger sequencing. We additionally sequenced the T3SS-1 ATPase from 8 additional isolates of *P. sneebia* originating from the hemolymph of wild caught *D. melanogaster* and found that they all have this stop codon (data not shown). While this does not mean all isolates of *P. sneebia* will have a stop codon in the ATPase of T3SS-1, it does suggest that this ATPase does not need to be functional for *P. sneebia* to cause an infection in *D. melanogaster*. None of the sequenced strains of *P. sneebia*, *P. rettgeri* and *P. alcalifaciens* showed evidence for the ability to intracellularly replicate in *D. melanogaster* cells *in vitro*, although *P. alcalifaciens* showed some evidence for being invasive and *P. rettgeri* and *P. sneebia* had evidence for resisting phagocytosis [18]. While a T3SS may be involved in these phenotypes for each species, it remains possible that these bacteria may have different intracellular replication or invasive phenotypes in *D. melanogaster* *in vivo* or in other hosts. Indeed, different strains of *P. alcalifaciens* have previously been shown to exhibit invasion of vertebrate cells [60,61].

4. Conclusions

We sequenced and compared the draft genomes of four species of *Providencia* that were all isolated from the hemolymph of wild caught *D. melanogaster* [17,18]. We found the core genome of these isolates to be about 60% of the total coding content for each genome, even after
inclusion of four isolates of Providencia originating from human feces. We found no genes that were specific to and universal in bacteria isolated either from D. melanogaster or from the human gut. Approximately 15% of each genome sequence consisted of genes unique to that isolate. These genes should explain variable phenotypes among the isolates, including metabolic differences [17] and variation in virulence towards D. melanogaster [18]. We found that each of these isolates has at least one type 3 secretion system. The unique genes of each genome are enriched for genes with the “pilus” GO term, suggesting variation in substrates that the bacteria adhere to. The T3SSs and the variety of adhesion molecules suggest that host cell contact is an important part of the virulence mechanisms for these Providencia.

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References


CHAPTER 4

CONCLUSION
This work establishes the basic differences in infection biology among multiple strains in the bacterial genus *Providencia* that were isolated from infected wild *Drosophila melanogaster*. Comparisons of several infection phenotypes of these *Providencia* gave insight into the possible methods these bacteria are using to be virulent towards *D. melanogaster* [1]. The phenotypes I measured created an infection profile for each strain and allowed me to determine that *Providencia sneebia* differs from the general pattern seen with the other strains. Most of the strains had a correlation between the phenotypes such that those that caused the most host mortality were also the most prolific and induced the most antimicrobial peptide transcription by the host. In contrast, *P. sneebia* causes death in about 90% of infected flies and replicates the most out of the strains but induces a lower amount of antimicrobial peptides expression than the less virulent strains. By coinfecting *D. melanogaster* with both *P. sneebia* and another strain which causes less mortality but greater immune response transcription, I was able to conclude that *P. sneebia* is able to actively avoid recognition by the fly immune system.

To determine the differences in genomic content of these *Providencia* strains, I sequenced, assembled and compared the draft genomes of four of them [2]. I found these genomes to have a large amount of homogeneity in gene content both among themselves and in comparison with four other *Providencia* genomes sequenced as part of the Human Microbiome Project. This suggests that additional *Providencia* strains and isolates will also be found to have a sizable core similarity. Each genome also has a repertoire of unique genes that are enriched for the pilus-related genes in each species. All sequenced *Providencia* genomes have a type 3 secretion system (T3SS) so their virulence differences towards *D. melanogaster* are not explained by just the presence of this known virulence factor. The presence of T3SS and
differences in adhesion molecules leads to the hypothesis that these bacteria require contact with host cells to be virulent, though each bacterium may be virulent towards different hosts.

There are still many unanswered questions about these particular host-pathogen interactions. For example, the location of the bacteria in *D. melanogaster* during infection is unknown. Knowing if the bacteria are located throughout the fly or if they colonize a specific location would provide more insight into their virulence mechanisms, particularly for *P. sneebia* and *Providencia alcalifaciens* as they both replicate to very high numbers during infection. The protocol that I developed to insert DNA into the genome of *Providencia rettgeri* could be used to insert GFP into the genome to then track the bacteria within the fly during infection [2]. Although this insertion protocol was unsuccessful with *P. sneebia*, it is possible that this protocol may also work for *Providencia burhodogranariea* and *P. alcalifaciens*, as I did not attempt genetic manipulations of these bacteria. The availability of the genome sequences allow for the location of gene insertions to be chosen to minimize the effect on other genes [2]. Since fluorescent bacteria can be seen through the fly cuticle and the observation of where the bacteria are located is non-destructive, infection within a single fly can be tracked for the whole course of infection [3]. I hypothesized in Chapter 2 that those flies that carry the largest bacterial burden are those flies that succumb to the infection [1]. Although not precisely quantitative, this method could be used to determine if those flies with the greatest amount of bacteria are in also those that die from infection. The comparison of the multiple strains will continue to aid in the determination of the infection biology of these strains with *D. melanogaster*.

Beyond just these specific host-pathogen pairs, the infection biology found with *D. melanogaster-Providencia* may be applicable to other host-pathogen pairs. Bacteria often horizontally transfer genes, particularly virulence related genes, therefore, any genes whose
function is identified in this system could be found in another pathogenic bacteria and be relevant to other host-pathogen pairs [4]. These Providencia strains can also be used to study the host side of the interaction, beyond just the transcriptional read out of the innate immune response. The variability in virulence across these closely related strains can provide comparisons to see how D. melanogaster responds to different pathogens.

References


APPENDIX

ATTEMPTED VERIFICATION OF

PUTATIVE VIRULENCE FACTORS
Abstract

Type 3 secretion systems (T3SS) have evolved to facilitate injection of bacterial proteins into eukaryotic cells in order to manipulate the host for the bacterial pathogen’s benefit. Two natural Drosophila melanogaster bacterial pathogens, Providencia sneebia and Providencia rettgeri, both contain T3SSs but have different infection phenotypes in the fly. I sought to generate mutant strains of both bacteria containing a deletion of a vital T3SS gene with the anticipation of determining the role of the T3SSs for each bacteria during infections in D. melanogaster. Although I was unable to complete the construction of these deletion mutants, I outline here the methodological steps taken towards making these strains. I also discuss what I hypothesize could have been the results from disabling the T3SS system and what the implication would have been for T3SS involvement in P. sneebia or P. rettgeri infections.
1. Introduction

Type 3 Secretion Systems (T3SS) are a complex of proteins involved in virulence of many different bacterial species, particularly in Gram-negative pathogens [1]. The T3SS proteins assemble to form a needle-like apparatus used to breach host cell membranes to inject proteins into the host cytosol. These injected proteins, called effector proteins, manipulate the host cell in a variety of ways to the benefit of the infecting bacterium. The genes encoding the T3SS needle apparatus proteins are located within a single operon, generating what is called a pathogenicity island. This genomic structure of T3SS genes allows all of the genes to be transferred to other bacteria through horizontal gene transfer. Effectors proteins can be contained within these pathogenicity islands, but more often they are located elsewhere in the genome [1]. As these islands contain many genes, it is not easy to delete all of them to implicate T3SS in virulence phenotypes. The machinery can instead be disabled by deleting a protein essential to the function of the needle, such as the ATPase that provides energy for secretion of the effector proteins [2]. The construction of pathogenic bacteria without a functional T3SS can allow for the dissection of how the T3SS is involved in the bacterial virulence.

I analyzed the infection pathology of four different species of the genus *Providencia* that were isolated as natural bacterial pathogens of *Drosophila melanogaster* in this same host and sequenced their genomes [3,4]. I found these bacteria to vary in their pathology during infection in *D. melanogaster* [3]. In particular, *Providencia sneebia* is able to kill 90% of infected flies within 48 hours while *Providencia rettgeri* only kills 40% of infected flies. I also found that *P. sneebia* replicates to significantly higher numbers during infection than *P. rettgeri* but causes a lower induction of *D. melanogaster* antimicrobial peptides [3]. By coinfecting flies with both of
these bacteria, I was able to conclude that the reason for *P. sneebia*’s lower immune induction is that it avoids recognition by the immune system [3].

In analyzing the genomes of these *Providencia* I found two different T3SS. The first one, T3SS-1, is found in 3 of the four species: *P. sneebia*, *P. rettgeri* and *Providencia alcalifaciens*. It is most likely that this island was acquired prior to speciation and was lost from the fourth species, *Providencia burhodogranariea* [4]. T3SS-1 is similar in all three species, as well as in the closely related *Proteus mirabilis*, in protein sequence, gene content, and gene order. Additionally, the island is found in similar genomic locations in all three *Providencia* [4]. The second T3SS, T3SS-2, is found in only *P. sneebia* and *P. burhodogranariea*. Although the T3SS-2 island is found in different locations in the genomes those two bacteria, the T3SS-2 island in each bacteria is very similar in protein sequence, gene content, and gene order [4].

The *P. sneebia* T3SS-1 ATPase has a mutation that creates a stop codon at codon 281 middle of the 420-codon gene, which was verified by Sanger sequencing. This suggests that T3SS-1 may not be functional in *P. sneebia* and therefore, further destruction of this ATPase may have no effect on virulence. The ATPase is the only known essential gene of T3SS, suggesting that *P. sneebia* lacks TTSS-1 function [2]. T3SS vary a lot in what triggers expression of both the apparatus forming genes and the effector protein genes. These external triggers can be things like temperature, contact with the host cell, environmental factors or a combination of these [5–7]. It is unlikely that the apparatus for T3SS-1 and T3SS-2 are being expressed at the same time. Therefore it is unlikely that the ATPase from T3SS-2 would be able to function with the rest of the T3SS-1 machinery to compensate for the loss of T3SS-1 ATPase function. Furthermore, the ATPases of T3SS-1 and T3SS-2 belong to different families which have been suggested to function in different ways. The T3SS-1 is part of the Inv-Mxi-Spa ATPase family
which commonly functions in cell invasion, while T3SS-2 is part of the Ysc ATPase family and is generally found in extracellular pathogens [1].

To gain insight into the functionality and involvement in virulence of the T3SS-2 and to verify that T3SS-1 is nonfunctional in P. sneebia, I sought to make strains that contain an in-frame deletion each ATPase. I also wanted to delete the ATPase present in the T3SS-1 island of P. rettgeri to determine the effect of the loss of its T3SS-1 on its virulence. I designed primers to create in-frame deletions of each ATPase. I intended to use suicide plasmids that would integrate these in-frame deletions into each genome via homologous recombination. I next planned to use negative selection to select for those bacteria that excised the plasmid from the genome. Those that excised the plasmid would be screened by PCR to identify bacteria that had the in-frame deletion instead of the reversion to the wild type configuration. After creating strains with the deleted ATPase, I anticipated doing infections in wild type D. melanogaster to see how the loss of each T3SS affected P. sneebia and P. rettgeri’s virulence by measuring host mortality, bacterial load and host antimicrobial peptide expression after infection.

2. Methods and Materials

2.1 Bacterial strains

The type strain of P. sneebia (DSM 19967) and the Dmel1 strain of P. rettgeri were used in this study [3,8]. The E. coli strain S17 was used during construction of the in-frame deletion plasmids. All three strains were grown in LB at 37°C with shaking.
2.2 Suicide plasmid

To do a knock out by homologous recombination, a plasmid must be selected which is unable to replicate within the bacterium. This ensures that descendent bacteria with the antibiotic resistance or other selective marker associated with the plasmid have an integration of the plasmid into the genome and are not maintaining the plasmid in the cytosol. I opted to use a plasmid with a R6Kγ origin because these plasmids can only replicate in host bacteria with a specific pi protein encoded by the *pir* gene [9]. To ensure that neither *P. sneebia* nor *P. rettgeri* contain the *pir* gene, I created databases of all of the Roche/454 sequencing reads with BLAST+ (version 2.2.25) and used blastx to search for similarity to the pi protein [4]. I found no evidence that *P. sneebia* or *P. rettgeri* have the *pir* gene. I choose the R6Kγ origin plasmid pLD55, which has a locus with multiple restriction sites, an origin for transfer through conjugation, and genes that allow for negative selection once integrated into the bacterial genome [9].

2.3 Creating the deletion

I designed 4 primers to create an in-frame deletion for each *ATPase* using the method of crossover PCR as described in Link et al. 1997 (Figure A.1) [10]. This method first creates two PCR products with complementary sequences at the 3’ end of one product (Product A in Figure A.1) and the 5’ end of the other (Product B in Figure A.1) which allows them to overlap in a second PCR resulting in the formation of one product to create the in-frame deletion (Figure A.1, steps 1 and 2 in Figure A.2). Two of the primers, L1 and R2, were designed on either side of the *ATPase* gene, placed equal distance from where the deletion will ultimately be (Figure A.1). These primers included different restriction enzyme cut sites at their 5’ ends, XhoI or NotI. Two internal primers, R1 and L2, were designed to create the in-frame deletion (Figure A.1). The
Figure A.1. Diagram of primer locations and steps in crossover PCR protocol.
Figure A.2. Flowchart of steps taken and planned for deletion of ATPases in *P. sneebia* and *P. rettgeri*. See Figure A.1 for reference of primer and product names.
right internal primer, R1, contained the last 57 bases at the 3’ end of the \textit{ATPase} while the left internal primer, L2, contained first 18 bases from 5’ end of the \textit{ATPase} and the complement of the 21 nucleotides of the 5’ end of the right internal primer, R1 (Figure A.1).

The deletion was created in a two-step crossover PCR process. The first round of PCRs was done using genomic DNA for the template to create either side of the deletion with the pair of internal and external primers, L1 with L2 to make Product A and R1 with R2 to make Product B (Figure A.1, steps 1 in Figure A.2). The second PCR uses the products of the first PCR as templates, Product A and Product B, with the outermost primers, L1 and R2 (Figure A.1, steps 1 in Figure A.2). The complimentary 21 bases at the 5’ ends of both internal primers allow the products of the first PCR step to anneal into a single molecule that contains the in-frame deletion and flanking sequence. See Table A.1 for sequences of primers used.

The first PCR step (step 1 in Figure A.2) was done using high fidelity iProof polymerase (Bio-Rad) in 50 µl reaction with 300 ng of genomic DNA. The PCR cycles were: 1) 98°C for 30 seconds, 2) 98°C for 7 seconds, 3) appropriate annealing temperature for 20 seconds, 4) 72°C for 20 seconds, 5) repeat steps 2-4 for 30 cycles, 6) 72°C for seven minutes. Annealing temperatures were determined using manufacturer’s suggested method (Table A.2). These products were then gel purified using the E.Z.N.A Gel Extraction Kit (Omega). The second PCRs (step 2 in Figure A.2), the crossover PCR, was done with a total volume of 100 µl divided into two separate tubes, with approximately 87 ng of each of the products from the first round in each tube using iProof polymerase. The same PCR cycles and purification methods were performed in a manner similar to those of the first round.
Table A.1. Primer sequences used for crossover PCR. Primer type designations refer to those used in Figure A.1.

<table>
<thead>
<tr>
<th>species</th>
<th>T3SS</th>
<th>primer type</th>
<th>sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. rettgeri</em></td>
<td>T3SS-1</td>
<td>L1</td>
<td>ttttctcagctcagctcattgcagattttg</td>
</tr>
<tr>
<td></td>
<td></td>
<td>L2</td>
<td>atctatggtgctagtaattgaactcaattttggtct</td>
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<tr>
<td></td>
<td></td>
<td>R1</td>
<td>tttcatagcctacatcagcagttcctcattttgctt</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R2</td>
<td>tttttgcggccgtgaatggcggagtagttttgtgg</td>
</tr>
<tr>
<td><em>P. sneebia</em></td>
<td>T3SS-1</td>
<td>L1</td>
<td>ttttctcaggtgctcagttgctcagttttttgat</td>
</tr>
<tr>
<td></td>
<td></td>
<td>L2</td>
<td>atcatttatgatcattgactcatcattttttgaacatcatcaattttttgctt</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R1</td>
<td>gatgctttcgtcattttccctttttgccgcttagtttaa</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R2</td>
<td>tttttgcggcccttctttttttctttttttttttttttgag</td>
</tr>
<tr>
<td><em>P. sneebia</em></td>
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<td>L1</td>
<td>ttttctcagttggtgcagataagctcagttttggat</td>
</tr>
<tr>
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<tr>
<td></td>
<td></td>
<td>R1</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>R2</td>
<td>tttttgcggcccttctttttttcattttttttttttttatcagtttttagt</td>
</tr>
</tbody>
</table>
Table A.2. Primer annealing temperatures used for crossover PCR. Primer type designations refer to those used in Figure A.1.

<table>
<thead>
<tr>
<th>species</th>
<th>T3SS</th>
<th>primer type pair</th>
<th>crossover PCR step</th>
<th>annealing temp</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. rettgeri</em></td>
<td>T3SS-1</td>
<td>L1-L2</td>
<td>1</td>
<td>62°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R1-R2</td>
<td>1</td>
<td>69°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>L1-R2</td>
<td>2</td>
<td>62°C</td>
</tr>
<tr>
<td><em>P. sneebia</em></td>
<td>T3SS-1</td>
<td>L1-L2</td>
<td>1</td>
<td>61°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R1-R2</td>
<td>1</td>
<td>68°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>L1-R2</td>
<td>2</td>
<td>66°C</td>
</tr>
<tr>
<td><em>P. sneebia</em></td>
<td>T3SS-2</td>
<td>L1-L2</td>
<td>1</td>
<td>59°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R1-R2</td>
<td>1</td>
<td>67°C</td>
</tr>
<tr>
<td></td>
<td></td>
<td>L1-R2</td>
<td>2</td>
<td>64°C</td>
</tr>
</tbody>
</table>
2.4 Restriction digestions of pLD55 and inserts

Digestions of gel purified products or plasmid pLD55 were performed with XhoI and NotI at 37°C for 2 hours, followed by a 20 minute incubation at 65°C to inactivate the enzymes (step 3 in Figure A.2). The products were then cleaned by gel purification from a fresh 1% TAE gel. The inserts were extracted using Purelink gel extraction kit (Invitrogen). The digested plasmid was extracted using SNAP gel purification kit (Invitrogen).

2.5 Ligations of inserts into pLD55

The purified digested products were run on a gel to help estimate the proper volumes to add to the ligation reaction in order to achieve an approximately 3:1 insert to vector ratio (step 4 in Figure A.2). Ligations were done in a volume of 10 µl at 22°C for two hours.

2.6 Transformation of constructs into E. coli

E. coli strain S17 cells were made chemically competent. Bacteria were grown in 8 ml of liquid LB until $A_{600}$ was between 0.2 and 0.4, indicating the bacteria were in log phase. The bacteria were then spun down in 5 separate tubes at 4°C at 7000 rpm for 10 minutes. The pellets were washed with 333 µl of ice cold 0.1M CaCl$_2$ and mixed into one tube. The E. coli were spun and washed again before the addition of 12 µl DMSO. After a 15 minute incubation on ice, 11 µl more of DMSO were added followed by another 15 minute incubation on ice. The bacteria were aliquoted into pre-chilled 1.5 mL tubes before being snap frozen in a dry ice and ethanol bath.

For transformation, DNA was added to gently thawed competent cells. The cells were then incubated on ice for at least 30 minutes, followed by a 30 second heat shock at 42°C, and immediately returned to ice (step 5 in Figure A.2). Next 200 µl SOC media was added to the
bacteria and they were shaken at 37°C for one hour. The transformation was then plated on LB media containing tetracycline and grown overnight at 37°C. The pLD55+deletion construct provides resistance to tetracycline, allowing selection of transformants.

2.7 Conjugation with *P. rettgeri*

*E. coli* strain S17 carrying the plasmid with the pLD55+deletion construct and *P. rettgeri* were grown separately in liquid LB, with antibiotics added to the *E. coli* containing medium to maintain the plasmid, until log phase (step 6 in Figure A.2). Bacteria were then spun down and washed twice with LB. The bacteria were gently pelleted a third time, and each pellet was resuspended in 100 µl LB and then mixed into the same tube. The 200 µl of mixed bacteria were pipetted onto a filter paper placed on top of a LB plate and incubated overnight at 37°C. The filter paper was then removed and vortexed in a tube with 1 ml of liquid LB. Multiple 50 µl aliquots of the bacteria-LB mixture were spread on LB plates that contained both tetracycline and erythromycin. *P. rettgeri* strain Dmel1 is naturally resistant to erythromycin while *E. coli* strain S17 is not so its presence inhibits growth of all *E. coli* donor strains. The tetracycline was present to select for those *P. rettgeri* that obtained genomic integration of pLD55+deletion.

2.8 Electroporation with *P. sneebia*

I tried the conjugation protocol described above for *P. rettgeri* with *P. sneebia* but was unsuccessful at getting the plasmids described here, or other plasmids, into *P. sneebia* with it. Instead, I developed a method for transformation by electroporation for *P. sneebia* (step 6 in Figure A.2). *P. sneebia* cells were made electrocompetent by growing at 37°C until mid log phase 500 mL of SOC media spiked with 5 mL of an overnight culture. The bacteria were then
chilled on ice at 4°C for one hour followed by centrifugation at 6000 rpm for 20 minutes. The bacteria were then washed in sterile cold water twice before being resuspended in 20 mL cold 20% glycerol and spun again at 8000 rpm for 10 minutes. The pellet was resuspended in 2 mL 20% glycerol and 30 µl was aliquoted into chilled 1.5 mL tubes to be flash frozen with an ethanol and dry ice bath. Electroporation was successful after adding around 100 ng of plasmid DNA to an electrocompetent *P. sneebia* aliquot and shocking the cells at 1.8 kV. After shocking, the bacteria were grown for an hour at 37°C in 1 ml SOC media before being plated on selective medium. This electroporation protocol was successful in getting the pEGFP plasmid (Clontech) into *P. sneebia* but not either of the plasmids designed here for creating a deletion of each T3SS ATPase.

I tried to an alternative protocol to make electrocompetent cells but this was also unsuccessful. In this protocol 2 ml of overnight *P. sneebia* culture were spun for 2 minutes at 8000 g at room temp. The cells were then washed in 2 ml of sterile 300 mM sucrose before being spun again. Cells were washed with 300 mM sucrose and pelleted again before being resuspended in 200 µl of 300 mM sucrose and placed on ice to await addition of DNA and electroporation.

2.9 Selecting and screening for the ATPase deletion

After transforming *P. rettgeri* and *P. sneebia* with the ATPase-deletion plasmid and selecting for genomic integration, I intended to cause the plasmid to excise from the genome and then screen for the in-frame deletion (steps 7 and 8 in Figure A.2). Bacteria with pLD55+deletion genomic integration would be grown on selective media containing fusaric acid which kills bacteria possessing the *tetAR* genes present on pLD55, therefore selecting for
survival of those isolates that successfully excised the plasmid from their genome [11,12]. Approximately half of the surviving bacteria would have the \textit{ATPase} deletion construct, while the other half would have reverted back to wild type, depending on which way the plasmid is excised from the genome. I planned screen isolates by PCR, using the outside most L1 and R2 primers, expecting a 1.5 kb product for the wild type bacteria compared to an 500 bp product in bacteria that have sustained the deletion (Figure A.1). After PCR verification of the deletion of the \textit{ATPase}, I intended use Sanger sequencing methods to sequence the region around and including the deletion to verify that an in-frame deletion had been created.

2.10 Infections with mutants in \textit{D. melanogaster}

Infections of \textit{D. melanogaster} with the \textit{ATPase}-deleted bacteria would be performed as previously described [3]. The phenotypes of mortality from infection, amount of bacterial replication during infection, and the antimicrobial peptide expression by the fly during infection would be measured.

3. Results

I was able to create the plasmids containing an in-frame deletion with flanking regions of each T3SS \textit{ATPase} on the pLD55 backbone. Using conjugation, I was able to insert the appropriate plasmid into the genome of \textit{P. rettgeri}. Although I made several attempts to get the applicable plasmids into \textit{P. sneebia} by both conjugation and multiple electroporation protocols, I was never successful at achieving genomic insertion. As I was unable to get the deletion construct into the genome of \textit{P. sneebia}, I was unable to create a strain with either T3SS \textit{ATPase} deleted to test the involvement of T3SS in \textit{P. sneebia} virulence. I did not continue on with the...
subsequent steps of creating the in-frame deletion for *P. rettgeri* as my main interest was in determining the role T3SS in *P. sneebia*’s virulence and *P. rettgeri* was being constructed as a comparison.

**4. Discussion**

The most likely explanation for why the *P. sneebia* electroporation protocol worked with the pEGFP plasmid but not with the plasmids created here for deleting each T3SS ATPase is that maintenance of pEGFP did not require genomic integration. It is possible that the plasmid did get within *P. sneebia* cells but was not integrated into the genome, and since the suicide plasmids were unable to replicate, no transformants were established. It is possible that a longer period of growth after electroporation prior to being plated on selective media could allow for genomic integration to happen. There could also be other barriers in *P. sneebia* to successfully integrating DNA into the genome, which would be harder to surmount.

Had I been successful in creating the T3SS ATPase deletions, I had some hypotheses as to the phenotypes that I would have seen during fly infections with these mutant bacteria. I hypothesize that deleting the ATPase from T3SS-1 in *P. sneebia* would have no effect on infection phenotypes since the endogenous gene already carries a premature stop codon. But I predict that deleting the ATPase in T3SS-2 would decrease the amount of *D. melanogaster* host mortality seen from infection. I also hypothesize that there would be a slight reduction in *P. rettgeri* induced mortality in *D. melanogaster* with the inhibition of proper T3SS-1 function. I think that both bacteria are using T3SS, T3SS-1 in *P. rettgeri* and T3SS-2 in *P. sneebia*, in their virulence towards *D. melanogaster* and that the difference seen in their infection phenotypes results from different effector proteins or from the presence of non-T3SS related proteins. The
idea that each bacteria is using their T3SSs in very different ways is supported by the varied functions that are suggested based on the different ATPase families each belong to. It is most likely that T3SS-2 is functioning extracellularly while T3SS-1 is functioning to get the bacteria within the host cells. Although this suggests that \textit{P. rettgeri} may be an intracellular pathogen, I did not find that \textit{P. rettgeri} has the ability to replicate within \textit{D. melanogaster} cells [3]. This may be because the \textit{P. rettgeri}’s T3SS-1 is not used to be an intracellular pathogen of \textit{D. melanogaster} or the proper signal was not present in the experimental setup as the transcription of T3SS apparatuses or secretion of certain effector proteins can be dependent on specific external triggers [5].

\textbf{Acknowledgements}

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\textbf{References}


