Pathogenicity of Serratia marcescens Strains in Honey Bees

Kasie Raymann,a,b Kerri L. Coon,b Zack Shaffer,b Stephen Salisbury,b Nancy A. Moranb

aDepartment of Biology, University of North Carolina, Greensboro, North Carolina, USA
bDepartment of Integrative Biology, University of Texas at Austin, Austin, Texas, USA

ABSTRACT Although few honey bee diseases are known to be caused by bacteria, pathogens of adult worker bees may be underrecognized due to social immunity mechanisms. Specifically, infected adult bees typically abandon the hive or are removed by guards. Serratia marcescens, an opportunistic pathogen of many plants and animals, is often present at low abundance in the guts of honey bee workers and has recently been isolated from Varroa mites and from the hemolymph of dead and dying honey bees. However, the severity and prevalence of S. marcescens pathogenicity in honey bees have not been fully investigated. Here we characterized three S. marcescens strains isolated from the guts of honey bees and one previously isolated from hemolymph. In vivo tests confirmed that S. marcescens is pathogenic in workers. All strains caused mortality when a few cells were injected into the hemocoel, and the gut-isolated strains caused mortality when administered orally. In vitro assays and comparative genomics identified possible mechanisms of virulence of gut-associated strains. Expression of antimicrobial peptide and phenoloxidase genes was not elevated following infection, suggesting that these S. marcescens strains derived from honey bees can evade the immune response in their hosts. Finally, surveys from four locations in the United States indicated the presence of S. marcescens in the guts of over 60% of the worker bees evaluated. Taken together, these results suggest that S. marcescens is a widespread opportunistic pathogen of adult honey bees and that it may be highly virulent under some conditions such as perturbation of the normal gut microbiota or the presence of Varroa mites that puncture the integument, thereby enabling entry of bacterial cells.

IMPORTANCE Recently, it has become apparent that multiple factors are responsible for honey bee decline, including climate change, pests and pathogens, pesticides, and loss of foraging habitat. Of the large number of pathogens known to infect honey bees, very few are bacteria. Because adult workers abandon hives when diseased, many of their pathogens may go unnoticed. Here we characterized the virulence of Serratia marcescens strains isolated from honey bee guts and hemolymph. Our results indicate that S. marcescens, an opportunistic pathogen of many plants and animals, including humans, is a virulent opportunistic pathogen of honey bees, which could contribute to bee decline. Aside from the implications for honey bee health, the discovery of pathogenic S. marcescens strains in honey bees presents an opportunity to better understand how opportunistic pathogens infect and invade hosts.

KEYWORDS Serratia marcescens, honey bee, opportunistic pathogen, virulence

Despite extensive recent research on the reasons for global honey bee colony decline, no single factor has yet been implicated as a cause. In fact, most evidence suggests that honey bee colony decline stems from multiple factors, such as climate change, pests and pathogens, pesticides, and loss of foraging habitat (1, 2). Although honey bees have many known pathogens, few bacterial pathogens have been identified. The best-known bacterial diseases are European foulbrood and American foulbrood.
brood, which are caused by the Gram-positive species *Melissococcus plutonius* and *Paenibacillus larvae* and result from infection of larvae that die in the hive and leave diagnostic evidence of disease (3). Recognized pathogens of adult honey bees include *Bacillus pulviferacens* (powdery scale disease) and two species of the genus *Spiroplasma* (May disease) (4). Adults with bacterial infections may be overlooked in many cases due to behaviors associated with social immunity, a phenomenon by which individual group members cooperate to combat disease transmission (5). For example, honey bees initiate precocious foraging when exposed to different stressors, including Varroa mites and *Nosema ceranae*, and specialized workers guard the nest entrance to attack or exclude infected nest mates (5). Therefore, it is possible that other bacterial pathogens of adult honey bees exist but have gone unnoticed because infected workers are largely excluded from the hive.

Opportunistic pathogens are organisms that can become pathogenic only in susceptible hosts, e.g., hosts with weakened immune systems or altered microbiome compositions (6). However, in most cases opportunistic pathogens coexist peacefully within their host and can live in several nonhost environments, making it difficult to identify them and determine the factors that lead to their pathogenicity. Many opportunistic pathogens are resistant to multiple antibiotics, and antibiotic treatment is often a precursor to infection (6). Honey bees experience many stressors that can disrupt their microbiota and/or alter their immune system, such as exposure to antibiotics (7, 8) and pesticides (9–11), increasing their risk of infection by opportunistic pathogens.

*Serratia marcescens* is a Gram-negative opportunistic pathogen of a wide range of animals, including humans and insects (12). In most animals, *S. marcescens* is virulent only when present in the bloodstream (12). In honey bees, *S. marcescens* has been detected in diseased larvae (13) and is sporadically found at low frequencies (typically <5% relative abundance) in the guts of adults, where it is considered a signifier of an atypical microbiome composition (7, 14–16). Recently, a *S. marcescens* strain (kz11) isolated from the honey bee gut was shown to cause increased mortality in adult bees following antibiotic or pesticide exposure (7, 17), and another strain, *S. marcescens* Ss1, was isolated from Varroa mites and the hemolymph of immobilized and dead bees, particularly in overwintered hives (18). Taking the data together, these studies suggest that *S. marcescens* is a common pathogen of honey bees. However, the virulence of honey bee-associated *S. marcescens* strains has not been experimentally tested.

Here we isolated and characterized three strains of *S. marcescens* from the guts of honey bees and compared them to Varroa/hemolymph-associated strain *S. marcescens* Ss1 (18). Using in vivo mortality assays, we showed that *S. marcescens* strains isolated from the gut and inoculated orally or injected into the hemolymph are highly virulent to honey bees. In contrast, we found that *S. marcescens* Ss1 exhibits virulence only when present in the hemolymph of honey bees, suggesting that it possesses mechanisms for virulence and infection that are different from those of the gut isolates. Using in vitro enzyme activity assays and comparative genomics, we identified potential virulence factors associated with the pathogenicity of gut-isolated strains. Furthermore, we investigated how *S. marcescens* infection impacts the honey bee immune response and found that honey bee-associated *S. marcescens* strains do not induce expression of antimicrobial peptides (AMPs) or phenoloxidase, suggesting that these strains possess mechanisms for evading the honey bee immune system. Overall, our results indicate that *S. marcescens* is an important and overlooked threat to honey bees.

**RESULTS**

*S. marcescens* is commonly found in the gut microbiome of honey bees. We performed 16S rRNA profiling of the gut microbial community in honey bees from four locations: a hive kept on the University of Texas at Austin (UT) campus (*n* = 26); a commercial hive in Florida (*n* = 11); an organic hive in Tennessee (*n* = 21); and a long-established feral colony near Moab, Utah (*n* = 9). We identified *S. marcescens* in honey bees from all locations (Fig. 1). The proportions of bees containing *S. marcescens* in their gut ranged from 22% to 100% (Fig. 1A). However, the relative abundance was
**Serratia marcescens** is a Pathogen of Honey Bees

FIG 1 Frequency of *S. marcescens* in the gut microbiome of honey bees and the phylogeny of gut isolates. (A) Relative abundances of *S. marcescens* in the gut microbiome of honey bees from four locations: Utah (UT), Texas (TX), Tennessee (TN), and Florida (FL). (B) Phylogenetic tree of the genus *Serratia*. The three strains described here (kz2, kz11, and kz19), the other honey bee-associated strain (Ss1), and the *Drosophila*-associated strain (Db11) are shown in color. The tree was constructed using concatenated sequences of 125 proteins encoded by single-copy genes and analyzed with PhyML (GTR plus Gamm4) with 100 bootstrap replicates. Bootstrap values are represented by circles at each node (black, >90; gray, >70 and <90; white, <70). For the full tree, see Fig. S1. The Venn diagram on the right shows the number of genes shared by kz2, kz11, kz19, and Ss1.
low (0.007% to 9.32%) in all S. marcescens-positive bees, suggesting that it can live as a bee gut commensal at low abundance (see Table S1 in the supplemental material). These results, coupled with previous gut microbiome studies of honey bees (7, 14–16), suggest that S. marcescens is frequently present at low abundance in the honey bee gut.

The genomes of three S. marcescens strains (kz2, kz11, and kz19) isolated from the guts of honey bees were sequenced using Illumina MiSeq 2X300, assembled, and annotated (for more information about the genomes, see Table S2 and Materials and Methods). The kz11 strain was reported in a previous study (7) but was not sequenced or characterized. Phylogenetic and average nucleotide identity (ANI) analyses confirmed that these strains belong to the species S. marcescens (Table S2). Only one other honey bee-associated S. marcescens strain (Ss1) has been previously sequenced and characterized (18). One of our gut-isolated strains (kz19) is closely related to the Ss1 strain (99% ANI). The two other strains (kz2 and kz11) belong to a different clade, which consists of a mixture of human and environmental isolates, and these two strains are only 95% identical to Ss1 or ks19 based on ANI (Fig. 1B; see also Fig. S1 in the supplemental material). The genomes of kz11 and kz2 are 100% identical to one another based on ANI, differing only in gene content represented by a few accessory genes (Fig. 1B). The three kz strains share over 4,000 genes with Ss1 but also possess 298 genes that are not present in the Ss1 genome (Fig. 1B; see also Data Set S1 in the supplemental material). All of the honey bee-associated strains were 95% identical to the Db11 strain (19), a spontaneous streptomycin-resistant mutant derived from strain Db10 isolated from Drosophila (20); Db11 falls within a different subclade of S. marcescens (Fig. 1B; see also Table S2).

S. marcescens is a pathogen of honey bees. Using in vivo virulence assays, we found that all three S. marcescens bee gut isolates, Db11, and Ss1 are pathogenic to honey bees. The gut-isolated strains and Db11 showed high virulence when bees were exposed orally or through injection into the hemolymph. However, Ss1 was virulent only when present in the hemolymph. Adult worker bees were orally exposed to S. marcescens strains kz2, kz11, kz19, Db11, and Ss1. Two routes were used for oral exposure: (i) 500 µl of a bacterial solution applied to sterile pollen at an optical density (OD) of 1 (feeding exposure) and (ii) 10 µl of a bacterial sugar syrup solution applied to the body of each bee at an OD of 0.5 (immersion, which results in ingestion when bees clean one another). For both methods of exposure, kz2, kz11, kz19, and Db11 significantly decreased the survival rate of exposed bees compared to control bees, whereas Ss1 did not result in increased mortality (Fig. 2; see also Fig. S2). Exposure via immersion resulted in a much higher mortality rate (40% to 60%) (Fig. 2A; see also Fig. S2A and B) than exposure via feeding (10% to 20%) (Fig. 2B; see also Fig. S2C and D).

In other insects, it has been shown that S. marcescens is virulent only if it colonizes the hemolymph (12). To test if this is true in bees, adult workers were orally exposed to kz2, kz11, kz19, and Ss1 strains containing an E2 crimson fluorescent protein on an RSF1010 broad-host-range backbone with spectinomycin resistance (56). In order to maintain the plasmid, bees were fed a sterile sucrose solution containing 120 µg/ml of spectinomycin during the experiment. Plasmid-containing strains, which are blue/purple in color, were used to screen for the S. marcescens strains. For 3 days following exposure, six live bees were sampled from each group. In addition, up to six newly dead bees, if applicable (see Materials and Methods), were sampled. The gut and hemolymph of each bee were extracted and plated on LB agar containing 120 µg/ml spectinomycin. After 24 h of incubation at 30°C, the presence or absence of each S. marcescens strain was determined by the presence of blue/purple colonies on the plates. On day 1 following exposure, strains kz2, kz11, and kz19 were detected in at least 50% of the guts of living bees, but the numbers of bees sampled that possessed these strains in their gut decreased on days 2 and 3 (Fig. 3A). In contrast, Ss1 was never isolated from the guts of living bees (Fig. 3A). With the exception of a single kz19-exposed bee, none of the S. marcescens strains were observed in the hemolymph of living bees (Fig. 3A). In
dead bees, kz2, kz11, and kz19 were isolated from both the gut and the hemolymph of almost all bees sampled on days 1 and 2. On day 3, kz2, kz11, and kz19 were detected in at least 50% of the guts of dead bees and in the hemolymph of some bees (Fig. 3B). Very few control and Ss1-exposed bees died during the experiment. Ss1 was not detected in the gut or hemolymph of living or dead bees on days 1 and 2, but Ss1 was detected in the gut and hemolymph of a single dead bee on day 3 (Fig. 3B). S. marcescens was never detected in the gut or hemolymph of control bees (Fig. 3).

We performed hemolymph injection experiments to further investigate the virulence of the S. marcescens strains. Within 20 h, all bees injected with ~10 cells of kz11, kz19, kz2, or Db11 died (Fig. 4A; see also Fig. S3). Injection with Ss1 also resulted in increased mortality, but the increase was less than that seen with the other S. marcescens strains. After 24 h, approximately 30% of Ss1-injected bees died (Fig. 4A). In a second experiment, injection with kz11, kz19, and kz2 again killed all bees within 20 h, but Ss1 exposure did not significantly increase mortality compared to controls (Fig. S3). In contrast, almost all bees that were injected with phosphate-buffered saline (PBS) only or with Escherichia coli strain K12 were still alive after 26 h (Fig. 4A; see also Fig. S3).
In order to determine how quickly the bee-associated strains replicate within the hemolymph, we evaluated the number of cells present in the hemolymph 3 and 6 h after the injections with ~10 cells. At 3 h postinjection, the CFU count was similar to the number of cells injected (Fig. 4B; ~10 cells per bee). At 6 h, the average number of CFU of kz2, kz11, and kz19 was $1 \times 10^3$ to $2 \times 10^3$. In contrast, the number of Ss1 CFU did not significantly increase from h 3 to h 6 (Fig. 4C). The hemolymph of control bees produced no CFU.

In rich media, Ss1 grows more slowly than kz strains (Fig. S4). The slow growth of Ss1 compared to other *S. marcescens* strains was also reported in reference 18. Thus, Ss1 may also replicate more slowly within the hemolymph, resulting in a delay in virulence.

*Serratia marcescens* infection does not cause an immune response in honey bees. The honey bee innate immune response to *S. marcescens* infection was evaluated by assessing expression levels of the four honey bee antimicrobial peptide (AMP) gene families (abaecin, apidaecin, defensin, and hymenoptaecin) and the gene encoding the melanizing enzyme phenoloxidase following exposure to each strain. Bees were either injected with ~10 cells or orally exposed using the immersion exposure method. The abdomen of the injected bees and the guts of the orally exposed bees were sampled 6 h after infection, and AMP and phenoloxidase gene expression were evaluated.

No significant changes in AMP expression were observed between bees injected with *S. marcescens* and control bees for any of the strains tested (Fig. 5A). However, bees injected with Ss1 showed increased expression of phenoloxidase compared to control bees or bees injected with kz21 or kz19 (Fig. 5A). Ss1-injected bees also showed higher expression of abaecin, defensin, and hymenoptaecin, but this difference was significant only in comparisons to bees injected with kz21 (Fig. 5A). Oral exposure of bees to different *S. marcescens* strains also generally failed to elicit significant changes.
Virulence-associated enzymatic activity is detected in gut-isolated *S. marcescens* strains from honey bees. Virulence is associated with several enzymatic activities, such as siderophore production, protease, chitinase, gelatinase, DNase, and hemolytic activity (22–27). Motility (e.g., swimming and swarming) has also been shown to play a role in virulence (28). We performed *in vitro* assays at three different temperatures (22°C [room temperature (RT)], 30°C, and 37°C) to determine if our strains and Ss1 possessed any of these activities. The three kz strains tested positive for all enzymatic and motility assays at 30 and 37°C after 24 and 48 h of incubation, with the exception of kz11, which exhibited hemolytic activity only after 72 h (Fig. S5). Protease activity, siderophore production, and gelatinase activity were observed for all kz strains at RT at 24 and 48 h (Fig. S5). In contrast, Ss1 exhibited DNase activity and swarm motility, the latter of which was observed only after incubation at 30 and 37°C (Fig. S5). DNase activity has previously been demonstrated in Ss1 (18).

*S. marcescens* gut isolates possess virulence genes that are missing from Ss1. Because Ss1 did not exhibit virulence when ingested by bees under our experimental conditions, we investigated the genes shared by all kz strains but absent from Ss1 to identify genes potentially associated with the virulence of the gut isolates. Of the 298 genes unique to the kz strains (Data Set S1), 216 (72%) were found in clusters (sets of three or more consecutive genes that display conserved synteny). Because Db11 was
also virulent to bees when ingested, we focused on the 13 clusters shared by all kz strains and the Db11 strain.

One of the largest clusters missing from Ss1 encodes flagellar proteins (Fig. 6). Ss1 is missing 36 genes from the flagellar gene cluster which are present in all the kz strains as well as Db11 (Fig. 6), indicating that Ss1 does not have the ability to form fully functional flagella. In our motility assays, Ss1 did not exhibit swim motility but appeared to exhibit some swarming motility (Fig. S5). It is possible that Ss1 is not actually swarming but uses another mechanism that does not require flagella to move across surfaces, such as gliding or twitching (32). Two other clusters missing from Ss1 include genes involved in iron regulation and siderophore biosynthesis (Fig. S6), which is consistent with the finding that Ss1 does not produce siderophores (Fig. S5). The functional roles of the other 11 gene clusters missing from Ss1 are less clear, as they mostly include genes encoding hypothetical proteins, transcriptional regulators, and transporters (Fig. S7).

**DISCUSSION**

Here we characterized the virulence of three *S. marcescens* strains isolated from the honey bee gut as well as *S. marcescens* Ss1 isolated from Varroa mites and the hemolymph of bees (18). We experimentally confirmed that Ss1 is virulent when present in the hemolymph, as suggested in a previous study (18). Moreover, we showed that the gut-isolated *S. marcescens* strains, when ingested and when injected into the body cavity, are highly virulent. We detected Ss1 in the gut of only one bee, which suggests that it is rarely capable of colonizing the gut, at least under our experimental conditions. We note that bees were exposed to very high doses of *S. marcescens* in our oral exposure experiments. In field-collected bees, *S. marcescens* is usually at low abundance, and virulence of gut-dwelling *S. marcescens* may depend on unusual conditions in which it becomes abundant in the gut.

The immersion method of oral exposure resulted in higher mortality than the
feeding method. Bees immersed in the inoculum solution immediately clean themselves and each other and thus are rapidly exposed to large doses. In contrast, the amount ingested via feeding varies, depending on the rate and extent of consumption of the sugar syrup. Also, the immersion method potentially enables *S. marcescens* to invade through other routes, such as the respiratory tracheae or punctures in the integument. For our oral exposure and hemolymph injection experiments, we tested *S. marcescens* virulence on workers from hives that had not been treated with chemicals for 2 years. The mortality rate of bees infected with *S. marcescens* was previously shown to be much higher following exposure to the antibiotic tetracycline (7).

The immune response of honey bees and other insects consists of three levels of resistance: physical barriers, cell-mediated immunity, and cell-free humoral immunity (33). AMPs and phenoloxidase are key elements of humoral immunity (34). In general, we did not observe an increase in expression of the genes encoding the four bee AMPs or phenoloxidase following *S. marcescens* infection, suggesting that these strains possess mechanisms for evading (not triggering) the honey bee immune response, such as are known in some other animal pathogens (35). This general lack of immune induction potentially reflects the prior upregulation of AMP and phenoloxidase genes in response to colonization by the native gut microbial community (36), preventing us from observing further upregulation. Interestingly, we did observe a decrease in the expression of two AMP genes following oral exposure to two of the gut-isolated *S. marcescens* strains, consistent with an ability of some strains to suppress immune responses under certain conditions. We did not directly measure AMP or phenoloxidase activity in the present study. However, in a previous study of immune responses to gut microbiota in honey bees (36), transcript levels correlated with corresponding AMP abundance. Further investigation will be necessary to elucidate the molecular mechanisms underlying how different *S. marcescens* strains interact with the bee immune system as well as the impact of *S. marcescens* infection on expression of bee immune factors over longer durations of infection or under conditions in which the native microbiota are disrupted.

![Flagellar components in honey bee-associated *S. marcescens*.](image-url) (A) The flagellar gene cluster. Genes shown in yellow are missing from the Ss1 genome but are present in k22, k211, k19, and Db11. Genes shared by all five strains are shown in dark gray. (B) Flagellar components of *Salmonella enterica* serovar Typhimurium. (Adapted from reference 29 with permission of the publisher.) Filled circles and boxes represent components present in Ss1. White circles and boxes represent components missing from the Ss1 genome. (C) List of accession numbers and annotations for genes shown in panel A.
Successful colonization and pathogenicity may depend on the gut microbiome composition. Although we found that the gut-isolated *S. marcescens* strains were sometimes virulent when restricted to the gut (Fig. 3), virulence may require unusually high titers or proliferation near the gut wall, as might occur when the microbiome is disrupted. In the colonization experiments, bees were exposed to spectinomycin, prior to exposure to *S. marcescens*. Increased susceptibility to *S. marcescens* infection following tetracycline treatment (7) provides evidence supporting this hypothesis. Moreover, some bee gut community members possess antibacterial weaponry, potentially contributing to bee defense against bacterial pathogens (37). For example, one of the core bee gut bacterial species, *Snodgrassella alvi*, possesses type VI secretion systems that deliver antibacterial toxins and that kill members of other bacterial species (37).

As in other animals (12), *S. marcescens* virulence in honey bees might correlate with the ability of the strains to colonize the gut and subsequently penetrate the gut wall and enter the body cavity, causing septicemia. We identified *S. marcescens* in the hemolymph of most dead bees orally inoculated with the gut-isolated strains, whereas it was rare in the hemolymph of live bees (Fig. 3). While this suggests that pathogenicity involves invasion of the body cavity via the gut, another possibility is postmortem proliferation in hemolymph. In contrast, Ss1, which was isolated from hemolymph of diseased honey bees (18), appears unable to colonize the gut, suggesting that its mode of infection is different from that of the gut-isolated strains.

Under conditions of injection into the hemolymph, 100% of bees injected with the gut-isolated *S. marcescens* strains or Db11 died within 20 h, and 30% of bees injected with Ss1 died within 26 h. The delay in mortality of bees injected with Ss1 may reflect its low replication rate in the hemocoel (Fig. 4). Despite this slower replication, our results confirm that Ss1 is lethal to honey bees when present in the hemocoel. The fact that Ss1 colonizes only the hemocoel supports the hypothesis that Ss1 is transmitted by mites (18). We did not test whether the gut isolates are transmitted by Varroa mites. However, wounds from mite bites could enable these strains to access the hemolymph, which would result in death within 1 day.

Comparing gene sets possessed by *S. marcescens* kz2, kz11, kz19, Db11, and Ss1 strains allowed us to identify candidate genes associated with the virulence of the gut isolates. Although kz19 is more closely related to Ss1, it shares 298 genes uniquely with kz2 and kz11 (Fig. 1), which had similar pathogenicity attributes in our experiments. Among these genes, the flagellar genes and genes involved in iron acquisition are promising virulence candidates, and both have been shown to be important for pathogenicity in other bacteria (28, 38–40). The loss of flagellar genes from Ss1 is particularly interesting, because this loss is frequently observed in obligate endosymbionts and intracellular pathogens (41). Flagella are costly to maintain as their production significantly decreases bacterial growth (42). Transmission of Ss1 by Varroa mites (18) might make motility unimportant for transmission, resulting in the loss of flagellar genes.

*S. marcescens* strains are widespread in honey bee guts, and our results show that oral exposure to these strains can lead to lethal infections. Disruption of the gut microbiome appears to facilitate invasion by *S. marcescens* (7). Because honey bees are frequently exposed to factors that could disrupt the gut microbiome, *S. marcescens* infection may be common and potentially contributes to colony losses. Entry of even a few cells into the hemocoel can be lethal, supporting the possibility that *S. marcescens* pathogenicity is exacerbated by Varroa mites (18). In addition, the discovery of *S. marcescens* strains that can be pathogenic in honey bees, which represent a tractable model system, presents an opportunity to study how and when opportunistic pathogens are able to invade and kill their hosts.

**MATERIALS AND METHODS**

**Survey of *S. marcescens* in the honey bee gut microbiome.** Honey bees from a hive located on the UT campus, a feral hive from Utah, a commercial hive from Florida, and an organic hive from Tennessee...
were dissected, and DNA was extracted using the phenol chloroform bead-beading protocol described in reference 43. Community profiling was performed using the V4 region of the 16S RNA gene with primers 515F and 806R as described previously (44). Reaction products were purified with Ampure XP beads (Beckman Coulter). The resulting amplicons were subjected to Illumina sequencing on the MiSeq platform (2 × 250 sequencing runs) at the Genome Sequencing and Analysis Facility (GSAF) at UT. Reads were processed in QIIME (45). FASTQ files were filtered for quality by allowing a minimum Phred quality score of Q20. Forward and reverse Illumina reads were joined, and chimeric sequences were removed using the usearch6.1 detection method. Operational taxonomic units (OTUs) were clustered at 97% with the UCLUST algorithm using the July 2017 release of the SILVA (46) reference database. Sequences that did not match the SILVA data set were subsequently clustered into de novo OTUs with UCLUST. Unassigned, mitochondrial, and chloroplast reads were removed from the data set. All OTUs present in less than 0.1% abundance were removed. The samples were rarefied at a depth of 15,000 reads per sample. For each bee, presence or absence of S. marcescens was evaluated based on the taxonomic assignment of the OTUs; e.g., if a bee had an OTU assigned to the species S. marcescens after filtering, it was marked as positive. The relative abundance of S. marcescens per bee was determined based on the percentage of S. marcescens reads among the rarefied 15,000 total reads per sample (see Table S1 in the supplemental material).

Isolation of Serratia marcescens from honey bees. Honey bee guts were surveyed for the presence of S. marcescens. Using workers from a package of bees from a treatment-free apiary, we extracted the guts of hundreds of bees (of these, 21 were sampled for 16S RNA gene sequencing; see Tennessee [TN] data in Fig. 1). Gut extractions were performed as described in reference 7. The extracted guts were homogenized individually using sterile pestles and kept in 20% glycerol at −80°C. The stocks were plated onto LB agar plates and incubated at 30°C for 24 h (resident gut bacteria do not grow on this medium). Bacterial colonies that grew within 24 h were picked and replated. The isolated bacteria were PCR amplified using universal 16S rRNA gene primers 27F (5′-AGAGTTTGATCCTGGCTCAG-3′) and 1507R (5′-TACCTTGTTACGACTTCACCCCAG-3′). The PCR products were submitted for Sanger sequencing at the GSAF at UT. Three strains were identified with >97% identity to S. marcescens based on a BLASTn search against the NCBI nonredundant database (https://www.ncbi.nlm.nih.gov/).

DNA extraction and genome sequencing. DNA was extracted using the phenol chloroform bead-beading protocol described in reference 43, with the following modifications: cetyltrimethylammonium bromide (CTAB) and 2-mercaptoethanol were not used, the samples were bead beaten for only 2 min total, and the lysate was incubated at 56°C for 1 h. The resulting DNA was then submitted for Illumina MiSeq sequencing at the GSAF at UT, resulting in 1,281,027 (strain kz2), 1,994,790 (strain kz11), and 915,651 (strain kz19) reads. Reads were assembled using CLC Workbench (see Table S2 for assembly details) with default parameters, producing a total of 20 (kz2), 21 (kz11), and 24 (kz19) contigs. Genomes were annotated with the NCBI Prokaryotic Genome Annotation Pipeline with GeneMark+ version 4.3 (https://www.ncbi.nlm.nih.gov/genome/annotation_prok/).

Phylogenetic analysis. A total of 102 Serratia genomes were downloaded from NCBI as well as a genome of Yersinia enterocolitica, which was used as an outgroup. Coding sequences were translated into protein sequences using an in-house script. Usearch (47) was used to identify orthologs (based on best reciprocal hits) with thresholds of 70% sequence identity and 80% length conservation. In-house scripts were used to identify single-copy protein families present in 100% of the genomes. A total of 125 single-copy gene families were identified. These protein sequences were aligned using MAFFT v7 (48) and concatenated. The concatenated alignment was used to build a maximum likelihood phylogeny using PhyML (49) with the following parameters: GTR, Gamma4, and 100 bootstrap replicates.

Survival assays. Workers of random ages were taken from a single hive on the UT campus. Bees were immobilized at 4°C and split into the following seven treatment groups, each containing about 200 workers: (i) control, (ii) E. coli K12 and S. marcescens, (iii) strain kz2, (iv) strain kz11, (v) strain kz19, (vi) strain S11, and (vii) strain Db11.

Feeding exposure. Each group was divided into 5 cup cages containing approximately 40 bees. A solution of bacterial cells in PBS (or PBS alone for controls) at an OD of 1 was applied to irradiated bee bread and placed in each cup cage. Bees were monitored for survival every day for 1 week. The experiment was then replicated using a second hive on the UT campus.

Immersion exposure. Groups of bees were immobilized at 4°C in a 50-ml Falcon tube, and 10 μl of a 1:1 solution of bacterial cells at an OD of 0.5 in PBS and sterile sucrose (or PBS and sucrose alone for controls) was added per bee. The tube was lightly shaken for 15 s before the bees were placed into cup cages. Each group was divided into 5 cup cages containing approximately 40 bees. Bees were monitored for survival every day for 1 week. The experiment was then replicated using a second hive on the UT campus.

Hemolymph injections. A PBS bacterial solution (1 ml) at an OD of 1 was serial diluted to 10−4, and 1 μl of this solution (approximately 10 bacterial cells) was injected into the abdomen of worker bees using a fine-tipped glass capillary needle. Control bees were injected with 1 μl of sterile PBS. Each group was divided into 4 cup cages with approximately 12 bees per cup. Bees were monitored for survival for 24 h, and the number of dead bees for each treatment group was recorded approximately every 2 h. During the experiments, bees were kept in a 35°C incubator with 95% humidity to mimic hive conditions. The experiment was then replicated using a second hive on the UT campus.

Hemolymph colonization. Workers of random ages were taken from a single hive on the UT campus. Bees were immobilized at 4°C and split into five groups consisting of a control and S. marcescens strains kz2, kz11, kz19, and S11. During the experiments, bees were kept in a 35°C incubator with 95% humidity to mimic hive conditions. For each bacterial treatment, a PBS bacterial solution at an OD of 1 (1 ml) was serially diluted to 10−4, and 1 μl of this solution (approximately 10 bacterial cells) was injected...
into the abdomen of 10 worker bees per group using a fine-tipped glass capillary needle. Control bees were injected with 1 μl sterile PBS. At 3 and 6 h postinjection, hemolymph was collected from the thorax of each bee using a fine-tipped glass capillary needle. Aliquots (0.5 μl) of undiluted and 1:100-diluted hemolymph were plated onto LB agar. After overnight incubation at 30°C, the number of CFU per agar plate was determined to calculate the number of *S. marcescens* cells per microliter of hemolymph.

**Gut colonization and septicemia tests.** An E2-Crimson fluorescent protein on an RSF1010 broad-host-range backbone with spectinomycin resistance (S6) was transformed into *S. marcescens* strains k22, k211, k219, and S51. In brief, an E. coli donor strain bearing pBTK570 was mixed in a 1:1 OD ratio with the recipient *S. marcescens* strain and incubated overnight on LB plates supplemented with diaminopimelic acid (DAP). Transconjugant *S. marcescens* strains were selected with 180 μg/ml spectinomycin, and visible E2-Crimson expression confirmed successful transconjugants.

Workers of random ages were taken from a single hive on the UT campus. Bees were immobilized at 4°C and separated into groups according to strain. Half of the bees were fed and hemolymph of each bee was extracted. In brief, guts and hemolymph from each bee were homogenized in 1 ml sterile PBS. Bees were then divided into 3 groups of approximately 20 bees per group. Bees were fed a sterile sucrose solution containing 120 μg/ml spectinomycin for 2 days prior to *S. marcescens* exposure as well as for all 3 days of the experiment. During the experiments, bees were kept in a 35°C incubator with 95% humidity to mimic hive conditions. Bees were exposed to the *S. marcescens* strains using the immersion method described above. Each day, six live bees were sampled from each group as well as six dead bees if applicable. Dead bees were sampled <2 h postmortem. In the case of survival assays, only five and six bees, respectively, died during the 3-day experiment. The gut and hemolymph of each bee were extracted. In brief, guts for individual bees were homogenized in 1 ml sterile PBS and plated on LB agar containing 120 μg/ml spectinomycin using an inoculation loop. The hemolymph was collected from each bee, and 0.5-μl aliquots of undiluted hemolymph were plated on LB agar containing 120 μg/ml spectinomycin. Plates were incubated at 30°C for 16 h and then screened for the presence of blue/purple colonies.

**Bacterial growth curves.** *S. marcescens* cells were cultured in 24-well plates containing LB medium. The plates were incubated in a plate reader (Tecan) at 30°C for 48 h. Optical density was measured at 600 nm every 0.5 h, and plates were shaken for 2 min before each measurement.

**Immune response to *Serratia marcescens* infection.** Worker bees of random ages were taken from a single hive on the UT campus. Bees were immobilized at 4°C and split into the following five groups: (i) control and *S. marcescens*, (ii) strain k22, (iii) strain k211, (iv) strain k219, and (v) strain S51. Half of the bees were fed (immersion exposure), and the other half were injected with the *S. marcescens* strains as described above under “Survival assays.” During the experiments, bees were kept in a 35°C incubator with 95% humidity to mimic hive conditions. Bees were exposed to the *S. marcescens* strains using the immersion method described above. Each day, six live bees were sampled from each group as well as six dead bees if applicable. Dead bees were sampled <2 h postmortem. In the case of survival assays, only five and six bees, respectively, died during the 3-day experiment. The gut and hemolymph of each bee were extracted. In brief, guts for individual bees were homogenized in 1 ml sterile PBS and plated on LB agar containing 120 μg/ml spectinomycin using an inoculation loop. The hemolymph was collected from each bee, and 0.5-μl aliquots of undiluted hemolymph were plated on LB agar containing 120 μg/ml spectinomycin. Plates were incubated at 30°C for 16 h and then screened for the presence of blue/purple colonies.

**Immune response to *Serratia marcescens* infection.** Worker bees of random ages were taken from a single hive on the UT campus. Bees were immobilized at 4°C and split into the following five groups: (i) control and *S. marcescens*, (ii) strain k22, (iii) strain k211, (iv) strain k219, and (v) strain S51. Half of the bees were fed (immersion exposure), and the other half were injected with the *S. marcescens* strains as described above under “Survival assays.” During the experiments, bees were kept in a 35°C incubator with 95% humidity to mimic hive conditions. Bees were exposed to the *S. marcescens* strains using the immersion method described above. Each day, six live bees were sampled from each group as well as six dead bees if applicable. Dead bees were sampled <2 h postmortem. In the case of survival assays, only five and six bees, respectively, died during the 3-day experiment. The gut and hemolymph of each bee were extracted. In brief, guts for individual bees were homogenized in 1 ml sterile PBS and plated on LB agar containing 120 μg/ml spectinomycin using an inoculation loop. The hemolymph was collected from each bee, and 0.5-μl aliquots of undiluted hemolymph were plated on LB agar containing 120 μg/ml spectinomycin. Plates were incubated at 30°C for 16 h and then screened for the presence of blue/purple colonies.

**Bacterial growth curves.** *S. marcescens* cells were cultured in 24-well plates containing LB medium. The plates were incubated in a plate reader (Tecan) at 30°C for 48 h. Optical density was measured at 600 nm every 0.5 h, and plates were shaken for 2 min before each measurement.

**Immune response to *Serratia marcescens* infection.** Worker bees of random ages were taken from a single hive on the UT campus. Bees were immobilized at 4°C and split into the following five groups: (i) control and *S. marcescens*, (ii) strain k22, (iii) strain k211, (iv) strain k219, and (v) strain S51. Half of the bees were fed (immersion exposure), and the other half were injected with the *S. marcescens* strains as described above under “Survival assays.” During the experiments, bees were kept in a 35°C incubator with 95% humidity to mimic hive conditions. Bees were exposed to the *S. marcescens* strains using the immersion method described above. Each day, six live bees were sampled from each group as well as six dead bees if applicable. Dead bees were sampled <2 h postmortem. In the case of survival assays, only five and six bees, respectively, died during the 3-day experiment. The gut and hemolymph of each bee were extracted. In brief, guts for individual bees were homogenized in 1 ml sterile PBS and plated on LB agar containing 120 μg/ml spectinomycin using an inoculation loop. The hemolymph was collected from each bee, and 0.5-μl aliquots of undiluted hemolymph were plated on LB agar containing 120 μg/ml spectinomycin. Plates were incubated at 30°C for 16 h and then screened for the presence of blue/purple colonies.

**Bacterial growth curves.** *S. marcescens* cells were cultured in 24-well plates containing LB medium. The plates were incubated in a plate reader (Tecan) at 30°C for 48 h. Optical density was measured at 600 nm every 0.5 h, and plates were shaken for 2 min before each measurement.
and 1 µl was spotted in the center of the plate using a sterile flat toothpick. Swim and swarm plates were incubated at RT, 30°C, and 37°C and assessed at 24 and 48 h.

**Gelatinase hydrolysis test.** Gelatinase culture tubes were prepared as follows: 5 g of peptone, 3 g of yeast extract, and 120 g of gelatin were dissolved in 1 liter of water and autoclaved; the autoclaved medium was then dispensed into culture tubes and cooled overnight at 4°C. The gelatin was stabbed (toothpick) with bacteria (bacteria at an OD of 1 in PBS) and incubated at RT, 30°C, and 37°C. At 24 and 48 h after incubation, the gelatin was cooled and the assay was read.

**DNase activity assay.** Plates were made using BD Difco DNase test agar with methyl green. *S. marcescens* strains (bacteria at an OD of 1 in PBS) were streaked onto DNase agar plates and incubated at RT, 30°C, and 37°C. DNase activity was assessed at 24 and 48 h.

**Siderophore detection.** Siderophore detection plates were made as follows. Chromeazurol S (CAS) (60.5 mg) was dissolved in 50 ml water, and 72.9 mg CTAB was dissolved in 40 ml water. Those two solutions were then mixed with 10 ml of a 1 mM FeCl₃ hexahydrate and 10 mM HCl solution and added to LB agar medium (the recipe for 1 liter was used, and the medium was dissolved in 900 µl of water). The pH was adjusted to 6.8 and then autoclaved. *S. marcescens* strains (bacteria at an OD of 1 in PBS) were streaked onto the siderophore detection plates and incubated at RT, 30°C, and 37°C. Siderophore production was assessed at 24 and 48 h.

**Chitinase activity assay.** Chitin powder (40 g) was dissolved in 500 ml of concentrated hydrochloric acid and continuously stirred at 4°C for 1 h. The hydrolyzed chitin was washed several times with distilled water to remove the acid and to bring the pH to the range of 6 to 7. The colloidal chitin was then filtered and stored at 4°C. Chitin plates were made as follows: 3 g of MgSO₄ heptahydrate, 3 g of (NH₄)₂SO₄, 2 g of KOH, 2 g of citric acid, 15 g of agar, 4.5 g of colloidal chitin, and 15 g of bromocresol purple were dissolved in 1 liter of water; 200 µl of Tween 20 was then added; the pH was adjusted to 4.7; and the medium was autoclaved. *S. marcescens* strains (bacteria at an OD of 1 in PBS) were streaked onto the chitin plates and incubated at RT, 30°C, and 37°C. Chitinase activity was assessed at 24 and 48 h.

**Comparative genomics.** Protein families were clustered using Silix (53) with 85% identity and 70% length conservation thresholds. The identified protein families were then aligned, and HMMER (54) profiles were constructed. The HMMER profiles were used to search a local database of the *S. marcescens* kk2, kk11, kk19, and Ss1 genomes in order to confirm their presence or absence in each genome. Among the 328 unique genes identified (see Data Set S1 in the supplemental material), missing gene clusters were further analyzed. Gene clusters were defined as three or more consecutive genes that displayed conserved genome synteny. All gene clusters unique to kk2, kk11, and kk19 were analyzed and visually inspected using Geneious version 11.0.4 (55). Gene clusters were subjected to local BLAST search against *S. marcescens* D011 using Geneious version 11.0.4 (55).

**Data availability.** The genomes of *S. marcescens* strains kk2, kk11, and kk19 are available on NCBI (accession numbers PQGJ00000000, PQGI00000000, and PQGK00000000). 16S rRNA gene reads are deposited with NCBI Sequence Read BioProject under project number PRJNA483763.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at [https://doi.org/10.1128/mBio.01649-18](https://doi.org/10.1128/mBio.01649-18).

**FIG S1**, PDF file, 0.2 MB.
**FIG S2**, PDF file, 0.8 MB.
**FIG S3**, PDF file, 0.1 MB.
**FIG S4**, PDF file, 0.5 MB.
**FIG S5**, PDF file, 12.5 MB.
**FIG S6**, PDF file, 0.1 MB.
**FIG S7**, PDF file, 0.5 MB.
**TABLE S1**, PDF file, 0.04 MB.
**TABLE S2**, PDF file, 0.8 MB.
**DATA SET S1**, XLSX file, 0.1 MB.

**ACKNOWLEDGMENTS**

We thank James Burritt for generously providing the *S. marcescens* Ss1 strain and giving constructive feedback on the manuscript. Jerry Shue provided honey bee samples from a feral hive in Utah.

Funding was provided by a U.S. NIH award (1R01GM108477-01) to N.A.M. and USDA NIFA fellowships (2017-67012-26088) to K.R. and (2018-67012-28009) to K.L.C.

**Note Added after Publication**

In the originally published version, reference 56 was omitted. The reference was added to the References list and is now cited on pages 4 and 12.
REFERENCES

1. Cox-Foster DL, Conlan S, Holmes EC, Palacios G, Evans JD, Moran NA, Quan P-L, Briese T, Horning M, Geiser DM, Martinson V, vanEngelsdorp D, Kalkstein AL, Drysdale A, Hui J, Zhai J, Cui L, Hutchison SK, Simons JF, Egholm M, Petitt JS, Lipkin WI. 2007. A metagenomic survey of microbes in honey bee colony collapse disorder. Science 318:283–287. https://doi.org/10.1126/science.1146498.

2. Potts SG, Biesmeijer JC, Kremen C, Neumann P, Schweiger O, Kunin WE. 2010. Global pollinator declines: trends, impacts and drivers. Trends Ecol Evol 25:345–353. https://doi.org/10.1016/j.tree.2010.01.007.

3. Genersch E. 2010. American Foulbrood in honeybees and its causative agent, Paenibacillus larvae. J Invertebr Pathol 103:510–519. https://doi.org/10.1016/j.jip.2009.06.015.

4. Evans JD, Schwarz RS. 2011. Bees brought to their knees: microbes affecting honey bee health. Trends Microbiol 19:614–620. https://doi.org/10.1016/j.tim.2011.09.003.

5. Cremer S, Armitage SAO, Schmid-Hempel P. 2007. Social immunity. Curr Biol 17:R693–R702. https://doi.org/10.1016/j.cub.2007.06.008.

6. Brown SP, Cornforth DM, Mideo N. 2012. Evolution of virulence in bacteria: the flagellar export apparatus functions as a protein-secretion system. Proc Natl Acad Sci U S A 96:6456–6461. https://doi.org/10.1073/pnas.96.11.6456.

7. Raymann K, Shaffer Z, Moran NA. 2017. Antibiotic exposure perturbs the gut microbiota of honey bees by antibiotic treatment could increase the honey bee's vulnerability to Nosema infection. PLoS One 12:e0187505. https://doi.org/10.1371/journal.pone.0187505.

8. Brandt A, Gorenflo A, Siede R, Meixner M, Büchler R. 2016. The neonicotinoid thiacloprid, imidacloprid, and clothianidin affect the immune responses of honey bees (Apis mellifera L.). J Insect Physiol 86:40–47. https://doi.org/10.1016/j.jinsphys.2016.01.001.

9. Brandt A, Gorenflo A, Siede R, Meixner M, Büchler R. 2016. Honey bee gut microbiota assessed using deep sampling from individual worker bees. PLoS One 7:e36393.

10. Cresswell JE. 2011. A meta-analysis of experiments testing the effects of a neonicotinoid insecticide (imidacloprid) on honey bees. Ecotoxicology 20:149–157. https://doi.org/10.1111/j.1539-7416.2010.00566.x.

11. Di Prisco G, Cavaliere V, Annoscia D, Varricchio P, Caprio E, Nazzi F, Kaska M, Lysenko O, Chaloupka J. 1976. Exocellular proteases of Serratia marcescens and their toxicity to larvae of Galleria mellonella. Folia Microbiol 21:465–473. https://doi.org/10.1007/BF02876938.

12. Grimont PA, Grimont F. 1978. The genus Serratia. Annu Rev Microbiol 32:103–121. https://doi.org/10.1146/annurev.mi.32.100178.001253.

13. El Sanousi SM, El Sarag MSA, Mohamed SE. 1987. Properties of Serratia marcescens isolated from diseased honeybee (Apis mellifera) larvae. Microbiology 133:215–219. https://doi.org/10.1099/13235.

14. Toft C, Fares MA. 2008. The evolution of the flagellar assembly pathway in bacteria. Front Microbiol 7:1255.

15. DanihlíkJ,AronsteinK,Petr ˇivalskýM.2015.Antimicrobialpeptides:akeycomponentofhoneybeeinnateimmunity. J Insect Physiol 54:123–136. https://doi.org/10.1016/j.jinsphys.2016.03.001.

16. Kaska M, Lysenko O, Chaloupka J. 1976. Exocellular proteases of Serratia marcescens and their toxicity to larvae of Galleria mellonella. Folia Microbiol 21:465–473. https://doi.org/10.1007/BF02876938.

17. Raymann K, Motta EVS, Girard C, Riddington IM, Dinser JA, Moran NA. 2017. New evidence showing that the destruction of gut bacteria by antibiotic treatment could increase the honey bee’s vulnerability to Nosema infection. PLoS One 12:e0187505. https://doi.org/10.1371/journal.pone.0187505.

18. Jezieryski D, Miska J, Wierzbicki J, Nowakowski R, Grillenberger M, Quan P-L, Briese T, Hornig M, Geiser DM, Martinson V, vanEngelsdorp D, Owens SM, Betley J, Fraser L, Bauer M, Gormley N, Gilbert JA, Smith G, Knight R. 2012. Ultra-high-throughput microbial community analysis on
Serratia marcescens Is a Pathogen of Honey Bees

the Illumina HiSeq and MiSeq platforms. ISME J 6:1621–1624. https://doi.org/10.1038/ismej.2012.8.

45. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Peña AG, Goodrich JK, Gordon JI, Huttley GA, Kelley ST, Knights D, Koenig JE, Ley RE, Lozupone CA, McDonald D, Muegge BD, Pirrung M, Reeder J, Sevinsky JR, Turnbaugh PJ, Walters WA, Widmann J, Yatsunenko T, Zaneveld J, Knight R. 2010. QIIME allows analysis of high-throughput community sequencing data. Nat Methods 7:335–336. https://doi.org/10.1038/nmeth.f.303.

46. Pruesse E, Quast C, Knittel K, Fuchs BM, Ludwig W, Peplies J, Glöckner FO. 2007. SILVA: a comprehensive online resource for quality checked and aligned ribosomal RNA sequence data compatible with ARB. Nucleic Acids Res 35:7188–7196. https://doi.org/10.1093/nar/gkm864.

47. Edgar RC. 2010. Search and clustering orders of magnitude faster than BLAST. Bioinformatics 26:2460–2461. https://doi.org/10.1093/bioinformatics/btq461.

48. Katoh K, Standley DM. 2013. MAFFT multiple sequence alignment software version 7: improvements in performance and usability. Mol Biol Evol 30:772–780. https://doi.org/10.1093/molbev/mst010.

49. Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, Gascuel O. 2010. New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. Syst Biol 59:307–321. https://doi.org/10.1093/sysbio/sys010.

50. Yang X, Cox-Foster DL. 2005. Impact of an ectoparasite on the immunity and pathology of an invertebrate: Evidence for host immunosuppression and viral amplification. Proc Natl Acad Sci U S A 102:7470–7475. https://doi.org/10.1073/pnas.0501860102.

51. Kuster RD, Boncrustiani HF, Rueppell O. 2014. Immunogene and viral transcript dynamics during parasitic Varroa destructor mite infection of developing honey bee (Apis mellifera) pupae. J Exp Biol 217:1710–1718. https://doi.org/10.1242/jeb.097766.

52. Evans JD. 2006. Beepath: an ordered quantitative-PCR array for exploring honey bee immunity and disease. J Invertebr Pathol 93:135–139. https://doi.org/10.1016/j.jip.2006.04.004.

53. Miele V, Penel S, Duret L. 2011. Ultra-fast sequence clustering from similarity networks with SiLiX. BMC Bioinformatics 12:116. https://doi.org/10.1186/1471-2105-12-116.

54. Johnson LS, Eddy SR, Portugaly E. 2010. Hidden Markov model speed heuristic and iterative HMM search procedure. BMC Bioinformatics 11:431. https://doi.org/10.1186/1471-2105-11-431.

55. Kearse M, Moir R, Wilson A, Stones-Havas S, Cheung M, Sturrock S, Buxton S, Cooper A, Markowitz S, Duran C, Thierer T, Ashton B, Meintjes P, Drummond A. 2012. Geneious Basic: an integrated and extendable desktop software platform for the organization and analysis of sequence data. Bioinformatics 28:1647–1649. https://doi.org/10.1093/bioinformatics/bts199.

56. Leonard SP, Perutka J, Powell JE, Geng P, Richhart DD, Byrom M, Kar S, Davies BW, Ellington AD, Moran NA, Barrick JE. Genetic Engineering of Bee Gut Microbiome Bacteria with a Toolkit for Modular Assembly of Broad-Host-Range Plasmids. ACS Synth Biol. 2018 May 18;7(5):1279–1290. https://doi.org/10.1021/acssynbio.7b00399.