

# Horizontal Gene Transfer to a Defensive Symbiont with a Reduced Genome in a Multipartite Beetle Microbiome

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**ABSTRACT** Symbiotic mutualisms of bacteria and animals are ubiquitous in nature, running a continuum from facultative to obligate from the perspectives of both partners. The loss of functions required for living independently but not within a host gives rise to reduced genomes in many symbionts. Although the phenomenon of genome reduction can be explained by existing evolutionary models, the initiation of the process is not well understood. Here, we describe the microbiome associated with the eggs of the beetle *Lagria villosa*, consisting of multiple bacterial symbionts related to *Burkholderia gladioli*, including a reduced-genome symbiont thought to be the exclusive producer of the defensive compound lagriamide. We show that the putative lagriamide-producing symbiont is the only member of the microbiome undergoing genome reduction and that it has already lost the majority of its primary metabolism and DNA repair pathways. The key step preceding genome reduction in the symbiont was likely the horizontal acquisition of the putative lagriamide *lga* biosynthetic gene cluster. Unexpectedly, we uncovered evidence of additional horizontal transfers to the symbiont's genome while genome reduction was occurring and despite a current lack of genes needed for homologous recombination. These gene gains may have given the genome-reduced symbiont a selective advantage in the microbiome, especially given the maintenance of the large *lga* gene cluster despite ongoing genome reduction.

**IMPORTANCE** Associations between microorganisms and an animal, plant, or fungal host can result in increased dependence over time. This process is due partly to the bacterium not needing to produce nutrients that the host provides, leading to loss of genes that it would need to live independently and to a consequent reduction in genome size. It is often thought that genome reduction is aided by genetic isolation—bacteria that live in monocultures in special host organs, or inside host cells, have less access to other bacterial species from which they can obtain genes. Here, we describe exposure of a genome-reduced beetle symbiont to a community of related bacteria with nonreduced genomes. We show that the symbiont has acquired genes from other bacteria despite going through genome reduction, suggesting that isolation has not yet played a major role in this case of genome reduction, with horizontal gene gains still offering a potential route for adaptation.

**KEYWORDS** *Burkholderia*, insects, metagenomics, natural products, symbiosis

**M**utualistic symbioses between animals and bacteria, widespread in nature, serve a variety of functions such as biosynthesis of nutrients not found in the host's diet (1, 2) and protection from predation (3–5) or infection (6–8). Such relationships exist on a continuous spectrum of dependency and exclusivity from the perspectives of both the host and the symbiont. Symbionts generally become obligate after a prolonged

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period of exclusive association with the host, and the symbionts that become obligate tend to carry out highly important functions for the host (9–11). For example, mitochondria and chloroplasts, organelles that are required for energy production and carbon fixation in eukaryotic and plant cells, originated from endosymbiotic capture of alphaproteobacteria and cyanobacteria ~1.2 billion years ago (Bya) and ~900 million years ago (Mya), respectively (12). The acquisition of these organelles allowed the diversification of eukaryotic species (13). More recently, aphids evolved to feed on plant sap depleted of several essential amino acids only through capture of an endosymbiont that can synthesize these nutrients, namely, *Buchnera aphidicola*, ~160 to 280 Mya (14, 15). In these cases, the microbial symbiont has lost the ability to live independently of the host, and the hosts are also dependent on their symbionts.

The mechanism by which symbionts become obligate is through loss of genes required for independent (but not host-associated) life, leading to an overall reduction in genome size (9–11). This gene loss is the result of relaxed selection of genes for functions provided by the host and also of increased genetic drift as a consequence of the presence of small effective populations in strict vertical transmission (16, 17). A general mutational bias toward deletion in bacteria (18) combined with many successive population bottlenecks that allow the fixation of slightly deleterious mutations (10) mediates general gene degradation and genome reduction in symbionts. Also, a low level of recombination between different strains is often thought to contribute to this process. While these mechanisms are largely thought to be nonadaptive, there are also examples of free-living bacteria with reduced genomes, where reduction is thought to be the result of strong selection for reduced metabolic expenditure (19–21). In symbionts, there is also evidence that increases in AT content and reductions in genome size could be selected for to reduce the metabolic burden on the host (22, 23). The early stages of this process are manifested by a proliferation of nonfunctional pseudogenes and a decrease in coding density in the genome (11) before the intergenic sequences are lost to eventually give tiny (<1-Mbp) genomes (9, 10). While there is a robust model for the evolutionary forces that drive this process once a symbiont becomes host restricted, it is unknown how free-living bacteria first transition to an endosymbiotic lifestyle and start on the road to genome reduction (10).

The beetle subfamily Lagriinae offers an opportunity to examine this issue. Various Lagriinae species have evolved special symbiont-bearing structures that serve to facilitate the vertical transmission of bacteria (24). Symbionts are stored extracellularly in two accessory glands associated with the female reproductive system, allowing contamination of the egg surface with symbionts (24). These symbionts later colonize three compartments forming from invaginations of the dorsal cuticle in the larva (24). Beetles are typically coinfectected with multiple symbiont strains related to the plant pathogen *Burkholderia gladioli* that are secreted onto the surface of eggs as they are laid (24). In the species *Lagria villosa*, a South American soybean pest, at least one symbiotic *B. gladioli* strain (Lv-StA) has been cultured and is still capable of infecting plants (24). The same strain produces antibacterial and antifungal compounds that can protect the beetle's eggs from infection (24, 25). This is consistent with the hypothesis that the *B. gladioli* symbionts evolved from plant pathogens to become beetle mutualists. However, in field collections of *L. villosa*, Lv-StA is found only sporadically and is never highly abundant (26). Instead, the most abundant strain identified is often the uncultured Lv-StB strain (26), which we previously found harbors a biosynthetic gene cluster (BGC) predicted to produce lagriamide, a defensive compound found in field egg collections (6). We previously found through metagenomic sequencing that the genome of Lv-StB was much smaller than that of Lv-StA, suggesting that it has undergone genome reduction (6). It would seem that, while *L. villosa* has multiple options for symbionts that produce potential chemical defenses, only a subset have specialized as obligate mutualists. The presence of multiple related strains in this system, with selective genome reduction of a single strain, could potentially shed light on why the genomes of some symbionts become reduced.

Here, we show that in the *L. villosa* microbiome, Lv-StB is uniquely undergoing

genome reduction, despite other community members possessing biosynthetic pathways for potentially defensive molecules. We also suggest that this process was likely driven not only by horizontal acquisition of the putative lagriamide pathway but also by loss of genes that limit cell division and translation and by gain of *zot*, encoding zonular occludens, a toxin also found in *Vibrio cholerae* that aids invasion of host membranes. Further, we present evidence that these horizontal gene transfers occurred concurrently with genome reduction, suggesting that complete genetic isolation is not a main driving force for the reduction process. (This article was submitted to an online preprint archive [27]).

## RESULTS AND DISCUSSION

**Selective genome reduction of strain Lv-StB in *Lagria villosa*.** We previously analyzed the metagenome of eight *L. villosa* egg clutches (6), using our binning pipeline Autometa (28). This method has the advantage that it can separate noncharacterized eukaryotic contamination from metagenomes, and it uses multiple factors (nucleotide composition, sequence homology, the presence of single-copy marker genes, and level of coverage) to accurately produce bins from individual data sets. Because we had implemented several bug fixes and small improvements to the pipeline since our original analysis, we re-ran Autometa on the same metagenomic assembly. Despite some minor differences, the new bins were broadly similar to our previous results (see Data Set S1A in the supplemental material), with 19 bins. As before, the Lv-StB bin had the highest coverage, at 1,977× (294 contigs;  $N_{50}$ , 8,138 bp; see Data Set S1B for details of other bins), such that the constituent contigs are unlikely to represent repeats from lower-coverage bins. We classified the bins according to a new standardized bacterial taxonomy utilized by the Genome Taxonomy Database (GTDB) that minimizes polyphyletic taxa and standardizes divergence between taxa of the same rank (29). Notably, the GTDB taxonomy reclassifies betaproteobacteria as being under class *Gammaproteobacteria*. By this classification, all bins were in class *Gammaproteobacteria*, in three different orders: *Betaproteobacterales*, *Pseudomonadales*, and *Xanthomonadales* (Data Set S1C). The most abundant bins were all in the family *Burkholderiaceae*, with the highest abundance corresponding to the Lv-StB strain previously found to harbor the putative lagriamide BGC (6). Interestingly, the average nucleotide identity (ANI) of Lv-StB to the reference *B. gladioli* genome in GTDB (strain ATCC 10248) is 85.7%, much lower than the 95% cutoff suggested for species identifications (30) (Data Set S1C). This divergence suggests that Lv-StB is a novel species in the genus *Burkholderia*, even though we previously classified it as *B. gladioli* on the basis of 16S rRNA gene sequence (6, 26), and therefore we refer to the strain here as “*Burkholderia* sp. Lv-StB.” Likewise, most bins were found to be novel species, with one (DBSCAN\_round2\_3) being divergent enough to be a representative of a novel genus in the family *Burkholderiaceae*. Notably, the cultured *B. gladioli* strain that we had previously isolated from *L. villosa* eggs, Lv-StA (6, 24) was not present in this metagenome.

The bins obtained had a range of different sizes (Table 1; see also Data Set S1B), which could be due to genome reduction or poor assembly and/or binning of a larger genome, which is often observed if there are many related strains in a metagenome (28). As part of the binning procedure, genome completeness was estimated based on the presence of 139 single-copy marker genes (31) (Data Set S1B). However, as some complete genomes of genome-reduced symbionts have low apparent completeness by this measure (32, 33), this figure cannot be used alone to determine the size of an incompletely assembled genome. Conversely, even the drastically reduced genomes of intracellular obligate insect symbionts have been found to almost universally maintain certain genes that we refer to here as “core genes,” involved in replication, transcription, protein folding/stability, tRNA modification, sulfur metabolism, RNA modification, and translation (9). We would expect, therefore, that a well-assembled reduced genome would contain a near-complete core gene set but not necessarily the whole set of 139 single-copy marker genes. Conversely, incompletely assembled genomes are likely to

**TABLE 1** Genome characteristics of *Burkholderia* symbiont Lv-StB, its relative *B. gladioli* Lv-StA, and other bins obtained from the metagenome

Genome	Coverage (×)	Size (% of closest relative) <sup>a</sup>	% of core genes	Category
<i>B. gladioli</i> Lv-StA	NA <sup>c</sup>	96.2	95.2	Nonreduced
<i>B. gladioli</i> sp. Lv-StB	1,977	23.5	85.7	Reduced
DBSCAN_cluster_round5_1	355	105	92.9	Nonreduced
DBSCAN_cluster_round2_3	298	74.7	95.2	Nonreduced
DBSCAN_cluster_round6_18	207	18.2	36.9	Incomplete
DBSCAN_cluster_round4_6	170	86.1	95.2	Nonreduced
DBSCAN_cluster_round43_4	142	4.74 <sup>b</sup>	38.1	Incomplete
DBSCAN_cluster_round8_1	94	39.6	25	Incomplete
DBSCAN_cluster_round3_0	90	86.1	81	Nonreduced
DBSCAN_cluster_round5_3	71	77.2	81	Nonreduced
DBSCAN_cluster_round1_2	68	128	92.9	Nonreduced
DBSCAN_cluster_round6_14	61	20.9	39.3	Incomplete
DBSCAN_cluster_round66_39	47	0.351 <sup>b</sup>	25	Incomplete
DBSCAN_cluster_round7_14	37	5.3	16.7	Incomplete
DBSCAN_cluster_round34_1	34	40.9	20.2	Incomplete
DBSCAN_cluster_round7_0	24	62.8	31	Incomplete
DBSCAN_cluster_round44_0	18	58.4	34.5	Incomplete
DBSCAN_cluster_round1_3	16	103	94	Nonreduced
DBSCAN_cluster_round4_0	8	102	89.3	Nonreduced
DBSCAN_cluster_round4_12	5	63.1	47.6	Incomplete

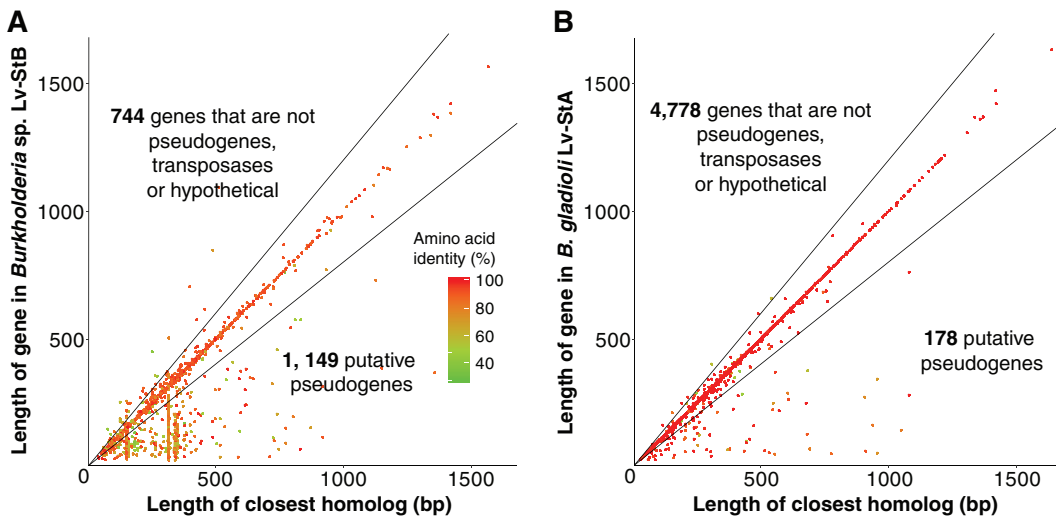
<sup>a</sup>Calculated relative to the genome of the closest relative identified by GTDB-Tk (see Data Set S1B).

<sup>b</sup>Calculated relative to the average size of 517 *Pseudomonas* genomes taken from the GTDB database.

<sup>c</sup>NA, not applicable.

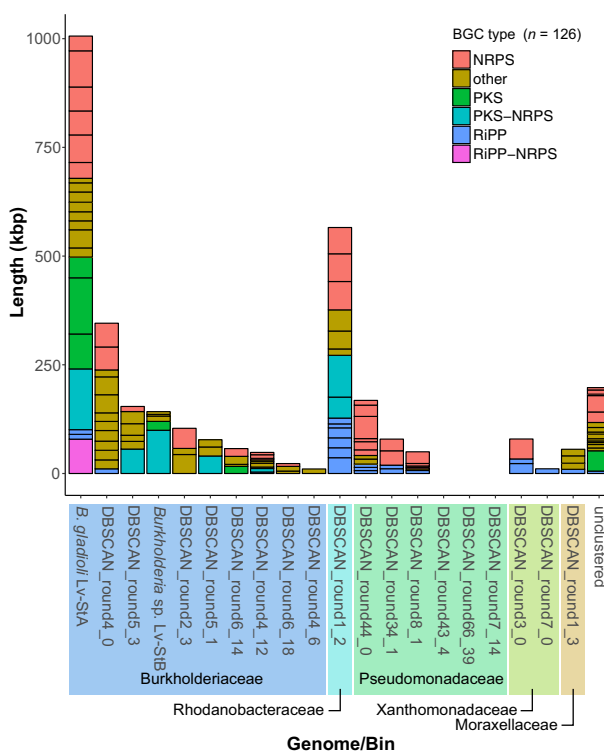
be missing a significant number of core genes that are required even in symbionts with highly reduced genomes. We examined the presence of core genes in all metagenomic bins, as well as in *B. gladioli* Lv-StA for comparison (Table 1; see also Data Set S1D). Nine bins were close in size to the genome of their respective closest relatives, while maintaining most core genes, and are classified as "nonreduced," and 10 bins were small but also lacked a significant fraction of core genes and are classified as "incomplete." Only the Lv-StB genome can be classified as reduced, on the basis of its reduced size compared to its close relative (2.07 Mbp, 23.5%) and maintenance of most core genes (85.7%). It is worth noting here that the "core" genes missing from the Lv-StB genome include genes essential for translation (*infB*, *trpS*, the tRNA-Met gene, and tRNA-Gln gene; Data Set S1D), which is probably explained by incomplete assembly of the draft-quality genome. The high level of maintenance of core genes nevertheless suggests that the assembly is missing a relatively small amount of the genome and that this genome is reduced in size. The Lv-StB bin exhibited additional features of genomes undergoing reduction, namely, reduced GC% content compared to the *B. gladioli* reference genome (58.7% versus 67.9%) and a proliferation of pseudogenes accounting for 45.29% of the annotated open reading frames (ORFs) (Fig. 1; see also Data Set S1C and E). Because of this, the Lv-StB genome exhibits low coding density (59.04%), and it also possesses a large number ( $n = 159$ ) of ORFs containing transposases. Both of these characteristics are hallmarks of symbionts in the early stages of genome reduction, whose genomes contain high numbers of pseudogenes and genome rearrangements (9). The Lv-StB genome also contains a low number of genes compared with its free-living relative, with 744 ORFs that are not pseudogenes, transposase genes, or hypothetical genes versus 4,778 such genes in *B. gladioli* Lv-StA.

**Diversity of biosynthetic gene clusters in the *L. villosa* microbiome.** Because we had previously isolated the nonreduced *B. gladioli* Lv-StA strain from *L. villosa* (6) and had found it to produce protective compounds despite having sporadic distribution and low abundance in field-collected beetles, we asked whether BGCs were a unique feature of Lv-StB in the metagenome or whether other community members have the biosynthetic machinery for potential chemical defenses. AntiSMASH (34) searches

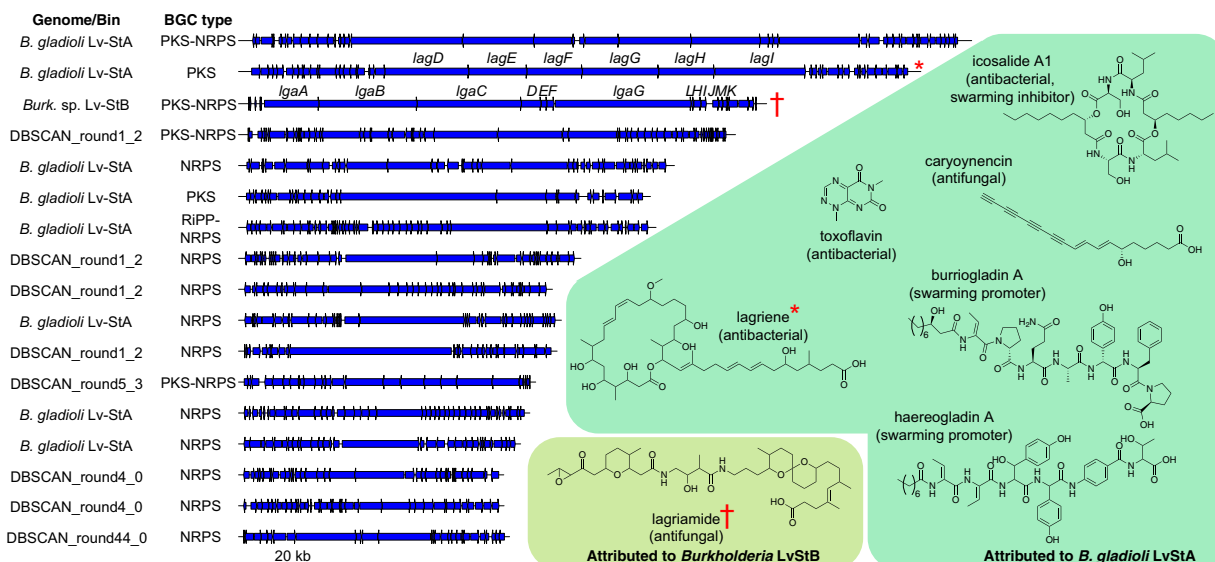


**FIG 1** Comparison of the lengths of genes in the *Burkholderia* sp. Lv-StB genome (A) and *B. gladioli* Lv-StA (B) with the closest homologs identified through BLAST searches against the NR database (see Materials and Methods). Genes which are less than 80% of the length of the closest relative (i.e., below the lower black line) are putatively assigned as pseudogenes, as described previously (4, 121). Note that two vertical groupings of pseudogenes in Lv-StB correspond to multiple copies of a 155-bp hypothetical gene and an IS5 family transposase.

revealed a total of 105 BGCs in this metagenome, as well as 21 BGCs in the *B. gladioli* Lv-StA genome (Fig. 2), with variable BGC content ranging from 0 to 566 kbp per bin (0 to 14 BGCs, with 16 BGCs in the unclustered bin), while the *B. gladioli* Lv-StA genome contained 1,006-kbp BGCs. This indicates that Lv-StB is not the only strain in the egg microbiome with the potential to produce complex natural products and that many



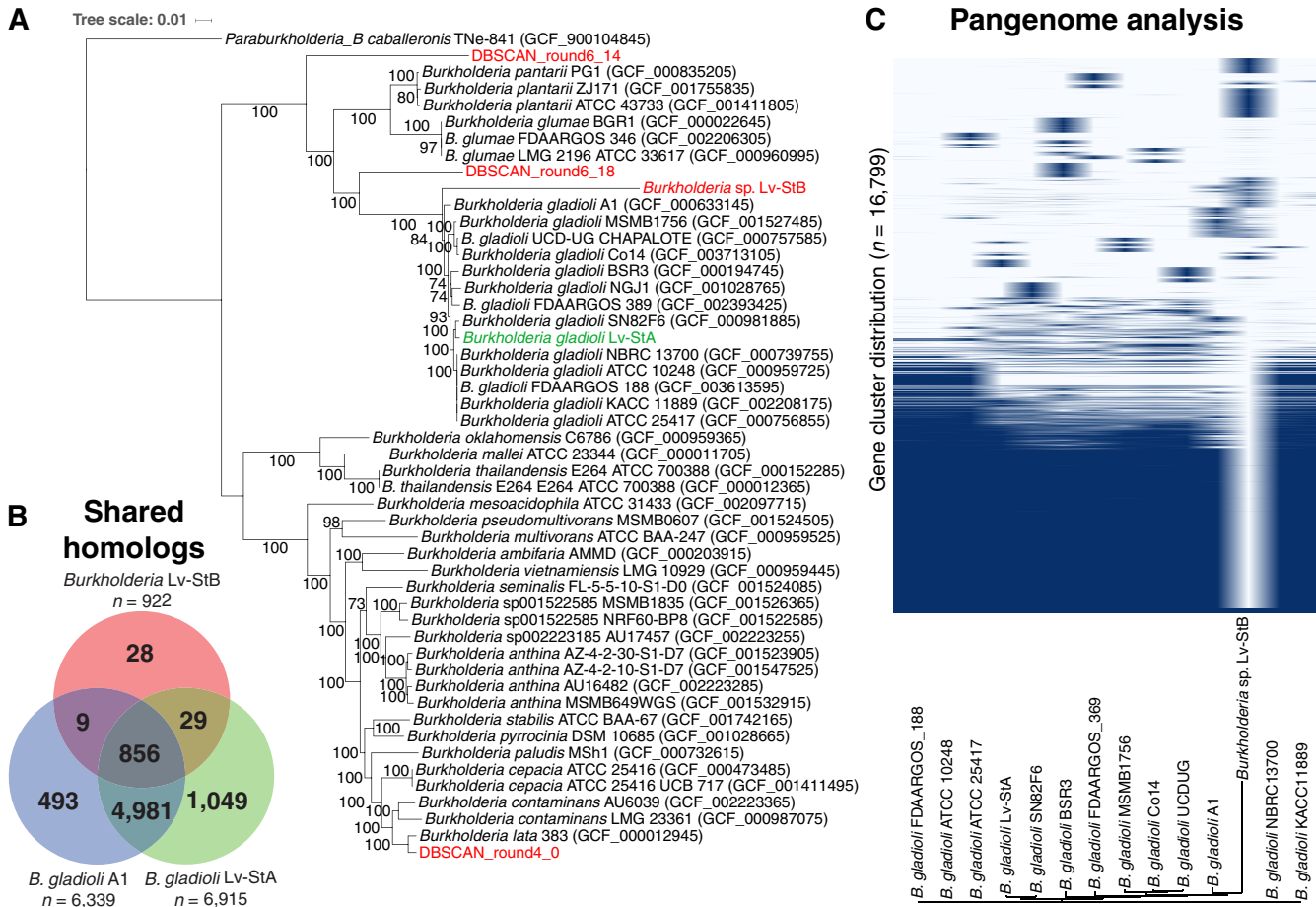
**FIG 2** Distribution of biosynthetic gene clusters (BGCs) among the *L. villoso* metagenome bins and the genome of *B. gladioli* Lv-StA. Colors indicate the type of BGC annotated by antiSMASH, simplified into polyketide synthase (PKS), nonribosomal peptide synthetase (NRPS), ribosomally synthesized and post-translationally modified peptides (RiPP), and “other” (126 identified) (34).



**FIG 3** Schematics of all BGCs with length greater than 50 kbp assembled from the *L. villosa* metagenome and in the *B. gladioli* Lv-StA isolate genome among 126 identified by antiSMASH (left). Shown on the right are structures of compounds putatively assigned to BGCs in the *B. gladioli* Lv-StA genome (24, 25) or putatively assigned to a BGC in the *Burkholderia* sp. Lv-StB genome (6). Structures highlighted with red symbols have been attributed to the indicated BGCs.

strains harbor multiple BGCs. In the metagenome, bins in the family *Burkholderiaceae* collectively contained the most BGCs by length (963 kbp), followed by a single bin in family *Rhodanobacteraceae* (DBSCAN\_round1\_2, 566 kbp, Fig. 2). This distribution suggests that although *Burkholderiaceae* appear to be an important reservoir of BGCs in the *L. villosa* egg microbiome, other groups have significant biosynthetic potential. Among the 126 BGCs detected in the metagenome and in the *B. gladioli* Lv-StA genome, there were 17 that were >50 kbp in length and were predicted to produce complex nonribosomal peptides or polyketides (Fig. 3). Two of these have been putatively assigned to production of the antibiotic lagriene in Lv-StA (24) and of the antifungal lagriamide in Lv-StB (6), whereas five other small molecules known to be produced by Lv-StA have been assigned to shorter BGCs (24, 25). We compared the 126 identified BGCs using BIG-SCAPE (35) and found only 7 examples of BGCs occurring in multiple strains, indicating that the biosynthetic potential in the metagenome and *B. gladioli* Lv-StA is largely nonredundant. Taken together, the data suggest that there is a large amount of undefined biosynthetic potential for production of potentially defensive small molecules in *L. villosa* symbionts, beyond *B. gladioli* Lv-StA and *Burkholderia* sp. Lv-StB.

**Divergence between Lv-StB and the closest free-living relative.** We constructed a phylogenetic tree of metagenomic bins assigned to the genus *Burkholderia* as well as *B. gladioli* Lv-StA, on the basis of 120 marker genes (Fig. 4A). This showed that *Burkholderia* sp. Lv-StB is most highly related to the *B. gladioli* clade but is divergent from it. We calculated genome-wide ANI values for pairs of *Burkholderia* genomes and found that *B. gladioli* strains shared between 97% and 100% ANI whereas *Burkholderia* sp. Lv-StB shared at most 85.79% ANI with *B. gladioli* A1 (Data Set S1F). During genome reduction, symbionts are known to undergo rapid evolution due to the loss of DNA repair pathways (as found in the Lv-StB genome; see below) and the relaxation of selection (9), and so the divergence of Lv-StB from *B. gladioli* may have been accelerated relative to free-living lineages. Genome-reduced symbionts have often been vertically transmitted for evolutionary timescales and across host speciation events, and therefore it is possible to calculate evolution rates where related symbionts occur in hosts with known divergence times inferred from fossil records. Such estimates in insect symbionts range widely over 3 orders of magnitude, but more recent ant and



**FIG 4** (A) Maximum-likelihood multilocus species tree of metagenomic bins classified in the genus *Burkholderia*, plus *B. gladioli* Lv-StB, using 120 marker gene protein sequences. Bootstrap proportions greater than 70% are indicated to the left of each node as a percentage of 1,000 replicates. Metagenomic bins are shown in red, while the *L. villoso*-associated isolate *B. gladioli* Lv-StA is shown in green. (B) Shared homologous gene groups in *Burkholderia* Lv-StA, *B. gladioli* A1, and *B. gladioli* Lv-StB, after discounting pseudogenes. Note that homologous groups are counted only once per genome (i.e., corresponding to collapsing paralogs), and therefore the indicated counts are lower than the absolute gene counts. Also, 492 genes in the Lv-StB genome were singletons and not included in homologous groups. (C) Hierarchical clustering of homologous gene clusters, showing presence and absence in *Burkholderia* sp. Lv-StB and closely related strains of *B. gladioli*.

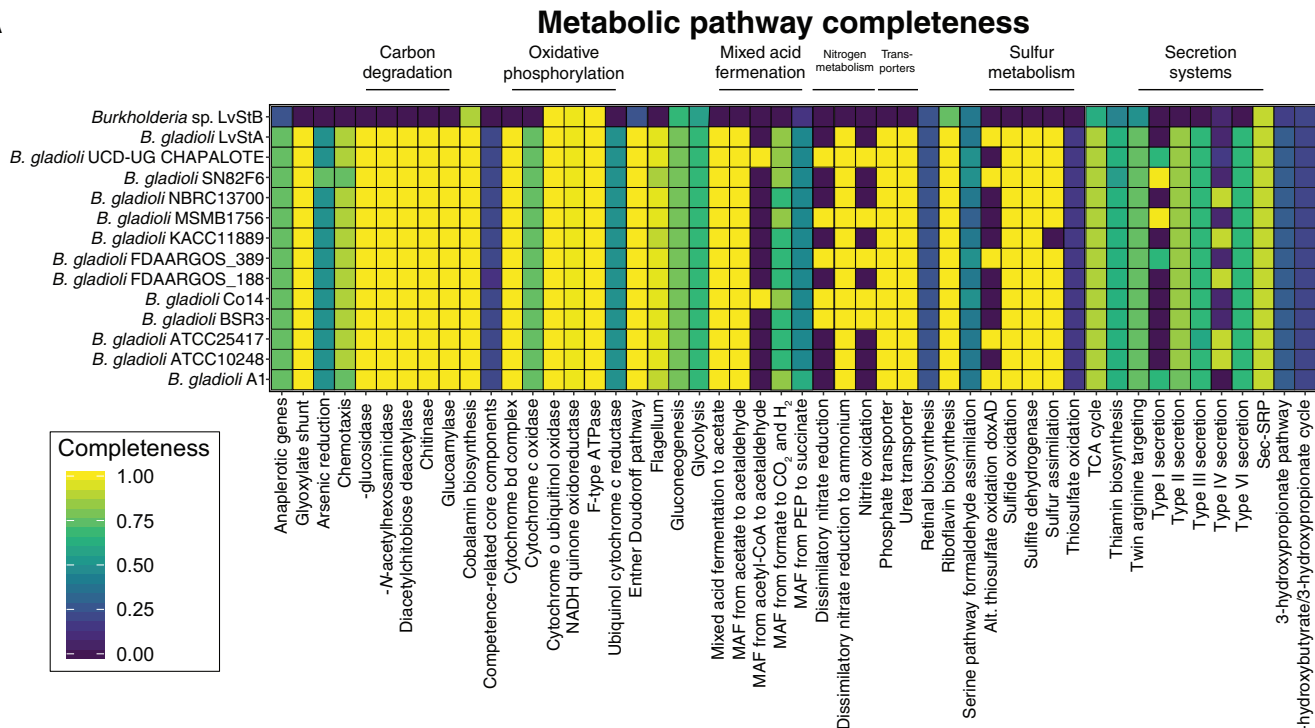
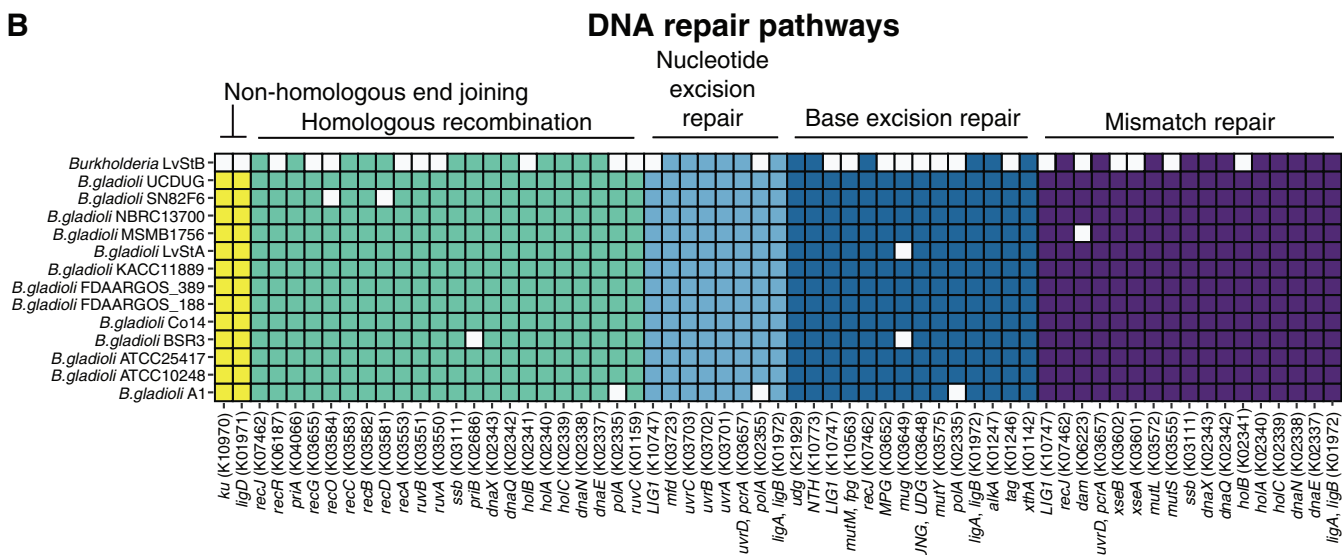
sharp-shooter symbiont lineages (established <50 Mya for “*Candidatus Baumannia cicadellinicola*,” *Blochmannia obliquus*, *Bl. pennsylvanicus*, and *Bl. floridanus*) show high rates of divergence per synonymous site per year (dS/t) of between  $1.1 \times 10^{-8}$  and  $8.9 \times 10^{-8}$  (36) (the divergence rates used here can be found in Table S1 in the supplemental material). Because of the large number of pseudogenes in the Lv-StB genome, we reasoned that it is likely to be a recent symbiont and therefore used these rates to estimate times of divergence between Lv-StB and *B. gladioli* A1. We found a dS rate of 0.5486 per site between these genomes and calculated divergence times of 6.15, 8.55, 6.93, and 49.76 My rates for *Bl. floridanus*, *Bl. pennsylvanicus*, *Bl. obliquus*, and “*Candidatus Baumannia cicadellinicola*” divergence rates, respectively (Table S1) (36). We should note here that these time estimates are very approximate and are possibly overestimates as symbiont evolution rates are likely not constant, with particularly rapid evolution occurring during lifestyle transitions (11). The range of these estimated divergence times suggests that the common ancestor of *Burkholderia* sp. Lv-StB and *B. gladioli* existed after the evolution of symbiont-bearing structures in Lagriinae beetles (see below).

We then sought to quantify the conservation of genes in Lv-StB compared to 13 related *B. gladioli* strains by identifying homologous gene groups in the entire set of nonpseudogenes in these strains with OMA (37). This pipeline aims to identify ortholo-

gous groups (OGs) while discounting paralogs among the genes in a given set of genomes (38). Of the 1,388 genes in Lv-StB that are not pseudogenes (Data Set S1B), 497 were not included in any OMA orthologous group (see below). A crude analysis of the OMA groups in Lv-StB, *B. gladioli* Lv-StA, and *B. gladioli* A1 revealed that Lv-StB retains a small subset of the groups found in both Lv-StA and A1 and has few unique groups (Fig. 4B), suggesting that Lv-StB has lost many of the genes conserved in *B. gladioli*. Consistent with this notion, we visualized the pangenome of *B. gladioli* and Lv-StB with Roary (39) (Fig. 4C) and found a large number of gene clusters that are conserved in *B. gladioli* but not Lv-StB. The gene clusters that are more variable among *B. gladioli* are also generally not found in Lv-StB. Conversely, there were gene clusters found in Lv-StB that are not present in *B. gladioli* strains, and these clusters may have been obtained by horizontal transfer after the divergence of Lv-StB or, alternatively, were lost in *B. gladioli*.

Remarkably, among the 1,149 pseudogenes detected in the Lv-StB genome, 976 were hypothetical and 129 were transposases, leaving only 44 that were recognizable (Data Set S1G). Pseudogenes were detected by identifying those that appeared truncated more than 20% compared to the closest BLAST hit (see Materials and Methods). While this methodology does not attempt to compare lengths to a consensus gene length (for example, to account for start site annotation variance) and while a firm cutoff for pseudogene length truncation has not been defined, the low number of recognizable pseudogenes suggests that we are not overannotating pseudogenes. The recognizable set of pseudogenes included a number of important genes in the categories noted to be depleted as described below. For instance, the DNA polymerase I (Pol I) gene (*polA*) appears to have been disrupted by a transposase, which is now flanked by two DNA Pol I fragments (E5299\_1120 and E5299\_01122). Likewise, the *uvrC* gene (E5299\_00503), a component of the nucleotide excision repair system (40), is also present as a truncated gene adjacent to a transposase gene. There were also pseudogenes involved in the Entner-Doudoroff and glycolysis energy-producing pathways (corresponding to phosphogluconate dehydratase [41] and glucokinase [42]), as well as purine biosynthesis (corresponding to phosphoribosylglycinamide formyltransferase [43] and phosphoribosylformylglycinamide synthase [44]). Interestingly, we found two pseudogenes that negatively regulate cell division and translation. Septum protein Maf is a nucleotide pyrophosphatase that has been shown to arrest cell division, especially after transformation or DNA damage (45). Deletion of the *Escherichia coli* gene for homolog YhdE increased the growth rate, while overexpression decreased the growth rate (46). Therefore, the loss of Maf in Lv-StB would be expected to increase the rate of cell division and reduce the conversion of nucleotides, which it probably obtains from the host, to the monophosphates. The gene for the energy-dependent translational throttle protein EttA was also found to be truncated. This protein slows translation through interacting with the ribosome in both the ATP- and ADP-bound forms (47, 48). Under energy-depleted conditions (i.e., high ADP levels), EttA was found to stabilize ribosomes and prevent commitment of metabolic resources; thus, the deletion mutant displayed reduced fitness during extended stationary phase (47). However, under circumstances where the host supplies ample nucleotides, the loss of *ettA* would be expected to increase translation rates.

**Degradation of primary metabolic pathways in *Burkholderia* sp. Lv-StB.** While the Lv-StB genome is of draft quality and thus may be missing some genes due to poor assembly, we found pervasive metabolic gaps across functional categories (Data Set S1A). Lv-StB is deficient in many metabolic pathways that are complete in related *B. gladioli* strains (Fig. 5), including the glyoxylate shunt (49) and various carbon degradation pathways as well as sulfur and nitrogen metabolism. Lv-StB appears incapable of making any of the following compounds due to the absence of several biosynthesis genes: thiamine, riboflavin, nicotinate, pantothenate, vitamin B12, and biotin. Likewise, there were deficiencies in amino acid biosynthesis (see Fig. S1 in the supplemental material). We predict that Lv-StB would be able to make chorismate, isoleucine, leucine,

**A****B**

**FIG 5** Completeness of metabolic and DNA repair pathways in *Burkholderia* sp. Lv-StB in comparison to closely related strains of *B. gladioli*. Note that presence/absence data shown here reflect detection in the draft-quality Lv-StB genome, and there is a possibility that genes are missing due to poor assembly. (A) Completeness of various metabolic pathways as determined by KEGG decoder. Note that categories that were not found in any of the examined genomes have been removed. MAF, mixed-acid fermentation; CoA, coenzyme A; TCA, tricarboxylic acid. (B) Presence (colored squares) and absence (white squares) of genes in the different DNA repair pathways in Lv-StB and related *B. gladioli* strains.

ornithine, proline, and threonine but likely lacks the ability to make aromatic amino acids, serine, methionine, lysine, histidine, cysteine, glutamine, and arginine due to the absence of several key genes in these pathways. The genome of Lv-StB also lacks genes involved in chemotaxis and flagella, suggesting that after the symbiont mixture is spread on eggs, the colonization of the dorsal cuticular structures in the embryo (24) does not require symbiont motility. Interestingly, the Lv-StB genome includes a trimeric autotransporter adhesin (TAA) related to SadA (50), which is involved in the pathogenicity of *Salmonella enterica* serovar Typhimurium by aiding cell aggregation, biofilm

formation, and adhesion to human intestinal epithelial cells. TAAs are found in *Proteobacteria* and consist of anchor, stalk, and head domains, among which the head forms the adhesive component (51). The bacterial honey bee symbiont *Snodgrassella alvi* is hypothesized to utilize TAAs in combination with other extracellular components during colonization of the host gut (52), and similar genes were identified in *S. alvi* symbionts in bumble bees and are predicted to perform a similar role (53). Therefore, this gene may play a role in the adhesion of Lv-StB cells to *L. villoso* eggs.

The Lv-StB genome is also missing several enzymes involved in glycolysis, most notably phosphoglycerate kinase and phosphoenolpyruvate carboxykinase (among others), which would suggest that Lv-StB has lost the ability to perform glycolysis. The loss of glycolysis is often compensated for by an alternative pathway, such as the pentose phosphate or Entner-Doudoroff pathway (54). This is not the case for Lv-StB, where both the glucose-6-phosphate dehydrogenase and 6-phosphogluconolactonase genes appear to be missing from the genome. The citrate cycle is largely complete, except that it is missing pyruvate carboxylase, the enzyme that converts pyruvate to oxaloacetate. However, phosphoenolpyruvate carboxylase is present in Lv-StB (E5299\_00983) and may alternatively be used for the production of oxaloacetate from exogenous phosphoenolpyruvate (55), as alternative pathways for the supply of oxaloacetate are also incomplete. Lv-StB is also missing all genes required to form cytochrome *c* oxidase and cytochrome *b-c* complexes. However, all genes required to encode NADH:quinone oxidoreductase (complex I), succinate dehydrogenase (complex II), and cytochrome *o* ubiquinol oxidase are present, along with all genes required for the F-type ATPase. The lack of complex III would likely result in a decreased rate of ATP production in Lv-StB as observed in fungi with alternative oxidases that bypass complexes III and IV (56). Lv-StB may be similar to the psyllid endosymbiont "Candidatus Liberibacter asiaticus," which has lost both key glycolysis genes and key glyoxalase genes and instead relies on the scavenging of ATP from the host (57).

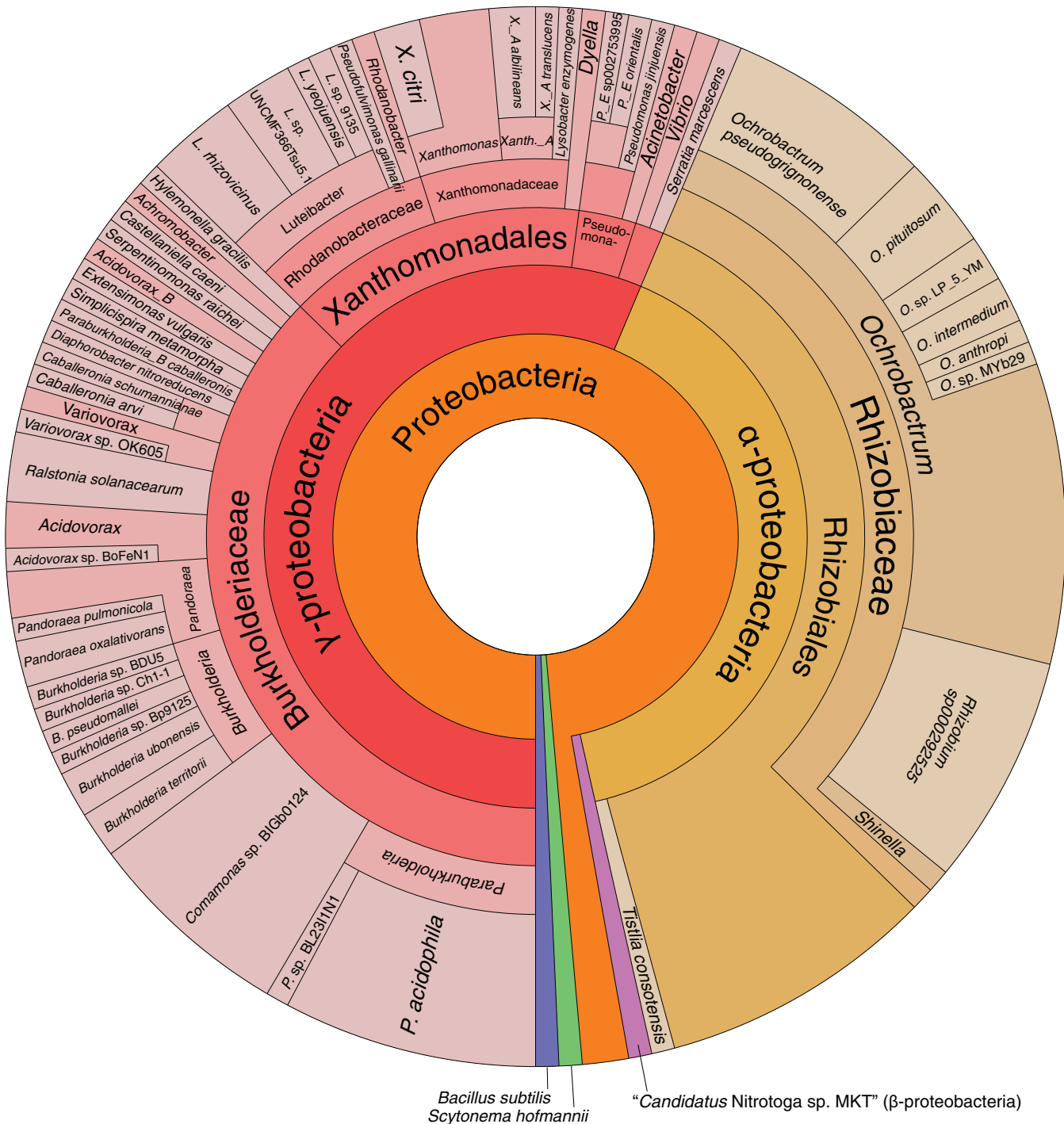
We also found many deficiencies in both the *de novo* and salvage nucleotide pathways (Fig. S2). In the pyrimidine biosynthetic pathway, genes for dihydroorotate (*pyrC*) (58) and orotate phosphoribosyl transferase (*pyrE*) (59) were missing, suggesting that Lv-StB cannot produce orotate from *N*-carbamoylaspartate and cannot create nucleotides from free pyrimidine bases. Ribonucleotide reductase (*nrdAB*) (60) and thymidylate synthase (*thyA*) (61) are present, suggesting that deoxypyrimidine nucleotides can be made from CTP. The deficiencies in purine synthesis were more profound. The majority of the genes involved in the *de novo* pathway (62) were missing (*purCDEFHLMT*, IMP dehydrogenase/*guaB*), except for those encoding adenylosuccinate lyase (*purrB*), adenylosuccinate synthase (*purA*), and GMP synthase (*guaA*). Lv-StB should therefore be able to make AMP from IMP and GMP from XMP (plus their deoxy analogs through ribonucleotide reductase) but cannot make purines *de novo*. We were also not able to find adenine phosphoribosyltransferase or hypoxanthine-guanine phosphoribosyltransferase (62), meaning that purine bases cannot be salvaged to make nucleotides.

The loss of so many seemingly vital pathways would suggest that several key nutrients are acquired from either the host or other microbes. There are a total of 56 intact genes associated with transport in the Lv-StB genome (Table S2), of which 12 are annotated as efflux/export proteins. A total of 32 genes were found to be associated with ABC-type transporters and are predicted to be involved in the transport of carbohydrates, lipids, lipoproteins, cations, sugars, and methionine. Other genes were also found that encoded transporters that were predicted to transport biopolymers, potassium, osmoprotectants, and fructose. Finally, only broad identification could be achieved for the remaining transporters, including those within the major facilitator superfamily (MFS), the resistance-nodulation-cell division superfamily (RND), the drug/metabolite transporter superfamily, and the HlyC/CorC family transporter. It is therefore possible that many nutrients that cannot be created *de novo* within Lv-StB, such as the amino acid methionine, may be acquired from external sources, such as is observed between *Buchnera* bacteria and their aphid hosts (63).

**Degradation of DNA repair pathways in *Burkholderia* sp. Lv-StB.** The genome of Lv-StB is missing many genes involved in DNA repair (Fig. 5B), similarly to other examples of genome-reduced symbionts (9). Compared to closely related *B. gladioli* strains, Lv-StB lacks genes in every repair pathway. In particular, the DNA polymerase I gene (*polA*), involved in homologous recombination, nucleotide excision repair, and base excision repair, is present only as two truncated pseudogenes (see above). Even though *polA* is involved in many different DNA repair pathways, it has been found to be nonessential in *E. coli* (64, 65), *B. pseudomallei* (66), and *B. cenocepacia* (67). In the homologous recombination pathway, Lv-StB lacks *recA*, *polA*, *rvuA*, *rvuB*, *rvuC*, and *recG*, all of which have been found to be essential for homologous recombination in *E. coli* (68). Likewise, Lv-StB is also missing *ku* and *ligD*, the two components of the nonhomologous end-joining pathway (69), suggesting that it cannot recover from double-strand breaks. Lv-StB lacks several DNA glycosylases in the base excision repair pathway which are responsible for removing chemically modified bases from double-stranded DNA (70). Some of these losses in Lv-StB simply reduce redundancy, but it has also lost the nonredundant glycosylase genes *mutM* and *mug*. The former recognizes 2,6-diamino-4-hydroxy-5-N-methylformamidopyrimidine (Fapy) and 8-hydroxyguanine (71), while the latter recognizes G:U and G:T mismatches (72) as well as epsilonC (73), 8-HM-epsilonC (74), 1,N(2)-epsilonG (75), and 5-formyluracil (76). Finally, in the mismatch repair system, Lv-StB is missing *mutS*, which is required for the recognition of mismatches in methyl-directed repair (77). In summary, Lv-StB is likely to be completely incapable of nonhomologous end-joining, homologous recombination, and mismatch repair, while being impaired in nucleotide excision repair and base excision repair due to the loss of DNA polymerase I and several DNA glycosylases.

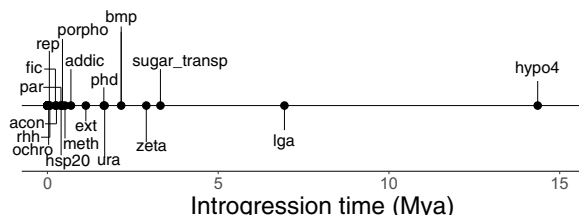
**Timing of horizontal acquisition of defensive and other genes in the Lv-StB genome.** We then attempted to identify genes in the *Burkholderia* sp. Lv-StB genome that are likely to have been acquired by horizontal gene transfer (HGT). A total of 497 intact genes were identified as unique to Lv-StB among 13 reference *B. gladioli* genomes (see Materials and Methods). Of these 497 genes, genes with no homologs in the NR database or genes that had homologs in *B. gladioli* genomes not included in this study were removed. A total of 148 genes appeared to be more closely related to species other than *B. gladioli* and may have been acquired through horizontal transfer (Fig. 6). A total of 79 and 57 genes appear to have been obtained from gammaproteobacteria and alphaproteobacteria, respectively, with one gene from a firmicute, one gene from a cyanobacterium, and three genes from phages (see Data Set S1G). The distribution is consistent with the notion that horizontal transfer occurs most frequently between closely related species (78). In particular, *Burkholderiaceae* species were the most frequent apparent donors of gammaproteobacterial proteins. Interestingly, the alphaproteobacterial genus *Ochrobactrum* (family *Brucellaceae* in the NCBI taxonomy and *Rhizobiaceae* in GTDB) was a major putative gene donor. This genus includes several symbionts of termites (79), army worms (80), weevils (81), and leeches (82, 83). In previous 16S amplicon investigations of *Lagria* beetles, *Ochrobactrum* spp. were often found (24, 26) (Fig. S3), suggesting that the donors of these genes could have also been associates of *L. villosa*. *Ochrobactrum* operational taxonomic units (OTUs) account for 5% to 20% of 16S rRNA gene reads, but this genus was not observed in the shotgun metagenome. However, disparities between 16S and shotgun metagenome abundances are not uncommon due to variable 16S copy numbers and primer and sequencing biases (84, 85). On the basis of the evidence for putatively horizontally transferred genes, we asked whether these could have contributed to the dominance of Lv-StB in *L. villosa* and set out to estimate the timings of horizontal transfer events.

Horizontally transferred genes are often detected on the basis of nucleotide composition differing from that of other genes in the genome (86). Such genes initially exhibit nucleotide composition consistent with the donor genome, which eventually normalizes to the composition of the recipient genome (87). The rate of this “amelioration” process ( $\Delta GC^{HT}$ ) has been modeled by Lawrence and Ochman (87), based on the



**FIG 6** Putative sources of genes unique to the *Burkholderia* sp. Lv-StB genome (compared to *B. gladioli* strains) based on hits to BLASTP searches. Darker colors denote basal taxonomic classifications, while lighter colors denote specific taxonomic classifications. Note that three proteins of putative phage origin (see Data Set S1H) are not included in the figure.

substitution rate ( $S$ ), the transition/transversion ratio ( $\kappa$ ), and the GC content of both the recipient genome ( $GC^{EQ}$ ) and the putatively horizontally transferred genes ( $GC^{HT}$ ), according to equation 1. By iterating this equation repeatedly until  $GC^{HT}$  equals  $GC^{EQ}$ , the time required from the present day to complete amelioration can be estimated. If the GC content of the donor genome is known, then equation 1 can be used in reverse to estimate the time since introgression. However, if the donor GC content is not



**FIG 7** Estimated introgression time for putative HGT gene sets, using the “*Candidatus Baumannia cicadellinicola*” substitution rates (36). Note that for clarity, the gene sets with estimated introgression time of <5,000 ya are not labeled. For these data and ages estimated with other substitution rates, see Data Set S1H. Abbreviations refer to groups of genes as outlined in Data Set S1H: acon, E5299\_02570–E5299\_2571; addic, E5299\_02248–E5299\_02250; meth, E5299\_02474–E5299\_02475; ochro, E5299\_01926–E5299\_01933; par, E5299\_02349–E5299\_02352; phd, E5299\_02420–E5299\_02423; porfo, E5299\_02300, E5299\_02302–E5299\_02303; rep, E5299\_02219–E5299\_02220; rhn, E5299\_01720, E5299\_01726; ura, E5299\_02533–E5299\_02435.

known, then the differing selection pressures on the first, second, and third codon positions can be exploited to estimate the introgression time. Because these positions have different degrees of amino acid degeneracy, they are subject to different degrees of selection, and therefore they ameliorate at different rates. As a consequence, Lawrence and Ochman (87) found that for genes in the process of amelioration, the relationship between overall GC content and the GC content at individual codon positions seen in genes at equilibrium (87, 88) (equations 2, 3, and 4) does not hold. So if equation 1 is applied in reverse separately for each codon position, the time since introgression can be inferred at the iteration yielding the minimum square difference from equations 2 to 4. Application of equation 1 also yields an estimate for the original donor GC.

$$\Delta GC^{HT} = S \times \frac{\kappa + \frac{1}{2}}{\kappa + 1} \times [GC^{EQ} - GC^{HT}] \quad (1)$$

$$GC_{1st} = 0.615 \times GC_{genome} + 26.9 \quad (2)$$

$$GC_{2nd} = 0.270 \times GC_{genome} + 26.7 \quad (3)$$

$$GC_{3rd} = 1.692 \times GC_{genome} - 32.3 \quad (4)$$

We identified groups of consecutive genes in the putative HGT set that could have been acquired together and used the method described above to estimate their introgression time (Data Set S1H). Among the 18 identified gene groups, 7 were found to have atypical GC content (defined by Lawrence and Ochman [87]), with GC% content that was either >10% lower or >8% higher at the first codon position or the third codon position, respectively, than that of the genome as a whole. The method described above was used to estimate the time of introgression for each gene group, using the *Bl. floridanus*, *Bl. pennsylvanicus*, *Bl. obliquus*, and “*Ca. Baumannia cicadellinicola*” divergence rates (see above and Fig. 7; see also Data Set S1H). The oldest horizontal transfer (1.79 to 14.36 Mya) was found to correspond to the “hypo4” group (E5299\_02488–E5299\_02489), representing two phage proteins, and the next oldest corresponded to the lagriamide BGC and neighboring genes (*lga*; 0.8 to 6.42 Mya; E5299\_00001–E5299\_00003, E5299\_00005–E5299\_00018). The *lga* BGC is predicted to have come from a high-GC organism, with original GC content of 72%. The closest relatives of many of the *lga* genes are found in *Pseudomonas* strains, which typically do not have GC content this high. However, as BGCs are often thought to be horizontally transferred (89), *Pseudomonas* may not be the direct source of *lga* in Lv-StB.

Several other HGT gene groups (sugar\_transp [E5299\_01760–E5299\_01761, E5299\_01763], bmp [E5299\_02444], ext [E5299\_02524–E5299\_02526], dinj [E5299\_02222, E5299\_02224], tonb [E5299\_00736–E5299\_00737], tonb2 [E5299\_02080]) were predicted to be involved in transport, with predicted substrates that included purine

nucleosides, trehalose, and vitamin B12. Of these, the sugar\_transp and bmp groups, predicted to be involved in importation of trehalose and purine, respectively, are relatively old (0.41 to 3.31 My and 0.27–2.16 My, respectively), and the groups likely involved in B12 importation (dijn, tonb, tonb2) are estimated to be <5,000 y old. Trehalose is the most abundant component of insect hemolymph, and was found to be provided by the host to its genome-reduced symbiont in a leaf beetle system (90). We also found HGT gene groups putatively involved in heat shock response (hsp20; 0.05 to 0.4 Mya) and DNA repair (ura [E5299\_02533–E5299\_02535], rep [E5299\_02219–E5299\_02220], ochro [E5299\_01926–E5299\_01933]). In the latter category, we identified a uracil-DNA glycosylase (in ura; 0.21 to 1.68 Mya), used in base excision repair in cases of deamination; RecB, used in homologous recombination (in rep; 0.01 to 0.055 Mya) (62); and YedK (in ochro; 0.01 to 0.035 Mya) (91), a protein used in the SOS response that binds to abasic sites in single-stranded DNA. The transferred transport functions and DNA repair proteins match functional categories that are currently lacking genes due to genome reduction (see above), and therefore the transfers could have been contemporaneous with the reduction process, acting as mechanisms compensating for lost functions.

One putative HGT toxin that is unique to Lv-StB in the metagenome and among *B. gladioli* strains is zonular occludens toxin (zot [E5299\_02544–E5299\_02545]; estimated to have been acquired <5,000 ya). The zot gene is responsible for the production of the zonula occludens toxin, a virulence factor which was initially identified in *Vibrio cholerae* and whose presence was found to lead to the disassembly of intracellular tight junctions and consequently to increased permeability of mammalian epithelium (92). Colocalized with the predicted zot gene was a gene encoding DUF2523, which we found often accompanied zot in our searches of the STRING database (93). Zot proteins have been identified in several strains of *Campylobacter* and have been shown to elicit an inflammatory response in intestinal epithelial cells (94, 95). Furthermore, a significant correlation was found between the presence of the Zot protein and hyperinvasive strains of *Neisseria meningitidis* (96). Potentially, zot may aid in the infection of the *L. villosa* embryonic structures through increasing permeability across the outer layers of the egg, although this remains speculative.

**The evolution of *Burkholderia* sp. Lv-StB.** In the 1920s it was observed (97) that beetles in the *Lagria* and *Cerogria* genera possessed structures now known to harbor *Burkholderia* symbionts in *L. villosa* and *L. hirta* (26). Other genera in the Lagriinae subfamily, such as *Adynata* and *Arthromacra*, do not have these structures. According to the tree found at [timetree.org](http://timetree.org) (98), *Lagria* and *Cerogria* diverged 55 Mya, and the common ancestor of *Lagria*, *Cerogria*, and *Adynata* existed 82 Mya (this region of the tree utilizes data from Kergoat et al. [99]). On the basis of these estimates, the symbiont-bearing structures in *Lagria* and *Cerogria* likely evolved between 82 and 55 Mya. Our analysis suggests that the divergence of *Burkholderia* sp. Lv-StB from *B. gladioli* occurred after that point (6.15 to 49.76 Mya). During that time, the genome of *Burkholderia* sp. Lv-StB became reduced, and it is likely dependent on the host due to deficiencies in energy metabolism and nucleotide biosynthesis. Notably, the profound metabolic insufficiencies and incomplete DNA repair pathways in Lv-StB are typical of symbionts with smaller genomes, such as “*Ca. Endolissoclinum faulkneri*,” an intracellular tunicate symbiont with a 1.48-Mbp genome and a similar number of genes (783) (100), estimated to have been a symbiont for at least 6 to 31 My (101). While the presence of *Burkholderia* sp. Lv-StB and of its defensive compound lagriamide has been shown to decrease the rate of fungal egg infection (6), the symbiont is not essential for beetle reproduction (24). Therefore, the relationship is facultative from the perspective of the host, while Lv-StB is in the process of becoming dependent on *L. villosa*. A central issue that we aimed to address in this work was that of how the genome of Lv-StB became reduced, when *L. villosa* appears to maintain multiple other nonreduced *Burkholderia* spp. and other symbionts.

It is clear from previous work that the *Lagria* symbionts related to *B. gladioli* evolved from plant-associated strains (26), likely transmitted to the insects from the plant environment. Intriguingly, there are closely related *Burkholderia* (now formally *Caballeronia*) symbionts of several plant species within the Rubiaceae and Primulaceae families which reside within the host's leaf nodules (102) and have genomes that are similar in size to that of *Burkholderia* sp. Lv-StB. In "Ca. *Burkholderia crenata*," hosted by the plant *Ardisia crenata*, there are a number of interesting parallels with *Burkholderia* sp. Lv-StB, such as the maintenance of secondary metabolite BGCs and loss of essential genes and DNA repair functions (103). Interestingly, lateral gene transfers of the secondary metabolite BGCs between different symbionts have occurred, likely after transition to a host-restricted lifestyle (104). It is possible that the evolution of the *L. villosa* symbiont resembles that of the leaf nodulating *Burkholderia* symbionts given the similarities in function, genome characteristics, and degrees of host dependence, although "Ca. *Burkholderia crenata*" has the smallest genome of leaf nodule symbionts and, in contrast to Lv-StB, maintains the capability of synthesizing most cofactors and vitamins.

For the *Lagria* host, the probable advantage of the consistent presence of *Burkholderia* or other bacteria is that of protection of eggs from infection, through the activity of small molecules made by its microbiome. The strains characterized here, as well as the previously isolated Lv-StA strain, were found to contain ample biosynthetic potential, and both Lv-StA and Lv-StB produce antifungal compounds that were found to protect eggs from fungal infection in laboratory experiments (26). And yet, Lv-StA is found only sporadically as a minor component of the microbiome in the field (6). It is probably advantageous for *Lagria* beetles to maintain a pool of facultative symbionts with different biosynthetic capabilities to allow fast adaptation to different environmental infection pressures (105). However, there may be less selection pressure on a facultative symbiont to stay associated with its host if it can also survive in the environment and infect plants.

The foundational event in the establishment of the symbiosis between Lv-StB and *L. villosa* was likely the acquisition of the *Iga* pathway, which putatively produces lagriamide, in a nonreduced ancestor of Lv-StB (6). We place this as the first event for four reasons. First, the *Iga* BGC is almost the oldest detectable horizontal transfer that survives in the reduced genome of Lv-StB. Second, we found little evidence that Lv-StB is capable of making metabolites of use to the host, indicating that the symbiosis is likely not based on nutrition. *L. villosa*'s diet of plant leaves may be nitrogen poor, with hard-to-digest plant cell wall components (106), but we did not find polysaccharide-degrading pathways or pathways corresponding to extensive biosynthesis of essential amino acids in the Lv-StB genome. Therefore, the *Iga* BGC is the oldest remaining feature that potentially increases host fitness. Third, the reduced coding density seen in the Lv-StB genome may be indicative of a recent transitional event (11), such as strict host association or a move to vertical transmission. Fourth, even though we found genes missing from all DNA repair pathways, which is thought to be a driver for increased AT content in symbiont genomes (9), and increases in AT content may have an adaptive component that reduces the metabolic costs associated with symbionts (23), the GC content of the Lv-StB genome is not very different from that of free-living *B. gladioli* strains in comparison to other "transitional" symbionts. For instance, "*Candidatus Pantoea carbekii*" has a reduced genome and, like Lv-StB, lacks full-length DNA polymerase I (107). The GC content of this strain is almost 30% lower than that of its closest free-living relatives (11), while the GC content of the genome of Lv-StB is ~10% lower than that of its closest relatives. Therefore, we propose that loss of DNA repair pathways and other genome degradation events in Lv-StB occurred relatively recently, after the acquisition of *Iga*.

It can be envisioned that *Iga* provided a sustained survival advantage in an environment where lagriamide reduced egg fungal infections and that there was positive selection in beetles that vertically transmitted *Iga*-bearing symbionts. It is also possible, although speculative, that production of lagriamide confers a competitive advantage

among the members of the microbial community in the beetle. In *L. villosa*, symbionts are stored extracellularly, and they are spread onto the exterior of eggs as they are laid. According to observations in the congeneric species *L. hirta*, the symbionts are assumed to first enter through the egg micropyle to reach the embryonic organs where they are housed throughout larval development (97). It is thus likely that only a few of the cells are vertically transmitted as a result of colonizing these structures, potentially providing the population bottlenecks that could have caused initial accumulation of deleterious mutations that started the process of genome reduction. Meanwhile, loss of certain proteins limiting growth rate (see above) may have been selected through the presence of increased Lv-StB populations and compound production. It is unknown to what extent Lv-StB is genetically isolated in the larval or adult host, but we found evidence of ongoing horizontal transfer events having occurred in the recent past, presumably through contact with a complex microbiome associated with *L. villosa* egg surfaces. These horizontal gene transfers likely happened concurrently with the ongoing genome reduction process and may have been compensatory for gene losses. There is some precedence for the idea of the existence of extracellular symbionts with profoundly reduced genomes (106, 108–110). For instance, the leaf beetle *Cassida rubiginosa* harbors “*Candidatus Stammera capleta*,” a symbiont with the smallest genome of any extracellular organism (0.27 Mbp), which provides pectinolytic enzymes to help break down the host’s leafy diet (106). In many of these cases, symbionts are stored as isolated monocultures within specialized structures in adult hosts, while vertical transmission is assisted by packaging symbiont cells in protective “caplets” attached to eggs (106) or in a “symbiont capsule” encased in chitin (108) or by their secretion in a galactan-based jelly ingested by hatched juveniles (110), although reduced-genome symbionts have also been known to be vertically transmitted by simple egg surface contamination (109), as in *L. villosa*. Because these examples represent advanced cases of genome reduction, it would be difficult to determine whether horizontal transfer events occurred before or during genome reduction, and none have been noted. The complexity of the *L. villosa* microbiome appears to represent a different case, as it afforded ample opportunity for horizontal gene transfer even while the genome of Lv-StB was actively undergoing reduction.

In addition to the leaf nodule *Burkholderia* symbionts (see above), horizontal acquisition of genes has been observed in two types of reduced-genome symbionts, namely, eukaryotic parasites in the genus *Encephalitozoon* (111) and *Acetobacteraceae* strains associated with the gut community of red carpenter ants (112). In both of these cases, symbiont genomes were similar in size to Lv-StB (~2 Mbp) but showed far greater coding density and fewer pseudogenes. Furthermore, both the *Encephalitozoon* and *Acetobacteraceae* strains were culturable, suggesting that they are facultative symbionts in a less advanced state of genome reduction than Lv-StB. The genome of Lv-StB appears to be different from the genomes of these examples, because there is evidence of recent horizontal transfers, even though genes required for homologous recombination are currently missing. Either the loss of homologous recombination was very recent or such transfers could have occurred in a RecA-independent manner. For example, plasmids could have been transferred into Lv-StB cells, followed by the RecA-independent transposition of genes to the chromosome (113, 114). Another potential mechanism is phage infection. For instance, it is known that the genomes of symbionts that switch hosts such as *Wolbachia* strains tend to contain a large amount of mobile DNA and phage remnants (115).

**Conclusions.** It is unclear whether Lv-StB will continue on the path of genome reduction to become drastically reduced with a <1-Mbp genome. Where symbionts are required for host survival and are genetically isolated within host cells or specialized structures, such a process appears to be irreversible and unstoppable (116, 117). However, a number of alternate fates could be envisioned for Lv-StB. With a complex microbiome, if ongoing gene losses in Lv-StB reduce its fitness past a certain point, then it could be replaced by another strain, potentially accompanied by horizontal transfer

of the *lga* pathway to a less reduced genomic chassis. Alternatively, horizontal transfers of genes to Lv-StB could lead to an equilibrium of gene loss and gain. Interestingly, it appears that until the present time, horizontal transfer had not occurred fast enough to prevent widespread loss of metabolism and DNA repair in the Lv-StB genome. The host could also evolve strategies to maintain increasingly genome-reduced Lv-StB, perhaps by selective extracellular partitioning and packaging for vertical transfer similarly to the examples outlined above. However, it is unclear whether such an evolutionary path would be favorable, given that the coinfection of multiple BGC-bearing symbiont strains could be advantageous in environments with variable pathogen pressures.

In summary, evidence gathered here suggests that the introduction of the lagrimamide BGC preceded and perhaps initiated genome erosion of Lv-StB, potentially through selection of beetles that transferred the symbiont vertically, leading to a population structure with frequent bottlenecks. Simultaneous advantageous gene acquisitions may have enabled the preferential survival of Lv-StB and its dominance in the adult host and the egg surface.

## MATERIALS AND METHODS

**Sequencing and assembly of *Burkholderia gladioli* Lv-StA genome.** Genome sequencing of the isolated *B. gladioli* Lv-StA strain was carried out using PacBio with single-molecule, real-time (SMRT) technology. For *de novo* assembly (carried out by Eurofins Genomics), the HGAP pipeline was used (Hierarchical Genome Assembly Process). Briefly, a preassembly of long and accurate sequences was generated by mapping filtered subreads to so-called seed reads. Subsequently, the Celera assembler was used to generate a draft assembly using multikilobase long reads, which in this case rendered full-genome closure. Finally, the Quiver algorithm was used to correct indel and substitution errors by considering the quality values from the bas.h5 files.

**Metagenomic binning and annotation.** Metagenomic assembly files were clustered into putative genomic bins using Autometa (Master branch—commit bbcea30) (28). Contigs with lengths shorter than 3,000 bp were excluded from the binning process, and a taxonomy table was produced. Contigs classified as bacterial were further binned into putative genomic bins using run\_autometa.py. Unclustered contigs were recruited into clusters using ML\_recruitment.py. Results were summarized using cluster\_process.py. The resultant genome bins were compared to earlier versions (6) using Mash version 2.1.1 (118), which hashes genomes to patterns of *k*-mers (sketching), allowing rapid calculations of distances between two sketches. All bins were sketched, and distances were computed in a pairwise fashion. Pairwise distances were visualized in R as a dendrogram and enabled the determination of equivalent old and updated putative genome bins between analyses. The updated putative genomic bins were annotated using Prokka version 1.13 (119), with GenBank compliance enabled. Reference genomes downloaded from NCBI were similarly annotated with Prokka in order to maintain consistency between data sets. Amino acid sequences of open-reading frames (ORFs) were further annotated using DIAMOND blastp version 0.9.21.122 (120) against the Diamond formatted NR database. The search was limited to returning a maximum of 1 target sequence, and the maximum number of high-scoring pairs per subject sequence was set to 1. Results were summarized in BLAST tabular format with qseqid (Query gene identifier [ID]), stitle (aligned gene ID), pident (percentage of identical matches), evalue (expected value), qlen (query sequence length), and slen (aligned gene sequence length) as desired parameter output. Pseudogenes were identified by finding Lv-StB genes that were more than 20% shorter than their respective BLAST matches. This criterion has been used previously to identify pseudogenes (101, 121).

Coding density was calculated as the sum of all protein-coding sequences (coding sequence) as a percentage of the sum of all contigs (total sequence). In cases where protein-coding genes were found to overlap, the length of the overlap region was counted only once. This calculation was performed for all binned genomes, both on the initial data sets (GenBank files generated during Prokka annotations) and on the edited data sets where pseudogenes had been removed. For the identification and count of genes encoding transposases and hypothetical proteins, protein-coding gene amino acid files (\*.faa) containing Prokka annotations were parsed for gene descriptions containing “transposase” and “hypothetical” strings.

**Taxonomic classification of genome bins.** Putative genome bins clustered from the *L. villosa* metagenomic data set were taxonomically classified using GTDB-Tk v0.2.2 (reference database gtdbtk.r86\_v2) with default parameters (29). GTDB-Tk identifies and aligns 120 bacterial marker genes per genome before calculating the optimal placement of the respective alignments in the precomputed GTDB-Tk reference tree which consists of 94,759 genomes (see Data Set S1B in the supplemental material). A species was assigned to a genome if it shared 95% or more ANI with a reference genome.

**Identification of “core” genes.** The set of “core” genes generally found in even the most reduced symbiont genomes was taken from the data provided in Table 2 of a previous report by McCutcheon and Moran (9). GFF files produced for *B. gladioli* Lv-StA and the metagenomic bins by Prokka were searched for the following gene designations: “*dnaE*,” “*dnaO*,” “*rpoA*,” “*rpoB*,” “*rpoC*,” “*rpoD*,” “*groL*,” “*groS*,” “*dnaK*,” “*mnmA*,” “*mnmE*,” “*mnmG*,” “*sufS*,” “*sufB*,” “*sufC*,” “*iscS*,” “*iscA*,” “*iscU*,” “*rluA*,” “*rluB*,” “*rluC*,” “*rluD*,” “*rluE*,” “*rluF*,” “*infA*,” “*infB*,” “*infC*,” “*fusA*,” “*tsf*,” “*prfA*,” “*prfB*,” “*frr*,” “*def*,” “*alas*,” “*gltX*,” “*glyQ*,” “*ileS*,” “*metG*,” “*pheS*,” “*trpS*,” “*valS*,” “*rpsA*,” “*rpsB*,” “*rpsC*,” “*rpsD*,” “*rpsE*,” “*rpsG*,” “*rpsH*,” “*rpsI*,” “*rpsJ*,” “*rpsK*,” “*rpsL*,” “*rpsM*,” “*rpsN*,”

"rpsP," "rpsQ," "rpsR," "rpsS," "rplB," "rplC," "rplD," "rplE," "rplF," "rplK," "rplM," "rplN," "rplO," "rplP," "rplT," "rplV," "rpaA," "rpbB," "rpgG," "rplM," "tRNA-Met," "tRNA-Gly," "tRNA-Cys," "tRNA-Phe," "tRNA-Lys," "tRNA-Ala," "tRNA-Glu," "tRNA-Pro," "tRNA-Gln," and "tRNA-Ile." The presence of single or multiple examples of these genes per genome/bin was tabulated in Microsoft Excel to produce Data Set S1D, and the percentage of the core gene set found in a genome/bin was used for Table 1.

**Annotation and analysis of BGCs.** Putative biosynthetic gene clusters were identified in all binned genomes using the AntiSMASH (122) docker image (image ID: 8942d142d9ac). HMMer analysis of the entire genome was enabled, and identified clusters were compared to both antiSMASH-predicted clusters, the MIBiG database, and secondary-metabolite orthologous groups. Similarities between identified putative biosynthetic gene clusters were assessed using BiG-SCAPE version 20181005 (35) in "glocal" mode.

**Construction of multilocus species tree.** Genomes of *Burkholderia* sp. Lv-StB and *B. gladioli* Lv-StA and binned genomes taxonomically classified within the *Burkholderia* genus, including DBSCAN\_round6\_14, DBSCAN\_round6\_18, and DBSCAN\_round4\_0, were uploaded to the AutoMLST website (123). A concatenated species tree was constructed in *de novo* mode (with default options), with the IQ-TREE Ultrafast Bootstrap analysis and ModelFinder options enabled.

**Calculation of average nucleotide identities.** The average nucleotide identities (ANIs) of *Burkholderia* sp. Lv-StB and *B. gladioli* were calculated in a pairwise manner using FastANI (124) against 45 *Burkholderia* reference genomes downloaded from NCBI. A total of 13 genomes shared over 95% ANI (Data Set S1F). These genomes were all identified as *B. gladioli* species and were used in downstream analyses.

**Quantification of divergence between *Burkholderia* sp. Lv-StB and *B. gladioli* A1.** Orthologous protein sequences were identified in nonpseudogene sequence files of *Burkholderia* sp. Lv-StB and 13 closely related genomes (*B. gladioli* Lv-StA, *B. gladioli* A1, *B. gladioli* UCDUG, *B. gladioli* FDAARGOS\_389, *B. gladioli* ATCC 25417, *B. gladioli* Co14, *B. gladioli* SN82F6, *B. gladioli* ATCC 10248, *B. gladioli* NBRC13700, *B. gladioli* FDAARGOS\_188, *B. gladioli* MSMB1756, *B. gladioli* BSR3, and *B. gladioli* KACC11889) identified through the ANI analyses using OMA version 2.2.0 (37). A subset (797 groups) of the resultant orthologous groups (OGs) was identified which included genes from all 14 genomes used in the analysis. Each set of OG sequences were aligned using MUSCLE v3.8.31 (125), and the corresponding nucleotide files were extracted and aligned against the amino acid sequences using the PAL2NAL docker image (image ID: ce3b1d7d83ab) (126) using codon table 11 and specifying no gaps, with paml as the output format. The resultant paml files were used to estimate pairwise *dS* (synonymous divergence rate), *dN* (nonsynonymous divergence rate), and kappa (transition/transversion ratio) values from comparisons between individual genes per orthologous group with codeml (127) in the PAL2NAL package. The following parameters were specified in the control file: runmode = -2 (pairwise), model = 0 (one), fix\_kappa = 0 (kappa to be estimated), and fix\_omega = 0 (estimate omega, where omega is the *dN/dS* ratio, with initial omega set to 0.2). Any orthologous gene sets that included genes that gave a *dS* value over 3 were removed from the analysis (128). Individual sequences from remaining OGs were then gathered into genome-specific files (i.e., all Lv-StB genes in all OGs were moved into an ordered Lv-StB.faa/.ffn file). Stop codons were removed from each nucleotide sequence. Sequences were then concatenated per genome to produce a single sequence per genome. The concatenated amino acid sequences and corresponding nucleotide sequences were aligned against one another using PAL2NAL as performed for individual genes. Pairwise estimations of *dS*, *dN*, and kappa were calculated as before using codeml. Additionally, the concatenated genes were analyzed a second time using an alternative control file, in which the model was set to 2. The likelihood ratio test value for comparisons between pairwise null and alternative likelihood scores was calculated ( $2 \times \text{Alt\_lnl} - \text{Null\_lnl}$ ) for Lv-StB relative to the 13 reference genomes and found to be 0 in all cases, indicating that the omega (*dN/dS*) ratio was consistent in comparisons between Lv-StB and the reference genomes. Individual *dS* values were used to estimate divergence between Lv-StB and the 13 reference genomes with divergence rates estimated using the method described previously by Silva and Santos-Garcia (36) (see Table S1 in the supplemental material) in the following equation: age of divergence (Mya) = *dS* ÷ divergence rate × 1,000,000. As Lv-StB shared the greatest ANI with *B. gladioli* A1, the kappa value found in comparisons between these two genomes (7.03487) was used for amelioration estimates.

**Pangenome analysis.** To assess the pangenome of Lv-StB and other *B. gladioli* genomes, GFF files generated by Prokka were analyzed using Roary (39), which identifies core and accessory genes per genome. Concatenation and alignment of orthologous genes were enabled in Roary and used to build a phylogenetic tree with FastTree version 2.1.10 (129). The resultant phylogenetic tree and presence/absence matrix of genes in all genomes were visualized with the roary\_plots.py script. Additionally, nonpseudogenes of all genomes were annotated against the KEGG database (130, 131) using kofamscan (132) with the output in mapper format. An overview of the completeness of general metabolic pathways was visualized using KEGG-Decoder (133) with kofamscan annotations. For specific pathways of interest (amino acids, DNA repair, nucleotide *de novo* biosynthesis), presence/absence matrices of genes per KEGG pathway entry were visualized in R version 3.6.0 using the tidyR, ggplot2, and viridis libraries.

**Identification of genes putatively acquired by horizontal transfer.** All amino acid sequence files representing nonpseudogenes of all genomes used in the ANI analysis (*B. gladioli* Lv-StA and 45 *Burkholderia* reference genomes) were concatenated and converted into a DIAMOND BLAST (120) database (build 125). The amino acid sequence files of nonpseudogenes in Lv-StB were then searched against this database. All genes that had no significant hit, or that had a significant hit but with a shared proportion of less than 50%, were considered unique to Lv-StB. Nonpseudogenes of Lv-StB that were not found to have an ortholog in the OMA analysis were used to validate this list. These unique genes were then compared to the NR database using DIAMOND blastp (as described above), and any genes that had

no significant hit were removed from the set of unique genes. Manual inspection of the remaining genes resulted in the removal of any genes that were closely related to *B. gladioli* genes (i.e., were found within *B. gladioli* genomes other than the ones investigated here). The remaining unique genes were consequently considered to represent genes potentially acquired via horizontal transfer. This list was expanded with other genes within Lv-StB, for which homologs in *Burkholderia* genomes could be found, but the closest match against the NR database belonged to a different genus. For example, E5299\_02249 of the “addic” group shared 53.1% sequence identity with a gene from *B. contaminans* strain LMG 23361 but shared a higher sequence identity level of 96.9% with *Ochrobactrum pituitosum*.

**Deamelioration of putatively horizontally transferred genes.** The method of Lawrence and Ochman (87) was implemented in Python and is available at [https://bitbucket.org/jason\\_c\\_kwan/age\\_horizontal\\_transfers.py](https://bitbucket.org/jason_c_kwan/age_horizontal_transfers.py). The script takes as input (i) an in-frame nucleotide FASTA file containing the sequences of putatively horizontally transferred genes, (ii) an in-frame nucleotide FASTA file containing a comparison set of gene sequences from the genome, (iii) a synonymous mutation rate in substitutions per 100 sites per million years, (iv) a nonsynonymous mutation rate in substitutions per 100 sites per million years, (v) a transition/transversion ratio ( $\kappa$ ), (vi) a step time in millions of years, and (vii) a maximum time to iterate to. The script outputs GC content of each codon position (plus overall GC) at each time point and reports the estimated age of the gene cluster corresponding to the iteration with the smallest sum of squared deviations from equations 2 to 4. The substitution rates used in our calculations were half of the divergence rates estimated by Silva and Santos-Garcia (36) and were as follows: for “Ca. Ba. cicadellinicola,” synonymous rate of 0.55 and nonsynonymous rate of 0.05; for *Bl. obliquus*, synonymous rate of 3.95 and nonsynonymous rate of 0.26; for *Bl. pennsylvanicus*, synonymous rate of 3.2 and nonsynonymous rate of 0.28; for *Bl. floridanus*, synonymous rate of 4.45 and nonsynonymous rate of 0.395. A value of 7.0348 was used for  $\kappa$ , previously calculated for *Burkholderia* sp. Lv-StB and *B. gladioli* A1 (see above). A step time of 0.005 My and a maximum time of 50 My were used in all calculations. The comparison gene set included only nonpseudogenes that were not identified as putative horizontally transferred genes.

**Microbial community analysis.** 16S rRNA amplicon sequence data sets analyzed via Qiime (134) as described previously (24) were reanalyzed using Mothur v.1.40.3 (135). Reads shorter than 200 bp and reads containing ambiguous bases or homopolymeric runs longer than 7 bases were removed from the data set. Chimeric sequences were identified using VSEARCH (136) and removed from the data set. Reads were taxonomically classified against the Silva database (version 132), and all reads classified as unknown, eukaryotic, or mitochondrial or as representative of chloroplasts were removed from the data set. Reads were aligned using the Silva database (v. 132) as the reference and clustered into operational taxonomic units (OTUs) at a distance of 0.03, an approximation used for bacterial species. Counts of OTUs per sample were generated, and the top 10 most abundant OTUs were plotted (see Fig. S3 in the supplemental material). The top 50 most abundant OTUs were queried against the “nt” nucleotide database using BLAST for taxonomic classification.

**Data availability.** The *Burkholderia gladioli* Lv-StA genome was deposited in GenBank under accession number WITE00000000. The metagenomic bins obtained in this study were deposited in GenBank under the following accession numbers: for *Burkholderia* sp. Lv-StB, WNDN00000000; for DBSCAN\_round1\_2, WNDQ00000000; for DBSCAN\_round1\_3, WNDP00000000; for DBSCAN\_round2\_3, WNDQ00000000; for DBSCAN\_round34\_1, WNDR00000000; for DBSCAN\_round3\_0, WNDS00000000; for DBSCAN\_round43\_4, WNNT00000000; for DBSCAN\_round44\_0, WNDU00000000; for DBSCAN\_round4\_0, WNDV00000000; for DBSCAN\_round4\_12, WNDW00000000; for DBSCAN\_round4\_6, WNDX00000000; for DBSCAN\_round5\_1, WNDY00000000; for DBSCAN\_round5\_3, WNDZ00000000; for DBSCAN\_round6\_14, WNEA00000000; for DBSCAN\_round6\_18, WNEB00000000; for DBSCAN\_round7\_0, WNEC00000000; for DBSCAN\_round7\_14, WNED00000000; for DBSCAN\_round8\_1, WNEE00000000. The raw Illumina reads used to construct the *L. villoso* egg metagenome were deposited to the SRA database under accession number PRJNA531449.

## SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

**FIG S1**, EPS file, 1.8 MB.

**FIG S2**, EPS file, 2.5 MB.

**FIG S3**, EPS file, 1.6 MB.

**TABLE S1**, DOCX file, 0.01 MB.

**TABLE S2**, DOCX file, 0.01 MB.

**DATA SET S1**, XLSX file, 0.2 MB.

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