Experimental-Evolution-Driven Identification of Arabidopsis Rhizosphere Competence Genes in Pseudomonas protegens

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ABSTRACT Beneficial plant root-associated microorganisms carry out a range of functions that are essential for plant performance. Establishment of a bacterium on plant roots, however, requires overcoming several challenges, including competition with neighboring microorganisms and host immunity. Forward and reverse genetics have led to the identification of mechanisms that are used by beneficial microorganisms to overcome these challenges, such as the production of iron-chelating compounds, the formation of strong biofilms, or the concealment of characteristic microbial molecular patterns that trigger the host immune system. However, how such mechanisms arose from an evolutionary perspective is much less understood. To study bacterial adaptation in the rhizosphere, we employed experimental evolution to track the physiological and genetic dynamics of root-dwelling Pseudomonas protegens in the Arabidopsis thaliana rhizosphere under axenic conditions. This simplified binary one plant/one bacterium system allows for the amplification of key adaptive mechanisms for bacterial rhizosphere colonization. We identified 35 mutations, including single-nucleotide polymorphisms, insertions, and deletions, distributed over 28 genes. We found that mutations in genes encoding global regulators and in genes for siderophore production, cell surface decoration, attachment, and motility accumulated in parallel, underlining the finding that bacterial adaptation to the rhizosphere follows multiple strategies. Notably, we observed that motility increased in parallel across multiple independent evolutionary lines. All together, these results underscore the strength of experimental evolution in identifying key genes, pathways, and processes for bacterial rhizosphere colonization and a methodology for the development of elite beneficial microorganisms with enhanced root-colonizing capacities that can support sustainable agriculture in the future.

IMPORTANCE Beneficial root-associated microorganisms carry out many functions that are essential for plant performance. Establishment of a bacterium on plant roots, however, requires overcoming many challenges. Previously, diverse mechanisms that are used by beneficial microorganisms to overcome these challenges were identified. However, how such mechanisms have developed from an evolutionary perspective is much less understood. Here, we employed experimental evolution to track the evolutionary dynamics of a root-dwelling pseudomonad on the root of Arabidopsis. We found that mutations in global regulators, as well as in genes for siderophore production, cell surface decoration, attachment, and motility, accumulate in parallel, emphasizing these strategies for bacterial adaptation to the rhizosphere. We identified 35 mutations distributed over 28 genes. All together, our results demonstrate the power of experimental evolution in identifying key pathways for rhizosphere colonization and a methodology for the development of elite beneficial microorganisms that can support sustainable agriculture.
Plants are associated with complex microbial communities assembled into a functional microbiome that safeguards optimal plant performance under harsh environmental conditions (1). The rhizosphere is a particularly interesting hot spot of plant-microbe interactions. Plants deposit up to 44% of their photosynthetically fixed carbon into the rhizosphere, fueling a specific microbial community (2). The microbial species pool in the bulk soil is the source from which the root microbiome is recruited, and plant genotype, immune responses, and environmental factors are postulated to affect this process (3–6). The establishment of beneficial microbial associations requires a high degree of coordination of both plant and microbial responses by means of a continuous molecular dialogue (7, 8). Plant-associated microorganisms can improve plant yield by protecting the plant from abiotic stresses (9), improving plant nutrition and growth (10–12), antagonizing soilborne pathogens (13), or stimulating plant immunity (14). To exert their beneficial effects on plant performance, bacteria must colonize the roots efficiently and establish significant populations. For example, bacterial population densities above $10^5$ cells per gram of root are required for efficient suppression of soilborne plant pathogens by *Pseudomonas* spp. (15, 16). Therefore, bacterial adaptation to the plant root environment may be essential for the successful implementation of microbiome services in agriculture in order to support plant health.

Among all root-dwelling organisms, fluorescent *Pseudomonas* spp. are well characterized in terms of the traits required for their growth in the rhizosphere and for the establishment of beneficial plant-microbe interactions (17, 18). To study bacterial traits involved in efficient root colonization, mutants defective in specific traits suspected of being involved in colonization are compared to the parental strains for their ability to colonize plant roots. Such studies have highlighted a range of bacterial traits involved in efficient root colonization, including flagella (19), surface lipopolysaccharides (LPS) (20), and amino acid synthesis (21). Using random mutagenesis and by determining the fitness of each mutant in competition with its parental strain in the rhizosphere, many other important traits for rhizosphere competence in *Pseudomonas* were discovered (17). Recently, random mutagenesis in *Pseudomonas capeferrum* WCS358 led to the identification of two genes that are important for gluconic acid (GA) biosynthesis. GA, in turn, is essential for the suppression of local, flagellin-induced root immune responses (22). Such suppression was shown to be important for rhizosphere competence, as GA-deficient mutants maintain reduced populations in the rhizosphere (22). In another recent study, genome-wide saturation mutagenesis of *Pseudomonas simiae* WCS417r revealed that 2% of the protein-coding genes are important for successful root colonization (23). Mutations that negatively affect rhizosphere competence and mutations that confer a competitive advantage were identified in this study. The identification of mutations that can lead to increased root colonization (23) suggests that there is room for improvement of bacterial fitness in the rhizosphere.

In the present study, we used an experimental-evolution approach (24) to study how *Pseudomonas protegens* CHA0 evolves during repeated colonization of the rhizosphere of the model plant *Arabidopsis thaliana*. The model biological control agent CHA0 displays broad-spectrum antagonistic activity against several plant-pathogenic fungi and bacteria (11), and its complete genome sequence is available (25). We performed highly controlled experimental evolution in a gnotobiotic and carbon-free sand system in which bacteria depend solely on the plant for their supply of carbon. Following inoculation and establishment on the roots, bacterial populations were transferred to new plants, and this cycle was repeated eight times. We hypothesized that the repeated colonization and establishment of the bacterial population on the plant root would create an environment in which selection pressure drives the accumulation of better colonizers. *In vitro* characterization of individual bacterial colonies from these populations combined with sequencing analysis led to the identification of...
several evolutionary trajectories involving 35 distinct mutations that impact social traits representing interpopulation communication and cooperation, carbon source utilization, motility, or biocontrol activity. By combining experimental evolution with whole-genome resequencing, we created a powerful screen for the identification of adaptive mutations with positive effects on rhizosphere colonization.

RESULTS

Mutational events in independent evolutionary lines. We previously studied five experimental-evolutionary populations, referred to as lines, of CHA0 evolving in the rhizosphere of Arabidopsis in a gnotobiotic system. Independent populations were introduced on the roots, and after 4 weeks of plant growth, the populations were transferred to new plants. This cycle of transferring was repeated eight times, and we performed extensive characterizations up until cycle 6 to account for feasibility (26). In short, for each line, after every cycle, we plated a fraction of the population on culture media and randomly picked 16 colonies for extensive phenotypic assessment of bacterial life history traits (26). In order to study adaptation at the genetic level, we selected six colonies from each line at cycles 2, 4, and 6, such that they represented most of the observed phenotypic diversity among the 16 colonies that were initially picked and characterized. These colonies, as well as six colonies from the ancestral population that was initially introduced on the roots, were resequenced to an average depth of 25-fold coverage (minimum, 10; maximum, 70) and used for the identification of single nucleotide polymorphisms (SNPs), as well as small and large insertions or deletions (indels). In total, we thus set out to acquire genetic data for 96 bacterial colonies (5 lines \( \times \) 3 cycles \( \times \) 6 colonies + 6 ancestral colonies). Unfortunately, we were unable to acquire sufficient sequencing data for two colonies from line 4 at cycle 4, yielding a final set of 94 (88 evolved, 6 ancestral) resequenced colonies. The six ancestral colonies were all identical, indicating that there was no genetic variation in the starting population and that all observed mutations were de novo mutations. In total, one or more mutations were detected in 64 evolved colonies, which collectively represent 73% of the 88 characterized bacterial colonies (see Table S1 in the supplemental material). We identified 5 synonymous substitutions, 20 nonsynonymous substitutions, and 4 deletions ranging in length from 1 bp to about 400 bp, distributed over 22 genes, and 6 additional mutational events located in intergenic regions (Table 1). Mutations located in intergenic regions possibly affect the transcription of nearby genes via affecting their regulatory protein binding sites and making subsequent changes to their promoter activity (27, 28). Several mutations were found to be clustered in select genes and/or regions in the CHA0 genome, e.g., those in the response regulator gacA gene (PFLCHA0_RS17965; GenBank accession no. NC_021237, 4,039,113–4,039,754) or in the OBC3 gene cluster involved in LPS biosynthesis (accession no. NC_021237, 2,173,707–2,196,028) (29), but the majority of the mutations were spread across the 6.1-Mbp CHA0 genome (Fig. 1). Functional characterization of the mutated genes by analyzing their cluster of orthologous groups (COG) annotation revealed that the majority of these genes are involved in transcription (COG term K), signal transduction mechanisms (COG term T), amino acid transport and metabolism (COG term E), and cell wall/membrane/envelope biogenesis (COG term M) (Fig. 1; Table 1).

Identification of potential root colonization genes. Bacterial genes that are involved in the colonization of plant roots can be revealed by identifying beneficial mutations that evolve during adaption of bacteria to the rhizosphere environment and have positive effects on root colonization. Over time, such mutants will outcompete the ancestral strain and become dominant in the bacterial population. The observation that only a limited number of mutations accumulated relative to the total number of genes within the genome across the entire experiment makes it highly unlikely that the same gene would acquire several changes by chance in independent evolutionary lines. Nevertheless, we observed recurrent mutations in several of the same genes and/or pathways (Table 1), which is a strong indication for adaptive evolution. Genes or pathways that acquired mutations in multiple independent CHA0 populations

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### TABLE 1 Mutations that occurred in 28 genes during Arabidopsis rhizosphere adaptation over five replicate CHA0 populations (lines)

| Locus tag* | Gene name | Description | COG\(^b\) | Line(s) | Total no. of alleles | Total no. of mutations or deletions in protein-coding genes | No. of intergenic mutations |
|-----------|-----------|-------------|-----------|--------|---------------------|------------------------------------------------|--------------------------|
|           |           |             |           |        |                     | Nonsynonymous | Synonymous | Deletions |                     |
| PFLCHA0_RS02080 | hutI | Imidazolonepropionase | F | 4 | 1 | 0 | 1 | 0 | 0 |
| PFLCHA0_RS03400 | accC | Biotin carboxylase, acetyl-CoA carboxylase subunit | I | 2 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS05510 | nudL | Coenzyme A pyrophosphatase/nudix hydrolase NudL | L | 4 | 1 | 0 | 1 | 0 | 0 |
| PFLCHA0_RS06125 | rpoS | RNA polymerase sigma factor RpoS | K | 5 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS08340 | flxQ | Sigma 54-dependent Fis family transcriptional regulator | T | 5 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS08490 | flhA | Flagellar biosynthesis protein FlhA | N | 4 | 1 | 0 | 1 | 0 | 0 |
| PFLCHA0_RS09880 | RS09880 | Glycosyltransferase (GT) | M | 5 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS09980 | oafA | O-antigen acetylase | I | 1, 3 | 2 | 1 | 0 | 1 | 0 |
| PFLCHA0_RS09992 | galE | UDP-glucose 4-epimerase | M | 2 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS11785 | RS1178S | LysR family transcriptional regulator | K | 4 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS11820 | RS11820 | Paal family thioesterase | Q | 4 | 1 | 0 | 1 | 0 | 0 |
| PFLCHA0_RS12070 | RS12070 | 2OG-Fe(ll) oxygenase superfamily | S | 5 | 1 | 0 | 1 | 0 | 0 |
| PFLCHA0_RS13000 | yvaQ2 | Methyl-accepting chemotaxis protein | NT | 2 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS14960 | tetR | TetR/AcrR family transcriptional regulator | K | 5 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS17350 | RS17350 | Methyltransferase domain-containing protein | H | 1 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS17965 | gacA | UvrY/SirA/GacA family response regulator transcription factor | K | 1, 2, 4 | 6 | 5 | 0 | 0 | 1 |
| PFLCHA0_RS18525 | RS18525 | ABC transporter substrate-binding protein | E | 5 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS21265 | prdS | RNA polymerase factors sigma 70 | K | 1 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS21275 | RS21275 | Transporter substrate-binding domain-containing protein | ET | 1 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS2185S | wtpM | Polysaccharide biosynthesis protein/NDP-sugar epimerase | GM | 1 | 1 | 0 | 0 | 0 | 0 |
| PFLCHA0_RS22600 | gacS | Hybrid sensor histidine kinase/response regulator | T | 3 | 1 | 1 | 0 | 0 | 0 |
| PFLCHA0_RS22950 | argT5 | ABC transporter substrate-binding protein/lysine-arginine-ornithine-binding periplasmic protein ArgT | ET | 4 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS25175 | mraZ | Division/cell wall cluster transcriptional repressor MraZ | K | 2 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS26215 | osmY | Osmotically inducible protein OsmY/BON domain-containing protein | S | 2 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS27515 | rpsH | 30S ribosomal protein 58 | J | 4 | 1 | 0 | 0 | 1 | 0 |
| PFLCHA0_RS30075 | sadB | Surface attachment defective (SadB) ortholog/HDOD domain-containing protein | T | 1 | 2 | 2 | 0 | 0 | 0 |
| PFLCHA0_RS30120 | tssM | Type VI secretion system membrane subunit TssM | S | 2 | 1 | 0 | 1 | 0 | 0 |
| PFLCHA0_RS31060 | nlpD | Lipoprotein NlpD/LppB/LysM domain-containing protein | D | 3 | 1 | 1 | 0 | 0 | 0 |

Total 28 35 20 5 4 6

*Corresponding updated GenBank CHA0 locus tags can be found at [https://doi.org/10.6084/m9.figshare.13295828.v2](https://doi.org/10.6084/m9.figshare.13295828.v2).

COG, cluster of orthologous groups ([103; https://www.ncbi.nlm.nih.gov/COG](https://www.ncbi.nlm.nih.gov/COG)).
included gacA and gacS, the OBC3 gene cluster (oafA; galE; PFLCHA0_RS09880 [29]), and a putative pyoverdine siderophore biosynthesis cluster (pvdS; PFLCHA0_RS21275).

Other genes, like sadB, were targeted more than once in the same population. Because these genes were repeatedly identified in the CHA0 evolution experiment, they can be
assumed to contribute significantly to bacterial fitness in the rhizosphere (Table 1; Fig. 2). Moreover, our findings suggest that the independent evolutionary lines converge on similar evolutionary trajectories involving overlapping biological processes and molecular mechanisms.

To illustrate the diverse evolutionary trajectories, we constructed phylogenetic trees of each replicate population (line) using all mutations that were detected (Fig. 3). In these lines, both the numbers and the depths of branches were very different (Fig. 3; Table 2). In lines 1 and 3, the populations were swept early by oafA mutants and later by specific gac mutants, while in lines 2 and 4, the coexistence of multiple genotypes characterized by gac mutations was observed, indicative of clonal interference between lineages (Fig. 3) (30). Notably, the relative abundance of each genotype in the population assumed to contribute significantly to bacterial fitness in the rhizosphere (Table 1; Fig. 2). Moreover, our findings suggest that the independent evolutionary lines converge on similar evolutionary trajectories involving overlapping biological processes and molecular mechanisms.

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FIG 3  Phylogenetic trees and Muller plots of five independently evolving *P. protegens* CHA0 populations. (Left) Phylogenies for 18 genomes from each population (16 for the population in experimental line 4), based on the sequential appearance of mutations, are shown. Synonymous mutations are underlined. Circles represent unique bacterial genotypes. The color of the circle fill represents the time point at which a genotype was first detected. OBC3 and gac mutants are highlighted in blue and purple, respectively. (Right) Muller plots depicting the dynamics of mutant alleles during the evolutionary experiment. The Muller plots show the estimated frequencies, by their heights, of 33 mutations in the respective population over 6 experimental cycles. Descendant genotypes are shown emerging from inside of their respective ancestors. The frequency of each mutation in the respective population can also be found in Table 2. Muller plots are prepared using ggmuller in R (102).
is estimated from a small set of sequenced individuals and therefore is expected to lack precision. The evolved isolates contain one to four mutations in general, and in each line, as was expected for independently evolving populations. As the frequency of each mutation (Fig. 3; Table 2) was determined from only six individuals, the evolutionary line is estimated from a small set of sequenced individuals and therefore is expected to lack precision. The HRM methodology allows for the accurate quantification of mutant frequencies across a wide range for all three mutations (P < 0.001) (Fig. S1). Using this method, we

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### Table 2 Nomenclature for CHA0 mutations and variants and their frequency/cycle in each respective population

| Line | Locus tag | Gene name | DNA sequence change<sup>a</sup> | Amino acid change (three-letter code)<sup>b</sup> | Standard nomenclature (used in text) | Frequency in the population (x/6)<sup>c</sup> | Cycle 2 | Cycle 4 | Cycle 6 | HRM |
|------|-----------|-----------|-------------------------------|-----------------------------------------------|---------------------------------------------|-----------------------------------------------|--------|--------|--------|------|
| 1    | PFLCHA0_RS090890 | adaA | c.1005C>A | p.Tyr335X | Y335X | 4/6 | 6/6 | 6/6 | |
| 1    | PFLCHA0_RS17350 | RS17350 | c.230delC | p.Ala77Ala,fsX14 | A77A,fsX14 | 2/6 | 6/6 | 6/6 | |
| 1    | PFLCHA0_RS17965 | gacA | c.145G>T | p.Asp49Tyr | D49Y | 2/6 | 4/6 | 2/6 | 9.2% | 19.1% | 46.7% | x |
| 1    | PFLCHA0_RS21855 | wbpM | c.235G>C | p.Gly79Arg | G79R | 1/6 | | | |
| 1    | PFLCHA0_RS21275 | RS21275 | c.233G>T | p.Arg78Leu | R78L | 1/6 | | | |
| 1    | PFLCHA0_RS21265 | pvdS | c.-337>C | NA | -337>C | 1/6 | | | |
| 1    | PFLCHA0_RS30075 | sadB | c.548G>C | p.Arg183Pro | R183P | 1/6 | | | |
| 1    | PFLCHA0_RS17965 | gacA | c.773T>A | p.Leu238Gln | L238Q | 2/6 | | | |
| 2    | PFLCHA0_RS09920 | gaiE | c.94G>A | p.Val32Met | V32M | 2/6 | 5/6 | 6/6 | |
| 2    | PFLCHA0_RS30120 | tssM | c.1336C>T | p.Leu446Leu | L446F | 1/6 | | | |
| 2    | PFLCHA0_RS03400 | accC | c.1237G>A | p.Glu413Lys | E413K | 4/6 | 6/6 | | |
| 2    | PFLCHA0_RS13000 | yvaQ2 | c.-9G>T | NA | -9G>T | 1/6 | | | |
| 2    | PFLCHA0_RS17965 | gacA | c.160G>T | p.Asp54Tyr | D54Y | 2/6 | | | 5.5% | 4.5% | 15.5% | x |
| 2    | PFLCHA0_RS22600 | gacS | c.80G>A | p.Gly27Asp | G27D | 1/6 | 4/6 | | | 3.8% | 19.9% | 75.6% | x |
| 3    | PFLCHA0_RS31060 | nlpD | c.590A>G | p.Gln197Pro | Q197P | 1/6 | | | |
| 4    | PFLCHA0_RS22950 | argT5 | c.120-533del | p.Lys40-Leu185del,fsX6 | ΔargT5 | 2/6 | | | |
| 4    | PFLCHA0_RS02080 | hutL | c.786C>T | p.Phe262Phe | P262P | 1/6 | | | |
| 4    | PFLCHA0_RS17965 | gacA | c.-40T | NA | -40T>A | 2/6 | 1/4 | | |
| 4    | PFLCHA0_RS17965 | gacA | c.112G>T | p.E38X | E38X | 2/6 | 3/4 | | |
| 4    | PFLCHA0_RS05510 | nudl | c.288C>T | p.Ser96Ser | S96S | 2/6 | 3/4 | | |
| 4    | PFLCHA0_RS11820 | RS11820 | c.33C>T | p.Ala11Ala | A11A | 1/6 | | | |
| 4    | PFLCHA0_RS08490 | fha | c.1176delT | p.His393Gln,fsX15 | H393Q,fsX15 | 3/4 | | | |
| 4    | PFLCHA0_RS11785 | RS11785 | c.766A | p.Ser256Cys | S256C | 3/4 | | | |
| 4    | PFLCHA0_RS27515 | rpsH | c.90G>C | NA | -90G>G | 1/4 | | | |
| 5    | PFLCHA0_RS18525 | RS18525 | c.254delT | p.Val85Gly,fsX40 | V85G,fsX40 | 1/6 | | | |
| 5    | PFLCHA0_RS06152 | ropS | c.193C>T | p.Gln65X | Q65X | 1/6 | 4/6 | | |
| 5    | PFLCHA0_RS08340 | fieQ | c.959G>A | p.Arg320Gln | R320Q | 1/6 | | | |
| 5    | PFLCHA0_RS14960 | tetR | c.381G>C | p.Tyr127X | Y127X | 2/6 | | | |
| 5    | PFLCHA0_RS09880 | RS09880 | c.246C>A | p.Tyr82X | Y82X | 1/6 | | | |
| 5    | PFLCHA0_RS12070 | RS12070 | c.463C>A | p.Glu155Lys | E155K | 1/6 | | | |

<sup>a</sup>DNA sequence change positions are relative to the cDNA (c). del, deletion. Mutations upstream of the transcription start site (TSS) are preceded by a dash to indicate the position relative to the TSS.

<sup>b</sup>Standard nomenclature for CHA0 mutations and variants (used in text).

<sup>c</sup>Frequency in the population (x/6).

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*DNA sequence change positions are relative to the cDNA (c). del, deletion. Mutations upstream of the transcription start site (TSS) are preceded by a dash to indicate the position relative to the TSS. 

*Stop codon (at its relative position in case of a shifted frame); fs, frameshift; p, amino acid substitution relative to the protein; NA, not applicable.

*Percentages represent the frequency of the selected (indicated by x) mutation in the population as determined by PCR-based high-resolution melt (HRM) analysis.
found that the population-level frequency of these three mutations in the respective evolutionary lines corresponded remarkably well with the frequency previously obtained from the cultured and sequenced isolates (Table 2; Fig. S2). These findings corroborate the culture-based quantification of mutant frequency, suggesting that they provide a reasonable measure for population-level frequency. Intriguingly, it can be seen that the \textit{gacS}\textsubscript{G27D} mutation reaches fixation after eight experimental cycles, while the \textit{gacA}\textsubscript{D54Y} and \textit{gacA}\textsubscript{D54Y} mutants stabilize at around 50% and 25%, respectively (Fig. S2). Stabilization of mutations is indicative of frequency-dependent (FD) selection putatively reinforced by clonal interference with coexisting lineages carrying beneficial mutations in \textit{sadb} (L258Q) and \textit{gacA} (G97S) (Fig. 3). FD selection describes the phenomenon that the fitness of a particular genotype or phenotype is dependent on its frequency. Such context dependency has been linked to cheating behavior in which microbial cells that lack or have limited production of certain costly compounds benefit from other cells that do produce these compounds. When a minimal amount of such a compound increases genotype or phenotype fitness, FD selection can occur, resulting in stabilization of the mutation frequency.

**Early adaptations are driven by cell surface-related genes.** In general, mutations that are fixed early on in the rhizosphere adaptation process tend to have a high selective advantage (31). Disruptive mutations in \textit{aafA}, resulting in premature stops halfway in the coding region, were detected as the first acquired mutations in two independent evolutionary lines (lines 1 and 3) and appear to have swept the population in the following generations (Table 2). \textit{aafA} is part of the O-polysaccharide (O-PS; O-antigen) biosynthesis cluster 3 (\textit{OBC3}) (29) and encodes an O-acetyltransferase, which is postulated to acetylate the O-antigen component of the outer membrane LPS (32). Another \textit{OBC3} mutation that accumulated early on in the rhizosphere adaptation process, \textit{galE}\textsubscript{E94G\rightarrow A}, leads to an amino acid substitution (V32M) in \textit{galE}’s product, and this mutation swept through the population in evolutionary line 2, reaching fixation in cycle 6 (Table 2; Fig. 3). \textit{galE} encodes an UDP-glucose 4-epimerase which is involved in O-antigen and LPS core biosynthesis (33–35). One colony with a mutation in a third \textit{OBC3} cluster gene, \textit{RS09880}, encoding a putative glycosyl transferase (GT), was found in cycle 6 of evolutionary line 5. Thus, in four out of the five evolutionary lines, mutations that likely affect bacterial LPS structure appeared during rhizosphere adaptation, and these mutations became dominant in three out of four evolutionary lines. These results strongly suggest that modifying bacterial cell surface structure is an important bacterial strategy in early adaptation to the rhizosphere.

**Adaptation driven by global regulators.** In the present study, six mutations were detected in \textit{gacA} in three out of five evolutionary lines, representing approximately 20% of all missense mutations. Notably, in evolutionary lines 2 and 4, multiple \textit{gacA} alleles accumulated, some of which were detected in early experimental cycles (Table 2). Additionally, a \textit{gacS} mutation accumulated in line 3. \textit{gacA} and \textit{gacS} encode the main constituents of the conserved GacA/GacS two-component regulator system, i.e., the hybrid sensor histidine kinase GacS and the cognate response regulator GacA (Fig. 2). In Gram-negative bacteria, activation of GacS results in cross-phosphorylation of GacA via phosphotransfer, which in turn leads to activation of the expression of the small RNA genes \textit{rsnY} and \textit{rsnZ} via its helix-turn-helix (HTH) domain-binding domain (36). In \textit{CHA0}, this regulatory pathway is known to control quorum sensing as well as secondary metabolism and stress resistance (37–40). In the closely related strain \textit{P. protegens} Pf-5, this pathway was shown to have a big impact on bacterial genome-wide gene expression, affecting the expression of more than 10% of the annotated genes (41). Similarly, \textit{gacA} mutants that arose on the edge of swarming Pf-5 colonies showed dramatically altered genome-wide gene expression patterns (42).

Including \textit{gacS} and \textit{gacA}, about half of the mutated genes in this study are global regulators or sigma factors (Table 1; Fig. 1). This high frequency suggests that global regulator-controlled networks are evolvable and play a major role in rapid bacterial adaptation. Pleiotropic adaptive mutations in global regulator genes have been shown to be important for bacterial adaption both in the laboratory (43, 44) and in natural
Bacterial motility. Bacterial motility is an important trait for rhizosphere competence, mediating colonization of distal parts of the root system (48), and both LPS and the GacS/GacA two-component regulator system are known to affect this trait. The O-antigen side chain of the LPS was reported to contribute to swimming and swarming motility in the plant-pathogenic bacterium Erwinia amylovora (49, 50). The GacS/GacA two-component regulator system controls bacterial motility, for example, by affecting the transcription of genes related to flagellum and biosurfactant biosynthesis (42, 48, 51). We also identified several other mutations across the various evolutionary lines that can be linked to bacterial motility. For instance, a disruptive mutation in the flagellar biosynthesis protein-coding gene flhA (H393Q/fsX15) that is involved in the biogenesis of the bacterial flagellum (Table 1) appeared in evolutionary line 4, reaching up to a predicted frequency of 75% (Table 2; Fig. 3). In P. fluorescens Pf-5 and in Pseudomonas aeruginosa PA01, FlhA is reported to be essential for swimming motility (41).

Furthermore, we identified one amino acid substitution, R320Q, in FleQ and two in SadB, R183P and L258Q, that based on sequence similarity to well-studied, homologous proteins in other bacteria can be linked to motility in addition to several other bacterial traits. FleQ is a $\sigma^{54}$-dependent Fis family transcriptional regulator which regulates flagellar motility and biofilm formation as well as Pel exopolysaccharide (EPS) production in response to cellular c-di-GMP levels in P. aeruginosa (52). P. protegens FleQ shares 84% sequence identity and 98% sequence coverage with P. aeruginosa FleQ, and like P. aeruginosa FleQ, it is comprised of the N-terminal flagellar regulatory FleQ domain (PF06490), a central AAA+/ATPase $\sigma^{54}$ interaction domain (PF00158), and a C-terminal Fis-type HTH DNA-binding domain (PF02954). The R320Q substitution that we identified here is found in the AAA+/ATPase $\sigma^{54}$-interacting domain in between the arginine (Arg) finger (amino acids 300 to 303) and a c-di-GMP-binding motif, ExxxR (amino acids 330 to 334) (52). The conserved arginine residues in FleQ, including the here-mutated Arg320, are thought to be important for protein oligomerization, and substitution of any of these residues abolishes ATPase activity in Vibrio cholerae EpsE completely (53). Finally, sadB encodes an HD-related (named after the conserved histidine [H] aspartic acid [D] doublet of predicted catalytic residues [104]) output domain (HDOD)-containing protein (Table 1) that shares 77% sequence identity and 99% sequence coverage with P. aeruginosa SadB. In P. aeruginosa, SadB stimulates Pel EPS production and the chemotaxis-like cluster CheIV, which in turn affects flagellar motility as well as biofilm formation (54). In Pseudomonas fluorescens F113, SadB and FleQ control flagellar motility, both dependent upon and independent of the GacS/GacA two-component regulator system (55, 56).

Parallelism of targeted mutations on this functional-motility pathway impelled us to assess bacterial motility and track its dynamics across all evolutionary lines. We selected all OBC3 and gac mutants as well as sadB, fleQ, and flhA mutants. Additionally, we included two gacA mutant progenitors with mutations in accC and RS17350, encoding a biotin carboxylase and a methyltransferase, respectively, plus two gacA mutant descendants with mutations in osmY and mraZ that encode an osmotically inducible protein and a transcriptional repressor, respectively (Table 1). Altogether, we assessed the swimming and swarming motility of strains with 17 distinct genotypes (Fig. 4; Fig. S3). We found that OBC3 mutants, the gacA progenitors with mutations in accC and RS17350, and the gacA descendants with mutations in osmY and mraZ were unaltered compared to their respective ancestors when both swimming and swarming motility were considered, with the exception of a small yet significant increase in the swimming motility of the oafA$^{Y335X}$ mutant. However, gac mutants themselves were significantly affected, both in swimming, which is generally enhanced, and in swarming, which is repeatedly decreased (Fig. 4). sadB mutants, like gac mutants, display enhanced swimming and worsened swarming compared to those of their respective progenitors. Oddly, both
Evolved, root-competent mutant *P. protegens* CHA0 strains are characterized by enhanced swimming and impaired swarming. The accumulative effects of parallel and sequential mutations on swimming motility (top panel) and swarming motility (Continued on next page)
the fleQ and the flhA mutant displayed severely reduced swimming and swarming motility, thus representing two examples of an alternative evolutionary route toward adaptation in the rhizosphere in which motility is reduced. Loss of motility might coincide with another trait in these cases, such as EPS production and/or biofilm formation, regulated via shared yet oftentimes opposing mechanisms. Also, the frequency of the fleQ mutation was low; i.e., only one out of six isolates from cycle 6 of experimental line 5 carried this mutation, and therefore, whether this mutant is truly beneficial is unclear. On the other hand, the flhA mutation was found in three out of the four cultured and sequenced isolates from cycle 6 of line 4 and therefore may be present in a significant proportion of the total population.

**Dynamics of global phenotypic change.** Since natural selection eventually operates at the phenotypic level, revealing bacterial global phenotypic evolutionary dynamics can help us to identify traits that are under selection. Moreover, beneficial genetic mutations can be predicted if they are linked to well-known root colonization traits. A broad range of 30 bacterial traits, including different aspects of bacterial life/history traits, were examined for the sequenced isolates, which allows for genotype-phenotype association analysis.

As shown in Fig. 5a, the 30 bacterial traits separated into four clusters that share similar patterns across the different mutant genotypes, and this clustering is supported by model-based clustering analysis (Fig. S4). The growth of the bacteria in 1/3-strength King’s B (KB) medium was positively correlated with siderophore and exoprotease production, tryptophan side oxidase activity, and growth inhibition of the bacterial plant pathogen *Ralstonia solanacearum*. Thus, this cluster, designated cluster 1, contains traits associated with bacterial social behavior related to microbe-microbe communication and cooperation, such as the production of public goods. In a principal-component analysis (PCA) (Fig. S5a), the first principal component (PC1) is strongly correlated with all seven traits, with a total explanation of 53.9% for all variables. Cluster 2 contains traits linked to carbon source utilization. For this cluster, PC1 is strongly correlated with all carbon source usage-related traits, with a total explanation of 83.9% for all variables (Fig. S5b). A third cluster was observed for the bacterial ability to form a biofilm, to produce indole-3-acetic acid (IAA), and to inhibit the growth of two fungal plant pathogens, with a total explanation of 82.4% by PC1 (Fig. S5c). Finally, cluster 4 contains all seven traits that are related to bacterial resistance to biotic and abiotic stresses. The PC1 is strongly correlated with all seven traits, with a total explanation of 53.9% for all variables (Fig. S5d). For these four clusters, the PC1 (or –PC1) value is used as a proxy to present the general performance of all the traits that clustered together.

As is shown in Fig. 5b, all five evolutionary lines showed parallel trends of accumulative declines in social traits. Mutations in *gacA, gacS, and rpoS* resulted in significantly decreased bacterial social traits, and this was also observed for the *RS17350* double mutant relative to the social traits of a strain with the background *gacA* mutation. *RS11785*, like *gacA*, encodes a LysR-type transcriptional regulator. In addition to the mutations in these global regulators, earlier mutations in *oafA, gaeL, and RS17350* encoding a putative methyltransferase, also resulted in a significant but relatively small decrease in social traits. This parallel decline of the bacterial...
social trait index suggests a negative selection of bacterial social behavior, especially the production of costly public goods, such as siderophores and exoproteases.

All traits related to the utilization of 14 different carbon sources, which were selected based on their reported presence in root exudates of Arabidopsis (57), grouped in one cluster (cluster 2) (Fig. 5a). This suggests that carbon source utilization is coregulated. Different mutations in gacS and gacA resulted in contrasting bacterial carbon source

FIG 5 Genotype-phenotype association analysis of evolved bacterial populations. Isolates without mutations or with only synonymous mutations were excluded for this association analysis. (a) Heatmap representing the performance of 30 life history traits of six ancestral and 63 evolved isolates. The ward. D2 method was used for hierarchical clustering, based on the dissimilarities of the set of objects being clustered. The four clusters depicted on top are model predicted. Each column illustrates one bacterial life history trait, indicated at the bottom of the figure. Each row represents an isolate, indicated by its sample ID number and mutational events. (b to e) Principal-component analysis was applied to generate a general symbolic index for each model-predicted cluster. The proxy values of each cluster were normalized; higher values represent better performance. To illustrate better the accumulative effects of each mutational event, ANOVA was applied to reveal the phenotypic change between each genotype and its identified nearest ancestor in each line (this was applied only for genotypes that were detected more than once \( n \geq 2 \)). Asterisks alongside the mutational events indicate significant differences (*, \( \alpha = 0.05 \); **, \( \alpha = 0.01 \); ***, \( \alpha = 0.001 \)). Colored circles represent bacterial genotypes, with different colors representing independent evolving populations, and the size denotes the number of replicates (n). The dashed line represents the average value of the ancestor.
utilizations. A mutation in the first N-terminal transmembrane domain of GacS (G27D) (Fig. 2) resulted in significant enhancement of carbon source utilization (Fig. 5a and c). A similar trend was observed for the gacAG97S mutant. In contrast, the majority of evolved genotypes, including most mutations in gacA and rpoS, showed a reduced ability to utilize carbon sources (Fig. 5a and c).

Traits in clusters 3 (Fig. 5d) and 4 (Fig. 5e), unlike in the previous two clusters, were more stable, as most genotypes behave like the ancestor. Only two gacA mutants, the gacAE38X and gacAY183S mutants, were associated with a significant decline of bacterial traits associated with biocontrol, i.e., antifungal activity and biofilm formation, while the rpoSQ65X mutation resulted in a significant increase in these traits. The double mutation RS11785S256CflhAH393Q.fsX15, in the background of the disruptive gacAE38X mutation (Fig. 5e), was the only mutational event that led to a significant increase of general resistance to various environmental stresses. flhA encodes the flagellar biosynthesis protein FlhA and is linked to bacterial motility. The LysR-type transcriptional regulator RS11785 comprises an N-terminal HTH DNA-binding domain (PF00126) and a C-terminal substrate-binding domain (PF03466). This substrate-binding domain is linked to aromatic compound degradation and resembles that of the type 2 periplasmic binding proteins, or PBP2s. PBP2s are responsible for the binding and uptake of several substrates, including polysaccharides and several amino acids. We used HHpred (58) for protein remote homology detection and three-dimensional structure analysis to assess the possible consequence of the S256C amino acid substitution. This analysis identified OxyR from *Escherichia coli* as the best template for modeling (E value, 2.9e–30; score,
OxyR represents the master peroxide sensor in Gram-negative bacteria and operates via intramolecular disulfide bond formation in the regulatory PBP2-like domain. RS11785 is unlikely to represent the master peroxide sensor in P. protegens CHA0 because another CHA0 protein encoded by PFLCHA0_RS030065 is much more similar to OxyR (89% pairwise sequence identity versus 29% for RS11785). Nevertheless, it is tempting to speculate that the serine (S, Ser)-to-cysteine (C, Cys) amino acid substitution (S256C) that we observed here might influence regulatory activity by altering intramolecular disulfide bond formation, as such bonds are governed by pairs of cysteine residues. Lastly, RS11785 bears resemblance (37% sequence identity, 93% sequence coverage) to the LysR-type PBP2 domain-containing regulator alsR, which regulates the activity of the acetoin operon (alsSD) in response to several signals, such as glucose and acetate, in Bacillus amyloliquefaciens. Acetoin (3-hydroxy-2-butanone, a precursor of 2,3-butanediol) can elicit induced systemic resistance (59) and is linked to general plant growth promotion in bacilli. In the absence of singular mutations in flhA and RS11785, the molecular mechanism underlying the observed enhanced environmental stress resistance in this double mutant remains to be clarified, but we hypothesize that altered regulatory activity by RS11785256C is the most likely cause.

**DISCUSSION**

The rhizosphere is a nutrient-rich environment for root-associated bacteria. However, to access the available nutrients, bacteria must overcome various challenges, including the plant immune system, the presence of competing and/or predatory microorganisms, and abiotic stresses. In the gnotobiotic binary system used in this study, we tracked changes in P. protegens CHA0 in the rhizosphere of Arabidopsis under reproducible and controlled conditions without interference of complex interactions with other microbes. Mutations affecting global regulators, bacterial cell surface structure, and motility accumulated in parallel in our evolutionary experiment, revealing at least three important strategies of bacterial adaptation to the rhizosphere.

**Global regulators and rhizosphere adaptation.** The GacS/GacA two-component regulator system controls the production of antimicrobial secondary metabolites, exoenzymes, and siderophores but also biofilm formation, stress responses, motility, and quorum sensing (13, 38, 41, 60, 61). In the present study, mutations in the GacS/GacA two-component regulator system caused dramatic changes in several bacterial phenotypic traits, including motility, carbon source utilization, and social traits (Fig. 4 and 5). The overall gain in swimming motility is not unexpected, as swimming motility, driven by the flagellum apparatus, has been repeatedly reported to be an important root colonization trait in root-associated bacteria, including several *Pseudomonas* spp. (17, 62–64). In line with this, genome-wide transposon disruption mutant analysis by Cole and coworkers showed that the majority of motility-related genes in *Pseudomonas simiae* WCS417 are positively associated with Arabidopsis root colonization (23).

In contrast with the observed enhanced swimming motility across the evolutionary selection lines, swarming motility was found to be severely hampered throughout, and this appears to be the case especially for gac mutants. Swarming, like swimming, is driven by the flagellum but in addition depends on the production of several compounds, including quorum-sensing molecules and biosurfactants. *P. protegens* CHA0 is a known producer of the biosurfactant orfamide A, and the GacS/GacA two-component system is known to be an important regulator for its biosynthesis (65). These results fit observations by Song and coworkers (42) in the closely related *P. protegens* strain Pf-5. During swarming motility, gac mutants that lack production of the surfactant orfamide emerged. These mutants cannot swarm, but they coswarm with orfamide-producing cells (42).

Remarkably, we also identified two completely nonmotile mutants in our experiments with disruptions in the fleQ and flhA genes. Possibly, these mutants adhere much better to the root surface or might be able to form a biofilm more rapidly or strongly, as these traits
are often inversely correlated with bacterial motility (66–68). Future experiments should reveal whether such a tradeoff between bacterial motility and root adherence indeed underlies our observations in this evolutionary experiment.

The decreased production of public goods, such as siderophores and exoproteases, which were observed in all independent evolutionary lines (Fig. 5), may be beneficial for bacterial fitness by saving energy and primary metabolites. For example, adaptation of *Pseudomonas aeruginosa* to the human host through mutations in regulators was accompanied by losses of siderophore production, secreted proteases, and biofilm formation (69, 70).

**Cell surface structures as bacterial adaptation targets.** Bacterial cell surface components are the first line of defense against environmental stress and interplay with hosts (71–73). LPS is a central outer membrane component for Gram-negative bacteria and exhibits structural adaptability that is contributed especially by its O-antigen part (71, 73). Bacterial O-antigen structure modification plays an important role in the evasion of host immunity (74) and has the potential to change host-bacterium interactions (71, 73, 75). In plant-pathogenic bacteria, LPS components are important virulence determinants (76, 77) that can activate a variety of defense-related responses (76, 78–80). In *P. fluorescens*, the O-antigen component has been implicated to induce systemic resistance in radish (81) and *Arabidopsis* (82).

We observed parallel mutations in genes that are involved in LPS biosynthesis and structure modification. In three out of the five evolutionary lines, the first fixed mutations were identified in *oafA* and *galE*, which are annotated as related to O-antigen biosynthesis and structure modification (Table 1). *oafA* encodes an O-acetyltransferase, which is postulated to modify the O antigen by acetylation (32). The enzyme GalE, UDP-galactose 4-epimerase, is involved in the interconversion of UDP-glucose to UDP-galactose, an essential intermediate for LPS core and O-antigen structures (33–35). Inactivation of *galE* in *Porphyromonas gingivalis* resulted in shortening of the O antigen (83), and in *Bradyrhizobium japonicum*, disruption of *galE* resulted in the complete absence of O antigen (33). Thus, it is tempting to speculate that evasion of the plant’s immune response plays a role in the adaptation of CHA0 to the rhizosphere of *Arabidopsis*.

In conclusion, the observed bacterial genetic and phenotypic adaptation dynamics emphasize important roles for global regulators, motility, and cell surface structure in bacterial adaptation to its host. The parallel emergence of mutations in similar genes resulted in specific fitness advantages for mutants in the rhizosphere, suggesting that this evolutionary process is driven by the rhizosphere environment.

**MATERIALS AND METHODS**

**Experimental setup.** We set up an experimental evolution experiment with *Arabidopsis thaliana* ecotype Col-0 as the host plant and *Pseudomonas protegens* CHA0 as the evolving bacterial strain. CHA0 (84) is a model strain originally isolated from roots of tobacco plants grown in soil naturally suppressive to black root rot (85). CHA0 was chromosomally tagged with green fluorescent protein (GFP) and a kanamycin resistance cassette (86) to enable consistent tracking of the strain and identification of contaminations. We previously described the setup of the evolutionary experiment in great detail (26). In brief, the ancestral bacterial population (10⁸ cells) was inoculated on *Arabidopsis* roots grown under gnotobiotic conditions inside ECO2 boxes in carbon-free silver sand. For each cycle, *Arabidopsis* seeds were surface sterilized using chlorine gas, germinated on modified Hoagland’s agar medium, and grown for 2 weeks until transplantation into the ECO2 boxes, each containing two plants (26). Inoculated 2-week-old seedlings were then grown for an additional 4 weeks, after which the root-associated bacteria were collected in 10 mM MgSO₄ and subjected to fluorescence-based cell counting by flow cytometry, yielding on average 10⁸ cells/root. Cells of the evolved bacterial populations (10⁶) were then transferred to new plants, and this cycle was repeated eight times. After each cycle, a small fraction of each population was plated on general-purpose, nonselective medium, 3 g/liter tryptic soy agar (TSA), to assess for contaminations and to verify that all colonies carried the GFP marker gene, as observed under UV light, on the one hand, and to select individual isolates for phenotypic characterization on the other hand.

We previously picked 16 random isolates from each of five experimental lines at cycles 2, 4, and 6 plus 16 isolates from the ancestor population, yielding a total set of 256 isolates (26). These 256 isolates were subsequently grouped into five distinct phenotypes based on their performance on a variety of bacterial life history traits (26). In the current study, we selected 6 of these 16 isolates per line, and per cycle, for genome sequence analysis. Isolates were selected to represent the breadth of phenotypic diversity observed previously (26). We set out to obtain genome sequences of 96 isolates in total, together with 6 isolates from the ancestor population, using the NextSeq-500 Illumina platform (2 × 75-bp paired-end reads). Sequencing of two isolates from line 4, cycle 4, however, failed, and thus a final set of 94
genomes were retrieved. We then used the snippy pipeline (https://github.com/tseemann/snippy), integrating of data generated by mapping the Illumina reads by BWA-MEM, variant calling by SAMtools and Freebayes, and variant impact analysis using SnpEff (87), to identify single nucleotide polymorphisms (SNPs) and small insertions and deletions (indels). Larger indels were identified by calculating the breadths of coverage of the mapped Illumina reads on the reference genome in a sliding window using bedtools (88). Regions with reduced coverage (<99%) were manually inspected in the Integrative Genome Viewer (IGV). Phylogenetic trees for each line were constructed manually with Illustrator based on all detected mutations, with the lengths of the branches representing the numbers of mutations. The genealogy and frequency of each lineage are shown in the Muller plots that are prepared with the R package ggmuller.

**Bacterial life history traits.** For the 94 sequenced isolates, a variety of bacterial life history traits reflecting various aspects of bacterial physiological processes were measured previously as part of all 256 isolates initially collected (26). Briefly, we monitored optical density (OD) at a wavelength of 600 nm to estimate the bacterial yield after 72 h of growth under different growth conditions in 96-well microplates. We measured bacterial growth yield, resistance to various stresses, including acidic (pH 5) and alkaline (pH 9) conditions, oxidative stress in 0.0025% H_2O_2, water potential stress (15% polyethylene glycol plates. We measured bacterial growth yield, resistance to various stresses, including acidic (pH 5) and alkaline (pH 9) conditions, oxidative stress in 0.0025% H_2O_2, water potential stress (15% polyethylene glycol 6000 [PEG-6000]), and salt stress (2% NaCl), and resistance to the antibiotics streptomycin (1 μg · ml⁻¹), tetracycline (1 μg · ml⁻¹), and penicillin (5 μg · ml⁻¹). In this study, stress resistance was defined as the ratio of bacterial growth in the stressed relative to that in the nonstressed control treatment.

**Bacterial carbon source utilization** was quantified as growth yield in modified Omston and Stanier (OS) minimal medium (89) supplemented with single-carbon sources that have been reported to be abundant in Arabidopsis root exudates (57). These included the following carbon sources: alanine, arabinose, butyrolactam, fructose, galactose, glucose, glycerol, glycine, lactic acid, putrescine, serine, succinic acid, threonine, and valine, which were added to a final concentration of 0.5 g · liter⁻¹. In addition, we measured bacterial auxin (indole-3-acetic acid, or IAA) production with a colorimetric test (90), iron-chelating ability using a modified chrome azuro 5 (CA5) assay (91), proteolytic activity by the adapted assay from Smeltzer et al. (92), tryptophan side chain oxidase activity using a colorimetric assay (93), and biofilm formation using a modified crystal violet staining assay (94). We measured the OD values reflecting the color intensities at specific wavelengths to quantify these traits. We further assessed bacterial antimicrobial activity by quantifying their effect on the growth of the fungi Vorticillium dahliae and Fusarium oxysporum and on the bacterium Ralstonia solanacearum.

**Motility assays.** Motility assays were undertaken in round petri dish plates containing Cook's Cytophaga medium (CC medium) (0.3% agar for swimming, 0.5% agar for swarming) (95), using typical swimming and swarming assays as described by Déziel et al. (96). All tested strains were grown on King's B medium agar plates for 24 h before inoculation. Swim and swarm plates were inoculated with the tested strains with a sterile toothpick. For swimming plates, the inoculum was introduced by gently piercing the agar such that the motility within the semisolid agar could be evaluated. For swarming plates, the inoculum was introduced on the agar surface, enabling visualization of motility across the agar surface. Both swimming and swarming plates were imaged after 18 h of incubation at 21°C with the right side up. The radii of swimming and swarming motility were determined from the photographs by ImageJ by examining the inner circular turbid zone inside the 0.3% agar for swimming and the outer circular zone on top of the 0.5% agar surface for swarming.

**Hierarchical and model-based clustering of bacterial traits.** (i) **Hierarchical clustering.** A heat-map to illustrate the association patterns of bacterial genotypes and their measured traits was constructed in R using the ggpplot2 package. Isolates without mutations or with only synonymous mutations were excluded from the association analysis. Hierarchical clustering was performed using the Ward.D2 method, which is based on the squared dissimilarities of the set of objects being clustered (97).

(ii) **Model-based clustering.** We applied a model-based clustering method to reveal the best-fitting structures of trait covariance patterns. For example, some traits might be either directly or indirectly regulated by the same gene, which is expected for global regulators, particularly those that can coregulate thousands of genes. We used the mclust package in R to run the model simulation (98). This method assumes that the input data are distributed as a mixture of two or more clusters. The advantage of the model-based clustering method is that it avoids heuristic assumptions and uses a soft assignment that every data point has a possibility of falling to each cluster, which facilitates the best clustering solution. The so-called Bayesian information criterion (BIC) was used to select the best model. A large BIC score indicates a better fit of the model.

This result is in line with the outcome of hierarchical clustering with adjusted Rand index (ARI) set as 1 and k set as 4 in ward.D2, as indicated in Fig. 5a. ARI is usually used to evaluate the match degree of a given clustering solution compared to the model-based clustering result, with 0 reflecting a random partition and 1 the boundary of accuracy of a certain clustering solution (98).

**Genotype-phenotype association analysis.** Bacterial traits within each model-predicted cluster have similar data distribution patterns and vary with the definition of the clustering method. Thus, we applied a linear-regression-based method, i.e., principal-component analysis (PCA), to reduce the dimensionality of data and generate a proxy for each model-predicted cluster. These proxies were later used as the x axis values in Fig. 5b to e. We applied the package ggbioplot in R to generate the PCA plots and PC1 index from the normalized data sets. The proxies were normalized for further analysis.

To examine the accumulative effects of each mutation on bacterial phenotype, analysis of variance (ANOVA) was used to compare cluster proxies of evolved genotypes with their direct ancestors. Only genotypes identified more than once (n ≥ 2) were included in this analysis.
Relative quantification of mutant frequency using HRM profile analysis. We used PCR-based high-resolution melting (HRM) profile analysis with integrated LunaProbes to quantify the ratio of mutant to wild type genotypes (99–101). The probes and primers used in this study are listed in Table S2. Primers were designed using Primer3. Probes were designed with the SNP located in the middle of the sequence, and the 3’ end was blocked by carbon spacer C-3. The primer asymmetry was set to 2:1 (excess primer, limiting primer) in all cases. Pre-PCR was performed in a 10-µl reaction system, with 0.25 µM excess primer, 0.125 µM limiting primer, 0.25 µM probe, 0.5 µM bacterial sample culture (with a 100-fold-diluted saved sample, the OD at 600 nm [OD600] is about 0.01), and 1× LightScanner master mix (BioFire Defense). Dimethyl sulfoxide (DMSO) at a final concentration of 5% was added to all reaction mixtures to ensure that the targeted melting domains were within the detection limit of the LightScanner (Idaho Technology Inc.). Finally, Milli-Q (MQ) water was used to supplement up to 10 µl of a 96-well black microtiter plate with white wells was used to minimize background fluorescence. Before amplification, 25 µl mineral oil was loaded in each well to prevent evaporation, and the plate was covered with a foil seal to prevent the degradation of fluorescent molecules. Amplification was initiated by a holding the plate at 95°C for 3 min, followed by 55 cycles of denaturation at 95°C for 30 s, annealing at 60°C for 30 s, and extension at 72°C for 30 s; the plate was then kept at 72°C for 10 min. After amplification, samples were heated in a thermal cycler (Bio-Rad) shortly to 95°C for 30 s to denature all double-stranded structures, followed by a rapid cooling to 25°C for 30 s to facilitate successful hybridization between probes and the target strands. The plate was then transferred to a LightScanner (Idaho Technology Inc.). Melting profiles of each well were collected by monitoring the continuous loss of fluorescence with a steady increase of the temperature from 35°C to 97°C, with a ramp rate of 0.1°C C/s. The relative quantification was based on the negative first derivative plots using MATLAB software. The areas of probe-target duplex melting peaks were auto-calculated with the AutoFit Peaks I Residuals function in PeakFit software (SeaSolve Software Inc.). The mutant frequency, X, was calculated using the equation $X = \frac{X_{\text{mutant}}}{X_{\text{WT}}}$, where WT is the wild type. To validate the HRM method, standard curves were generated by measuring mixed samples with the following known proportions of mutant templates: 0%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, and 100%. Measurements for each sample were done in triplicate. The linear-regression formula of each mutant between actual frequencies and measured frequencies is shown in Fig. S1. The high $R^2$ values and slope values of these equations that were nearly equal to 1 confirmed that the HRM method can accurately detect a mutant’s frequency in a mixed population.

Data availability. The P. protegens CHA0-GFP reference genome is deposited on GenBank under accession no. RCSR00000000.1. Raw sequencing data used in this study are deposited in the NCBI database under BioProject no. PRJNA473919. A conversion table for the CHA0-GFP–to–CHA0 gene annotations, including recent NCBI accession codes, is available at the following URL: https://doi.org/10.6084/m9.figshare.13295828.v2.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, JPG file, 0.1 MB.
FIG S2, JPG file, 0.1 MB.
FIG S3, JPG file, 0.4 MB.
FIG S4, JPG file, 0.1 MB.
FIG S5, JPG file, 0.1 MB.
TABLE S1, DOCX file, 0.02 MB.
TABLE S2, DOCX file, 0.01 MB.

ACKNOWLEDGMENTS

This work was supported by China Scholarship Council fellowships (to E.L. and H.Z.), a postdoctoral fellowship of the Research Foundation Flanders (FWO 12B8116RN) (to R.D.J.), and a European Research Council advanced grant (269072 to C.M.J.P.).

E.L., H.Z., A.J., C.M.J.P., P.A.H.M.B., and R.D.J. designed experiments; E.L., H.Z., and H.J. performed experiments; E.L., H.Z., and R.D.J. analyzed data. All authors discussed the results and contributed to the final manuscript.

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