RESEARCH ARTICLE

Re-evaluation of a Tn5::gacA mutant of Pseudomonas syringae pv. tomato DC3000 uncovers roles for uvrC and anmK in promoting virulence

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Abstract

Pseudomonas syringae is a taxon of plant pathogenic bacteria that can colonize and proliferate within the interior space of leaf tissue. This process requires P. syringae to rapidly upregulate the production of virulence factors including a type III secretion system (T3SS) that suppress host defenses. GacS/A is a two-component system that regulates virulence of many plant and animal pathogenic bacteria including P. syringae. We recently investigated the virulence defect of strain AC811, a Tn5::gacA mutant of P. syringae pv. tomato DC3000 that is less virulent on Arabidopsis. We discovered that decreased virulence of AC811 is not caused by loss of GacA function. Here, we report the molecular basis of the virulence defect of AC811. We show that AC811 possesses a nonsense mutation in anmK, a gene predicted to encode a 1,6-anhydromuramic acid kinase involved in cell wall recycling. Expression of a wild-type allele of anmK partially increased growth of AC811 in Arabidopsis leaves. In addition to the defective anmK allele, we also show that the Tn5 insertion in gacA exerts a polar effect on uvrC, a downstream gene encoding a regulator of DNA damage repair. Expression of the wild-type anmK allele together with increased expression of uvrC fully restored the virulence of AC811 during infection of Arabidopsis. These results demonstrate that defects in anmK and uvrC are together sufficient to account for the decreased virulence of AC811, and suggest caution is warranted in assigning phenotypes to GacA function based on insertional mutagenesis of the gacA-uvrC locus.

Introduction

Pseudomonas syringae is a plant pathogenic bacterium that infects a broad range of host species including numerous agronomically-important food crops and ornamental plants. Due to its genetic tractability, well-defined infection cycle, and ability to infect model plants such as Arabidopsis, P. syringae has been adopted as a model organism for studying the genetic bases of bacterial pathogenesis [1]. P. syringae can infect most aerial plant tissues, but is commonly
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studied for its ability to colonize and infect leaves. Upon introduction to a plant host, *P. syringae* is capable of persisting as an epiphyte on leaf surfaces [2]. Under favorable environmental conditions, *P. syringae* swims into the apoplast, or interstitial space, of leaf tissue through stomata or wounds in the leaf surface. Once in the leaf interior, *P. syringae* rapidly proliferates to a high density and causes visible chlorosis as well as water-soaked necrotic lesions on infected tissue.

In order to colonize leaf tissue and cause disease, *P. syringae* must overcome both preformed and induced plant host defenses. A primary line of defense encountered by *P. syringae* are pattern recognition receptors (PRRs) on the surface of plant cells [3]. PRRs monitor the extracellular space for the presence of conserved microbial features and, upon activation, rapidly initiate defense responses to limit pathogen growth. To counteract PRR-activated defenses, *P. syringae* deploys the type III secretion system (T3SS), a syringe-like translocon that delivers bacterial effector proteins directly into plant cells. Within the host cytosol, many of these effector proteins inhibit PRRs or their downstream signaling cascades, thereby suppressing host defenses and promoting *P. syringae* growth within the apoplast [3]. Mutant strains of *P. syringae* that lack a functional T3SS are unable to grow to high levels in host tissues and fail to cause disease.

Although the T3SS is an essential component of *P. syringae* virulence, other factors are also necessary for full infection. These T3SS-independent factors include small molecule toxins that disrupt host immunity [1], enzymes that detoxify plant defense compounds [4], and extracellular proteases [5], and catalases [6] that modify the apoplast environment to enable bacterial survival. Physiological attributes, such as flagella and pili that facilitate bacterial motility, have also been shown to play important roles in the progression of infection [7–8]. Furthermore, signaling pathways that coordinate the expression of these diverse virulence factors are also fundamental to the development of bacterial plant disease [9]. Recent analyses of the *P. syringae* transcriptome during host infection identified various other metabolic, transport, and abiotic stress pathways that are differentially expressed throughout various stages of plant infection [10–12] and that may potentially contribute to pathogenesis. Given the complex, multigene process of adapting to the host environment, it is likely that many additional unknown genetic factors are required for *P. syringae* to successfully grow within the apoplast and cause disease.

GacS/A is a highly conserved two component system that is hypothesized to function as a global regulator of diverse cellular processes including motility, exopolysaccharide biosynthesis, exoenzyme secretion, and T3SS deployment [13–15]. In *Pseudomonas syringae* pv. *tomato* DC3000, a model pathogen capable of infecting both Arabidopsis and tomato, GacS/A has been characterized as a positive regulator of T3SS expression and virulence. Predictive models were developed primarily on the basis of characterizations of a single mutant strain AC811 that contains a Tn5 transposon insertion within the gacA open reading frame [16]. In a recent study, we re-examined phenotypes of AC811 and discovered that, contrary to previous reports, this mutant strain has increased, rather than decreased, expression of T3SS genes during infection of Arabidopsis [17]. Furthermore, we demonstrated that decreased growth of AC811 within Arabidopsis leaves was not due to loss of GacA function. Together, these data indicated that AC811 possesses a T3SS- and GacA-independent defect that compromises its virulence.

In this work we investigated the genetic basis of decreased virulence of strain AC811. We demonstrate that a defective allele of *anmK*, as well as a polar effect of Tn5 insertion on *uvrC* expression, are sufficient to account for the decreased growth of AC811 in Arabidopsis leaves. Together, these data demonstrate that *anmK* and *uvrC* are required for maximum virulence of DC3000 in Arabidopsis. In addition to virulence phenotypes, we also show that *uvrC*, but not *gacA*, is required for siderophore production by DC3000. These findings highlight potential
challenges with assigning functions to GacA based solely on analysis of transposon insertional mutants, and underscore the importance of complementation of GacA-associated phenotypes.

**Results**

**Decreased relative growth of AC811 occurs in arabidopsis leaves but not in King’s B medium**

AC811 is unable to grow to high levels or cause disease to Arabidopsis [16–17]. To determine whether AC811 has a general growth defect, we cultured DC3000, AC811 and a DC3000 ΔgacA deletion strain in King’s B (KB) broth at 28˚C, a temperature frequently used for overnight culturing of *P. syringae*. At 28˚C, growth rates of AC811 and DC3000 ΔgacA were not significantly different from that of DC3000 (S1A Fig). We also tested growth of these strains in KB broth at 21˚C, the same temperature we use to maintain Arabidopsis plants during infection assays, and observed no growth defect for either AC811 or DC3000 ΔgacA (S1B Fig). These data indicate that AC811 does not have a general growth defect, and suggest that growth of this strain may be specifically hindered within the leaf apoplast environment.

**A nonsense mutation in anmK contributes to decreased virulence of AC811**

Decreased virulence of AC811 in the apoplast of Arabidopsis leaves is not due to loss of gacA [17]. We hypothesized that another mutation may be responsible for the decreased virulence of AC811. We used the Illumina HiSeq platform to sequence the genomes of strain AC811 and its DC3000 parental strain, enabling us to search for single nucleotide polymorphisms (SNPs) that were unique to AC811 [18]. We identified a total of 120 high confidence SNPs in AC811 relative to the published DC3000 reference genome (S1 Table). A total of 77 of these SNPs are unique to AC811, as determined based on their absence in the genome sequence of the parental strain. Of these, two SNPs were identified as the most likely to significantly impact protein function and cause the AC811 mutant phenotype. One of these unique SNPs results in a nonsense mutation in the first third of the coding region of PSPTO_0606, encoding a predicted 1,6-anhydromuramic acid kinase (AnmK). We sequenced the *anmK* locus of AC811 and its DC3000 parental strain, confirming that the nonsense mutation is present in AC811 but absent in the parental strain (Fig 1A). We also identified a SNP exclusive to AC811 that causes a frameshift in PSPTO_1067, a predicted glycosyltransferase. All other SNPs unique to AC811 were either predicted missense mutations, synonymous mutations, or mutations within intergenic regions, and were not predicted to have severe effects on protein function (S1 Table).

To assess whether the mutations in *anmK* and PSPTO_1067 contribute to loss of AC811 virulence, we individually cloned the DC3000 alleles of *anmK* and PSPTO_1067 into pME6010, introduced the resulting constructs into DC3000 and AC811, and infected Arabidopsis leaves with these strains by syringe-infiltration. AC811 carrying *anmK*-pME6010 grew to significantly higher levels relative to AC811, indicating that introduction of the DC3000 allele of *anmK* partially complemented the ability of AC811 to grow in planta (Fig 1B). However, growth of AC811 *anmK*-pME6010 was not fully restored to DC3000 levels, suggesting that other factors may contribute to AC811 *in planta* growth (Fig 1B). Expression of *anmK* did not significantly impact the growth of wild type DC3000 (Fig 1B). DC3000 is a highly virulent pathogen that can achieve high levels of growth, which could mask any potential growth-promoting effects of *anmK*. To test this, we introduced *anmK*-pME6010 into a virulence-attenuated DC3000 ΔavrPtoΔavrPtoB strain [19]. Similar to DC3000, we detected no *anmK*-dependent increase in the growth of this strain in Arabidopsis leaves (S2 Fig). No increase in growth was measured for AC811 carrying PSPTO_1067::pME6010, suggesting that the
frameshift mutation of this gene does not negatively impact AC811 virulence (S3 Fig).

Together, these data indicate that a nonsense mutation in anmK contributes to the decreased virulence of AC811.

Tn5 insertion in gacA disrupts the expression of downstream uvrC

We reasoned that the insertion of Tn5 into the gacA open reading frame in AC811 may have a polar effect on nearby genes. In DC3000, gacA is predicted to be part of an operon including uvrC and pgsA [13] that is conserved among γ-proteobacteria [14–15, 20–22] (Fig 2A). The uvrC gene encodes a component of the dinucleotide excision repair (NER) complex involved in remediating DNA damage [23], whereas pgsA is predicted to encode a protein involved in phospholipid biosynthesis [13]. In E. coli, a similarly structured operon can be transcribed...
either as a polycistronic transcript, or as separate monocistronic transcripts due to promoter elements located within the gacA open reading frame [24].

We used quantitative RT-PCR (qRT-PCR) and gene-specific primers to measure the abundance of gacA-, uvrC- and pgsA-containing transcripts in AC811. As expected, we could not detect gacA-containing transcripts from AC811 or the DC3000 ΔgacA-1 deletion strain (Fig 2B). Transcripts containing uvrC were also significantly decreased in AC811 bacteria relative to levels in DC3000 (Fig 2B). In contrast, the abundance of pgsA-containing transcripts was not significantly altered in AC811 (Fig 2B). We also measured uvrC and pgsA expression in DC3000 ΔgacA-1 and found that the abundance of uvrC transcripts was similar to DC3000 levels, whereas the average abundance of pgsA transcripts was reduced approximately 2-fold, although this reduction was not significantly different from pgsA levels in either DC3000 or AC811 (Fig 2B). For all genes assessed, virtually no transcripts were detected from negative control reactions in which cDNA was mock synthesized in the absence of reverse transcriptase (S4 Fig). Based on these data, we conclude that the Tn5 insertion in gacA decreases the expression of uvrC but not pgsA. Furthermore, these data indicate that deletion of the entire gacA open reading frame does not exert a significant polar effect on the expression of downstream genes.

We additionally assessed whether polycistronic transcripts containing both gacA and uvrC are altered in AC811. To investigate, we performed qRT-PCR with primers that span the gacA-uvrC junction and measured a significant decrease in these transcripts in AC811 (S5B Fig). We also detected transcripts containing the uvrC-pgsA junction by qRT-PCR, but did not observe
any significant change in the abundance of these transcripts in either AC811 or DC3000 ΔgacA-1 (S5B Fig).

Expression of uvrC and anmK fully rescues the growth defect of AC811 in arabidopsis

To determine whether decreased uvrC expression contributes to the virulence defect of AC811, we cloned uvrC into broad host range vector pME6010 and introduced this construct into both DC3000 and AC811. We then syringe-infiltrated these strains into Arabidopsis leaves and observed that AC811 carrying uvrC::pME6010 grew to significantly higher levels than AC811 carrying empty pME6010 (Fig 3A). Similar to our results with anmK expression, uvrC::pME6010 did not alter growth of DC3000 WT (Fig 3A) or DC3000 ΔavrPtoΔavrPtoB (S2 Fig) in Arabidopsis. To alleviate the polar effect of Tn5 on uvrC, we used allelic exchange to delete both gacA and Tn5 from the AC811 genome, generating an AC811 ΔgacA strain. Expression of uvrC in AC811 ΔgacA was restored to DC3000 levels as determined by qRT-PCR (S5C Fig). Growth of AC811 ΔgacA was significantly higher relative to AC811 (Fig 3B) but was not fully restored to DC3000 or DC3000 ΔgacA levels.

We next introduced a plasmid with the DC3000 allele of anmK into AC811 ΔgacA. The introduction of anmKDC3000 was sufficient to fully restore the growth of AC811 ΔgacA to DC3000 levels in Arabidopsis leaves (Fig 4). Disease symptoms caused by AC811 ΔgacA carrying anmKDC3000, such as chlorosis of infected leaves, also appeared similar to those produced by DC3000 infection (S6 Fig). Based on these data we conclude that decreased uvrC expression and loss of AnmK function are together causal for decreased virulence of AC811 in Arabidopsis leaves.

Decreased uvrC expression reduces siderophore production in DC3000

The biosynthesis and extracellular secretion of iron-chelating siderophores is positively regulated by GacS/A in certain pseudomonads [25], but suppressed by GacS/A in others [26–27]. To investigate if GacA regulates siderophore production in DC3000, we incubated DC3000, AC811 and DC3000 ΔgacA on CAS blue agar to detect siderophore production. We observed a significant reduction in siderophore production by AC811 (Fig 5A and 5B). However, no decrease in siderophore levels was observed for DC3000 ΔgacA, indicating the decrease in siderophore production by AC811 is gacA-independent (Fig 5A and 5B). Expression of uvrC in AC811 significantly increased siderophore production by AC811 to near DC3000 levels (Fig 5C). Furthermore, siderophore production by AC811 ΔgacA was indistinguishable from DC3000 (Fig 5B). Together, these data indicate that, similar to DC3000 growth in the leaf apoplastic, polar effects of Tn5 on uvrC expression influence the levels of siderophore production.

Discussion

Isolating and characterizing mutants that have loss-of-virulence phenotypes is fundamental to understanding virulence mechanisms of P. syringae and other bacterial pathogens. However, laboratory manipulation of bacterial genomes can present various challenges. Common techniques used to generate mutants, such as Tn5 insertional mutagenesis, carry the risk of disrupting expression of surrounding genes. Furthermore, accumulation of SNPs or other mutations within the genomes of laboratory strains may further confound the analysis of mutant phenotypes. Here, we investigated the genetic defects of AC811 and demonstrated that a polar effect of the Tn5 transposon, as well as a second-site mutation are causal for decreased virulence of AC811. In this case, these unintended genetic defects led to the discovery of two genes that positively regulate DC3000 virulence. However, these gacA-independent effects also
Fig 3. Growth of AC811 in Arabidopsis is enhanced by eliminating polar effects of Tn5:gacA insertion on uvrC expression. (A) Arabidopsis leaves were syringe-infiltrated with DC3000 or AC811 carrying either pME6010 empty vector or pME6010::uvrC as indicated. Graphed are means ± SE of leaf bacteria measured by serial dilution plating; n = 9. Data are pooled from three independent experiments. (B) Arabidopsis leaves were syringe infiltrated with DC3000 WT, AC811, ΔgacA-1 or AC811 ΔgacA. Graphed are means ± SE of leaf bacteria measured by serial dilution plating; n = 14. Abbreviation dpi is days post-infection. Data are pooled from four independent experiments. Small-case letters in (A) and (B) denote statistical groups determined by ANOVA with multiple pairwise t-test comparisons and Tukey’s post-hoc HSD analysis, p < 0.05.

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Fig 4. Growth of AC811 in Arabidopsis is fully restored by eliminating Tn5 effects on uvrC and by expression of a wild-type anmK allele. Arabidopsis leaves were syringe-infiltrated with DC3000, AC811 or AC811 ΔgacA carrying either pME6010 empty vector or pME6010::anmK. Graphed are means ± SE of leaf bacteria enumerated by serial dilution plating; n = 3. Abbreviation dpi is days post-infection. Data are pooled from four independent experiments; n = 12. Statistical groups were determined by ANOVA with multiple pairwise t-test comparisons and Tukey’s post-hoc HSD analysis, p < 0.05.

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contributed to confusion regarding the role of gacA in DC3000 virulence, and highlight the need for caution when interpreting phenotypes of mutant strains, as well as the need to confirm that the casual lesion is indeed within the gene of interest. In this work, whole genome sequencing of AC811 proved to be an effective approach for identifying an off-target genetic defect. With the ever-decreasing cost of high-throughput DNA sequencing, we anticipate that whole genome sequencing of mutant strains may become a routine and necessary practice in the laboratory for confirming the results of bacterial genetic studies.

We demonstrated that a lesion in anmK (PSPTO_0606) causes a reduction in in planta growth of AC811. The anmK gene is predicted to encode for anhydro-1,6-muramic acid kinase, an enzyme that catalyzes the phosphorylation of muropeptides generated during the natural recycling of bacterial cell wall peptidoglycan [28]. AnmK has not been previously identified as necessary for virulence of P. syringae, and the specific role of this enzyme during host infection remains unknown. In E. coli, loss of AnmK function is known to disable
metabolic reuptake of peptidoglycan. This disruption of cell wall regeneration leads to the phosphorylation-dependent retention of muropeptides in the bacterial cytosol, preventing the stress, such as high cellular osmolarity induced by water stress or cell envelope damage inflicted by antibiotics [29]. AnmK-mediated cell wall recycling could be required for the rapid proliferation of DC3000 following colonization of the host apoplast. Alternatively, abnormal accumulation of muropeptides in the extracellular space may disrupt DC3000 cellular stress signaling, possibly affecting the ability of DC3000 to tolerate stressors of the plant environment such as immune defenses and/or water stress.

Our results suggest that UvrC may play a role in regulating DC3000 virulence. As a subunit of the conserved dinucleotide excision repair (NER) complex, UvrC catalyzes the excision of DNA lesions, contributing to bacterial survival under UV- or mutagen-induced stress [30]. Accordingly, gacA mutants in various bacterial species show UV-hypersensitivity, likely due to negative polar effects on uvrC expression [15, 20, 22, 31–33]. While UV stress tolerance is known to impact epiphytic survival in Pseudomonas spp. [34], it is unknown how UvrC may contribute to growth of bacteria such as DC3000 that are primarily endophytic and are unable to persist epiphytically on the leaf surface. Similar to our findings, a study of both polar and nonpolar gacA mutants in the animal pathogen Pseudomonas aeruginosa revealed that negative effects on uvrC expression, rather than inactivation of gacA, were associated with a loss of virulence in Bombyx mori larvae [33, 35]. Furthermore, an uvrC mutant of the human pathogen Yersinia enterocolitica has been shown to cause reduced expression of inv, a gene required for colonization of the intestinal lymphoid tissues, although possible polar effects on gacA expression in this mutant could not be ruled out [36]. Interestingly, normal inv expression by the Y. enterocolitica uvrC mutant could be restored by expression of either SspA or ClpB, both of which function as regulators of stress tolerance in E. coli [32–35]. These results suggest that UvrC may broadly contribute to virulence in a range of pathogens, potentially through an unknown role in the cellular stress response. In DC3000, UvrC may regulate tolerance of environmental stressors encountered in the leaf apoplast enabling bacterial survival and proliferation in host tissues.

Throughout the γ-proteobacteria, gacA is located in an operon with uvrC and psgA. In numerous species, the translation start site of uvrC overlaps the stop codon of gacA [21–22, 31, 37]. Previously, gacA was reported to be expressed as a single gene in DC3000 based on detection of a ~0.75 kb transcript by northern blot with a gacA sequence-specific probe [16]. Our qRT-PCR data demonstrate that transcripts containing gacA-uvrC and uvrC-psgA junctions are also produced in DC3000, indicating that gacA is expressed as part of a larger operon structure than previously reported. Consistent with this operon model of gacA and uvrC expression, we observed a significant reduction in uvrC and gacA-uvrC transcripts in AC811 due to polar effects of the Tn5 insertion in gacA. Although we also detected bicistronic uvrC-psgA transcripts, both psgA- and uvrC-psgA-containing transcripts were not significantly altered in abundance in AC811 (Fig 4 and S4 Fig). In E. coli, monocistronic uvrC transcripts and bicistronic uvrC-psgA transcripts are produced by promoter sequences within the gacA coding region [24] and these promoter elements are conserved within DC3000 gacA based on our sequence analysis. Therefore, uvrC-psgA transcription in AC811 may occur through promoter elements in gacA that are downstream of Tn5 insertion, allowing for expression of psgA to remain unperturbed. Notably, expression of uvrC was restored to wild type levels in our AC811 ΔgacA strain, indicating that promoter elements upstream of gacA are sufficient to drive the expression of uvrC even in the absence of gacA. Despite the conserved genomic arrangement of gacA and uvrC, the functional relationship between these genes (if any)
remains unknown. Further analysis of the mechanistic role of UvrC during \textit{P. syringae} infection may provide important insights into the functional significance of the \textit{gacA-uvrC} association.

Our results also uncovered a role for \textit{uvrC} in siderophore production by DC3000. Siderophores are a class of fluorescent iron-chelating compounds that are secreted by bacteria into the extracellular space during iron-limiting growth conditions. Upregulation of \textit{uvrC} expression under iron-limiting conditions has been reported in certain bacterial species [38], although this effect is not observed in all cases [39]. Both \textit{uvrC} expression and siderophore production are induced in biofilm-forming cells of \textit{Mycobacterium smegmatis}, though it is unknown whether \textit{uvrC} influences siderophore production or is simply concomitantly expressed [40]. Siderophores contribute to the virulence of many bacterial pathogens. However, a DC3000 mutant that is unable to synthesize or import siderophores remains fully virulent on host tomato plants [41]. Therefore, although both decreased virulence and siderophore production are observed with AC811, it is unlikely that these two phenotypes are causally-related. In addition to identifying UvrC as a regulator of siderophore production, our results also demonstrate that GacA is dispensable for this phenotype. Regulation of siderophore biosynthesis by GacS/A has been described in various Pseudomonads, though whether GacS/A functions as a positive or negative regulator of siderophore production appears to vary across species [25–27]. As many previous studies have relied on insertional mutants to assess the function of GacA, in some cases it may be necessary to re-assess these mutant strains to determine if off-target effects on \textit{uvrC} expression may be responsible for \textit{gacA}-associated phenotypes.

\textbf{Materials and methods}

\textbf{Bacterial strains and growth conditions}

\textit{P. syringae} strains stored at -80°C were streaked from 25% glycerol stocks onto King’s B medium (KBM) 1.5% agar plates. The KBM agar was supplemented with 50 μg/mL rifampicin and antibiotics (10 μg/mL tetracycline, 20 μg/mL gentamycin, 50 μg/mL chloramphenicol) as necessary. For all experiments \textit{P. syringae} were grown at room temperature on KBM agar for 2 days prior to use. \textit{E. coli} strains were grown at 37°C in Luria Broth (LB) supplemented with antibiotics (5 μg/mL tetracycline, 25 μg/mL gentamycin, 30 μg/mL chloramphenicol) as necessary.

\textbf{Construction of \textit{anmK} and \textit{uvrC} complementation plasmids}

The Gibson assembly method was used to clone the open reading frames of \textit{uvrC} and \textit{anmK} into broad host range vector pME6010 under control of a constitutive kanamycin promoter [42]. Primers 6010-F1/6010-R1 and 6010-F2/6010-R2 (S2 Table) were used to PCR amplify pME6010 as 3.0-kb and 5.5-kb DNA fragments. Primer sets 6010-\textit{uvrC}-F/6010-\textit{uvrC}-R and 6010-\textit{anmK}-F/6010-\textit{anmK}-R (S2 Table) were used to amplify \textit{uvrC} and \textit{anmK}, respectively, from DC3000 genomic DNA. PCR products of \textit{uvrC} or \textit{anmK} were then mixed with the 3.0- and 5.5-kb fragments of pME6010 and assembled using the NEBuilder HiFi mix (NEB). Plasmids were transformed into \textit{E. coli} DH5a competent cells (NEB) by heat shock. Tc\textsuperscript{R} clones were screened for the presence of the \textit{uvrC} or \textit{anmK} insert by colony PCR with primer sets 6010-\textit{uvrC}-F/6010-\textit{uvrC}-R or 6010-\textit{anmK}-F/6010-\textit{anmK}-R, respectively. Constructs were confirmed by Sanger sequencing. Plasmids were conjugated into DC3000 by tri-parental mating using an \textit{E. coli} helper strain carrying pRK600 [43].
Generation of gacA deletions in DC3000

The previously described ΔgacA::pK18mobsacB suicide vector [17] was conjugated into DC3000 by tri-parental mating using an E. coli helper strain carrying pRK600 [43]. Merodiploid colonies were passaged 1–3 days in KB broth with 50 μg/mL rifampicin prior to counterselection on KB agar with 15% sucrose. Sucrose-resistant colonies were plated on KB agar with 50 μg/mL rifampicin supplemented with or without selective antibiotics. PCR reactions with primers gacA-F and gacA-R (S2 Table) were used to screen genomic DNA isolated from kanamycin-sensitive colonies for deletion of gacA. To generate a ΔgacA deletion in AC811 (KanR), the ΔgacA deletion cassette from pK18mobsacB was sub-cloned into pT18mobsacB (Addgene #72648, TcR) using the Gibson assembly method. Primers pT18gacA-F and pT18gacA-R (S2 Table) were used to PCR amplify the ΔgacA deletion cassette from pK18mobsacB. Primers pT18-F and pT18-R (S2 Table) were used to PCR amplify the entire backbone of pT18mobsacB. PCR products were assembled into an intact plasmid using NEBuilder HiFi mix (NEB). Plasmids were transformed into E. coli DH5a competent cells (NEB) by heat shock. Clones were screened for the presence of the ΔgacA deletion cassette by colony PCR with primers M13F and M13R, and constructs were confirmed by Sanger sequencing.

Measurement of bacterial growth in culture

P. syringae were scraped from agar plates and inoculated into 3 mL cultures of KB broth containing 50 μg/mL rifampicin. Cultures were grown at 28˚C overnight in a shaking incubator until cultures reached an optical density at 600 nm (OD600) of ~1.0. Overnight cultures were then diluted to OD600 = 0.05 in 30 mL of KB broth containing 50 μg/mL rifampicin in 250-mL volume flasks. Cultures were maintained at 28˚C or 21˚C with shaking. One mL samples were removed from each flask every 2 hrs for 12 hours and a spectrometer was used to measure the OD600.

In planta bacterial growth assays

Arabidopsis thaliana Col-0 seeds were sterilized, stratified and sown onto MS agar plates as described previously [9]. After two weeks of growth, seedling were transplanted to Sunshine mix soil (Sun Gro Horticulture) in 24-well flats. Plants were maintained in a 22˚C, 10-hour day growth chamber before and during the infection. To prepare the inoculum, P. syringae were scraped from the surface of KB agar and vortexed in 1 mL of sterile H2O. Cells were then washed twice and pelleted between each wash by centrifugation at 21,000 x g for 1 min, followed by resuspension of bacterial pellets in 1 mL H2O. Bacterial suspensions were diluted with H2O to an OD600 of 0.001 (~1 x 10⁶ colony-forming units/mL). A needle-less syringe was used to infiltrate bacteria into leaves of five week-old plants. Three to five leaves were infected per plant, and individual plants were infected with only a single strain of bacteria. At times indicated, a cork borer was used to remove 0.2 cm² discs from infected leaves. For each sample, three leaf disks from a single plant were pooled into a single microcentrifuge tube and homogenized in 500 μL H2O by grinding the leaf tissue with a plastic pestle. The resulting homogenate was serially-diluted and pipetted in 10 μL volumes onto the surface of KB agar containing 50 μg/mL rifampicin. After 24 hours bacteria, agar plates were viewed under a stereomicroscope to visualize and enumerate individual colonies.

Genome sequencing and SNP calling

The Qiagen DNeasy Blood & Tissue Kit Genomic DNA was used to extract genomic DNA from DC3000 and AC811. The Nextera XT library prep kit was used to prepare sequencing
libraries from each genomic DNA sample. The resulting libraries were sequenced (150 bp paired end) on a single lane of an Illumina HiSeq 3000 instrument at the Center for Genome Research and Biocomputing at Oregon State University. Illumina sequencing reads are deposited in the National Center for Biotechnology Information (NCBI) short read archive (SRA) under BioProject # PRJNA574776. Reads were aligned to the *Pseudomonas syringae* pv. tomato DC3000 reference genome (NCBI: 223283), using bowtie2 version 2.3.2 with the option “—local” [44]. Alignments were converted to bam format using samtools version 0.1.19 and read groups were added using Picard tools version 2.0.1 [45] (Picard Tools, 2015). GATK version 3.7 HaplotypeCaller with the options ‘-ERC GVCF -ploidy 1’ was used to call variants, and the data were then combined using GenotypeGVCFs [46]. The program snpEff version 4.3T was used to predict effects of SNPs on protein function [47].

**Quantification of bacterial gene expression by qRT-PCR**

Bacteria were grown overnight at 20˚C in KB broth containing 50 μg/mL rifampicin to an OD$_{600}$ = ~1.0. Cells were centrifuged for 10 min at 21,000 g and the resulting bacterial pellets were frozen in liquid nitrogen and stored at -80˚C prior to use. RNA was extracted from the bacterial pellets and 500 ng converted into cDNA in 25 μL reactions as described previously [9]. Quantitative RT-PCR (qRT-PCR) reactions were performed in 10 μL volumes using 5 μL of SsoAdvanced Universal SYBR Green Supermix (Bio-Rad), 4 μL of a mix containing 0.5 μM of each transcript-specific primer (S2 Table), and 1 μL of cDNA. SYBR Green fluorescence from each qRT-PCR reaction was monitored using the C1000 Thermal Cycler with CFX96 Real-Time System (Bio-Rad). Relative abundance of transcripts was calculated relative to *gyrA* using the formula Transcript Abundance = PCR efficiency$^{(Ct_{gene}-Ct_{gyrA})}$ [48]. PCR efficiency values were calculated for each qRT-PCR reaction using LinRegPCR [49].

**Detection of siderophore production**

Chrome azurol (CAS) blue agar plates [50] containing 2% (w/v) sucrose, 0.2% asparagine, 0.1% KH$_2$PO$_4$, 0.05% MgSO$_4$·7H$_2$O, 0.1 M PIPES salt, and 1.5% agar were prepared. *P. syringae* were scraped from the surface of KB agar and suspended in 1 mL of sterile H$_2$O by vortexing. Cells were then washed three times with 1 mL of sterile H$_2$O. Bacterial suspensions were diluted with H$_2$O to an OD$_{600}$ of 1.0 and 5 μL spotted at the center of each plate. Plates were incubated in the dark at room temperature for 48 hr prior to imaging. Radii of the diffuse orange halos that appear in the agar were measured starting from the edge of the bacterial colony.

**Supporting information**

**S1 Fig. gacA mutants exhibit wild type levels of growth in culture.** Time-course of bacterial growth at 28˚C. DC3000, AC811 and ΔgacA-1 were inoculated at OD$_{600}$ = 0.05 into KB broth and incubated at (A) 28˚C or (B) 21˚C with shaking. Graphed are means ± SE of OD$_{600}$, n = 3. Data are pooled from three independent experiments. (TIF)

**S2 Fig. Expression of uvrC or annK does not alter growth of DC3000 ΔavrPtoΔavrPtoB.** Arabidopsis leaves were syringe-infiltrated with DC3000 ΔavrPtoΔavrPtoB carrying empty pME6010 (EV), pME6010::uvrC, or pME6010::annK as indicated. Abbreviation dpi is days post-infection. Graphed are means ± SE from data pooled from two independent experiments; n = 6. (TIF)
S3 Fig. Expression of PSPTO_1067 does not increase growth of AC811. Arabidopsis leaves were syringe infiltrated with DC3000 and AC811 carrying either pME6010 empty vector or pME6010::PSPTO_1067 plasmid. Graphed are means ± SE of bacterial growth as determined by serial dilution plating, n = 3. Small-case letters denote statistical groups determined by ANOVA with multiple pairwise t-test comparisons and Tukey’s post-hoc HSD analysis, p < 0.05. Data are representative of two independent experiments.

(TIF)

S4 Fig. Reverse transcriptase-dependent quantification of gacA operon-associated transcripts. (A) RNA was extracted from DC3000, AC811, and ΔgacA-1, and cDNA synthesis performed with (+RT) or without reverse transcriptase (-RT). Shown is quantitative RT-PCR analysis of gacA, uvrC, and pgsA transcripts in these samples. Graphed are means ± SE of uvrC and pgsA transcript abundance normalized to gyrA from corresponding +RT samples and calculated relative to transcript levels measured in DC3000 (+RT). Data are pooled from two independent experiments with two technical replicates each; n = 4. Data for +RT samples are the same as shown in Fig 2B. (B) Transcript abundance of gacA, gacA:uvrC, and uvrC:pgsA junctions from DC3000 cDNA synthesized with (+RT) or without (-RT) reverse transcriptase. Data are pooled from two independent experiments with two technical replicates each; n = 4.

(TIF)

S5 Fig. Polar effects of Tn5 on uvrC and bicistronic gacA-uvrC and uvrC-pgsA transcript abundance. (A) Schematic of the predicted gacA-uvrC-pgsA operon in DC3000, with vertical bars indicating predicted translation start sites of uvrC and pgsA. Shading and numbering indicates regions targeted by qRT-PCR, with amplicons designated as follows: (1) gacA:uvrC; (2) uvrC; (3) uvrC:pgsA. (B) Abundance of uvrC transcripts was assessed by qRT-PCR using a gyrA reference gene as previously described. Values are normalized to DC3000. Graphed are means ± SE from data pooled across two independent experiments with two technical replicates each; n = 4. ns = no significant difference based on t-test (p > 0.05). (C) qRT-PCR analysis of transcripts containing gacA-uvrC or uvrC-pgsA junctions. Graphed are means ± SE from data pooled across two independent experiments with two technical replicates each; n = 4. **p < 0.01; ns = no significant difference (p > 0.05).

(TIF)

S6 Fig. DC3000 and an AC811 ΔgacA anmK+ strain cause similar disease symptoms during infection of Arabidopsis. Arabidopsis leaves were syringe-infiltrated with DC3000, AC811, or AC811 ΔgacA carrying pME6010::anmK. Infected plants were photographed at 3 days post-infection (dpi). Circular punches on infected leaves are from sampling for bacterial cfu enumeration. Images are representative of symptoms observed in three independent experiments.

(TIF)

S1 Table. Single nucleotide polymorphisms identified by Illumina sequencing.

(XLSX)

S2 Table. Sequences of oligonucleotide primers used in this study.

(XLSX)

S3 Table. List of strains used in this study.

(XLSX)
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