A cooperative virulence plasmid imposes a high fitness cost under conditions that induce pathogenesis

Thomas G. Platt*, James D. Bever and Clay Fuqua

Department of Biology, Indiana University, 1001 East Third Street, Jordan Hall 142, Bloomington, IN 47405, USA

Harbouring a plasmid often imposes a fitness cost on the bacterial host. Motivated by implications for public health, the majority of studies on plasmid cost are focused on elements that impart antibiotic resistance. Plasmids, however, can provide a wide range of ecologically important phenotypes to their bacterial hosts—such as virulence, specialized catabolism and metal resistance. The Agrobacterium tumefaciens tumour-inducing (Ti) plasmid confers both the ability to infect dicotyledonous plants and to catabolize the metabolites that plants produce as a result of being infected. We demonstrate that this virulence and catabolic plasmid imposes a measurable fitness cost on host cells under resource-limiting, but not resource replete, environmental conditions. Additionally, we show that the expression of Ti-plasmid-borne pathogenesis genes necessary to initiate cooperative pathogenesis is extremely costly to the host cell. The benefits of agrobacterial pathogenesis stem from the catabolism of public goods produced by infected host plants. Thus, the virulence-plasmid-dependent costs we demonstrate constitute costs of cooperation typically associated with the ability to garner the benefits of cooperation. Interestingly, genotypes that harbour derived opine catabolic plasmids minimize this trade-off, and are thus able to free-load upon the pathogenesis initiated by other individuals.

Keywords: cooperation; public good; Ti plasmid; trade-off; Agrobacterium tumefaciens; greenbeard

1. INTRODUCTION

Many ecologically important functions—including virulence, antibiotic resistance, bacteriocin production and specialized catabolism—are conferred by genes commonly found on bacterial plasmids. These self-replicating DNA elements tend to carry non-essential genes and can be transmitted vertically, and often horizontally, between host cells [1]. In many cases, the benefits that plasmids confer are only realized under certain environmental conditions [2]. In contrast, plasmids may be costly to their host cell regardless of the environment, because at a minimum their maintenance requires their replication and partitioning during each bacterial generation [3]. Indeed, numerous studies have found that harbouring a plasmid reduces the fitness of host cells [4–6]. The carriage cost of plasmids is centrally important to their population dynamics [2,7]. However, probably motivated by public health implications, most studies have measured carriage costs associated with plasmids that confer antibiotic resistance to host cells. Indeed, relatively few studies have examined the cost of harbouring plasmids conferring other ecologically important functions (see [8,9] for exceptions). Natural plasmids are often relatively large, have low copy number and maintain long-standing associations with bacterial lineages [1]. Each of these factors is likely to lead to the evolution of limited costs associated with these plasmids. Despite this, as well as the fact that these plasmids frequently confer important phenotypes such as the ability to infect plant or animal hosts, few studies have examined how the costs associated with these plasmids shape their population dynamics.

The tumour-inducing (Ti) plasmid of Agrobacterium tumefaciens confers the majority of virulence functions underlying crown gall disease of dicotyledonous plants [10]. Plant-associated cues stimulate agrobacterial cells harbouring this large (approx. 200 kb) plasmid to infect hosts using a Ti plasmid encoded type IV secretion (T4S) system that delivers copies of a subset of Ti plasmid genes (carried on the transferred DNA or T-DNA) into the plant genome [10]. The infection process is predominantly mediated by a large set of more than 30 different Ti plasmid virulence (vir) genes, including those that form the T4S system and others that drive T-DNA replication and processing. Following transformation of the plant host, T-DNA genes are recognized by the plant’s expression machinery and expressed strongly within the plant nucleus. Several of these genes increase pools of the plant hormones, auxin and cytokinin, resulting in a rapid proliferation of the infected cells and thereby tumour development [10]. Other T-DNA genes direct the synthesis and release of metabolites called opines—a suite of specialized nutritional resources that cells use via catabolic genes that, importantly, are also encoded on the Ti plasmid [10]. Most opines are unusual ligation products of amino acids with small organic acids or sugar phosphates that can supply the infecting bacteria with carbon, nitrogen and phosphorus.

In this study, we examine the costs associated with the A. tumefaciens Ti plasmid. We find that these costs...
probably vary over space and time as a consequence of
dependence on local environmental conditions. We
observed a significant cost associated with carriage of the
Ti plasmid when either carbon or nitrogen availability lim-
ited population carrying capacity, but not when these
resources were abundant. We also demonstrate that the
expression of Ti-plasmid-borne pathogenesis genes
imposes a dramatic fitness cost on host cells. While it is
not unusual for plasmids to have costs, it is striking that
in this system the costs associated with the Ti plasmid
are costs of cooperation. Agrobacterial pathogenesis is
cooperative with the actions of few individuals infecting
the host plant, leading to the availability of a public good
resource [11,12]. Cells that harbour a Ti plasmid suffer a
modest cost of carrying the Ti plasmid, as well as a high
cost of infecting plant host cells, but are able to garner
the benefits stemming from the catabolism of the public
good nutrients produced by infected plant cells. In
nature, some agrobacterial genotypes have an opine cata-
bolism system that enables the bacteria to catabolize
niche, some agrobacterial genotypes have an opine cata-
bolism system that enables the bacteria to catabolize

2. MATERIAL AND METHODS
(a) Strains, plasmids and growth conditions
All strains and plasmids used in this study are described
in the electronic supplementary material. We purchased
reagents, antibiotics and media components from Fisher
Scientific (Pittsburgh, PA), Sigma-Aldrich (St Louis, MO)
and New England Biolabs (Ipswich, MA). We obtained
oligonucleotide primers from Integrated DNA Technolo-
gies (Coralville, IA) and used QIAquick Spin kits (QIAGEN,
Valencia, CA) for nucleic acid purification. DNA
manipulations were performed using standard protocols
[15]. Plasmids were transferred into Agrobacterium strains via
either conjugation or electroporation using standard approa-
ches [16,17]. Unless specified otherwise, A. tumefaciens
strains were grown in AT minimal media supplemented
with 27.5 mM glucose and 15 mM (NH₄)₂SO₄ (ATGN [18]) and
incubated at 28 °C in a rotary aerator or on 1.5 per cent
agar plates. Unlike the original recipe, we omitted
FeSO₄•H₂O from our AT minimal media [19]. Escherichia
coli strains were grown in LB media at 37 °C on a shaker
platform or 1.5 per cent agar plates. Agrobacterium tumefaciens
strains TGP110 and TGP114 are spontaneous nalidixic acid
resistant mutants of 15955 and TGP101, respectively,
which contains the cured vector. We confirmed that we successfully cured
A. tumefaciens strain 15955 of pTi15955 using a modified Eckhardt method
[22] to evaluate the large plasmid content of our cured
strain relative to that of its parent strain, as well as R10,
KYC55 and SA122 as controls.

(b) Curing the tumour-inducing plasmid
In order to isolate the impact of the Ti plasmid on the growth
and competitive ability of bacterial cells, we generated a
plasmid-free derivative of A. tumefaciens strain 15955 that
differs from the parent strain only in that it lacks
pTi15955. We cured the Ti plasmid from 15955, using the
approach described by Uraji et al. [20]. We introduced the
curing vector pTP101 into the 15955 genetic background
via conjugation with an E. coli S17-1 Apir donor. Transconju-
gants of A. tumefaciens were selected for by their ability to
grow on ATGN supplemented with kanamycin and ampicil-
lin. The curing vector was constructed by cloning the entire
vegetative replication region of pTi15955 (repABC) into
pCF117, which contains a counterselectable sacRB gene
 conferring sucrose sensitivity. The two plasmids, pTP101
and pTi15955, use the same replication and partitioning
machinery, and are therefore incompatible [20,21]. As a con-
sequence, transconjugants that contain both plasmids give
rise to cells that lack one or the other plasmid. Populations
of these cells were subsequently screened for a lack of octo-
pine catabolism, consistent with a loss of pTi15955, by
patching colonies onto AT minimal media supplemented
with 3.25 mM octopine (Sigma; discontinued) as the sole
source of carbon and nitrogen. We then counterselected
against the sacRB gene on pTP101 by plating populations
of derivatives incapable of octopine catabolism onto AT
minimal media supplemented with 0.5 per cent sucrose
and 15 mM (NH₄)₂SO₄ to obtain a markerless derivative
of 15955 that has been cured of the Ti plasmid and also
the curing vector. We confirmed that we successfully cured
15955 of pTi15955 using a modified Eckhardt method
[22] to evaluate the large plasmid content of our cured
strain relative to that of its parent strain, as well as R10,
KYC55 and SA122 as controls.

(c) Characterizing resource-limited growth
We evaluated the population yield of A. tumefaciens strain
15955 in response to increasing amounts of glucose as the
sole carbon source or ammonia as the sole nitrogen source.
To do this, we prepared A. tumefaciens 15955 inoculum
free of nitrogen and carbon sources by growing this strain
to mid-log phase, harvesting cells via centrifugation and
washing them seven times with 78.6 mM KH₂PO₄ buffer
(pH 7.0). The washed cells were then diluted to an optical
density at 600 nm (OD₆₀₀) of 0.005 in the appropriate
media. Glucose was the sole source of carbon in AT minimal
media supplemented with 15 mM (NH₄)₂SO₄ and 0, 1, 3, 6,
12, 17 or 27.5 mM glucose. Analogously, ammonium was
the sole source of nitrogen in AT minimal media supplemented
with 27.5 mM glucose and 0, 0.04, 0.08, 0.16,
0.3, 0.6, 1.2, 2.4, 5, 10 or 20 mM ammonia. We monitored the
OD₆₀₀ of these cultures until their population density, as
monitored by optical density, ceased to increase.

(d) Measuring the cost of harbouring the tumour-
inducing plasmid in competition
In order to assess the fitness cost associated with bearing the
Ti plasmid, we grew pTi⁺ cells and pTi⁻ cells together under a variety of environmental conditions. pTi⁺
and pTi⁻ cells were competed in standard ATGN media as well as
AT minimal media modified to an acidic pH. This latter
media contained standard AT salts, 27.5 mM glucose,
15 mM (NH₄)₂SO₄ and 78.6 mM KH₂PO₄ (pH 5.6), and
was buffered to pH 5.6 by 2-(N-morpholino)ethanesulfonic
(MES). Under the stress of these acidic conditions, agrobac-
terial populations grow markedly slower than they grow
under standard, pH neutral ATGN conditions. We also
competed pTi⁺ and pTi⁻ cells under carbon- and nitrogen-
limiting conditions. The carbon-limiting media was AT
minimal media supplemented with 1 mM glucose and
30 mM ammonia, while the nitrogen-limiting media was AT

minimal media supplemented with 27.5 mM glucose and 1.2 mM ammonia.

All competitions were inoculated with approximately $6 \times 10^8$ cells, with approximately half $pTi^+$ and half a nearly isogenic $pTi$-cured derivative. Half of the competitions for each environmental condition competed 15955 ($pTi^+$) against TGP114 ($pTi^+$; NalR), while the other half of the replicates competed TGP110 ($pTi^+$; NalR) and TGP101 ($pTi^+$), thereby controlling for possible effects of the antibiotic resistance label. For all competition experiments, conjugation and loss of the plasmid is very rare under the tested environmental conditions [16,20]. We prepared carbon- and nitrogen-free inocula of each strain as described earlier. These cells were then used to inoculate 2 ml of appropriate fresh media to an approximate OD$_{600}$ of 0.005. The competition cultures were then incubated at 28°C for 24 h, after which all mixed cultures were sub-cultured 1:100 into 2 ml of the appropriate fresh media and incubated as before. We repeated this sub-culturing for a total of five passages. We plated a dilution series of each mixed culture at the start of the experiment, as well as after the initial, first passage and fifth passage populations reached stationary phase, onto ATGN and ATGN supplemented with 50 µg ml$^{-1}$ Nal so that we could estimate the density and frequency of both strains present. After the fifth passage reached stationary phase, a subset of the mixed cultures were allowed to incubate for an additional 24 h and then plated as described before.

(c) Virulence gene expression

Ti plasmid borne vir genes are expressed in response to acidic conditions, low phosphate levels and, most critically, the presence of plant-produced phenolics, such as acetosyringone (AS) [23]. To confirm that the vir genes are expressed in response to the resource-limiting conditions used in our experiments, we measured the expression levels driven off the virB promoter. We electroporated pSW2091 into 15955 to obtain a reporter of virulence gene expression. This plasmid contains a copy of lacZ fused to the virB promoter ($P_{virB}$:lacZ), such that β-galactoside levels—as measured by standard assays using cleavage of the colorimetric substrate o-nitrophenyl-β-D-galactoside—correspond to levels of expression driven from this promoter [24]. We grew populations of 15955 pSW2091 to mid-log phase and washed the cells seven times with 20 mM MES pH 5.6, each time collecting the cells by centrifugation. We used these washed cells to inoculate to an OD$_{600}$ of 0.005 media that contained AT minimal media salts, 20 mM MES (pH 5.6). As described earlier, we estimated the frequency and density of each cell type in each mixed culture at the start of the experiment, as well as after the initial and first passage populations reached stationary phase.

(g) Statistical analysis

For both competition experiments, the relative fitness of $pTi^+$ cells ($W_{pTi}$) was estimated as the ratio of the number of doublings by $pTi^+$ cells to that of $pTi^-$ cells over the course of the experiment [4]. Costs associated with the Ti plasmid (c) were estimated by subtracting one from the relative fitness of $pTi^+$ cells competing with $pTi^-$ cells ($1 + c = W_{pTi}^{-}$). These fitness and plasmid cost data were analysed using Proc GLM based on SAS software. The model included both the environmental condition and the NalR marker orientation as factors. Including the latter factor allowed us to remove variance associated with the marker and thereby isolate the effect of virulence gene expression by comparing least square (LS) means of the different media treatments.

3. RESULTS

(a) Curing the tumour-inducing plasmid

Experimental examination of the growth and fitness consequences of bearing a plasmid requires the comparison of strains that differ only in the presence or absence of the plasmid. Studies that do not carefully make such isogenic derivatives are prone to falsely attributing the effects of other factors to the plasmid. We generated these strains by curing A. tumefaciens 15955 of $pTi$ using plasmid incompatibility. The unmarked derivative we obtained is nearly isogenic with the 15955 parent strain, differing only in that it lacks the Ti plasmid. The absence of the Ti plasmid was confirmed with Eckhardt gel analysis, and the inability to PCR amplify several Ti plasmid specific sequences distributed around the plasmid, in parallel with reactions that did amplify sequences from the 15955 parental strain. We also confirmed that our cured derivative is unable to induce tumourigenesis in a potato-disc virulence assay [25] and cannot activate a bioreporter of acyl-homoserine lactones [26], another Ti plasmid encoded activity. Both of these phenotypes are consistent with the absence of the Ti plasmid. This $pTi$-cured derivative was designated as TGP101.
glucose as the sole source of carbon (quadratic regression, \( \beta_{\text{glucose}} = 0.18, t = 21.61, p < 0.001 \)) and \( \text{(b) ammonium as the sole source of nitrogen (quadratic regression, } \beta_{\text{ammonium}} = 0.28, t = 44.3, p < 0.0001 \)). In both cases, this effect eventually levelled off, suggesting that carbon (quadratic regression, \( \beta_{\text{glucose}} = -0.004, t = -13.9, p < 0.0001 \)) or nitrogen (quadratic regression, \( \beta_{\text{ammonium}} = -0.009, t = -28.3, p < 0.0001 \)) availability no longer limited population growth. OD_{600} was measured 45 h after inoculation. Values represent mean \( \pm \) s.e. of three replicates. All means are statistically different from one another in the glucose dose response \( (p < 0.05) \). However, the 0, 0.04, 0.08 and 0.16 mM ammonia treatment means are not significantly different from one another, while all other means in \( \text{(b)} \) are significantly different \( (p < 0.05) \).

**Figure 1.** Agrobacterium tumefaciens strain 15955 achieved higher population density in response to higher levels of \( \text{(a) glucose as the sole source of carbon (quadratic regression, } \beta_{\text{glucose}} = 0.18, t = 21.61, p < 0.001 \)) and \( \text{(b) ammonium as the sole source of nitrogen (quadratic regression, } \beta_{\text{ammonium}} = 0.28, t = 44.3, p < 0.0001 \)). In both cases, this effect eventually levelled off, suggesting that carbon (quadratic regression, \( \beta_{\text{glucose}} = -0.004, t = -13.9, p < 0.0001 \)) or nitrogen (quadratic regression, \( \beta_{\text{ammonium}} = -0.009, t = -28.3, p < 0.0001 \)) availability no longer limited population growth. OD_{600} was measured 45 h after inoculation. Values represent mean \( \pm \) s.e. of three replicates. All means are statistically different from one another in the glucose dose response \( (p < 0.05) \). However, the 0, 0.04, 0.08 and 0.16 mM ammonia treatment means are not significantly different from one another, while all other means in \( \text{(b)} \) are significantly different \( (p < 0.05) \).

**b) Resource-limited growth**

We monitored the population growth of 15955 under a variety of environmental conditions supplied with increasing amounts of glucose as the sole carbon source. We also performed an analogous experiment varying the concentration of ammonia as the sole nitrogen source. The carrying capacity of 15955 populations increased with increasing availability of glucose and ammonia, respectively (figure 1). These effects started to plateau at glucose or ammonia levels greater than approximately 5 mM (figure 1).

**c) Fitness cost of harbouring the plasmid**

The fitness of cells harbouring the Ti plasmid depended upon the environmental conditions in which they competed with \( \text{pTi}^- \) cells \( (F_{3,39} = 18.3, p < 0.0001) \). Over the course of approximately 35 generations in carbon- or nitrogen-limiting media, \( \text{pTi}^- \) cells began to overtake the \( \text{pTi}^+ \) cells in the population, regardless of which cells were resistant to nalidixic acid. This indicates that cells harbouring the Ti plasmid were at a competitive disadvantage to cells lacking the plasmid under both carbon- and nitrogen-limiting environmental conditions (figure 2). However, we did not observe this when these nutrients were abundant at both neutral and stressful, low pH (figure 2). We also observed that the frequencies of the two strains did not significantly change over the course of 24 h in stationary phase in any of the tested media \( (F_{2,23} = 0.65, \text{n.s.}) \). This suggests that the advantage that \( \text{pTi}^- \) cells manifested when resources were limiting stems from differences between the strains during lag and/or log population growth phases.

**d) Regulation of virulence gene expression**

Induction of the \( \text{vir} \) genes by the plant phenolic acetosyringone (AS) is well established \[23\]. Our experiments required modifying standard \( \text{vir} \) induction media in order to achieve either carbon-limiting or nitrogen-limiting conditions. Doing this involved increasing the availability of phosphate and decreasing the availability of glucose and ammonia. We confirmed that the \( \text{vir} \) genes are strongly induced in our carbon- and nitrogen-limiting modified induction media. When both carbon \( (F_{1,9} = 160.2, p < 0.001) \) and nitrogen \( (F_{1,9} = 1282.92, p < 0.001) \) were limiting, the presence of AS strongly induced the expression of virulence genes as measured by the activity of a \( \beta\text{-galactosidase} \) fusion (figure 3).

**e) Fitness cost of expressing plasmid-borne cooperative virulence genes**

The growth rate of clonal \( \text{pTi}^- \) 15955 populations slowed dramatically \( (F_{1,3} = 20.24, p < 0.05) \) when the environment contained AS. Under \( \text{vir} \)-inducing conditions, 15955 populations took nearly twice as long to double in size (figure 4). In contrast, populations of
Cost of cooperative virulence plasmid  T. G. Platt et al.  1695

Figure 3. Virulence gene expression is tightly regulated by environmental stimuli. The presence of the plant phenolic acetosyringone (AS) strongly promoted expression of the *virB* promoter in pTi⁺ cells (15955 pSW209ΔΩ) when both carbon (p < 0.05) and nitrogen (p < 0.05) is limiting under acidic, low-phosphate conditions. Values represent mean ± s.e. of five replicates. Filled bars, with AS; open bars, without AS.

Figure 4. The presence of the plant phenolic acetosyringone (AS) was associated with a large increase in the mid-log phase doubling time of pTi⁺ populations but not pTi⁻ populations. Environmental conditions are limiting carbon (3 mM glucose), low phosphate (500 μM), acidic pH (pH 5.6) either with (filled bars) or without (open bars) 100 μM AS present. Values represent mean ± s.e. of three replicates. Different letters denote treatment means that are significantly different (p < 0.05).

TGP101, which lack the Ti plasmid and therefore are unable to induce expression of the Ti plasmid virulence genes in response to the plant phenolic, were not affected for growth by the presence of AS (F₁,₅ = 0.11, n.s.). Furthermore, after approximately only 12 generations under *vir*-inducing conditions, pTi⁻ cells were much more common than pTi⁺ (15955) cells in populations initially composed of equal numbers of each type of cell. Under *vir*-inducing conditions, cells harbouring the Ti plasmid were at a striking competitive disadvantage to pTi⁻ cells regardless of nutrient levels (figure 5). In this experiment, we did not observe a significant cost to harbouring the Ti plasmid when AS was absent under carbon-limiting or nitrogen-limiting conditions (figure 5). This cost was observable in similar experiments that ran for approximately 35 generations (figure 2). This suggests that 12 generations is insufficient to observe the modest cost of harbouring the Ti plasmid when expression of the *vir* genes is not induced or that the differences between the media used for these experiments affect the carriage cost of the Ti plasmid.

4. DISCUSSION

Bacterial plasmids frequently impose a fitness burden on their host cells [4–6]. Most empirical estimates of these costs have focused on plasmids that impart antibiotic resistance on their hosts, while the costs of other important plasmids such as those conferring virulence, bacteriocin and catabolic functions have received much less empirical attention (see [8,9] for exceptions). In this study, we demonstrated that a natural, low-copy-number, ecologically important plasmid—the *A. tumefaciens* Ti plasmid—is associated with two kinds of fitness costs. We observed a relatively modest cost to harbouring the plasmid when few of its genes are induced under either nitrogen- (LS mean c ± s.e. = -0.028 ± 0.005) or carbon- (−0.044 ± 0.005) limiting conditions. We also demonstrated a more severe burden on host cells associated with the expression of plasmid-borne genes underlying virulence that did not significantly depend on nutrient availability (nutrient replete: LS mean c ± s.e. = -0.500 ± 0.087; nitrogen-limiting: -0.389 ± 0.101; carbon-limiting: -0.410 ± 0.092).

(a) Costs and plasmid population dynamics

Fitness costs associated with plasmids are thought to reflect the consequences of metabolic drain from the expression of plasmid genes and/or the plasmid having adverse effects on host physiology [27]. These costs can vary depending on external environmental conditions [28,29], the genetic background of the cell that harbours them [27] or the population frequency of the plasmid [8].
In this study, we demonstrated that there is a fitness cost associated with the carriage of the *A. tumefaciens* virulence plasmid under resource-limiting conditions. In contrast, we did not observe this cost when cells harbouring the Ti plasmid competed against plasmidless cells in a resource-abundant environment. The resource dependence of this cost is consistent with the carriage cost of the Ti plasmid stemming from the Ti plasmid imposing a metabolic drain on the host cell. Similar results have also been reported for the antibiotic resistance plasmid pB15 in *E. coli*, where reduced conjugation rates in high-carbon environments were associated with lower plasmid burden on the host cell [29].

Previous work has found that the long-term association of plasmid and chromosomal genes can diminish the costs associated with plasmids owing to changes in either the plasmid or the chromosomal genetic content [4,27,30]. In this study, we showed that despite the long-term association between pTi5955 and the 15955 chromosomal background, there continue to be fitness costs associated with harbouring this plasmid. However, we also note that these costs were not apparent under resource-replete conditions and thus were not readily apparent in laboratory experiments where saturating nutrients mask the cost of the plasmid. In most natural terrestrial environments, bacteria are resource-limited, and thus the costs of the Ti plasmid are likely to be manifested. Like many natural plasmids, the Ti plasmid exhibits several phenotypes consistent with selection to minimize the burden that it imposes on its host cell, also promoting the fitness of the plasmid itself because it benefits from vertical transmission during host cell division [29,31,32].

Most of the genes on the Ti plasmid are known to be tightly regulated according to biotic or abiotic stimuli such that these gene products are only produced when needed [33]. Natural plasmids are often maintained at a low copy number and typically have stability systems promoting efficient partitioning into both daughter cells, so that they are not lost during cell division [34]. The low copy number of these plasmids is thought to minimize their burden on host cells, thereby promoting the vertical transmission of the plasmid [31,32].

Despite the Ti plasmid generally having a low copy number, expressing genes only when induced by proper environmental stimuli and having a long-standing association with agrobacterial genetic backgrounds, it still imposes a significant fitness burden on host cells when resources are limiting and few plasmid-borne genes are expressed (figure 2). Ti plasmids belong to the repABC plasmid family, whose plasmids are widely distributed among *a*-proteobacteria, but particularly common in the Rhizobiales, the order that includes *Agrobacterium, Rhizobium* and *Brucella*, among others [21]. As with the Ti plasmid, these plasmids tend to be found at low copy number within host cells. Stable vertical transmission of these plasmids depends upon an efficient partitioning system involving the RepA and RepB proteins, which physically links replication of the plasmid to the process of cell division [35]. In addition, recent work has identified a toxin-antidote system on pTiC58 that further enhances the stability of this Ti plasmid [36]. Cells that lose the plasmid after segregation are often killed owing to poor stability of the antidote and unconstrained toxin activity [37]. These replication, partitioning, toxin-antidote and perhaps other stability systems can slow the generation of pTi cells from pTi, thereby helping to account for how the Ti plasmid is maintained despite the carriage costs that it imposes on its host cell. Ironically, they may also contribute to the burden that the Ti plasmid imposes on its agrobacterial host because these systems must be expressed each generation in their role of stabilizing the inheritance of the plasmid during cell division [3].

We also show that the expression of the plasmid-borne virulence genes was associated with a dramatic reduction in fitness. Consistent with previous studies on other plasmids [8,38], pTi cells were at a much greater competitive disadvantage to pTi cells when environmental conditions stimulated the expression of plasmid-borne virulence genes than when these genes were not induced (figure 5). Agrobacterial virulence is likely to be a metabolically expensive behaviour because it involves the construction of the elaborate T4S apparatus necessary for agrobacterial pathogenesis. These macromolecular bacterial secretion machines span the inner membrane, periplasm and outer membranes of the bacterial cell and are composed of numerous protein subunits. The T4S system encoded by the vir genes is composed of many copies of 11 distinct proteins [39]. During infection, the T-DNA and several effector proteins, which modify plant cell physiology and promote T-DNA integration, translocate into the plant host cell through the channel formed by this complex structure [10], allowing for subsequent genetic transformation of the plant cell. Additionally, vir-induction has been shown to be associated with an increase in Ti plasmid copy number, potentially adding to the cost associated with harbouring the plasmid [40,41]. This increase in Ti plasmid copy number, in addition to the metabolic drain of producing T4S machinery and associated functions, may contribute to the high plasmid cost that cells experience under vir-inducing conditions. In light of this high fitness cost, it is sensible that the expression of these genes is tightly regulated by external environmental cues associated with the presence of a potential plant host. This regulation minimizes the plasmid’s burden on the host cell, thereby benefiting the plasmid in terms of the fitness it gains through vertical transmission during host cell reproduction [31,32].

Expression of the plasmid conjugal machinery used to transfer the plasmid between bacterial cells can also impose a dramatic fitness burden on host cells [29,32,38]. This cost establishes a trade-off between horizontal and vertical transmission of a plasmid because conjugation of the plasmid comes at the expense of lower cell division rates, and thus a lower efficiency of plasmid vertical transmission [32]. The Ti plasmid is able to colocalize new agrobacterial genetic backgrounds via conjugal transfer of the plasmid; however, under our experimental conditions, the Ti plasmid conjugal machinery is not expressed—and therefore does not contribute to the plasmid associated costs that we observed. The conjugation of the Ti plasmid is highly regulated such that it only occurs in the disease environment where opines are abundant [16] and in which pTi cells have a competitive advantage [42].

The cost associated with carrying a plasmid plays a central role in its population dynamics [2,7,29]. The fitness costs that we have demonstrated in this study probably
factor into the observation that the majority of natural agrobacterial isolates lack a virulence-conferring Ti plasmid [43,44], despite the plasmid’s high stability [21], ability to conjugate into novel backgrounds [16], and benefits associated with phenolic resistance [45] and opine catabolism [42].

(b) Cooperation and plasmid evolutionary dynamics

While the expression of the genes underlying agrobacterial pathogenesis is very costly, its benefits stem from the Ti plasmid mediated catabolism of a public good—specialized catabolites produced by infected plant tissues [46,47]. Access to these opines depends on a cell expressing opine catabolic genes that are carried on the Ti plasmid. Thus, the Ti plasmid is a greenbeard wherein the benefits of the cooperative act of pathogenesis are intrinsically directed towards genetically similar cells that tend to have both virulence and opine catabolic genes due to their efficient coinheritance during vertical or horizontal transmission of the Ti plasmid [11,12]. Greenbeards were once thought to be rare and improbable; however, several examples have been identified in recent years [12]. The agrobacterial greenbeard provides a clear example with a particularly well-characterized genetic basis, making this an ideal model system for studying the evolutionary dynamics of greenbeard cooperation. Towards this goal, we take advantage of this knowledge and this system’s genetic tractability to experimentally dissect the costs of cooperation for this greenbeard system.

The greenbeard-like recognition intrinsic to agrobacterial pathogenesis restricts the availability of opines as a nutritional source. However, once opines become available in the disease environment, there is opportunity for selection favouring cheating agrobacterial genotypes that do not pay the substantial costs of being cooperative but are able to access its benefits (figure 5). These cheaters may either arise de novo within the disease environment or colonize from outside sources. The high cost of expressing vir genes (figure 5) establishes a strong selective pressure favouring mutants that suppress expression of these genes, while maintaining the ability to catabolize opines using other plasmid genes. Many natural isolates of agrobacteria harbour opine catabolic plasmids, which confer the ability to degrade opines but not the ability to infect host plants [13,14]. In addition to these agrobacterial cheaters, several non-agrobacterial soil microbes have also evolved the ability to degrade opines [14,48]. Thus, virulent agrobacteria face both intraspecific and interspecific competition for opine resources from agrobacterial cheaters and non-agrobacterial opine catabolizers, respectively, in the soil.

Genes underlying public good cooperation are over-represented on mobile genetic elements [49]. One possible explanation for this is that horizontal transmission of these genes into non-cooperative individuals helps stabilize cooperation by forcing potential freeloaders to bear the costs of cooperation [50]. This is akin to the idea that the horizontal transmission of public good cooperation genes effectively increases the relatedness at these loci by causing neighbouring cells to share these genes [49,51,52]. Because chromosomal genes are only vertically transmitted, they do not experience this effect; consequently, chromosomal- and plasmid-borne genes are expected to be in conflict over how cooperatively the cell should behave [49]. Interestingly, the ability of conjugation to force agrobacterial cheaters to cooperate [50] is constrained by the fact that these avirulent, opine catabolic mutants carry an incompaticable plasmid that can prevent the cooperative plasmid from being introduced by conjugation [53]. Thus, the persistence of virulent agrobacteria despite these potential competitors suggests that crown gall communities are sufficiently spatially structured such that they are able to at least transiently access the opine benefits of pathogenesis such that the trait can be favoured by kin selection [11,53].

The ability to infect hosts often comes at a cost for pathogens [50,54]. This cost can manifest itself in many ways, including the expression of virulence factors. Interestingly, many forms of virulence involve cooperative behaviour on the part of infecting pathogens [50,55]. This high cost, coupled with the opportunity of freeloading, establishes a selective pressure favouring cheating strains that avoid the costs of pathogenesis but are able to reap its benefits. Consistent with this, Fortin et al. [56] reported the origin and rapid spread of avirulent mutants in laboratory cultures of A. tumefaciens growing under vir-inducing conditions. In some cases, these avirulent mutants no longer harboured the Ti plasmid present in the parent strain, while others appeared to contain a modified form of the plasmid that had undergone deletion events that abolish or diminish the ability to respond to the plant phenolic cues that trigger wild-type vir gene expression [56]. Similar observations have also been made with some agrobacterial strains growing in the presence of infected apple trees [57].

Several studies indicate that selection favouring cheaters can drive the evolution of less virulent Pseudomonas aeruginosa strains that fail to produce at least some important virulence factors [58–60]. Moreover, several studies have found that avirulent E. coli and Shigella flexneri strains often arise and spread within populations growing under conditions inducing virulence expression [61,62]. These avirulent mutants can result from loss of the entire plasmid [61], deletion of virulence-determining regions of the plasmid [61] or integration of the plasmid into the chromosome, which results in lower expression of virulence genes [62]. Interestingly, the Shigella virulence plasmid counteracts the selective pressure favouring the loss of the plasmid using a toxin-antidote system that leads to the death of daughter cells that do not inherit the plasmid [63]. These results parallel the observation that the Ti plasmid is rapidly lost or undergoes virulence-disabling deletion events under laboratory conditions [56]. Our results suggest that at least one source of the selective pressure favouring the spread of these avirulent mutants stems from the high costs associated with the expression of the genes underlying pathogenesis.

The population dynamics and maintenance of bacterial plasmids depend on the costs they impose and benefits they confer on the cells that host them. The costs (figures 2 and 5) and benefits [42] associated with the A. tumefaciens Ti plasmid, as with other plasmids, are environmentally context-dependent. Consequently, the outcome of competition between pTi<sup>+</sup> and pTi<sup>−</sup> agrobacteria varies with the environmental conditions in which they are competing. This genotype-by-genotype-by-environment interaction
suggests that the Ti plasmid may be subject to selective pressures that vary over space and time. Because of the benefits of opine catabolism, disease environments are likely to be sources for pathogenic agrobacteria [42]. In contrast, non-disease environments are predicted to be sinks for virulent agrobacteria owing to Ti-plasmid-associated costs when resources are limiting.

We thank Anna Larimer, Elise Morton, Sean Curtis, Peter Merritt and Mike Hiebing for conceptual feedback or technical assistance throughout this study. Helpful conversations with Stephen Farrand, Spencer Hall, Michael Hynes and Jeff Smith also improved this paper. T.G.P. was funded by the NSF (DEB-0608155) and the NIH (R01 GM092660).

REFERENCES

Cost of cooperative virulence plasmid  
T. G. Platt et al.  


38 Haft, R. J. F., Mittler, J. E. & Traxler, B. 2009 Competition favours reduced cost of plasmids to host bacteria. ISME J. 3, 761–769. (doi:10.1038/ismej.2009.22)


42 Platt, T. G., Fuqua, C. & Bever, J. D. Submitted. Resource competition and competitive dynamics shape the benefits of public goods cooperation in a plant pathogen.


