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Title: Positive selection inhibits gene mobilisation and transfer in soil bacterial communities

Authors: James P. J. Hall\textsuperscript{1,2,*}, David Williams\textsuperscript{3}, Steve Paterson\textsuperscript{3}, Ellie Harrison\textsuperscript{1}, Michael A. Brockhurst\textsuperscript{1,*}

Affiliations:
\textsuperscript{1}Department of Animal and Plant Sciences, University of Sheffield, Western Bank, Sheffield, S10 2TN, United Kingdom
\textsuperscript{2}Department of Biology, University of York, York, YO10 5DD, United Kingdom
\textsuperscript{3}Institute of Integrative Biology, University of Liverpool, Biosciences Building, Liverpool, L69 7ZB, United Kingdom

*Correspondence to: j.p.hall@sheffield.ac.uk, m.brockhurst@sheffield.ac.uk

Main text:
Horizontal gene transfer (HGT) between bacterial lineages is a fundamental evolutionary process that accelerates adaptation. Sequence analyses show conjugative plasmids are principal agents of HGT in natural communities. However, we lack understanding of how the ecology of bacterial communities and their environments affect the dynamics of plasmid-mediated gene mobilisation and transfer. Here we show, in simple experimental soil bacterial communities containing a conjugative mercury resistance plasmid, the repeated, independent mobilisation of transposon-borne genes from chromosome to plasmid, plasmid to chromosome, and, in the absence of mercury selection, interspecific
gene transfers from the chromosome of one species to the other via the plasmid. By reducing conjugation, positive selection for plasmid-encoded traits, like mercury resistance, can consequently inhibit HGT. Our results suggest that interspecific plasmid-mediated gene mobilisation is most likely to occur in environments where plasmids are infectious, parasitic elements rather than those where plasmids are positively selected, beneficial elements.

Conjugative plasmids — semi-autonomous mobile genetic elements that transfer between bacteria — are key agents of horizontal gene transfer (HGT)\(^1,2\), facilitating rapid bacterial adaptation by spreading ecologically important traits between lineages\(^3\). The physical movement or duplication of genes (gene mobilisation) between chromosomes and plasmids, and their subsequent transfer between hosts, can be decisive in microbial evolution, and has facilitated acquisition of antimicrobial resistance\(^4,5\) and emergence of virulent pathogens\(^6,7\). Such mobilisation can be facilitated by transposable elements (TEs), which encode enzymes (transposases) allowing transfer of genes between replicons\(^2\). However, the effects of ecological factors and natural selection on gene mobilisation and subsequent HGT is unclear. This is particularly the case for natural environments with a high degree of spatial structure\(^8\), which is rarely captured by experimental studies. An outstanding question is how positive selection for plasmid-encoded traits, like resistance genes, affects the ability of that plasmid to spread genes through a community. Although positive selection can favour HGT by benefitting bacteria which have acquired the plasmid (transconjugants), it can also prevent HGT by killing or inhibiting growth of potential plasmid recipients\(^9,10\).
Experimental evolution of bacterial communities is a powerful tool for exploring the evolutionary dynamics of plasmid-mediated HGT, bridging the gap between simplified short-term laboratory studies \(^{11,12}\) and comparative genomics of natural populations \(^{4,5,13,14}\). To investigate how positive selection for plasmid-borne genes and the presence of an alternative host species interact to determine plasmid dynamics, we established communities of the common soil bacteria \textit{Pseudomonas fluorescens} and \textit{Pseudomonas putida}, either alone or in co-culture, in sterilized soil microcosms, which offer a complex, spatially-structured, and experimentally tractable environment \(^{15-17}\). Communities began with the 307-kb conjugative plasmid pQBR57 at ~50% frequency; this plasmid was isolated from agricultural soil and carries a 7-kb mercury resistance (Hg\(^R\)) TE, Tn5042. We also established control communities without pQBR57. Microcosms were supplemented with mercuric chloride to 16 µg/g Hg(II), or an equivalent volume of water (0 µg/g Hg(II)). This level of mercury contamination, similar to that found in industrial or post-industrial sites \(^{18}\), selects for specific Hg\(^R\) but does not necessitate it \(^{16}\).

Six replicate populations for each combination of treatments were grown for ~440 generations. Analysis of plasmid frequency dynamics showed that pQBR57 was generally maintained by \textit{P. fluorescens} and lost by \textit{P. putida}, but persisted in \textit{P. putida} when co-cultured with \textit{P. fluorescens}, due to interspecific ‘source-sink’ plasmid transfer \(^{17}\). To investigate consequent effects on gene mobilisation and transfer we sequenced clones from the beginning and the end of the experiment, and used the Bacterial and Archaeal Genome Analyser (BAGA) pipeline to identify structural variations \(^{19}\).
Results:

We detected multiple, independent gene mobilisation events between plasmid and chromosome in each species. Strikingly and unexpectedly, we also identified numerous interspecific transfers of chromosomal genes in the co-cultured treatments, facilitated by pQBR57 (Fig. 1). Gene transfer from *P. fluorescens* to *P. putida* was exemplified by a previously-unannotated *P. fluorescens* TE, Tn6291, in *P. putida* plasmids from 2/6 co-cultured communities (Fig. 1, replicates b and f). Subsequent PCR analyses found Tn6291 in *P. putida* clones from 2 further co-cultured communities (replicates d and e). Tn6291, a 22 kb TE carrying 25 predicted open reading frames and located between 2060105 and 2082440 in the *P. fluorescens* SBW25 reference sequence (ENA identifier AM181176, part of genomic island SBW_GI-1 (ref. 20)), carries an array of cargo genes with putative cytochrome c/d oxidase functions. The presence of Tn6291 in *P. putida* indicates these genes mobilised from the *P. fluorescens* chromosome onto pQBR57, and subsequently transferred between species via conjugation. All Tn6291+ *P. putida* clones were also positive for pQBR57, suggesting Tn6291 remains plasmid-borne in these isolates. Indeed, we detected plasmid-borne Tn6291 in 4/6 0 µg/g Hg(II) single-species *P. fluorescens* populations, and BLAST analysis shows a similar transposon in another soil Pseudomonad, *P. syringae* pv. *syringae* B301D (ref. 21), suggesting Tn6291 mobilises readily.

We also detected gene transfer from *P. putida* to *P. fluorescens*. *P. fluorescens* clones from 3/6 co-cultured populations had acquired the well-described *P. putida* TE Tn4652 (ref. 22) in their chromosomes, with plasmid-borne Tn4652 also present. Tn4652 is a 17
kb TE closely related to the Tn4651 toluene degradation transposon and encodes various putative enzymes including a diacylglycerol kinase and a sulfatase. Tn4652 mobilisation to the plasmid occurred readily, with events already detectible in the ancestral \textit{P. putida} clones used to inoculate the soil and begin the experiment (see Materials and Methods). PCR analysis of clones obtained over the course of the evolution experiment detected Tn4652+ plasmids in \textit{P. fluorescens} as early as transfer 3, however Tn4652 insertion in the \textit{P. fluorescens} chromosome was only detected later, after transfer 41 (Supplementary Table 2). In all cases, Tn4652 inserted in a region of the \textit{P. fluorescens} chromosome with atypical sequence composition, likely to be recently acquired DNA. Importantly, interspecific transfer of chromosomal TEs via the plasmid was only detected in populations grown without positive selection for the plasmid (0 µg/g Hg(II)). The amount of plasmid conjugation occurring, and thus opportunities for interspecific gene transfer, is likely a function of the densities of plasmid bearers and recipients. By killing potential plasmid recipients, mercury selection reduces encounters between plasmid donors and recipients, and therefore conjugation. Indeed, short-term experiments examining pQBR57 transfer (Figure 2) showed reduced effects of conjugation on plasmid dynamics when the plasmid was under selection, implying limited gene exchange. Together, these data suggest that positive selection for plasmid-borne resistance genes reduced the ability of that plasmid to facilitate HGT of chromosomal genes.
While mercury selection reduced TE transfer between species, we detected frequent mobilisation of the Hg\textsuperscript{R} TE Tn5042 from pQBR57 to the chromosome. Single-species \textit{P. putida} populations tended to lose the plasmid\textsuperscript{17}, and sequences show that under mercury selection this was facilitated by acquisition of chromosomal Tn5042. To track the acquisition of chromosomal Tn5042 by \textit{P. putida} populations, we designed PCR primers targeting ‘focal’ Tn5042 insertions (i.e. insertions detected in the end-point genome sequences) in eight different \textit{P. putida} populations, and applied these to clones collected across the experiment. As with single-species \textit{P. putida} populations, chromosomal Tn5042 was readily acquired by co-cultured \textit{P. putida}, although these populations tended to also maintain pQBR57 (Figure 3). Similarly, chromosomal Tn5042 was detected in the \textit{P. fluorescens} chromosome, which maintained the plasmid, despite its redundancy\textsuperscript{17}. These findings suggest that long-term plasmid maintenance largely depends on community context and on compatibility between plasmid and host, and, provided there are no restrictions on recombination of plasmid genes into the chromosome, is unlikely to be secured by positive selection for accessory genes alone\textsuperscript{17,25,26}.

Tn5042 also mobilised in the mercury-free treatments: we detected three instances of Tn5042 multiplying on plasmids, and one instance of Tn5042 copying to the \textit{P. fluorescens} chromosome (which occurred by transfer 35, see Figure 3). Tn5042 insertions sometimes occurred multiple times in a lineage — in co-cultured \textit{P. putida} with 16 µg/g Hg(II), one clone (from replicate c) ultimately carried six copies. Although Tn5042 copy number increased in some clones from the mercury-free treatment, we detected more copies in clones evolved under mercury selection ($Z = -5.4404$, $p < 0.0001$, $n = 48$, Exact
General Independence Test). We did not detect any Tn5042 loss. For *P. putida*, Tn5042 tended to insert in a ~10 kb region near the origin of replication, while Tn5042 tended to insert in *P. fluorescens* near or inside Tn6291 (detected in 4/12 populations under mercury selection), in three cases representing the *de novo* formation of a composite resistance transposon. Here, Tn5042 became part of the cargo of Tn6291, broadening opportunities for spread, because subsequent events favouring Tn6291 mobilisation (perhaps different to those of Tn5042) will cause co-mobilisation of Tn5042 and its Hg\(^R\) genes. The pervasive mobility of Tn5042 supports a model in which TEs exploit plasmids to rapidly spread in the natural environment\(^6\), consistent with sequence analysis suggesting Tn5042 was acquired relatively recently by pQBR57 (ref. 16).

Surprisingly, we found plasmid size generally increased, primarily due to TE accumulation. Plasmid size in one clone evolved in 0 µg/g Hg(II) increased by over 10% compared with the ancestor (Figure 1). Increased plasmid size is expected to contribute to increased cost of plasmid carriage\(^{27}\), however these results suggest such costs are negligibly small, and may be outweighed by transposition rates and/or general plasmid cost amelioration\(^{26}\).

**Discussion:**

The central role of HGT in adaptation is increasingly apparent, as ever-wider sequencing of isolates reveals the dynamic nature of microbial genomes\(^{4,5,28}\). Between-species transfer of chromosomal genes occurred only where plasmid-encoded mercury resistance was not under positive selection and the plasmid persisted instead as an infectious
element. Bacterial genome evolution is determined by the interaction between selection and recombination — here, we observed that recombination indeed makes an increased contribution to genome evolution when selection is relaxed. The transferred genes were part of the ‘accessory’ genome, which can vary even between closely-related strains and is often more strongly associated with ecological niche than phylogenetic lineage. In this case, the transferred genes were located on TEs and putative transposases could be identified. This is relevant because TEs can transfer between replicons at a high rate, providing an efficient platform for the movement of genes between chromosomes and conjugative elements. Plasmids and TEs have a close — even symbiotic — relationship. TEs can comprise a substantial fraction of a plasmid genome and where their genes are under positive selection they can boost the fitness of the plasmids that carry them due to genetic linkage. Similarly, unless they encode their own conjugative machinery, TEs must collaborate with elements such as conjugative plasmids to access new hosts; indeed models suggest that conjugative plasmids are required for TE survival and spread. Transposase activity can be affected by stress, for example nutritional deprivation or oxidative damage, and one intriguing possibility is that stresses caused by plasmid acquisition could signal to a TE that a vehicle had arrived, triggering transposition and thus increasing rates of exchange from the chromosome to that plasmid. At least one Pseudomonad TE has been shown to increase activity following conjugation.

Our results provide rare direct experimental evidence of pervasive plasmid-mediated gene mobilization, transfer and acquisition in a simple soil microcosm community. This
has profound implications for the spread of accessory genes in natural communities. Consistent with our findings, two recent studies of resistance plasmids in hospital outbreaks indicate that TE mobilisation dominates plasmid evolution. Furthermore, both studies suggest that plasmids may have acquired TEs outside of patients, i.e. in the environment, where they are less likely to experience direct antibiotic selection. HGT vastly expands the evolutionary opportunities available to bacteria, allowing species to draw upon a collective mobile gene pool: our data indicate that environmental and ecological factors will be key modulators of the rate and extent of HGT in natural communities. HGT, particularly of antibiotic resistance and virulence genes, poses a major health concern and understanding the ecology of HGT-mediated bacterial evolution will be crucial to predicting and designing interventions to prevent and mitigate such threats.

**Materials and methods:**

**Experimental design**

The evolution experiment, described previously, was designed to understand the effect of an alternative host species on plasmid population dynamics and evolution. The experiment used *Pseudomonas fluorescens* SBW25 and *P. putida* KT2440 — representative soil *Pseudomonas* species, a widespread and naturally co-occurring genus — and the 307-kb HgR plasmid pQBR57 which was isolated from the same geographic site as *P. fluorescens* SBW25 (ref. 16). Cultures were grown at 28°C and 80% relative humidity in soil microcosms consisting of 10 g twice-autoclaved John Innes No. 2 potting soil, supplemented with 900 µl sterile H2O or 900 µl HgCl2 solution. We used a fully-
factorial design with two levels of mercury treatment (0 µg/g, or 16 µg/g); two levels of
plasmid treatment (pQBR57+ starting with pQBR57-bearers at 50% frequency, or
plasmid-free starting without plasmid); and three levels of culture treatment (single-
species *P. fluorescens*, single-species *P. putida*, or co-culture with each species starting at
50% frequency). Six independent biological replicates (‘populations’) were initiated for
each treatment, consistent with previous evolution experiments \(^{42,43}\) and sufficient to
detect differences in population dynamics between the treatments \(^{17}\). Each replicate was
initiated from independent single colonies, and populations were blocked by replicate to
minimize confounding effects. The experiment was not blinded. To control for marker
effects, replicates a–c used gentamicin-labelled (Gm\(^R\)) *P. fluorescens* and streptomycin-
labelled (Sm\(^R\)) *P. putida*, whereas replicates d–f used Sm\(^R\) *P. fluorescens* and Gm\(^R\) *P.
*putida*. Samples of culture (100 µl soil wash) were serially transferred into fresh soil
microcosms containing either H\(_2\)O or HgCl\(_2\) every four days for 65 transfers (estimated as
~440 generations \(^{17}\)); this was decided before the experiment to be broadly consistent
with other plasmid experimental evolution studies \(^{42,43}\). At 16 points during and at the end
of the experiment samples were spread on selective media to isolate clones, which were
archived for subsequent analysis. After 65 transfers, a random number generator was
used to select one plasmid-bearing one plasmid-free clone (where present) from each
pQBR57+ population for DNA sequencing. If plasmid-free or plasmid-bearing clones
were present throughout the experiment but not at transfer 65, a clone from transfer 59
were used (this was the case for plasmid-bearing *P. putida* from single-species 16 µg/g
replicate f, and plasmid-bearing *P. putida* from co-cultured 16 µg/g replicates a and c).
We also sequenced ancestral clones, and three (single-species) or two (co-culture) clones
from plasmid-free treatments to test for mutations occurring in the absence of plasmid. No Tn6291, Tn4652, or Tn6290 activity was detected in the plasmid-free treatments.

**DNA sequencing and analysis**

DNA was extracted using the QIAGEN DNeasy kit, prepared using the TruSeq Nano DNA Library Preparation Kit (350 bp insert size), and sequenced on the Illumina HiSeq platform. Reads trimmed using Cutadapt (version 1.2.1) and Sickle (1.200) were analysed using the Bacterial and Archaeal Genome Analyser pipeline, which uses the bwa short read aligner and calls variants using the Genome Analysis Toolkit HaplotypeCaller.

To identify structural variation (deletions, duplications and TE insertions) in the re-sequenced clones we used the BAGA module Structure, which uses a threshold ratio of non-proper to proper paired reads to identify putative genome disruptions. Reads mapping to putative disruptions were re-assembled using SPAdes, and contigs were aligned with the reference to identify structural variants. We also used two complementary approaches to identify structural variation: Breakdancer, and custom scripts that examined coverage for characteristic direct repeats introduced by TE insertion (increase in coverage of ≥25% over a <30 bp region, compared with neighbouring positions). These different approaches were broadly consistent, and all putative structural variants were examined using the Integrated Genome Viewer (IGV). Because of differences between ancestral clones and the sequenced reference genome, variation appearing in all samples (including the ancestor) was removed from the analysis. In addition, apparent variation in hard-to-map regions (identified in an examination of parallel mutations in IGV) was considered unreliable and excluded (Supplementary Table 2).
3). We also examined putative SNV called near TE insertions, and removed these manually if miscalled. Representative TE insertions were tested by PCR on clones, and in all cases yielded products of the anticipated size.

Sequence analysis of ancestral clones revealed that in three cases, pQBR57 had acquired a TE before the experiment was initiated, indicated by dotted lines in Figure 1. In the Sm\(^R\) P. fluorescens ancestor, pQBR57 had acquired Tn6290 at position 164349–164354. This event likely occurred in P. putida UWC1 during preparation as a donor for transfer of pQBR57 into P. fluorescens \(^16\) because Tn6290 is present in P. putida UWC1 and not in P. fluorescens SBW25. In Sm\(^R\) P. putida, pQBR57 had acquired Tn4652 at 152552–152558 while in Gm\(^R\) P. putida pQBR57 had acquired Tn4652 at 162797–162802. These events may have either occurred in the donor P. putida UWC1 strain or in the recipient P. putida KT2440 strain, as both contain identical copies of Tn4652. In any case, TE insertion must have occurred rapidly as our stocks were all prepared from single colonies and ancestral pQBR57 contains neither Tn6290 nor Tn4652 (ref. 16) Tn4652 insertion into resident plasmids is consistent with previous work which found Tn4652 in pQBR plasmids pQBR55 and pQBR44 (ref. 16) presumably after acquisition by P. putida UWC1 (ref. 49).

**PCR analysis of clones**

We tested archived clones for TE insertions by PCR. Standard reactions were performed using GoTaq Green Master Mix (Promega), 0.4 \(\mu\)M each primer (Supplementary Table 4), and 0.2 \(\mu\)l archived culture on a program of 95°C for 5 minutes, followed by 30 cycles
of 95°C for 30 seconds, 58°C for 30 seconds and 72°C for 1 minute, followed by a final extension of 72°C for 5 minutes. Tn6291 was detected in reisolated *P. putida* clones, and parallel reactions using primers targeting the *P. fluorescens* 16S rDNA locus were performed to rule out the presence of contaminating *P. fluorescens*.

**Statistical analyses**

To analyse the number of Tn5042 insertions in 0 µg/g and 16 µg/g Hg(II) we used the R package ‘coin’ to perform an Exact General Independence Test. To avoid pseudoreplication with populations from which >1 sample was sequenced, we analysed the mean number of Tn5042 insertions per species per population. To analyse the effect of mercury on conjugation dynamics we performed an Exact General Independence Test on plasmid distribution between donor and recipient after 5 transfers.

**Data availability**

Short read data is available at the European Nucleotide, Archive Project Accession PRJEB15009. Data presented in Figures 1–3 are on Dryad Digital Archive doi:10.5061/dryad.6gf28.

**Code availability**

The Bacterial and Archaeal Genome Analyser (BAGA) is available online at https://github.com/daveuu/baga. Representative scripts used to analyse our data are on Dryad Digital Archive doi:10.5061/dryad.6gf28.
References:


37. Baharoglu, Z., Bikard, D. & Mazel, D. Conjugative DNA Transfer Induces the


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**Author contributions:** JPJH, EH, and MAB designed the study; JPJH collected data; JPJH, DW and SP analysed the data. JPJH and MAB drafted the manuscript. All authors discussed results and commented on the manuscript.
Author information: The authors declare no competing financial interests.

Correspondence and requests for materials should be addressed to
j.p.hall@sheffield.ac.uk or m.brockhurst@sheffield.ac.uk.

Figure Legends:

Figure 1. Evolved clones show extensive within- and between-species gene mobilisation. Each panel shows events detected in evolved *P. putida* (left, light green) and *P. fluorescens* (right, light blue), with changes in associated pQBR57 (if present) shown below. In each panel, six concentric lanes a–f indicate independent populations. One clone was sequenced from each population for each species, except where both plasmid-bearing and plasmid-free genotypes were detected. In this case, we sequenced one clone of each, with the plasmid-bearing clone indicated by the inner set of symbols in that lane (Supplementary Table 1). Duplicative insertions of large TEs are indicated by filled triangles coloured according to TE type (see key) and connected to ancestral positions (indicated by open triangles) by an arrow describing direction of duplication. Dotted lines indicate insertions that occurred before the evolution experiment was initiated (see Methods). Insertions of smaller insertion sequence (IS) elements are in black. Black bars indicate large deletions, and yellow bars (panel C, replicate e) indicate large tandem duplications. Scale is given in Mbp, and replicons are scaled to the same size for clarity. (A) Clones evolved in single-species populations with 0 µg/g Hg(II). (B) Clones evolved in single-species population with 16 µg/g Hg(II). In panels A and B, lines indicate the
physical separation of the two species. (C) Clones evolved in co-cultured populations with 0 µg/g Hg(II). (D) Clones evolved in co-cultured populations with 16 µg/g Hg(II).

Figure 2.
Plasmid dynamics are altered under positive selection. Top row: plasmid-bearing (‘donor’) and plasmid-free (‘recipient’) P. fluorescens were mixed in approximately equal ratios and cultured for 5 transfers in 0 µg/g Hg(II) microcosms. Densities of donors (dotted line) and recipients (solid line) and their plasmid statuses (donor, yellow fill; recipient, green fill; filled areas are overlaid) were estimated each transfer by plating onto selective media and replica plating onto Hg(II) where appropriate. Each panel represents an independent population. Bottom row: as top row except with 16 µg/g Hg(II).

Conjugation makes a reduced contribution to plasmid dynamics under 16 µg/g Hg(II) (Z = 2.88, p = 0.002, n = 12, Exact General Independence Test). These results are similar to those reported by Stevenson et al.10, showing that this pattern holds in soil microcosms.

Figure 3. Spread of chromosomally-acquired mercury resistance. (A) Frequency dynamics of focal Tn5042 insertions in P. putida chromosomes under 16 µg/g Hg(II) were tracked from transfer 9 (when insertions were first detected) to transfer 65. For each timepoint in each population, presence and frequency of the focal insertion was tested by PCR on ~30 clones using primers bridging the transposon and the chromosome (this giving a 95% chance of detecting a subpopulation comprising 10% of the total); other Tn5042 insertions were identified previously as pQBR57– merA+ clones17. Plasmid-bearing clones were identified previously17. Frequencies of different genotypes are
indicated by filled stacked areas. Each panel represents an independent separate population: top row, single-species populations a, b, c, e; bottom row, co-cultured with P. *fluorescens* populations c, d, e, f. (B) Frequency dynamics of focal Tn5042 insertion in *P. putida* (replicate b).
0 µg/g Hg(II) 16 µg/g Hg(II)

P. put. P. flu.

Tn4652 Tn5042 Tn6290 Tn6291

pQBR57 in P. put. pQBR57 in P. flu.

A B C D

single-species culture co-cultured