The Gossip Paradox: why do bacteria share genes?

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Abstract

Bacteria, in contrast to eukaryotic cells contain two types of genes: chromosomal genes that are fixed to the cell, and plasmids that are mobile genes, easily shared to other cells. The sharing of plasmid genes between individual bacteria and between bacterial lineages has contributed vastly to bacterial evolution, allowing specialized traits to ‘jump ship’ between one lineage or species and the next. The benefits of this generosity from the point of view of both recipient and plasmid are generally understood, but come at the expense of chromosomal genes in the donor cell, which share potentially advantageous genes with their competition while receiving no benefit. Using both continuous models and agent based simulations, we demonstrate that ‘secretive’ genes which restrict horizontal gene transfer are favored over wide range of models and parameter values. Our findings lead to a peculiar paradox: given the obvious benefits of keeping secrets, why do bacteria share information so freely?

1 Introduction

Evolution, that blind process of inheritance and selection, is both a process of constant refinement and optimization, and also a process of adaption; while in stable environments the species best able to make efficient use of available resources may drive others to extinction, in more variable environments it is those species most able to change and adapt which will flourish.

This tension between efficiency and adaptability can be seen most clearly in bacterial genomes, where ‘core’ genes are stored on bacterial chromosomes, while ‘accessory’ genes are commonly found on mobile gene elements, such as plasmids: small stable loops of DNA, independent from the host chromosome.

Core genes code for critical metabolic pathways, cell division and motility, while accessory genes code for more esoteric capabilities— for example resistance to heavy metals [53], uncommon metabolic pathways[50], virulence factors [39], or antibiotic resistance [9]. Genes stored on mobile gene elements can be lost, rearranged, and most importantly for our purposes, passed from one bacteria to another via the process of Horizontal Gene Transfer (HGT).
Research into HGT over the past decades has fundamentally altered our understanding of bacterial evolution. Rather than a gradual accumulation of mutations within a single continuous clonal lineage, HGT allows entire gene modules to be transferred from one genome to another, fulfilling much the same role as sexual recombination. Bacterial genomes are less a tapestry which must be altered one thread at a time, but instead a patchwork quilt, with some estimates [13] suggesting that upwards of 80% of bacterial genes have undergone HGT at some point in their history. While the historical impact of HGT is enough to earn great scientific and philosophical interest, the role of HGT in the spread of antibiotic resistance and bacterial virulence factors also fuels significant medical and practical concerns in our understanding of HGT [64, 7, 47].

HGT can take one of several pathways (see Fig. 1). It can take place via transformation: one bacterial cell taking up genetic material from its environment [29]. The process of transduction takes place when HGT takes place via an invading viruses. By chance, a virus may accidentally package host DNA inside a viral capsid before lysing a cell, eventually leading to DNA from one host being inserted into the next. Most strikingly of all, accessory genes can be shared via bacterial conjugation, in which one cell bridges the gap to another, duplicates its plasmids, and funnels these copies across. For the interested reader seeking further detail we recommend Eberhard’s classical review paper [15], and the more recent overview of Thomas and Nielson [57].

Conjugation is extensively regulated, both from the point of view of the donor and recipient. On the side of the donor cell, plasmid bound quorum sensing genes determine the appropriate density of donor cells for a particular environment [32] and intricate restriction-modification mechanisms (what might be thought of as the ‘bacterial immune system’) act to suppress foreign gene elements [44], though under what conditions a such foreign elements are accepted and when they are repressed remains unclear. On the side of the recipient cell, individual plasmids exclude incoming plasmids from the same ‘incompatibility group’, as these are likely to disrupt plasmid replication and regulatory processes [17]. Conjugation rate depends both on the plasmid in question, but also on the bacterial background these plasmids inhabit [20].

Bacterial conjugation is, for many reasons, somewhat odd. In particular, it involves one bacteria supplying another bacteria with (presumably) beneficial genetic material, often at reasonable expense to itself [14]. From the point of view of the plasmid genes being shared, this expense makes sense- the time and resources involved create more copies of the plasmid and allow it to ride over selective sweeps [4] or spread to new species or environments [6]. From the point of view of chromosomal genes in the recipient cell, the benefits are also clear; if the plasmid is beneficial the receiving bacteria and its descendants gain in fitness. Given the sometimes heavy burden imposed by newly arrived plasmid genes [69, 43, 1], the cost of accepting new plasmids can not be assumed to be trivial. Acceptance of an arriving plasmid can viewed as an act of ‘trust’, but not one of ‘generosity’ (at least, to the extent that such intentionalistic language can be applied to bacterial behavior). Though interesting in its own right, this question of trust will not be the focus of our present work.
Figure 1: Modes of Horizontal Gene Transfer. Left: HGT via transformation. After a cell dies, the lysing cell spills fragments of genes into the environment. Later, some other cell collects one of these fragments, incorporating the gene fragment into its genome. Middle: HGT via transduction. A phage infects a cell, hijacking the cellular machinery in order to produce further phages. In the process, one of the new phages is accidentally constructed containing a piece of the original cell’s DNA, rather than the phage’s. Later, this defective phage inserts the DNA fragment into a new cell. Right: HGT via conjugation. A cell contains DNA on a plasmid. This plasmid codes for bacterial conjugation, leading to the construction of a ‘pilus’. The donor cell binds to another bacteria, duplicates the plasmid DNA, and sends it through the pilus to its new host. Afterwards, the two cells go their separate ways.
While the benefits of HGT to the plasmid are clear, and the benefit to the receiving cell is at least somewhat understandable, from the perspective of the chromosomal genes in the donor cell, conjugation would appear to be strictly harmful. Not only does conjugation take significant time and risk infection by certain types of phages [9], in sharing advantageous DNA, a bacteria runs the risk of not only helping a rival in the present, but of generating a superior lineage and rendering its own descendents obsolete. In some cases, the recipient cell may even be of a completely different species altogether [39].

While previous authors have studied the peculiar evolutionary forces resulting from HGT from the point of view of bacterial plasmids—asking such questions as ‘how are deleterous genes not lost?’ and ‘why are beneficial genes not incorporated into chromosomal genomes more directly?’ [22], in this work we instead consider HGT from the point of view of those chromosomal genes left behind. Our goal in what follows is to investigate and demonstrate the tension between the interests of chromosomal and plasmid bound genes, particularly with respect to ‘preferred’ conjugation rate.

In section 2 we make use of a deterministic model to study the evolutionary dynamics in the simple case of one plasmid and two possible conjugation rates; we find that bacterial strains which suppress plasmid conjugation out compete their more generous competitors. We extend this model in section 3 in order to consider longer evolutionary time spans. In section 4 we construct a more detailed agent based model, and through simulation demonstrate how genes that limit plasmid conjugation gain a long term evolutionary advantage over a wide parameter range, even when accounting for space. In section 5 we build further upon this model, and discuss the similarities and differences between HGT and previous evolutionary ‘paradoxes’; namely the emergence of altruism and the evolution of sex. In both cases we find that classical resolutions to these past paradoxes fail to stabilize plasmid sharing.

2 A Single Selective Sweep

Our goal throughout this article will be to explore the evolutionary pressures experienced by chromosomal genes and how they interact with more mobile plasmids. While the course of evolution is governed by many factors, in this section our goal is to construct the simplest possible deterministic model in order to study the interaction between mobile and ‘static’ genes, and build our intuition, before moving on to more complex modelling approaches.

Consider a population of bacteria, each of which is either positive (⊕) or negative (⊖) for some beneficial plasmid. In addition, each bacteria possess either a generous (G) or secretive (S) gene on their chromosome, which share plasmids via conjugation at rate \( c \), or at some drastically reduced rate \( \epsilon c \). Note that this genotype determines a bacteria’s propensity to donate plasmids to others; all bacteria are equally capable of receiving plasmids. Here \( \epsilon c \) accounts for both imperfect conjugation suppression on behalf of our chromosome, and also ‘plasmid leakage’ via the unregulated...
HGT processes of transduction or transformation. Plasmid positive bacteria have some fitness \( f_\oplus \), while plasmid negative bacteria have fitness \( f_\ominus \). See figure 2 for a schematic of this set up.

Written as chemical equations we have:

\[

g_\pm \xrightarrow{f_\pm} 2g_\pm, \quad s_\pm \xrightarrow{f_\pm} 2s_\pm, \quad (1)
\]

\[
\begin{align*}
g_\oplus + s_\ominus & \rightarrow g_\oplus + s_\oplus, \\
g_\ominus + g_\oplus & \rightarrow 2g_\ominus, \\
s_\ominus + g_\oplus & \rightarrow s_\ominus + g_\oplus, \\
s_\oplus + s_\ominus & \rightarrow 2s_\ominus. \quad (2)
\end{align*}
\]

We assume a ‘death rate’ such that total population is held constant; \( g_\oplus + g_\ominus + s_\oplus + s_\ominus = 1 \) at all times.

Written as a series of differential equations we have:

\[
\begin{align*}
g_\oplus' &= (f_\oplus - \bar{f})g_\oplus + (c g_\oplus + ccs_\oplus)g_\ominus, \\
s_\ominus' &= (f_\ominus - \bar{f})s_\ominus + (c g_\ominus + ccs_\ominus)s_\oplus, \\
g_\ominus' &= (f_\ominus - \bar{f})g_\ominus - (c g_\ominus + ccs_\ominus)g_\oplus, \\
s_\oplus' &= (f_\ominus - \bar{f})s_\oplus - (c g_\ominus + ccs_\ominus)s_\ominus. \quad (4)
\end{align*}
\]

Here \( \bar{f} = f_\oplus (G_\oplus + S_\ominus) + f_\ominus (G_\ominus + S_\oplus) \) refers to the average fitness of the current population. The first term in each derivative refers to changes in population due to population growth and death/displacement. The second term in each derivative gives the demographic change caused by plasmid conjugation. These equations, can be viewed as filling similar role to the standard evolutionary replicator equations [40, 11], although HGT somewhat complicates this picture.

Suppose, we add a single beneficial plasmid to an otherwise plasmid negative population. This plasmid may find itself in either a generous or secretive bacteria. Due to its higher fitness, the plasmid is going to ‘sweep’ through, becoming ubiquitous in our population (a so called “selective sweep”). An illustration of two such selective sweeps is given in figure 3. Given the interaction between plasmids and chromosomes, we wish to determine how the balance between generous and secretive chromosomes changes as the new plasmid reaches saturation in our population. Stated mathematically, we wish to determine \( G_\oplus(\infty) \) and \( S_\oplus(\infty) \), given our parameters and initial conditions.

For this task, it proves convenient to make use of a slight relabeling of variables. Consider the population of mutants, \( M \), which are always plasmid positive, and either all generous or all secretive; \( M \) represents the original mutant bacteria and all its direct descendants. We also consider a “Wild type” population \( W \), which can be plasmid positive \( W_\oplus \) or plasmid negative \( W_\ominus \). The wild type population consists of some mixture of generous and secretive chromosomal genes. Because \( 1/n (G_\oplus/S_\oplus) = 0 \) we know that the relative proportions of generous and secretive genes remains fixed for \( W_\oplus \). Because \( W_\ominus \) grows proportional to \( W_\ominus \) in its first term, and \( W_\ominus \) in its second, and because \( W_\oplus \) initially has the same \( G/S \) ratio as \( W_\ominus \), it must inevitably continue with this same ratio throughout the entire course of our ODE. Thus, the relative ratio of generous and secretive chromosomal genes are fixed in the wild type population.
Figure 2: Our four ‘variants’, $G_\oplus, G_\ominus, S_\oplus$ and $S_\ominus$. ‘Generous’ variants are depicted in blue (left), ‘secretive’ in green (right). Plasmids are actively spread by $G_\oplus$ type bacteria, who posses both the plasmid and the chromosomal gene inclined to spread it. $S_\oplus$ type bacteria share the plasmid at a much lower rate, potentially close to zero. All variants replicate according to their fitness, with plasmid carrying cells assumed to have $f_\oplus > f_\ominus$. The case of burdensome plasmids, or plasmids with varying fitness, we ignore for the time being.

Figure 3: Solving equations 4, we observe an incoming plasmid arrive and become ubiquitous in the population (a selective sweep). As it does so, the balance between generous and secretive genes shifts. $c = 0.01$, $\epsilon = 0.01$, $f_\oplus = 1.01$, $f_\ominus = 1$ (Left) Initial plasmid positive population is $G_\oplus = 10^{-3}, S_\oplus = 0, G_\ominus \approx 0.9, S_\ominus = 0.1$. (Right) Initial plasmid positive population is $G_\oplus = 0, S_\oplus = 10^{-3}, G_\ominus \approx 0.9, S_\ominus = 0.1$. In the right hand panel, black diamonds represent the predicted final values using the approximation eq.19, while circles represent predictions made by solving eq. 16 using Newton’s method. In the left hand panel, these values are indistinguishable and the symbols are placed one on top of the other. Overall, when starting with a generous mutant (left), the levels of generous individuals increases from 0.899 to 0.9035, a 0.5% increase. When starting with a secretive mutant (right), the secretive population increases from 0.100 to 0.3955, a four-fold increase.
Stated algebraically, we have:

\[ \begin{align*}
G &= \rho W + \chi M, \\
S &= (1 - \rho) W + (1 - \chi) M, \\
G &= \rho W, \\
S &= (1 - \rho) W.
\end{align*} \tag{5} \]

Where \( \rho = \frac{G - (0)}{G - (0) + S - (0)} \) and \( \chi = \frac{G - (0)}{G - (0) + S - (0)} \). This partitioning of the population is algebraic rather than physical, but allows us to simplify our equations:

\[ \begin{align*}
\dot{M} &= (f - \bar{f}) M, \\
\dot{W} &= (f - \bar{f}) W - (c_M M + c_W W) W, \\
\dot{W} &= (f - \bar{f}) W + (c_M M + c_W W) W.
\end{align*} \tag{6} \]

Here \( c_M \) and \( c_W \) represent the conjugation rates of the mutant and wild type populations, and are given by \( c_M = \chi c_G + (1 - \chi) c_S \) and \( c_W = \rho c_G + (1 - \rho) c_S \).

Figure 4: A graphical illustration of our change of variables. Presence/absence of beneficial plasmid is denoted by bright/dark colours. secretive/generous chromosomes are denoted by green and blue (respectively). Over time descendants of our initial mutant displaces the wild type population, while simultaneously, HGT via conjugation supplies the wild type population with the beneficial plasmid, leaving the relative abundance of the chromosomal genes in the wild type population unchanged.

As before, the total population is held constant, \( M + W + W = 1 \), and \( \bar{f} \) is the mean fitness of the current population, \( \bar{f} = f_M M + f_W W + f_W W \).

Combining the definition of \( \bar{f} \) with population conservation, we find that \( (f - \bar{f}) = (f - \bar{f}) W = \Delta f W \) and \( (f - \bar{f}) = (f - \bar{f})(1 - W) \). Similar use of the total population equation gives

\[ \begin{align*}
\dot{M} &= \Delta f W M, \\
\dot{W} &= -\Delta f (1 - W) W - (c_M M + c_W (1 - M - W)) W, \\
&= -\Delta f + (1 - W) W - (c_M - c_W) M W.
\end{align*} \tag{7} \]

Dividing the equation \( \dot{W} \) by \( \dot{M} \) eliminates time dependence:

\[ \frac{\dot{W}}{\dot{M}} = \frac{\Delta f + c_W (1 - W)}{\Delta f M} - \frac{c_M - c_W}{\Delta f}. \tag{8} \]
The above can be rearranged to give a D’Alembert equation \(^{27}\) of the form:

\[
W_\oplus = \frac{(W_\odot(M) + \beta)M}{\alpha} + 1,
\]

where \(\alpha = (\Delta f + c_w)/\Delta f\), \(\beta = (c_m - c_w)/(\Delta f)\) and \(W_\odot\) is the derivative of \(W_\oplus\) with respect to \(M\) (as opposed to \(t\)). This equation permits solutions of the form:

\[
W_\odot(M) = 1 + kM^\alpha + \frac{\beta M}{\alpha - 1},
\]

where \(k\) is a constant of integration. Given the initial conditions are \(M(0) = M_0 \ll 1, \ W_\oplus = 0, \ W_\ominus = 1 - M_0\). This leads to:

\[
W_\odot(M_0) = 1 - M_0 = 1 + kM_0^\alpha + \frac{\beta M_0}{\alpha - 1},
\]

\[
\left(-1 - \frac{\beta}{\alpha - 1}\right)M_0 = kM_0^\alpha,
\]

\[
k = -\left(1 + \frac{\beta}{\alpha - 1}\right)M_0^{\alpha-1}.
\]

The final mutant population is found by noting that \(W_\oplus \to 0\) as time proceeds, and hence

\[
W_\odot(M_\infty) = 0 = 1 + kM_\infty^\alpha + \frac{\beta M_\infty}{\alpha - 1}.
\]

If we assume that \(M_0\) and \(M_\infty\) are of the same order of magnitude, then we see that for \(M_0 \ll 1\), the term \(\frac{\beta M}{\alpha - 1}\) is negligible compared to the +1 term unless \(\alpha \approx 1\). Hence

\[
M_\infty \approx (-1/k)^{1/\alpha},
\]

\[
\approx \left(1 + \frac{\beta}{\alpha - 1}\right)^{-1/\alpha}M_0^{1-1/\alpha},
\]

\[
\approx \left(\frac{c_m}{c_w}\right)^{\frac{-\Delta f}{W}}M_0^{\frac{c_w}{c_m}+\frac{\Delta f}{W}}.
\]

In cases where this formula suggests \(M_\infty\) is large enough such that the approximation \(M_\infty \ll 1\) fails, it is possible to instead use Newton’s method to solve eq. \(^{16}\) numerically. Figure 3 gives results using both methods.

For any beneficial plasmid, we can now determine the eventual population share that a mutant will have after one selective sweep, based on its initial fitness advantage, \(\Delta f\), its propensity to conjugate and share plasmids, \(c_m\), and the rate at which the (potentially mixed) wild type population shares plasmids \(c_w\).

Evolution however, particularly of plasmids, does not consist of one selective sweep, but of many. This leads naturally to the question of ‘how do conjugation suppressing genes fare over many selective sweeps?’.

8
3 Evolutionary Trajectory

Let us now consider how the balance of chromosomal genes evolves over many selective sweeps. Let $s_i$ refer to the fraction of the population containing $S$-type genes at the start of the $i^{th}$ selective sweep. Each time a beneficial plasmid arrives (through mutation or immigration), it is acquired by an individual, ‘the mutant’, which will be secretive with probability $s_i$ and generous with probability $1 - s_i$. If the mutant is secretive, then by the end of the evolutionary sweep, secretive individuals will occupy $s_{i+1} = M_\infty + (1 - M_\infty)s_i$ of the population. In contrast, if a generous individual is chosen the final population will have $s_{i+1} = (1 - M_\infty)s_i$ - note that $M_\infty$ in each case will not be identical, as $M_\infty$ is calculated using (amongst other things) the conjugation rate of the mutant (see eq. 19).

Figure 5 shows lineages with low conjugation rate taking over a population; this take over is initially very slow (taking thousands of evolutionary sweeps), accelerating as secretive genes take over and overall conjugation rate is reduced.

![Figure 5](image)

Figure 5: With each selective sweep the balance of the population changes. Over many sweeps, this leads to the fixation of lineages which are less inclined to share plasmids. However, for large population sizes (Left) this generally takes a very very long time, indicating that this effect is probably entirely overpowered compared to the many many other effects influencing evolution. (Right) we repeat the same, but with smaller population size/ larger $M(0)$, so as to make the individual jumps more visible.

In a population with a generally high rate of conjugation ($c_w \approx \Delta f$), almost any non zero $c_w$ will quickly lead to plasmids spreading through the population, hence the advantage enjoyed by secretive lineages is minor. If conjugation rates in a population are already low secretive chromosomal genes enjoy a far greater advantage, reaching high population density after only a few selective sweeps. Regardless of the exact parameters however, given sufficient time, secrecy is favored.
4 An Agent based model

The simplified evolutionary dynamics considered in the section indicate that chromosomal genes which repress plasmid spread gain an evolutionary advantage (however slight) over long evolutionary time frames. While useful for the sake of analytic tractability and building intuition, this simplistic model ignores many important features, several of which may contribute to stabilizing plasmid sharing. In order to test the robustness of our previous findings, let us now consider a more detailed “agent based” model. We will start by constructing a ‘baseline’ model, and then will consider a number of extensions.

Consider a population of bacteria on an \(N\) by \(N\) grid with periodic boundary conditions. Each grid cell contains a single bacteria which is either generous (\(G\)), secretive (\(S\)) or dead (\(X\)). In nature, plasmids exist in a number of ‘incompatibility groups’ \([56, 12, 49]\) - plasmids belonging to the same incompatibility group make use of the same regulatory proteins, and thus interfere with one another’s reproduction, driving one another to extinction whenever they exist within the same cell. Plasmids from different incompatibility groups co-exist with no such interference. In our model, we assume that each living bacteria possess one of \(k\) possible plasmids from each of \(q\) incompatibility groups. If there are (for example) \(q = 3\) incompatibility groups, the state of a single grid cell is given by a vector \(v\) of the form \(v = \{G, 2, 5, 3\}\).

Individual cells die at some death rate \(d(v, t)\), and reproduce at some reproduction rate \(r(v, t)\), both of which depend on the plasmids contained within the cell, and the current time. Dying cells will transition to the state \(v = \{X, -, -, -\}\). Reproducing cells will select a direction, and duplicate their state into the adjacent cell in the chosen direction, assuming the target cell is empty. If the selected cell is occupied, replication fails; this gives a crude representation of competition.

Plasmid mutation within cells occurs at some low constant rate \(m = 10^{-5}\); cells undergoing mutation select a plasmid group randomly, from 1 to \(q\), and set its value to some random value from 1 to \(k\). This mutation process is included so as to prevent plasmid extinction and ensure a minimal level of variance in the population.

Finally, cells conjugate at a rate \(c(v)\); for the purposes of our current exploration, conjugation rate depends only on chromosomal genes, with default rates \(c(G) = 0.1, c(S) = 10^{-3}\) and \(c(X) = 0\). A conjugating cell selects a neighboring gridcell uniformly at random, and replaces one of the neighbors plasmids with its own value. If the selected gridcell is empty, conjugation fails. While it is well known that many plasmids possess surface exclusion mechanisms in order to prevent invasion by incompatible plasmids \([17]\), these exclusion mechanisms are only partially effective \([16]\). For the purposes of our current investigation, we assume that \(c(v)\) is the conjugation rate after taking pre-existing plasmid exclusion mechanisms into account.

All events are assumed to be exponentially distributed, and we simulate the entire system using Gillespie’s Algorithm. Simulation code is available via Github \([\cdot]\).

A schematic illustration of this model is given in figure \([\cdot]\).
Figure 6: (Left) An N by N grid ($N = 5$) with periodic boundary conditions containing ‘generous’ type bacteria (blue cricles), secretive bacteria (green octagons) and empty space. In this example, each bacteria contains a plasmid belonging to $k = 2$ incompatibility groups (one group represented by triangles, the other by squares). Each plasmid comes in one of $q = 3$ different variants (variants distinguished by colour). (Right) The system evolves according to four mechanisms: death, birth, conjugation and mutation.

4.1 Base case

The agent based model described above is rather detailed. For the sake of concreteness, let us begin our investigation by considering the case where each bacteria contains only a single plasmid ($q = 1$), and this plasmid comes in one of $k = 4$ varieties. Assume $N = 64$. Each gridcell initially has a 54% chance of containing generous bacteria, 6% secretive bacteria and 40% empty space. The plasmid type for each bacteria in the initial population is selected uniformly at random.

At time $t = 0$, the simulation selects $d(x,t)$ and $r(x,t)$ uniformly at random in the ranges $(0.35 - 0.42)$ and $(1 - 1.3)$ respectively, for each plasmid state. At time $t = \tau = 50$, and every $\tau$ minutes thereafter, new death and replication rates are selected for each plasmid state. We refer to each such time window as an ‘epoch’ and refer to $\tau$ as the ‘epoch time’. This approach gives a rudimentary approximation of the manner in which plasmid fitness changes with variable environmental conditions.

Illustrations of the systems state throughout one simulation run are given in figure [7]. Simulation is allowed to run until either $t = 20,000$, or $G$ or $S$ is driven to extinction.

We run 100 simulations using the parameter values described. In 47 out of 100 simulations, despite their low initial population, secretive chro-
Figure 7: (Above), illustrations of the full system state at a variety of times. Pixel colours give the identity of the corresponding plasmid, with bright colored nodes being generous, and darker nodes secretive. (Graphs, top) Every $\tau$ time steps, new $r(v)$ and $d(v)$ values are selected, leading to different ‘fitness’ values $r(v)/d(v)$ for each of the four plasmid types. (Graphs, middle) The abundance of each plasmid strain rises and falls throughout the simulation. (Graphs, bottom) Here we graph the abundance of generous and secretive chromosomal genes (bright and dark, respectively). Over time, secrecy is favored.

mosomes reach fixation in the population, driving generous chromosomes to extinction. This rate is nearly five times higher than we would expect for a neutral mutation, which would reach fixation with 10% probability (as they represent 10% of the initial non-empty population). Generous chromosomes reach fixation in 53 of 100 simulations; all simulations complete before $t = 20,000$. In cases where secretive chromosomes reach fixation in the population, the average fixation time is $t_{fix} = 3024$ (60.5 epochs). The average time until secretive chromosomes reach extinction time is $t_{ext} = 1203$ (24 epochs). What these results indicate is that simple ‘network reciprocity’ (discussed later) is insufficient to stabilize conjugation, at least, not for the simple 2d lattice considered here.

4.2 Exploring parameter space, and environmental heterogeneity

With the base case now covered, let us now consider a range of different simulation conditions. Our aim in what follows is to determine the robustness of our previous results, and determine which aspects of the system
(if any) affect fixation probability of secretive genes. In all simulations, we assume \( N = 64 \), and initially gridcells are empty with probability 40% probability, generous with 54% probability and secretive with 6% probability. Table 1 summarizes the results of this section.

We start with two ‘sanity tests’ on our simulation. First, we consider a ‘control condition’, in which \( c(S) = c(G) = 0.1 \). In this neutral case we expect a fixation probability for \( S \) precisely equal to the initial population fraction, 66\%. In practice, we observe 12% fixation, 81% extinction, and 7% timeout (Scenario # B, table 1). This is consistent with what we would expect by chance. Next, in order to test sensitivity to simulation geometry, we reuse the same parameters as the base case simulation, but allow bacteria to conjugate and replicate diagonally, such that each cell is adjacent to 8 others rather than 4 (scenario C). No substantial change in fixation probability is observed; this is reassuring, as it indicates that our results are not overly sensitive to arbitrary choices made in the construction of the simulation.

With these basic tests out of the way, let us now explore variations of the more biologically meaningful parameters. We consider increasing/decreasing epoch times to \( \tau = 250 \) and \( \tau = 5 \) (corresponds to considering a more or less stable environment), decreasing/increasing conjugation rates to \( c(S) = 10^{-5} \) and \( c(S) = 10^{-2} \) (stricter and looser plasmid suppression, respectively), and setting the mutation rate a factor of 30 higher or lower. In all cases, fixation probability for \( S \)-type genes remains high (scenarios DEFGHI).

We also consider the case of either increasing death rate to \( d(v) = (0.75 - 0.90) \), dramatically reducing population density, or decreasing it to \( d(v) = (0.05 - 0.06) \), drastically increasing population density. In the former case, the fixation probability of \( S \) type chromosomes drops to 11% – close to what we would expect in the neutral case. This suggests that when populations are sparse enough to render plasmid conjugation rare and ineffective, plasmid suppressing genes have no evolutionary effect, either positive or negative (scenario J). In the low death rate/high density case, the most common result of simulations is timeout; no meaningful conclusions can be drawn in this parameter regime (scenario K).

If we simplify our model by assuming full population density, and that all death events are immediately followed by a corresponding birth event (treating our system as a death-birth Moran process on a graph \[38, 35\]) simulation speed can be significantly increased, and the timeout condition at \( t = 20,000 \) can be removed. In this case fixation occurs in 46% of cases, close to our base rate. When fixation does occur, the average time period is \( t \approx 44,000 \). Extinction occurs by \( t \approx 17,400 \) on average. Taken together, these simulations indicate that the advantage of secretive type chromosomal genes is exceptionally robust to changes in parameter values. If we desire a system that will stabilize sharing of plasmids, simple changes to parameter values are insufficient.

In natural settings, plasmids frequently contain resistance \[8\] and metabolic \[50\] genes adapted for specific environments. For this reason we might expect plasmids to have particular evolutionary importance in and around transition regions or boundaries between different environments \[23\]. In order to explore the effects of environmental variability, we extend our
basic model to one in which each gridcell is assigned an ‘environment’ parameter (either $A$ or $B$). Death and reproduction functions are replaced with environment specific functions $r_A(v), r_B(v), d_A(v), d_B(v)$; with each epoch, the rate functions associated with either $A$ or $B$ (but not both) are changed. In order to explore the effects of variable environment, we consider four different environmental geometries: block, checkerboard, random and gradient (see figure 8). Death and replication rates in the gradient geometry are given as position dependent linear combinations of those found in $A$ and $B$ (hence $r(v) = xr_A(v) + (1 - x)r_B(v)$ for some position dependent $x$ in the range $[0,1]$. Regardless of the environmental condition considered, fixation is observed to occur in roughly $40 - 50\%$ of all cases (scenarios KLMN). Hence, it would appear that environmental heterogeneity is also insufficient to stabilize plasmid sharing.

![Figure 8: Four possible environmental geometries; from left to right we have ‘block’, ‘checkerboard’, ‘random’ and ‘gradient’ type geometries. For each block, checkerboard and random conditions, cells experience one of two death and reproduction rates ($r_a(v)$ or $r_b(v)$) depending on the color of their gridcell. This allows us to simulate the behavior of cells around boundary regions (of various shapes). Block geometry assigns environment $A$ to all cells in the left half of the simulation and environment $B$ to the right half. Checkerboard assigns opposite environment conditions to adjacent cells. In the random condition each gridcell has a $50\%$ chance of having each environment type, each gridcell is determined independently. In the gradient condition, $r(v)$ and $d(v)$ are a linear combinations of the death and reproductive rates that would be experienced in ‘pure’ environments.

5 Relation to past paradoxes

The paradox posed so far by the maintenance of conjugation is by no means the first paradox in the study of evolutionary dynamics, nor is it likely to be the last. Two paradoxes of the past, namely, the evolution of sex, and the evolution of altruism (also called the paradox of cooperation), bare striking resemblance to the current conundrum. Let us take a brief detour to examine each of these paradoxes, their resolutions, and the similarities and contrasts to the question currently under study.
5.1 Can recombination preserve HGT?

In the study of the evolution of sex, simple modelling suggests that asexual mutants would possess two major advantages over variants that reproduce sexually. Firstly, all members of the asexual species are able to reproduce, in contrast, for sexual species, only the female population can reproduce (the cost of males) [37]. At the individual level, asexual individuals are able to pass on 100% of their genes to each offspring, while sexual individuals pass on only 50% of their genes (the cost of meiosis) [62, 63]. Classically, this paradox is resolved by considering ‘the evolution of evolvability’ [28]. For asexual species, if beneficial mutants arise independently in two different lineages, one will inevitably drive the other to extinction, a process known as clonal interference. Clonal interference severely limits the speed of evolution, especially for large population size [18]. In contrast, sexual recombination causes beneficial genes to accumulate. Mating allows genes initially present in separate lineages to come together in a single organism, both advantages can be retained, and the rate of evolution scales with population size [15, 58].

Similar to sex, HGT allows genes to accumulate, and recombine. In contrast to sex, in which all genes have (approximately) equal chance of transfer and recombination, for bacterial conjugation, plasmid bound genes and other mobile gene elements gain the benefits of recombination, while core chromosomal genes do not. This leads to the question; are the benefits of recombination enough to stabilize plasmid sharing in a bacterial population? In order to examine this we consider a collection of cells each containing \( q = 3 \) plasmids, each from a different incompatibility group. For this simulation, we assume that individual conjugation events transport only a single plasmid, and thus, at the boundary between two clonal lineages, conjugation will quickly lead to a wide variety of different plasmid combinations, if cells are generous. This effect will be significantly reduced for secretive variants, leading to increased clonal interference.

In order to study the multi-plasmid case, we must first determine how \( r(v) \) is defined as a function of the three plasmid values \( v_1, v_2, v_3 \). We consider three possible cases, in order of increasing complexity: in the first case (scenario Q), we assume that \( r(v) \) for each possible plasmid combination is selected in the range \( (1 - 1.3) \) entirely independently, and that new \( r(v) \) are selected in each epoch. The fitness of the combination \([G, 1, 1, 1]\) and that of \([G, 1, 1, 2]\) are entirely independent of one another. One can think of this as being a ‘complex’ genospace, the value of each plasmid varies based on the presence and absence of other plasmids. In the second case (scenario R), \( r(v) \) is formed as a linear combination \( r(v) = 1 + 0.1r_1(v_1) + 0.1r_2(v_2) + 0.1r_3(v_3) \). In this case \( r_i(v_i) \) are selected in the range \((0, 1)\). Each plasmid contributes to fecundity independently, and bacteria can succeed by ‘optimizing’ for each plasmid incompatibility group independently. Finally, we consider the hybrid case, in which half of the fitness is determined via each of the previous two methods (scenario S). In order to observe the benefits of recombination and the cost clonal interference, mutants must arrive frequently enough such that multiple mutant strains are competing at any one time. In order to achieve this, we assume \( N = 128 \) and \( m = 10^{-4} \) for this batch of simulations.
In all three cases we observed significantly more fixation events than the control case (Scenario B), 45%, 47% and 51%, respectively. Fixation and extinction times are comparable to the base case (Scenario A). Taken together these results indicate that recombination is also insufficient to maintain plasmid sharing (at least at the scale simulated here). The evolutionary pressure on the static chromosomal genes is too great.

5.2 Is HGT a public good?

If the advantages of recombination are not enough to stabilize plasmid sharing, let us turn our attention to another evolutionary paradox, and its resolutions. The second major paradox in the study of evolutionary dynamics is the emergence and maintenance of cooperation: slime molds cooperate in order to form stalks and bud \[45\], wolves hunt in packs \[10\], meerkats keep watch, and humans cooperate on vast and complex scales spanning the entire globe \[60\] and beyond \[2\]. Cooperation is powerful, and yet at every turn, it is the organism that most benefits *its own* lineage that evolution will select. The lazy wolf, the cowardly meerkat, the spore that forces itself into the fruiting body rather than the stalk. Evolution is a game with winners and losers, not one that played for fun. Given the ever present advantage of selfish behavior, how then is cooperation maintained? - this is the question posed in the study of cooperation and public goods.

Numerous answers to this question have been proposed \[41\], and at it would seem possible that several of the mechanisms that stabilize cooperation might also stabilize HGT. Much like cooperative and altruistic behavior, HGT involves one individual paying some ‘cost’ for the benefit of another; how this benefit is distributed between the receiving cell and the plasmid itself is unknown, and likely to vary wildly between contexts. Unlike classical public goods interactions however, HGT involves the transfer not of resources, of *information*, a collection of genes that will potentially benefit the recipient not only in the immediate future, but for generations to come. When estimating the ‘value’ of this gift, it is unclear what time horizon is appropriate.

With this differences and similarities in mind, let us now consider a variety of mechanisms that have been proposed for stabilizing cooperation, and how they might apply in the context of HGT. We note that there remains some debate in the literature as to the level of overlap between these mechanisms \[34, 5\]. Here, we make no claims as to the distinctness or similarities between alternative evolutionary mechanisms, and instead simply err on the side of inclusivity whenever doing so appears physically appropriate in the bacterial context.

The most well known and well understood mechanism for stabilization of cooperation of evolutionary time scales is kin selection \[19\]. Kin selection suggests that the benefits of cooperation are disproportionately directed at ones relatives. Cooperative genes thus persist because they inevitably end up helping *other copies* of those same cooperative genes. This is in some sense similar to parental care, albeit on a wider scale. In the context of HGT, kin selection plays a strange role: those individuals most closely related to a plasmid donor are also those most likely to *already*
posses a given plasmid, and hence gain no benefit. However, if a related individual does not posses a plasmid that the donor bacteria is currently benefiting from, they are also more likely than other bacteria to benefit, due to the plasmid and their chromosome being more closely adapted to one another [14, 51]. The kin selection hypothesis is further hampered in the bacterial context, as there is little evidence that bacteria are able to track their kin (beyond spatial proximity), unlike vertebrate animals. Given that our previous simulations already include a spatial component, it would appear kin selection via spatial associativity is ineffective.

A second resolution that has been proposed to the paradox of cooperation, is reciprocity; that is to say, an individual who pays a cost today may well be on the receiving end of generosity tomorrow. Reciprocity comes in a variety of forms. Direct reciprocity involves two parties directly benefiting one another, such as plants supplying root fungi with sugars, in exchange for key nutrients [31, 25]. Indirect reciprocity takes place when individuals provide benefit too, and gain benefit from a wider community; I am willing to help you because I trust that someone else will help me [22, 43]. Finally, ‘Network’ reciprocity involves repeated interactions between neighboring individuals; these individuals are also likely to be related to the donor. Neither direct, indirect nor network reciprocity would appear relevant in the context of bacterial conjugation. To the best of our current knowledge, bacteria are incapable of tracking the complex reputational networks required for indirect reciprocity, nor are they selective in who they donate to. Because export of plasmids depends on a variety of protein complexes within the donating cell [54], plasmid donation is an explicitly ‘single directional’ process, and hence direct reciprocity would also appear unlikely. Network reciprocity is already baked into the agent based model we are using, and has proven ineffective at preventing the invasion of secretive chromosomal genes.

One final resolution to the paradox of cooperation is the hypothesis of ‘group selection’. Under this model, individuals are separated into $M$ groups, and while selfish genes have the advantage on an individual level within each group, groups with a greater proportion of cooperative individuals are more likely to spread and divide than groups containing more selfish individuals. Unlike reciprocity and kin selection, this hypothesis is particularly suited to the microscopic world of bacteria. In order to test group selection in the context of HGT, we consider an alternative simulation, in which each column of the grid is considered one ‘group’, and individuals may reproduce and conjugate to any cell within their column. Group selection can be implemented in a variety of different ways; for a classical review, see Wade 1978 [61]). For the sake of this initial exploration, we mimic Traulsen and Nowak’s minimal model [58]; cells duplicate within a group according to their fitness, expanding into the free space available, until eventually the group reaches size $N$. At this stage, with each duplication event, the group will either split with probability $w$, or a randomly selected group member will perish with probability $1-w$. When a group splits, one of the other groups is selected at random and is eradicated, and each member of the splitting group either migrates to the new space, or stays put (50% probability of each outcome). Unlike previous simulation scenarios, death is not explicitly modeled, and is treated
as a downstream consequence of either group splitting or individual replication. Conjugation is allowed to occur within groups, but not between groups; for the sake of conjugation, all groups are considered well mixed.

In terms of simulation, we consider two separate scenarios. In both cases, we assume $w = 10^{-4}$ (rare group splitting). For scenario U, we assume initial conditions 54% generous, 6% secretive (as previously). This results in generous chromosomes overtaking the population 91% of the time, and secretive chromosomes reaching fixation 9% of the time, a result in line with neutral selection. In order to investigate this further, we also consider the same simulation with ‘balanced’ initial conditions with initial populations of both generous and secretive chromosomes at 30% (scenario V). Once again we find results consistent with neutral selection. When plasmids are unable to transmit between groups, the advantage of hoarding them is neutralized, and fixation probability reflects initial population fractions almost perfectly.

As is clear from these experiments, the selective advantage of conjugation suppressing genes is robust across a wide range of parameter regimes, including regimes that have previously been found to stabilize cooperation or sexual reproduction. Fixation probability is only reduced back to the 10% probability expected in for neutral evolution in circumstances where plasmid transmission is in some sense disabled (group isolation, sparse populations). Generous chromosomes are not found to have an evolutionary advantage for any of the scenarios considered.

We also observe that once fixation of secretive chromosomes is likely, fixation probability itself is relatively unaffected by parameter values. The general uniformity in fixation probability across scenarios would appear to indicate that extinction probability is governed almost entirely by local processes, and that larger scale and longer time frame effects (environment, mutation, epoch time), have little to no effect on the local dynamics. Given that extinction (when it occurs) happens 2-3 times faster than fixation, this may be indicative that fixation dynamics of secretive chromosomes are governed primarily by the probability of ‘early extinction’, and that once early extinction is avoided, fixation occurs with high probability. The one exception to this ‘uniform fixation probability’ is scenario F, in which we consider $c(S) = 10^{-2}$, that is to say significantly weakened HGT suppression reduces the evolutionary advantage of. As might be expected in this case fixation of secretive genes is reduced. See table 1 for a summary of these results.

6 Conclusion

Horizontal gene transfer amongst micro-organisms is a significant contributor to the vast complexity and variety of life we see in the world today. It allows for the sharing of resistance[53, 5] novel metabolic pathways[50], and virulence factors[50], and is critical both to our understanding of life, and our forays into medicine. HGT is also something of a mystery, complicating our understanding of the evolutionary ‘tree’. Plasmids have been described as ‘paradoxical’ in the literature [22], with Harrison and Brockhurst observing that costly plasmids should be purged via purifying
| Scenario | Simulation description | Prob. fix. | Prob. ext. | mean $t_{fix}$ | mean $t_{ext}$ |
|----------|------------------------|------------|------------|---------------|---------------|
| A        | Base case              | 41%        | 53%        | 3024          | 1203          |
| B        | Control condition, \(c(S) = c(G) = 0.1\). | 04%        | 87%        | 17182         | 3062          |
| C        | Diagonal travel        | 34%        | 66%        | 2327          | 952           |
| D        | Epoch time \(\tau = 5\) | 47%        | 53%        | 4776          | 2216          |
| E        | Epoch time \(\tau = 250\) | 44%        | 55%        | 4488          | 2007          |
| F        | Weak Plasmid Restriction \(c(S) = 10^{-2}\) | 32%        | 68%        | 4314          | 1507          |
| G        | Strong Plasmid Restriction \(c(S) = 10^{-5}\) | 46%        | 51%        | 4299          | 1778          |
| H        | High Mutation rate \((\times 30)\) | 45%        | 55%        | 4878          | 1816          |
| I        | Low mutation rate \((\div 30)\) | 43%        | 56%        | 4081          | 2470          |
| J        | Sparse population      | 11%        | 89%        | 1595          | 440           |
| K        | Dense population       | 0%         | 42%        | -             | 11687         |
| L        | Dense population (Death-birth process) | 46%        | 54%        | 44433         | 17400         |
| M        | Environment: 2 blocks  | 41%        | 58%        | 4826          | 2565          |
| N        | Environment: Checkerboard | 43%        | 56%        | 4836          | 1555          |
| O        | Environment: Random    | 47%        | 51%        | 4462          | 2021          |
| P        | Environment: Gradient  | 41%        | 59%        | 4205          | 1791          |
| Q        | Multiplasmid: linear sum \(f = f_a(v_a) + f_b(v_b) + f_c(v_c)\) | 45%        | 49%        | 4600          | 2007          |
| R        | Multiplasmid: Independent \(f = f_{abc}(v_a, v_b, v_c)\) | 51%        | 46%        | 4630          | 1540          |
| S        | Multiplasmid: Hybrid case \(f = f_{abc}(v) + f_a(v_a) + f_b(v_b) + f_c(v_c)\) | 44%        | 51%        | 4452          | 1810          |
| T        | Multiplasmid: \(q = 3, k = 25\) | 47%        | 52%        | 3106          | 1213          |
| U        | Group selection.       | 9%         | 91%        | 7721          | 2438          |
| V        | Group section, balanced initial conditions | 52%        | 48%        | 5400          | 5850          |

Table 1: Table of results of simulations. For each scenario, we give a brief description, fixation and extinction probability (of secretive chromosomal genes), and the mean time to fixation or extinction. By default, initial population densities are 6% secretive bacteria, 54% generous (with the exception of scenario V, where we have 30%, 30%. Because not all simulations reach fixation before time out, probabilities do not add to 100%. Aside from scenario L have a time cut off of \(t = 20,000\).
selection, while beneficial plasmid genes would be expected to integrate 
into the chromosome, rendering the plasmid bound copy redundant. In 
this work, we present another paradox, namely the question of why the 
sharing of plasmids itself is not selected against.

Many past mathematical models have explored the dynamics of HGT 
and plasmid conjugation from the point of view of the plasmids themselves[52, 
55, 20, 36, 21, 4, 24]. However, plasmids are only one half of the evolu-
tionary story. In this work, we have explored a number of models of HGT, 
foocusing instead on the evolutionary dynamics of chromosomal DNA. We 
are able to show that for a wide array of modelling assumptions and 
parameter values, chromosomal genes that restrict bacterial conjugation 
rates have an evolutionary advantage. Even when deliberately construct-
ing models which have classically been shown to favor cooperation, we 
find that selection for generous chromosomal genes is at best neutral, and 
in the vast majority of cases, selected against. With each change of envi-
ronment or selective sweep, bacteria which horde advantageous plasmids 
to themselves fare better than those which share freely; this occurs despite 
ignoring the non-trivial costs [9, 48, 26] of conjugation.

So where does this leave us? Horizontal gene transfer and conjuga-
tion are ubiquitous in real world biology. As always, where modelling and 
observation disagree, it is the model which is lacking. There is no point 
shaking our fists at nature for disobeying the equations. Rather, our hope 
in this paper is to point out that bacterial generosity in the sharing of 
plasmids can not be taken for granted, especially when viewed from the 
point of view of individual chromosomal genes, which may benefit signif-
icantly from restricting plasmid spread. Regulatory mechanisms control-
lung bacterial conjugation serve two masters, and it is easy to produce 
circumstances where chromosomal and plasmid bound genes experience 
very different evolutionary pressures. Unlike classical ‘public goods’ in-
teractions, where the good being shared is a transient physical resource, 
the sharing of plasmids involves the transfer of information, information 
which provides benefits over an evolutionary timescale. Unlike the clas-
sical question of sexual recombination, plasmid conjugation breaks any 
notion of symmetry between genes, partitioning the genome into mobile 
genes that benefit disproportionately from recombination, and immobile 
genes, which do not.

In terms of resolving this paradox, the work here has been exploratory 
rather than exhaustive, and there remain a significant number of avenues 
to explore. Here we study only the evolution of chromosomal genes, and 
ignore plasmid-chromosome co-evolution; it is entirely plausible that fast 
evolution of plasmids may be enough to escape any ‘regulatory restriction’ 
chromosomal genes might impose. The cost of restricting conjugation it-
self may be high enough such that any such genes will be quickly purged 
from the ‘core genome’ of a species as non-essential. When plasmids code 
for genes promoting public goods (such as siderophore production [33]) 
then sharing of plasmids may be directly beneficial to the donating cell. 
What concentration of mutualistic plasmid genes would be needed to 
stabilize HGT via this mechanism is as yet unknown. Alternative mod-
els of group selection may yet demonstrate the evolutionary advantage 
of generosity, even if the models considered here do not, or alternatively,
justification may be found for reconsidering kin selection, or some other hypothesis that we have discounted as improbable in the current context. Regardless of the exact models considered by future researchers, it is our hope to encourage discussion, and draw attention to the largely neglected role of chromosomal genes in the study of bacteria conjugation. As demonstrated by those smallest of creatures that fill the world around us, with free sharing and recombination of information, answers to near any problem can be found.

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