The dynamics of gene duplication and transposons in microbial genomes following a sudden environmental change

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A variety of genome transformations can occur as a microbial population adapts to a large environmental change. In particular, genomic surveys indicate that, following the transition to an obligate, host-dependent symbiont, the density of transposons first rises, then subsequently declines over evolutionary time. Here, we show that these observations can be accounted for by a class of generic stochastic models for the evolution of genomes in the presence of continuous selection and gene duplication. The models use a fitness function that allows for partial contributions from multiple gene copies, is an increasing but bounded function of copy number, and is optimal for one fully adapted gene copy. We use Monte Carlo simulation to show that the dynamics result in an initial rise in gene copy number followed by a subsequent fall-off due to adaptation to the new environmental parameters. These results are robust for reasonable gene duplication and mutation parameters when adapting to a novel target sequence. Our model provides a generic explanation for the dynamics of microbial transposon density following a large environmental changes such as host restriction.

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I. INTRODUCTION

Biological evolution involves a variety of genome transformations, which include point mutation, homologous recombination, gene duplication, and horizontal gene transfer [1,3], and which involve a variety of mobile genetic elements [4,6]. Understanding the influence that each of these mechanisms and elements exerts on the process of evolution is one of the current frontiers in biology [7]. Transposons are one such genetic element—capable of copying and pasting segments from one location to another within a genome, they provide vehicles for gene duplication [8,9].

Transposons are copied and inserted across genomes through a variety of mechanisms [8,9]. Non-conservative transposons multiply within a genome by replicating themselves elsewhere. Many code for the proteins that copy and insert themselves throughout the genome. Other transposable elements are more passive, relying on proteins from other transposons in order to proliferate.

Insertion sequences, or IS elements, are a particular type of non-conservative transposon, which contain their own proteins for replication but typically do not contain any additional proteins [10,11]. However, when two IS elements are near each other along a genome—for example, flanking both sides of a gene—they may form a composite transposon that includes the host genes sandwiched between the IS elements [8,10]. These composite transposons have been associated with the evolution of virulence factors that affect infection and severity of diseases [12,13], and have been implicated in large chromosomal rearrangements [14].

Due to the conserved tendency of the genes required to replicate and insert an IS element, it is possible to estimate the IS element density within a genome through sequence analysis. In order to probe the evolutionary dynamics of microbial genomes, Moran and Plague estimated the IS density in Bacteria following host-restriction, the transition to becoming an obligate, host-dependent, organism such as a gastrointestinal symbiont [15]. They found that on average initial host restriction was followed by a sharp increase in IS density but that at long times the IS density generally declined to near zero. This overall pattern has been more directly validated in a few taxonomic lineages by tracking the evolution of particular closely related genomes [16,17]. This pattern may be sufficiently widespread, and if so, it should have a generic explanation, independent of the particular organisms or environment, and this is what we seek to provide in the present article.

In order to explain the trend in IS density following host-restriction, we focus on the role of IS elements as vectors for gene amplification through their roles in composite transposons [8,10]. Several studies have indicated the importance of gene amplification in the rapid evolution of Bacteria [18,19]. Duplicated genes provide a basis for the evolution of novel function [20,21], and have been implicated in the evolution of new organismal forms [22] and lineages [23]. Gene duplication events have been invoked in medically important traits and diseases in humans [24], including various forms of cancer [25,26], as well as in the expansion of gene families such as the globins [27] and the DNA replication processivity complex subunits [28]. Even whole genome duplications have been observed in many organisms including yeast [29].
small flowering plants [30], and pufferfish [31, 32].

If gene duplications confer an adaptive advantage to their hosts, then one might expect a concomitant proliferation in the IS elements that provide the mechanism for gene duplication—especially when adapting to a new environment. Conversely, in relatively consistent environments gene duplications confer no advantage to a well-adapted organism and we might anticipate selection pressure for a decrease in IS density. Thus, our strategy for interpreting the genomic trends reported by Moran and Plague [15] is to better understand the evolutionary dynamics of gene duplication, and thence to infer the corresponding dynamics of the IS elements.

In order to probe the link between gene duplication and adaptation to a novel environment, we model genome dynamics with gene duplication. Previous evolutionary modeling has largely focused on the process of mutation and recombination [33, 34], large scale genome duplications [35, 36], or static features such as copy number distribution [37]. Here we quantify a continuous selection mechanism for the evolution of novel genes put forth by Bergthorsson et al. [21]. In our model, we consider a protein encoded by a gene that has multiple activities—for example, enzymatic activity or non-specific binding. As shown schematically in Fig. 1, the initially deleterious effect of a gene product may subsequently become beneficial to the organism when exposed to different environmental conditions, as documented, for example, in the growing body of literature on non-specific interactions involving proteins [38].

There are then two mechanisms by which an organism may increase a particular protein activity—efficiency and expression. In the first mechanism, a gene may undergo mutations that result in a more effective protein, i.e., a gene with some small activity which is favorable to the organism can undergo mutations that subsequently enhance that activity. In the second mechanism, increasing the level of gene expression—for example, by creating additional copies of a gene within the genome—also positively impacts the total activity of a given gene product.

The purpose of this work is to show that adaptation to a large environmental change provides a sufficient explanation for both the short term increase and the long term decrease in IS density following host restriction—as outlined in Fig. 2. The rapid adaptation to a new environment results in a large number of gene duplications in adapting novel functions, these changed conditions result in an enhancement of the IS element density within the genome. Free-living bacteria experience a fluctuating environment, which results in the maintenance of IS density. Host-restriction represents a large change in environmental conditions, resulting in an initial boom in IS density, such as that seen in the recent obligate organisms. However, a host also represents a stable and consistent environment. The lack of any need for rapid adaptation leads to near-zero IS density by a process of slow decay.

This paper is organized as follows. In Section II, we present our quantitative model of Bergthorsson et al.’s
proposed mechanism for the emergence of novel genes under continuous selection [21]. Results of our simulations are presented in section [II]. Finally, in section [V], we describe the biological interpretation of our work, showing how our results are consistent with the trends observed by Moran and Plague [15].

II. MODEL OF CONTINUOUS SELECTION WITH GENE DUPLICATION

We consider a population of replicating cells whose replication rates depend on the genes within their genomes. We express the probability of the kth cell replicating according to a fitness function \( F \).

\[
F_k = \min \left( \sum_{j=1}^{n} g_j, 1 \right) - n\mu + \eta_d
\]  

(1)

where \( n \) represents the number of total genes in the genome, \( \mu \) denotes the fitness penalty arising from each additional gene, and \( g_j \) denotes the positive fitness contribution of the \( j \)th gene within the genome. The parameter \( \eta \) represents an assumed additive Gaussian noise with standard deviation \( d \). Introducing additive noise allows us to vary the strength with which the organismal fitness is coupled to the genome, allowing us to take into account the effect of environmental fluctuations as well as mutation in the parts of the genome that are unrepresented in this simple model. Note that the decoupling noise we have introduced here differs from the demographic noise intrinsic to a population [39]. In section [VII] we will discuss the functional form of the noise in more detail. The value of \( g \) depends inversely on the Hamming distance \( d \) between the gene sequence \( S_i \) and some target sequence \( T_j \), i.e.,

\[
g_j = 1 - \frac{d(S_j, T_j)}{N} = 1 - \frac{\sum_{i=1}^{N} \delta(S_{ji}, T_i)}{N}
\]  

(2)

where \( N \) represents the number of letters in each sequence (both \( S_i \) and \( T_j \), \( d(S_i, T) \) represents the Hamming distance between \( S_i \) and \( T_j \), and \( S_{ji} \) and \( T_i \) represent the \( i \)th letter of \( S_i \) and \( T_j \), respectively.

These parameters are intended to caricature a biological process whereby a novel beneficial functionality captured by target sequence \( T \) would provide an overall improvement in fitness given by \( g_j(S_i = T_j) = 1 \). Partially matching sequences also provide some of the catalytic activity necessary for the new function, yielding a partial benefit in proportion to the homology between the gene sequence \( S_i \) and the target sequence \( T_j \), in accordance with the continuous selection model of Berghorsson et al. [21]. The total benefit of a set of genes must also be less than the maximum benefit arising from a single gene. Therefore, each gene has a fixed cost \( \mu \) that represents the deleterious effect of non-specific interactions of the product coded for by a gene. For simplicity, we assume that each gene in the genome is expressed equally without regard to the more detailed considerations of gene regulation. This simplification has its basis in recent work that has indicated that gene copy number is positively correlated with gene expression level [40].

The scheme outlined above clearly sets the optimum at a the one gene solution, with the gene matches the target sequences. This assumption comes from the biological considerations of the need to minimize the deleterious non-specific interactions while reaching a certain level of functional activity for a given biomolecule. Note that while the optimum is chosen by design, the behavior of the system as it evolves in time is not. We do not a priori know whether the optimum behavior will be to duplicate or to not duplicate. After all, the long term advantage is for the single gene case, and it might be superfluous to duplicate genes only the then try to reduce their number. Furthermore, if we introduce high gene duplication rates, will competition suffice to overcome the duplication rate and drive the reduction in the number of genes? Or will the behavior of the system depend on the careful tuning of these parameters? Constructing a simple model enables us to answer these questions and to probe the viability of the scheme we outline in Fig. 2.

Genes are allowed to evolve by spontaneous point mutation, internal duplications, or deletions. Spontaneous mutation occurs by replacing a letter at a particular position within a gene, chosen at random from a uniform distribution, and replacing that letter with a randomly chosen one from an alphabet of size \( c \). Gene duplications and deletions, just as with spontaneous mutations, occur on randomly chosen genes within the population. Duplications are modeled as insertion events, and do not overwrite existing genes. Fig. 3 outlines the processes that lead to gene duplication and deletion.

Initially, all organisms contain a single identical copy of a randomly chosen gene. At each time \( t \), the fitness of a random cell \( k \) is selected out of a fixed population size and its fitness \( F_k \) from Eq. 1 is measured. This number then defines the probability that this cell will replicate at time \( t \). If it is decided that the organism replicates, it then overwrites a different randomly chosen organism in the population—the so-called roulette scheme [41]. Thus, organisms are on average being selected for higher replication rates (defined by \( F \)). One generation is defined as the time it takes for half the total population to undergo a growth attempt. A random update scheme governs which organisms will attempt to replicate.

III. SIMULATION RESULTS

We characterize the model of continuous selection with
gene duplication described in Section II and contrast it with a model of continuous selection with no gene duplication. In order to do so, we assign a randomly chosen target sequence $T$ and a randomly chosen gene sequence that is initially fixed in the population (i.e., zero diversity at $t = 0$). We then allow the system to evolve under a selection pressure described by Eq. (1).

As shown in Fig. 4, the average number of genes in a genome $\langle n \rangle$ increases sharply at the onset through gene duplication. As time passes and the individual genes become better adapted toward the target sequence, the number of genes then begins to decrease. In the long time limit, the gene number seemingly asymptotes to a small number greater than 1.

These changes in gene number are accompanied by changes in the individual gene scores or gene fitness $g$. Fig. 5 plots the average organismal fitness $\langle F \rangle$ as a function of time and confirms that gene duplication is enhancing the initial rate of adaptation (fitness increase) by providing a means for the organism to amplify the benefit of a gene.

In contrast to the case of organismal fitness, average gene score or fitness $\langle g \rangle$ does not necessarily increase with gene number $N$. In principle, there is a tension between the greater mutation rate that larger copy number engenders and the lesser effect on fitness from an individual gene. By plotting the average gene score $\langle g \rangle$ in Fig. 6 we see that, on average, genes initially adapt faster toward the target sequence $T$ with gene duplication. It thus appears that the primary effect comes from mutation of the gene copies provided by gene duplication, which leads to additional diversity in comparison to the case without gene duplication.

Gene duplication enhances the effect of point mutation by amplifying its effect. While the effect of a point mutation on a single gene may relatively small when compared to the noise $\eta$, when that point mutation is duplicated numerous times so is its effect on the fitness of the organism. This can mean the difference between being a mutation that is effectively washed out by noise or strongly selected. Fig. 7 shows that the relative speed-up from gene duplication becomes more dramatic as the magnitude of the noise increases. Notice how the point at which the single gene case crosses over the gene duplication case shifts further and further to the right with increasing $d$.

These results support the proposed mechanism for evolution of novel proteins proposed by Bergthorsson et al [21]. In particular, the primary steps of gene duplication to enhance expression, followed by the slower mutation and selection of gene with better catalytic properties then the original, and finally reduction of gene copies all appear to have been captured by this simple model. Note that in the long time limits we tested, the average gene number $\langle n \rangle$ remained above unity, probably reflecting the fact that there is a low probability of simultaneous beneficial mutations that would allow for a favorable gene deletion. Also, the standard deviations in Figs. 4 5...
FIG. 5: (Color online). Average organismal fitness \( \langle F \rangle \) with and without gene duplication. With gene duplication the initial rate of adaptation is faster than in the single gene case. However, at long times the single gene case results in genes that are closer to the target sequence \( T \). Parameters are the same as given in Fig. [4].

and [6] hint at the important role of population variance in adaptation [42, 43]. In particular, notice that the variance in gene number and in gene fitness in Figs. [4] and [6] respectively, both play a role in the additional population variance seen in Fig. [5]. This results in a organismal fitness variance that is greater in the case with gene duplication than without despite a larger gene variance in the single gene case.

FIG. 6: (Color online). Average gene fitness \( \langle g \rangle \) with and without gene duplication for simulations with the fitness function given by Eq. [1]. With gene duplication the initial rate of gene adaptation is faster than in the single gene case. However, at longer times the single gene case results in genes that are closer to the target sequence \( T \). Parameters are the same as given in Fig. [4]. Note that the horizontal axis scale differs from that shown in Fig. [4].

FIG. 7: (Color online). Average gene fitness \( \langle g \rangle \) with and without gene duplication for noise parameter \( d = 0.0, 0.2, 0.4, \) and 0.6 (from top to bottom). With gene duplication the initial rate of gene adaptation is faster than in the single gene case. However, at longer times the single gene case results in genes that are closer to the target sequence \( T \). Other parameters are the same as given in Fig. [4].

IV. PARAMETER CHOICE

In order to interpret these simulation results in terms of their impact on biological evolution, it is important to consider how to map the parameters from simulation to those of real biology. A direct mapping where simulation models biological processes at biological rates would be one particular solution. However, the microbial world is one of very large numbers, making this approach unfavourable. Thus, it is often best to turn to other approaches such as scaling analysis in order to estimate how a system will behave at a parameter setting that is far from computationally tractable.

The parameters chosen for the simulations described above are from matching those of real biology. Realistically, a microbe contains around 1000 genes, and the rates of mutation \( 10^{-9} \) and gene duplication \( 1 \) given by Ref. [21] describe per organism rates per generation. Our simulation is intended to model a particular slice of the genome, representing how one particular gene within the cell might be subject to particular conditions and subsequently gene family expansion and contraction within a population. Thus, the biologically relevant regime would be a mutation rate of \( 10^{-12} \) with duplication rates of \( 10^{-3} \). This is fairly far from our choice of \( 10^{-2} \) and 1 for these two parameters, respectively. The rates of these basic parameters influence the timescales within the simulations. Thus we can expect that realistic parameter range simulations would require at least 10 orders of magnitude longer simulation times based on the mutation rates alone. Even then, these simulations are still cannot be considered realistic in light of microbial populations that easily number in the billions.
V. ALTERNATE FITNESS FUNCTIONS

The above model and results represent approximations to a possible biological mechanism for generation of novel gene functions. We have attempted to keep our model close to that of Bergthorsson et al.\cite{21} in order to understand the plausibility of the gene duplication mechanism they propose. Any study of plausibility, however, should also contain some measure of sensitivity analysis. The sensitivity of the model to specific choice of parameters is already discussed above. This same sensitivity check should also be applied to our modeling choices.

In order to probe this, we put forth the following fitness function $F^2$.

$$F^2_k = \min \left( \sum_{j=1}^{n} g_j^2, 1 \right) - n \mu + \eta_d\ (3)$$

Eq. (3) alters the linear dependence of individual gene fitnesses given by Eq. (2) to a squared dependence. This
TABLE I: Summary of number of simulation runs with fitness function $F^2$ that reach the target sequence. In order to compute this, the average gene fitness was taken at the end of each run (10,000 generations) and were considered to have evolved sufficiently toward the target sequence if $\langle g \rangle > 0.5$. The genomes are initialized identically, with each initial gene being chosen at random. As the initial number of genes grows, the probability of a single gene being of sufficient benefit to outweigh it’s cost increases. Thus, the probability of success with gene duplication increases with initial gene number accordingly. However, in the case without duplication, since these genomes can neither duplicate nor delete genes, the extra initial genes only provide an extra fitness cost. Over 99% of the cases were $\langle g \rangle > 0.9$ or $\langle g \rangle < 0.01$.

| Initial Gene Number | With Duplication | Without Duplication |
|---------------------|-----------------|-------------------|
| 1                   | 418             | 1000              |
| 2                   | 858             | 956               |
| 3                   | 991             | 977               |
| 4                   | 1000            | 902               |
| 5                   | 1000            | 888               |
| 10                  | 1000            | 0                 |

Particular fitness function was chosen because it as we increase the power to which the individual fitness $g$ is raised, we weaken the effect of the continuous selection proposed by Bergthorsson et al [21]. In other words, since squaring smaller numbers reduces their value by a greater fraction than for larger numbers closer to 1, we have are modeling a weaker initial catalytic “side-effect” than in Eq. [1]. Indeed, sometimes, the benefit of the initial genes outweighs their associated fitness costs, leading to a number of simulations that never improve in fitness. Table I shows the number of cases out of 1,000 that reach the target sequence by the end of the run.

Figs. 11, 12, and 13 show the behavior of the model under the fitness function given by Eq. [3]. All three plots show qualitatively similar properties to those given in section III for the fitness function in Eq. [1]. Specifically, the rise and fall in gene number in the case with gene duplication in Fig. 11 and the faster initial rises in organismal and gene fitness in Figs. 12 and 13 closely resemble the results of the previous model.

Increasing the power to which $g$ is raised in the fitness function to 3,

$$F^3_k = \min \left( \sum_{j=1}^{n} g_j^m, 1 \right) - n\mu + \eta_d$$

further reduces the effect of continuous selection, but again results in qualitatively similar plots, though with a larger fraction of cases with gene duplication that do not adapt toward the target gene (data not shown).

Ultimately, this reduction of effect continues as we raise the power of $g$ in the fitness function

$$F^m_k = \min \left( \sum_{j=1}^{n} g_j^m, 1 \right) - n\mu + \eta_d$$

until the benefit from an additional gene will only outweigh the cost for genes that exactly match the target sequence $T$. Retaining results that are qualitatively similar to those presented in Section III requires some form of continuous selection that allows for fitness contributions from multiple genes, supporting the hypothesis that these features play a key role in the evolution of novel gene function [21]. Removing either of these two attributes entirely nullifies the benefits of gene duplication (data not shown). However, our results do not depend very strongly on the strength of either of these features, and our findings outlined in Section III do not appear sensitive to the specific form of the fitness function chosen, as long as they fall within a class of bound fitness functions that allow multiple gene contributions and some form of continuous selection.

VI. DISCUSSION

Significantly, the results for the case with gene duplication outperformed the single gene case. Continuous selection, especially in the absence of noise, provides a means for rapid uphill adaptation. Given that selection
pressure on any one gene is weaker when there are multiple copies, adaptation of genes could have arguably been slower with gene duplication. We did not observe this to be the case and found that gene duplication has an initial advantage over the single gene case. Note that this is for the specific case where there is a large difference between initial and the target sequences and not true for the case where only smaller adjustments are required to reach the target sequence. Also, we did not observe monotonic gene duplication with a non-zero gene penalty. Beneficial genes were able to proliferate, eventually resulting in a reduction in the average number of genes per genome. That gene number rises and falls according to fitness, overriding the duplication and deletion rates for a broad range of parameters, shows us that the selective advantage, as we represent it here, is enough to overcome these intrinsic rates. If it were otherwise, we would not be able to posit gene amplification as the driving dynamic behind changes in IS density.

A. IS Density Following Host-Restriction

IS elements and gene duplications go hand-in-hand with one another. IS elements copy and paste segments of the genome from one location to another location and thus are vehicles of gene duplication [8, 9]. Enhancing gene duplication rates allows an organism to take better advantage of the mode of adaptation described in this work.

Host-restriction is defined as the process by which a previously free-living organism becomes an obligate organism, i.e., the transition from a more independent organism to an organism dependent on a host for survival. Genome comparisons among *Buchnera* indicate that this process involves an initial period of massive deletions, large scale rearrangement, and the proliferation of repetitive elements followed by extreme stability and a slow loss of additional genes [16]. The pattern of repetitive element proliferation is consistent with Fig. 4 in which we see an initial spike in IS density (given by gene number $g$) followed by a slow decrease.

Wide surveys of genomes reveal that organisms with recently formed obligate associations show an increased level of IS density in comparison to free-living organisms [15]. Conversely, ancient obligate organisms generally show a much lower IS density in comparison to free-living organisms [15]. In the context of our model, as IS elements proliferate, they grow in number and overall density within a genome. The pattern of boom and bust in IS density seen in the literature [15, 17] corresponds to a cycle of rapid adaptation to a new environment. The level of transposon density in ancient obligate organisms is then predicted by the long-time asymptotics of the simulations in Fig. 4. Note that we do not see a reduction from 2 genes to 1 on the timescales of our simulation due to the fact that this requires a combination of point mutations to occur in order to increase fitness, which then makes the $n = 2$ to $n = 1$ transition very rare and slow. On the other hand, this qualitatively matches the finding that only ancient obligate organisms (i.e., after long times) exhibit exceedingly low transpo-
to a random sequence at randomly selected intervals according to a 0.04 probability of change per generation. The curve for constant environment is comprised of the averaged data from Fig. 4 and is plotted here for comparison purposes. Other parameters are the same as given in Fig. 4. Notice that the number of genes in the changing environment are generally larger than in the case for a constant environment.

FIG. 14: (Color online). Gene number $\langle n \rangle$ with gene duplication under changing and constant environments. The fitness function is given by Eq. 1. Example trajectory for changing environments was plotted by changing the target sequence $T$ to a random sequence at randomly selected intervals according to a 0.04 probability of change per generation. The curve for constant environment is comprised of the averaged data from Fig. 4 and is plotted here for comparison purposes. Other parameters are the same as given in Fig. 4. Notice that the number of genes in the changing environment are generally larger than in the case for a constant environment.

B. The Role of Noise

The additive noise present in the fitness function given by Eq. 1 plays an important role in determining the relative advantage of gene amplification over single gene evolution. In other words, noise determines how advantageous gene duplication will be for a population. The greater the noise, the more difficult the evolutionary problem of finding the optima becomes. At the same time, these more difficult evolutionary problems are particularly suited for gene duplication.

Consider for a moment the role and source of noise. Noise represents the coupling between the genome and the organismal fitness—the more noise, the weaker the coupling, and vice-versa. In principle, noise can arise from several sources. For instance, environmental fluctuations may destroy an organism and kill without regard for the organismal phenotype. Conversely, it is possible that every organism has an almost equal probability of reproducing at any given timepoint (even though on average the fitter organism will still retain a fixed advantage given by the noiseless fitness function). Nonetheless, this change still dramatically alters the timescale on which selection acts. Noise is thus not a proxy of environmental harshness, but instead of the sensitivity of selection, or selectivity.

Selection favors phenotypes that grow, survive, and reproduce more prolifically than their neighbors. If small changes in genotype result in large changes to the survival and reproduction of the organism, then the coupling between genotype and selection can be said to have a stronger effect than noise. Note that the overall rates of reproduction do not matter, but instead, competition is the dominant factor. Selection pressure is not a proxy for the harshness of the environment. An environment that kills indiscriminately is just as selective as an environment that allows indiscriminate growth within a finite capacity.

We must now differentiate between the selectivity (or conversely, the noise) of the system from the average environmental conditions or directionality of selection. When the environment changes on longer time scales, the direction of selection changes. We say that an environment is relatively stable or consistent when the average environment remains essentially constant over time with small fluctuations. However, these environmental fluctuations are not the same as noise in our model, which represents small scale fluctuations that essentially randomize an organism’s probability of survival or reproduction. Thus, we regard the host-restriction that necessitates rapid adaptation to the host environment as different from the selectivity of the environment.

Indeed, in our model the noise or selectivity of the system remains an important, but separate, contributor. An alternate explanation for the boom in IS density following host-restriction can be seen in Fig. 7. As the noise parameter $d$ increases, the advantage of gene duplications increases. Thus, it is possible that simply by entering
a noisier environment one should see an increase in the number of IS elements. However, this explanation cannot account for the later decrease in IS density in the ancient obligate organisms.

VII. CONCLUSION

We have presented a model for the dynamics of a population of microbial genomes following a change in environmental conditions. Our results indicate the advantages of higher IS density in accelerating the process of adaptation to different environmental conditions, and the ensuing decrease in IS density during subsequent reestablishment of the environment. This corroborates evidence from observational bioinformatics [16] that indicates increased IS density following host-restriction—a large environmental change. Although our discussion has primarily focused on host-restriction, genomic surveys can track transposon number along environmental gradients, for example as a function of depth in the ocean [14] [15], and our results should be relevant to interpretation of these data [16].

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