Experimental Evolution as a High-Throughput Screen for Genetic Adaptations

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ABSTRACT Experimental evolution is a method in which populations of organisms, often microbes, are founded by one or more ancestors of known genotype and then propagated under controlled conditions to study the evolutionary process. These evolving populations are influenced by all population genetic forces, including selection, mutation, drift, and recombination, and the relative contributions of these forces may be seen as mysterious. Here, I describe why the outcomes of experimental evolution should be viewed with greater certainty because the force of selection typically dominates. Importantly, any mutant rising rapidly to high frequency in large populations must have acquired adaptive traits in the selective environment. Sequencing the genomes of these mutants can identify genes or pathways that contribute to an adaptation. I review the logic and simple mathematics why this evolve-and-resquence approach is a powerful way to find the mutations or mutation combinations that best increase fitness in any new environment.

KEYWORDS evolutionary biology, genomics, population genetics

Experimental evolution is a method that is gaining in popularity because of several inspiring successes and because high-throughput sequencing can reveal the genetic basis of the evolutionary process (1, 2). An underappreciated benefit of these studies is their ability to identify mutants in genetic pathways that underlie functions of interest. Thus, the experimental evolution of large populations of microbes can and should be viewed as a powerful genetic screen for adaptations. I outline why the processes underlying most evolution experiments are practically straightforward and why evolved mutants can provide valuable insight into the biology of the microbe under study and its response to environmental pressures.

Experimental evolution is not a new method, dating back to the 1870s when Dallinger conducted continuous culture experiments at steadily increasing temperatures (3). It also still underlies a large fraction of vaccine development, in which viruses are propagated in alternative hosts or cell lines to render them nonpathogenic but still immunogenic (4). Likely the best known is the Long-Term Evolution Experiment (LTEE) in which 12 Escherichia coli populations have been propagated daily for more than 30 years (5). Many remarkable studies of the LTEE have taught us a great deal about the evolutionary dynamics of adaptation and revealed numerous adaptive phenotypes, including gains in cell size, metabolic capacity, thermal tolerance, life history parameters, and above all, competitive fitness. Yet for the first 13 years of the LTEE, the genetic sources of adaptation remained unknown because the causative mutations did not affect the candidates chosen for Sanger sequencing. The first few mutations were discovered with higher-throughput screens of metabolism, expression microarrays, and by probes targeted to mobile insertion sequences (6–8). The power of each of these
Evolution experiments are influenced by all population genetic forces: selection, genetic drift, mutation, and recombination. In conducting a genetic screen, we can usually ignore the process of recombination, given that we are mostly interested in “first-step” adaptive mutants arising in different clones. Given that mutations occur inevitably, the relative balance of the remaining forces is governed by the effective population size, $N_e$, and the strength of selection, $s$, acting on these mutations (Table 1). $N_e$ is a property that is typically much less than the census population size ($N$) at any given time and is biased toward bottlenecks that occur during transfers. In practice, microbial populations are transferred at sizes ranging from a single cell to $\sim 10^8$ individuals or more. Meanwhile, values for $s$ that have been measured during experimental evolution range from $\sim 0.01$ (e.g., a minor improvement in resource uptake [6]) to $\sim 4$ (9) (involving the absolute gain in ability to colonize a new niche), with smaller values more common. Mutants with $s < 0.01$ are likely much more abundant but remain largely undetected because they are lost by drift or outcompeted by more beneficial mutants in large experimental populations, a process known as clonal interference (Table 1) (2, 10). However, technologies that rarify cells and limit clonal interference could detect even smaller fitness effects (13, 42).

When evolving populations are maintained under conditions in which the product of these two properties $N_e s$ is clearly greater than 1 ($N_e s \gg 1$), then selection becomes the dominant force in the population (2, 21, 43). To understand these properties more practically, consider a typical serial dilution evolution experiment in which bacterial populations are grown from a single clone in medium that supports a population of $10^8$ cells/ml (Fig. 1). Let us also assume that the ancestor is not pre-adapted to this environment so there is opportunity for improvement. Population growth on the first day involves $\sim 10^6$ cell divisions and generates $\sim 10^5$ mutations given the approximate per-genome, per-generation mutation rate of $10^{-3}$ (11, 12). Selection could act upon any of these $\sim 10^5$ mutations, and the probability that any one of them would reach a detectable frequency (say, 0.01) depends on the mutant’s frequency and its selective value. So, the earlier a mutation occurs on this first day of population expansion, the more likely it will reach high frequency. Yet most mutations arise in the final division of population growth, i.e., when the population increases from $5 \times 10^7$ to $10^8$ cells/ml involving $5 \times 10^7$ cell divisions. The key point is that after this first growth cycle, there are already many mutations present in the population, but nearly all are very rare, having arisen in the last generation (16).

In a side note, in considering these estimates, one may wonder if some sites or genome regions may be more mutation-prone than others, thus increasing the number of certain mutations. While mutation rates do vary among sites, such rate variation typically involves fold changes ($2 \times$ to $5 \times$) and rarely more than an order of magnitude (10×) (11, 12). These processes can be influential over hundreds of generations but...
typically contribute little over short time scales because they would add only a handful of new mutations to a sample of $\sim 10^5$. Nonetheless, if these mutation hot spots encode a trait under strong selection, such as phase variation of membranes affecting antibiotic resistance (14, 17, 18), they may greatly accelerate the evolutionary rate.

On the next day of the experiment, we transfer some fraction of the first population to fresh medium, say 1/100, and growth resumes (Fig. 1). For example, $10^6$ cells/ml harboring $\sim 10^3$ mutations (simple dilution) now regrow to reach $10^8$ cells/ml, which involves another $10^6$ cell divisions ($10^5$ divisions to reach $2 \times 10^6$ cells/ml, $2 \times 10^6$ divisions to reach $4 \times 10^6$ cells/ml, and so on) and produces another $10^5$ mutations. Despite the many individual cell divisions, it is important to realize that the number of generations the entire population undergoes during the next cycle is $\log_2 (10^8/10^6) = 6.67$ generations. This is the number we commonly reference in describing the duration of an evolution experiment over any interval. It is also noteworthy that this demographic measure of generation number does not depend on the population size but rather on the dilution itself. Under these conditions, the $N_e$ of a population cycling between $10^6$ and $10^8$ cells/ml each day is $\sim 2 \times 10^6$ (Table 1).

Now, revisiting the condition in which the force of selection dominates, $N_e s \gg 1$, any mutation with a selective coefficient $s$ much greater than $1/N_e$ is likely to rise in frequency by selection provided it becomes established in the population. By establishment (Table 1), we mean the mutant must also be lucky enough to reach a frequency where it escapes genetic drift and then is guided deterministically by selection (Fig. 1). This frequency is typically $1/s$ individuals (2, 43). Thus, a fairly typical big-benefit mutation that increases fitness by 10% ($s \sim 0.1$) (19, 20) exceeds the general $N_e s$ threshold by 4 orders of magnitude. If its population size reaches a mere 10 individuals, it is nearly guaranteed to be governed by selection and is likely to completely take over (fix) the population in the absence of competition. However, should this big-benefit mutation cooccur in a population with an even more beneficial mutation ($s = 0.2$ or more), it may lose out to this superior competitor by the process of clonal interference (Table 1 and Fig. 2). In fact, even though beneficial mutations may amount to only 1% of all mutations (or less), large populations guarantee that many

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**FIG 1** Design of a typical evolution experiment with microbial populations. A single colony is isolated from an agar plate and used to inoculate a tube of growth medium. Every 24 h for days or weeks, 1/100 of the population is transferred to fresh medium enabling regrowth. Multiple replicate populations are typically established by the same protocol and can be tracked by plating, quantitation of genetic markers, or whole-population genomic sequencing.
beneficial mutations will arise and compete under these conditions, with only the best of them ever reaching a detectable frequency of 0.01 or more (Fig. 2) (2, 19, 21).

As the experiment progresses, suppose mutants with some potentially adaptive phenotype (a change in colony morphology indicative of altered secretions, growth parameter, metabolic capacity, host colonization, etc.) are observed in a population after seven transfers or ~46 generations. You pick these mutants to subculture them, and their phenotypes appear heritable. What is the probability that these mutants involve adaptations driven by selection and did not become frequent by the random forces of mutation and drift? Virtually certain. Here is why: any given mutation with no effect on fitness (and likely no phenotype) will be governed by genetic drift and will reach a potentially detected frequency of 0.1 in the following number of generations, where $g$ is the number of generations, $p$ is the mutant frequency, and $\mu$ is the mutation rate (22):

$$g = \frac{-\log_2 p}{\mu} = \frac{-\log_2 0.1}{10^{-3}}$$

This would equal 3,322 generations, or 499 days in this experimental design. This is the estimated time required for any mutation to reach a frequency of 10% by genetic drift alone. For a mutant to become detected 100 times faster than this in your evolution experiment (and this is not uncommon), the mutant must have been subject to strong positive selection because it is adaptive.
As evolution experiments become commonplace for many microbial species in a range of environments, reports of parallel mutations—even identical nucleotide changes in the same gene—are becoming more frequent (15, 23). This observation is often met with the question—is this particular site hypermutable? We need to consider three more questions. First, can a high mutation rate alone drive a mutation to high frequency? Second, what role might selection have played? Third, under what conditions would parallelism occur?

Let us suppose that this position is hypermutable, not the usual $1 \times 10^{-10}$ mutations/bp/generation but 100-fold more mutable at $1 \times 10^{-8}$ mutations/bp/generation. Would a high mutation rate alone cause it to rise to high frequency in multiple populations?

Using the same equation above, a specific mutation would reach a frequency of 10% by drift alone by $\frac{-\log_{10}0.1}{10} = 33,219,289$ generations, which is clearly implausible, so mutation pressure alone cannot lead to parallelism. However, let us imagine this mutant is strongly selected with $s = 0.2$, which is much greater than the mutation rate. It will rise to 10% frequency in $\frac{-\log_{0.2}0.1}{0.2} = 16.6$ generations, or less than 3 transfer days, which is fast! Selection can cause a highly beneficial mutation to sweep very rapidly, in the typical time required for an efficient laboratory genetic screen.

However, the probability that the same mutation becomes rapidly detected in another population depends on whether this same mutation actually occurs in time. The evolution experiment we have outlined here generates ~$10^5$ new mutations per day, with only $10^3$ on average surviving each bottleneck. The probability of this same mutation even recurring is the approximate per-site mutation rate of $2 \times 10^{-10}$, which will occur on average in $10^8$ cell divisions/day $\approx 39$ days or 263 generations. Site-specific parallelism is therefore still highly improbable. However, strongly selected mutations are much more likely to survive the bottleneck and may also become enriched both by demography (becoming slightly more common in the transfer by chance). Estimating the exact probability of parallelism is not straightforward because it depends on these chance effects as well as on the distribution of effects of other selected mutations within the population, but in essence, if the same mutation does indeed occur by chance in another population and escapes drift and outcompetes other mutants, it will also rapidly reach this frequency of 0.1.

To sum up, any single mutation rising from a frequency of $10^{-7}$ to $10^{-1}$ in an evolution experiment lasting weeks or months (100 to 1,000 generations) must be under positive selection, and equally important, the contribution of other population genetic processes is negligible. Parallelism at the level of nucleotides, amino acids, or even genes in replicate populations adds overwhelming evidence of strong selection on that target.

Why are these inferences useful? Given that the force of selection dominates evolution experiments conducted in large populations, mutants rising to high frequency must have acquired adaptive traits in the selective environment. Furthermore, the large population sizes typically used in these experiments often generate multiple beneficial mutants that compete with one another (Fig. 2), meaning that those that ultimately become detectable (e.g., as colonies on a plate) are among the most fit available. Genome sequencing of these mutants can therefore be used as a forward genetic screen for traits that enhance fitness in any environment, including new host organisms (9, 15, 24–27). Sequencing multiple mutants from independent populations or diverse samples of the evolved populations can expand the sample of adaptive mutations. Any parallelism at the level of gene or pathway provides powerful inference that alterations in this pathway provide an ideal adaptive solution. As one example from our screen of adaptations to biofilm growth, we observed four independent D652N mutations in the response regulator protein WspE in different populations of *Burkholderia cenocepacia* selected for only 32 to 64 generations (15). These mutants produced a characteristic wrinkly colony morphology and increased biofilm. No specific
features of the nucleotide sequence indicate that this site is hypermutable, but this remarkable parallelism is consistent with the fitness advantage of \( s \approx 1.0 \) that rapidly displaces the ancestor (15). These results also demonstrate that this aspartate residue (and not the adjacent aspartate at position 653) that is outside the primary receiver domain is functionally important for signaling that the cell is in contact with the surface (28, 44). Another example of an evolution experiment serving as a genetic screen involved \( E. coli \) selected for growth in test tubes in minimal medium containing glucose (19). Some of the most fit mutants acquired partial loss-of-function mutations in yfgA and prc, genes that encode a cytoskeletal protein and a protease involved in peptidoglycan synthesis. We reasoned that both mutations could result in the synthesis of a more limited peptidoglycan layer to facilitate transport but also alter cell shape, which we indeed observed as the production of more spherical cells (19). Put simply, this evolve-and-resquence approach is likely to find the best mutations (or mutation combinations) to respond to any selective challenge and can inform how the affected proteins function and enable the mutant to outcompete its ancestor.

Researchers with diverse interests have recognized this opportunity, leading to an explosion of the kinds of questions addressed by experimental evolution and two successful American Society for Microbiology (ASM) meetings focused on the subject. For example, experimental evolution has been used to discover: (i) how bacteria evolve resistance to antibiotics (29, 30) or environmental pollutants (31), (ii) how microbes evolve mutualistic or antagonistic social interactions with other microbes (32–35), (iii) how bacteriophage undergo host range expansion (36, 37), (iv) how microbes adapt to current hosts (38) or novel hosts (9), and (vi) how pathogens evolve during infections (39). These imaginative studies have identified new pathways that reveal how these microbes adapt to new conditions and some recurring themes, such as the selection for mutations in regulators that produce multiple adaptive traits at once. With the wealth of knowledge and experience from almost 150 years of evolution experiments and an influx of creative young investigators, the future of using evolution experiments to reveal unknown mechanisms of adaptation could not be brighter.

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REFERENCES


