Historical contingency and the evolution of a key innovation in an experimental population of *Escherichia coli*

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The role of historical contingency in evolution has been much debated, but rarely tested. Twelve initially identical populations of *Escherichia coli* were founded in 1988 to investigate this issue. They have since evolved in a glucose-limited medium that also contains citrate, which *E. coli* cannot use as a carbon source under oxic conditions. No population evolved the capacity to exploit citrate for >30,000 generations, although each population tested billions of mutations. A citrate-using (Cit⁺) variant finally evolved in one population by 31,500 generations, causing an increase in population size and diversity. The long-delayed and unique evolution of this function might indicate the involvement of some extremely rare mutation. Alternately, it may involve an ordinary mutation, but one whose physical occurrence or phenotypic expression is contingent on prior mutations in that population. We tested these hypotheses in experiments that “replayed” evolution from different points in that population’s history. We observed no Cit⁺ mutants among 8.4 × 10¹² ancestral cells, nor among 9 × 10¹² cells from 60 clones sampled in the first 15,000 generations. However, we observed a significantly greater tendency for later clones to evolve Cit⁺, indicating that some potentiating mutation arose by 20,000 generations. This potentiating change increased the mutation rate to Cit⁺ but did not cause generalized hypermutability. Thus, the evolution of this phenotype was contingent on the particular history of that population. More generally, we suggest that historical contingency is especially important when it facilitates the evolution of key innovations that are not easily evolved by gradual, cumulative selection.

At its core, evolution involves a profound tension between random and deterministic processes. Natural selection works systematically to adapt populations to their prevailing environments. However, selection requires heritable variation generated by random mutation, and even beneficial mutations may be lost by random drift. Moreover, random and deterministic processes become intertwined over time such that future alternatives may be contingent on the prior history of an evolving population. For example, multiple beneficial mutations will arise in some unpredictable order (1, 2), and those that are substituted first may differ from others in their pleiotropic effects and epistatic interactions (3, 4), thus constraining some evolutionary paths while potentiating other outcomes (5–9). These accidents of history may even determine the survival or extinction of entire lineages, given the capricious and sudden nature of some environmental changes (10–12).

Stephen Jay Gould maintained that these historical contingencies make evolution largely unpredictable. Although each change on an evolutionary path has some causal relation to the circumstances in which it arose, outcomes must eventually depend on the details of long chains of antecedent states, small changes in which may have enormous long-term repercussions (13–15). Thus, Gould argued that contingency renders evolution fundamentally quirky and unpredictable, and he famously suggested that replaying the “tape of life” from some point in the distant past would yield a living world far different from the one we see today. Simon Conway Morris countered that natural selection constrains organisms to a relatively few highly adaptive options, so that “the evolutionary routes are many, but the destinations are limited” (16). He and others point to numerous examples of convergent evolution as evidence that selection finds the same adaptations despite the vagaries of history. Evolution may thus be broadly repeatable, and multiple replays would reveal striking similarities in important features, with contingency mostly confined to minor details (16–19).

Of course, replaying life’s tape on the planetary scale is impossible, but careful experiments can examine the role of contingency in evolution on a more modest scale (15, 20, 21). To address the repeatability of evolutionary trajectories and outcomes, the long-term evolution experiment (LTEE) with *Escherichia coli* was started in 1988 with the founding of 12 populations from the same clone (2). These populations were initially identical except for a neutral marker that distinguished six lines from six others. They have since been propagated by daily 1:100 serial transfer in DM25, a minimal medium containing 25 mg/liter glucose as the limiting resource (2, 22). Environmental conditions have been controlled, constant, and identical for all 12 lines. To date, each population has evolved for >44,000 generations, and samples have been frozen every 500 generations, providing a rich “fossil record” (23). Moreover, these samples remain viable, allowing us to perform simultaneous measurements and other experiments with bacteria from different generations. The founding strain is strictly asexual, and thus populations have evolved by natural selection and genetic drift acting on variation generated solely by spontaneous mutations that occurred during the experiment. Thus, the LTEE allows us to examine the effects of contingency that are inherent to the core evolutionary processes of mutation, selection, and drift.

Previous analyses of this experiment have shown numerous examples of parallel phenotypic and genetic evolution. All twelve populations underwent rapid improvement in fitness that decelerated over time (2, 3, 22, 23). All evolved higher maximum growth rates on glucose, shorter lag phases upon transfer into fresh medium, reduced peak population densities, and larger average cell sizes relative to their ancestor (22–26). Ten populations evolved increased DNA supercoiling (27), and those populations examined to date show parallel changes in global adaptation | experimental evolution | mutation | selection


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gene-expression profiles (4, 28, 29). At least three genes have substitutions in all 12 populations (30, 31), and several others have substitutions in many populations (27–30), even though most loci harbor no substitutions in any of them (32). At the same time, there has also been some divergence between populations. Four have evolved defects in DNA repair, causing mutator phenotypes (3, 33). There is subtle, but significant, between-population variation in mean fitness in the glucose-limited medium in which they evolved (2, 23). In media containing other carbon sources, such as maltose or lactose, the variation in performance is much greater (34). And while the same genes often harbor substitutions, the precise location and details of the mutations almost always differ between the populations (27–31).

Throughout the duration of the LTEE, there has existed an ecological opportunity in the form of an abundant, but unused, resource. DM25 medium contains not only glucose, but also citrate at a high concentration. The inability to use citrate as an energy source under aerobic conditions has long been a defining characteristic of E. coli as a species (35, 36). Nevertheless, E. coli is not wholly indifferent to citrate. It uses a ferric dicitrate-sodium citrate transport system for iron acquisition, although citrate does not enter the cell in this process (37, 38). It also has a complete tricarboxylic acid cycle, and can thus metabolize citrate internally during aerobic growth on other substrates (39). E. coli is able to ferment citrate under anaerobic conditions if a cosubstrate is available for reducing power (40). The only known barrier to aerobic growth on citrate is its inability to transport citrate under aerobic conditions (41–43). Indeed, atypical E. coli that grow aerobically on citrate (Cit⁺) have been isolated from agricultural and clinical settings, and were found to harbor plasmids, presumably acquired from other species, that encode citrate transporters (44, 45).

Other findings suggest that E. coli has the potential to evolve a Cit⁺ phenotype. Hall (41) reported the only documented case of a spontaneous Cit⁺ mutant in E. coli. He hypothesized that some complex mutation, or multiple mutations, activated cryptic genes that jointly expressed a citrate transporter, although the genes were not identified. Pos et al. (43) identified an operon in E. coli K-12 that apparently allows anaerobic citrate fermentation, and which includes a gene, citT, encoding a citrate–sodium citrate antiporter. High-level constitutive expression of this gene on a multicopy plasmid allows aerobic growth on citrate, but the native operon has a single copy that is presumably induced only under anaerobic conditions.

Despite this potential, none of the 12 LTEE populations evolved the capacity to use the citrate that was present in their environment for over 30,000 generations. During that time, each population experienced billions of mutations (22), far more than the number of possible point mutations in the ~4.6-million-bp genome. This ratio implies, to a first approximation, that each population tried every typical one-step mutation many times. It must be difficult, therefore, to evolve the Cit⁺ phenotype, despite the ecological opportunity. Here we report that a Cit⁺ variant finally evolved in one population by 31,500 generations, and its descendants later rose to numerical dominance. The new Cit⁺ function has been the most profound adaptation observed during the LTEE and has had major consequences. As we will show, the population achieved a severalfold increase in size. Moreover, a stable polymorphism emerged, with a Cit⁻ minority coexisting with the new Cit⁺ majority. Interestingly, the population that evolved the Cit⁺ function is not one that had previously become hypermutable. It is also intriguing that this key innovation evolved so late in the experiment, given that the rate of fitness improvement had declined substantially in all of the populations (3, 23).

The long-delayed and unique evolution of the Cit⁺ phenotype might indicate that it required some unusually rare mutation, such as a particular chromosomal inversion, that does not scale with typical mutation rates. Alternatively, the occurrence or phenotypic expression of the mutation that generated the Cit⁺ function might depend on one or more earlier mutations, such that its evolution was contingent on the particular history of that population. Contingent adaptations should tend to be complex and require multiple steps, some of which might not be beneficial, at least not uniquely so given other advantageous paths. Otherwise, cumulative selection would predictably favor the same steps, and the evolutionary path should be repeatable (18). Contingent adaptations should thus display two characteristics. First, independent origins should be rare, because the same historical sequences would rarely recur (19). Second, significant time-lags should occur between the presentation of ecological opportunities or challenges and the evolution of those traits that confer adaptation to those circumstances (46).

Can the hypothesis of contingent adaptation be rigorously tested in the case of the evolution of the Cit⁺ function? And can this hypothesis be formally distinguished from the alternative explanation that this new function required an unusually rare mutation, but one that was not historically contingent? The answer to both questions is yes, owing to certain features of bacteria in general and the LTEE in particular that allow us quite literally to replay evolution from various points in a population’s history. We will first describe the emergence of this new function, and then present our experiments to distinguish between the hypotheses of mutational rarity and historical contingency in the origin of this key innovation.

**Results**

**Evolution of Cit⁺ Function in Population Ara-3.** The LTEE populations are transferred daily into fresh medium, and the turbidity of each is checked visually at that time. Owing to the low concentration of glucose in DM25 medium, the cultures are only slightly turbid when transferred. Occasional contaminants that grow on citrate have been seen over the 20 years of this experiment. These contaminated cultures reach much higher turbidity owing to the high concentration of citrate in the medium, which allows the contaminants to reach high density. (When contamination occurs, the affected population is restarted from the latest frozen sample.) After ~33,127 generations, one population, designated Ara-3, displayed significantly elevated turbidity that continued to rise for several days (Fig. 1).

A number of Cit⁺ clones were isolated from the population and checked for phenotypic markers characteristic of the ancestral E. coli strain used to start the LTEE: all were Ara⁻, T5-sensitive, and T6-resistant, as expected (2). DNA sequencing also showed that Cit⁺ clones have the same mutations in the pykF and nadR genes as do clones from earlier generations of the Ara-3 population, and each of these mutations distinguishes this population...
from all of the others (30). Therefore, the Cit\(^+\) variant arose within the LTEE and is not a contaminant.

The evolved Cit\(^+\) variant grows to high density in DM0 (a citrate-only medium), produces vigorous colonies on minimal citrate (MC) agar plates, and causes a positive color change on Simmon’s citrate agar, all of which indicate that it can use citrate (Fig. 2). Also, growth on citrate is inhibited by the citrate analog 5-fluorocitrate (data not shown), as was observed for the Cit\(^+\) variant. Cit\(^+\) cells undergo a period of rapid growth on glucose that is followed by slower growth on citrate (Fig. 2).

The hypothesis of clonal interference implies that competition the emerging Cit\(^+\)/H11001 analog 5-fluorocitrate (data not shown), as was observed for the Cit\(^+\) variant. Cit\(^+\) cells were found in the samples taken at 30,000, 30,500, or 31,000 generations. Cit\(^+\) cells constituted \(\approx 0.5\%\) of the population at generation 31,500, then 15\% and 19\% in the next two samples, but only 2\% at generation 33,000. It appears that the first Cit\(^+\) variant emerged between 31,000 and 31,500 generations, although we cannot exclude an earlier origin. The precipitous decline in the frequency of Cit\(^+\) cells just before the massive population expansion suggests clonal interference (47), whereby the Cit\(^-\) subpopulation produced a beneficial mutant that outcompeted the emerging Cit\(^+\) subpopulation until the latter evolved some other beneficial mutation that finally ensured its persistence. The hypothesis of clonal interference implies that the early Cit\(^+\) cells were very poor at using citrate, such as mutations that improved competition for glucose could have provided a greater advantage than did marginal exploitation of the unused citrate.

Indeed, the Cit\(^+\) clones isolated from generations 32,500 and earlier grow much more slowly on citrate than those from 33,000 generations and later. After depleting the glucose in DM25, the earliest Cit\(^+\) clones grow almost imperceptibly, if at all, for many hours before they begin efficiently using the citrate (data not shown), whereas later Cit\(^+\) clones switch to growth on citrate almost immediately (Fig. 2). Thus, the population expansion between generations 33,000 and 33,500 (Fig. 1) was triggered by one or more mutations that improved citrate utilization, rather than by the growth of the original Cit\(^+\) mutant. This finding also raises the question of whether weak Cit\(^+\) mutants might exist in any other LTEE population. We therefore screened the other 11 populations, in most cases by using samples from generation 41,500. None of 220 cultures inoculated with heterogeneous population samples grew in glucose-free DM0, nor did any of \(\geq 3,500\) clones show a positive reaction on Christensen’s citrate agar.

Historical Contingency in the Evolution of Cit\(^+\). We performed three experiments to test whether the evolution of the Cit\(^+\) function was low but constant over time. Under the historical-contingency hypothesis, the probability of this transition increased when a mutation arose that produced a genetic background with a higher mutation rate to Cit\(^+\).

In our first experiment, we performed the replays under the same conditions as the LTEE. We isolated three random clones from each of twelve time points, from the ancestor to 32,500 generations (Table 1), and obtained neutral Ara\(^+\) mutants of each clone to embed as protection against accidental cross-contamination during the experiment (2). In total, there were 72 replay populations, six from each generation, each founded by a single clone. These populations evolved for \(\approx 3,700\) generations, and they were checked visually each day for the elevated turbidity indicative of the Cit\(^+\) phenotype. We also tested samples on MC and Christensen’s citrate agar plates every 250 generations, with incubation for up to a week. New Cit\(^+\) variants emerged in four replay populations, all founded by clones from later generations of the original population (Table 1). These Cit\(^+\) variants emerged between 750 and 3,700 generations of the replay experiment.

Our second experiment also looked for Cit\(^+\) mutants derived from clones sampled at various times in the history of population Ara-3. This time, however, we incubated large populations of Cit\(^+\)/H11001 cells on MC plates, enabling us to test more clones and more cells of each clone. We also allowed a long incubation time to facilitate the growth and detection of very weak Cit\(^+\) mutants, as well as mutations that might occur as cells sat starving on the plates. In this experiment, \(\approx 3.9 \times 10^8\) cells of each of the same 68 clones used in the first replay experiment were spread on each of five MC plates, and these 340 plates were then incubated for 59 days. Five plates produced Cit\(^+\) mutants, and all used clones from generations 32,000 or 32,500 of the original population (Table 1). None of the particular clones that evolved Cit\(^+\) in this experiment did so in the first one, although there was overlap in the generations from which those clones were sampled.
The third replay experiment was similar in design to the second, but on a larger scale. We isolated 20 clones from each of 13 time points in the history of population Ara-3, again through 32,500 generations. We generated and tested 10 replicate cultures of each evolved clone and 200 replicates of the ancestor. Each culture grew to \( \sim 1 \times 10^{10} \) cells, which were pelleted by centrifugation, spread on an MC plate, and incubated for 45 days. In total, \( \sim 4 \times 10^{13} \) cells were tested for their ability to use citrate in this experiment. Cit\(^+\) mutants emerged from eight cultures representing seven clones (Table 1), with one clone yielding mutants in two replicate cultures. Four clones that produced Cit\(^+\) mutants came from generations 31,000 and later, two were from generation 27,000, and one (the one that produced two mutants) was from generation 20,000. We found no Cit\(^+\) mutants among any of the 200 ancestral cultures, nor among any of the other 600 cultures that used clones isolated before generation 20,000.

Interestingly, 7 of the 8 plates that yielded a mutant Cit\(^+\) colony produced multiple colonies, including one with 137 colonies. This pattern illustrates the “jackpot” effect discovered by Luria and Delbrück (48), and it implies that mutations arose during the population growth before plating on MC agar. On the other hand, the Cit\(^+\) colonies were not observed until at least 8 days of incubation and, in one case, they were first seen after 28 days. These late appearances suggest that the mutations to Cit\(^+\) occurred after plating. One possible explanation for this apparent discrepancy is that the mutants grow very slowly but, in fact, they typically produce visible colonies in only 2 days when restested on MC plates. Another potential explanation is that the high density of Cit\(^-\) cells on the plates interfered with the growth and detection of emerging Cit\(^+\) colonies. To test this possibility, we seeded dense Cit\(^-\) cultures with a few Cit\(^+\) cells before plating on MC agar, but the Cit\(^+\) colonies were seen only after 2–3 days. The rapid growth of Cit\(^+\) colonies occurred even when the Cit\(^+\) cells had grown on glucose, and not on citrate, before plating. These results imply that mutations to Cit\(^+\) occurred after cultures were plated on MC agar. This conclusion, taken together with the jackpot distribution of Cit\(^+\) mutants, indicates that the phenotypic change required two mutations, one of which occurred during the culture growth before plating and the other after plating.

Statistical Analysis of the Replay Experiments. All three experiments show the same tendency for Cit\(^+\) variants to evolve more often from clones sampled in later than earlier generations of population Ara-3 (Table 1). To calculate the significance of these data, we performed Monte Carlo resampling tests (shuffling without replacement) by using the Statistics101 Re-sampling Simulator version 1.0.6 (www.statistics101.net). For each experiment, we compared the observed mean generation of those clones that yielded Cit\(^+\) variants to the mean expected under the null hypothesis that clones from all generations have equal likelihood. The null thus corresponds to the rare-mutation hypothesis laid out in the Introduction. We ran one million resampling iterations for each experiment. The deviations from the null expectations range from marginally to highly significant in the three experiments, and in all cases they support the historical-contingency hypothesis, according to which clones from later generations have greater propensity to evolve the Cit\(^+\) phenotype (Table 2). Although the third experiment was the largest, it was the least significant, owing primarily to the production of two Cit\(^+\) mutants by a 20,000-generation clone. We also used the Z-transformation method (49) to combine the probabilities from our three experiments, and the result is extremely significant (\( P < 0.0001 \)) whether or not the experiments are weighted by the number of independent Cit\(^+\) mutants observed in each one. Furthermore, the potentiation effect in later generations is underestimated by these tests, because the number of cells trended lower in later-generation cultures owing to the evolution of larger cells that reach lower population density (24, 25, 41, 50).

These analyses compel us to reject the hypothesis that a rare mutation could have produced a Cit\(^+\) variant with equal probability at any point in the LTEE. Some unusually rare mutation

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<th>Table 1. Summary of replay experiments</th>
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<td><strong>First experiment</strong></td>
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<th>Table 2. Statistical analyses of three replay experiments</th>
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<td><strong>Mean generation of clones yielding Cit(^+)</strong></td>
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<td>Monte Carlo ( P ) value</td>
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Clones were incubated.

other DNA activity probably occurred during the many days that
through 15,000 generations; and because some cell turnover and
distribution based on the uncertainty in
confidence interval (CI) can also be calculated from the Poisson
mutation rate to Cit
in the third replay experiment, the upper bound on the ancestral
found among another 9.0
/H11003
mentally measured. It is also probably far too high because no
4). To the best of our knowledge, this value is the lowest upper
clone, and 280 for the ancestor. Potentiated and ancestral
ground. We prepared 40 replicate cultures for each potentiated

Effect. Given the fraction of cultures that produce no mutants
(P0) and the number of cells per culture (N), one can estimate the
mutation rate as \( \mu = -(\ln P0)/N \). The corresponding 95% confidence interval (CI) can also be calculated from the Poisson
distribution based on the uncertainty in
which is much greater
than the uncertainty in N. By comparing the mutation rate to
Cit
between clones, one can in principle quantify the potenti-
tion effect in the evolved genetic background. However, we
cannot unambiguously identify which evolved clones remain
nonpotentiated owing to the very low mutation rate to Cit
even in the potentiated clones. We also cannot separate potentiated
and nonpotentiated clones by generation because some gener-
alional samples may be polymorphic.

We therefore performed an independent series of fluctuation
tests using seven clones that yielded Cit
mutants in at least one
replay experiment. These additional tests permit an unbiased
estimate of the mutation rate to Cit
in the potentiated back-
ground. We prepared 40 replicate cultures for each potentiated
clon, and 280 for the ancestor. Potentiated and ancestral
cultures had, on average, 1.1 and 1.5 \times 10^{10}
cells, respectively, which were harvested and incubated on MC agar plates for 45
days.

None of the ancestral cultures yielded any Cit
mutants. We
can nonetheless calculate an upper limit to the mutation rate by
noting that the Poisson distribution has a 5% probability of
yielding zero events when the expectation is three. With no more
than three mutations among the 8.4 \times 10^{12}
cells tested here and
in the third replay experiment, the upper bound on the ancestral
mutation rate to Cit
is 3.6 \times 10^{-13}
per cell per generation (Fig. 4). To the best of our knowledge, this value is the lowest upper
bound ever reported for a mutation rate that has been experi-
mentally measured. It is also probably far too high because no
mutations were actually observed for the ancestor, nor were any
found among another 9.0 \times 10^{12}
cells of 60 clones sampled
through 15,000 generations; and because some cell turnover and
other DNA activity probably occurred during the many days that
plates were incubated.

Even among the potentiated clones, the rate of mutation to
Cit
is extremely low. Cit
mutants arose in 2 of the 280 new
cultures, giving an estimate of 6.6 \times 10^{-13}
for the mutation rate,
with the 95% CI extending from 7.9 \times 10^{-14}
to 2.4 \times 10^{-12}
(Fig.
4). (Although the upper bound of the CI for the ancestor
overlaps the lower bound of the CI for the potentiated clones,
that upper bound does not overlap the point estimate for the
potentiated clones, indicating a significant difference and adding
further support to the replay experiments.) The potentiated
background thus increases the mutation rate to Cit
at least 2-fold, and probably much more. However, even this
potentiated value represents an unusually low mutation rate. A
typical mutation rate in E. coli is \approx 5 \times 10^{-10}
per base pair per
generation (51). Such a low rate suggests that the final mutation
to Cit
is not a point mutation but instead involves some rarer
class of mutation or perhaps multiple mutations. The possibility
of multiple mutations is especially relevant, given our evidence
that the emergence of Cit
colonies on MC plates involved
events both during the growth of cultures before plating and
during prolonged incubation on the plates.

Another issue is whether the potentiating effect might indicate
the evolution of a generalized hypermutability in population
Ara-3. Previous surveys of mutation rates in the 12 LTEE lines
found that 4 had become mutators by generation 20,000, al-
though population Ara-3 retained the low ancestral mutability
(3, 33). To investigate this issue further, we performed another
series of fluctuation tests using the same seven potentiated
clones to determine their rate of mutation from Ara
- to Ara
+, which serves as a proxy for the background mutation rate (33). This mutation reverts a phenotype that was knocked out during
the derivation of a predecessor to the ancestor of the LTEE (2,
52). Sequence analysis indicates that most Ara
+ mutants have a
mutation from GAC to GGC at codon 92 of the araA gene,
restoring that codon to its distant ancestral state. Forty cultures
of each clone were grown in DM1000 (1 g of glucose per liter),
and the cells were spread on minimal agar plates containing
arabinose. At the same time, 280 cultures of the ancestor were
tested in the same way. The ancestral mutation rate to Ara
+ is
2.3 \times 10^{-10}
per cell per generation calculated by using the
method (95% CI from 1.8 \times 10^{-10}
to 2.9 \times 10^{-10}), a typical rate
for point mutations. The rate for potentiated clones is also 2.3 \times 10^{-10}
(95% CI from 2.0 \times 10^{-10}
to 2.7 \times 10^{-10}); in fact, these
estimates are within 1% of each other (Fig. 4). We conclude that
general hypermutability is not responsible for the elevated
mutation rate to Cit
in the potentiated clones.

Frequency-Dependent Selection Maintains Ecological Diversity. To
this point, we have examined the evolutionary origin of the Cit
function in population Ara-3. We now turn to its ecological
consequence. The Cit
phenotype did not achieve fixation
during the population expansion but, instead, Cit
 cells persisted
as a minority. When we mixed Cit
and Cit
clones at different
initial frequencies, they stably coexisted over many serial trans-
fers (Fig. 5). In these mixtures, the Cit
cells gradually
approached an equilibrium frequency of roughly 1%, regardless of
their initial frequency. We saw a transient jump in the frequency of the Cit⁻ subpopulation on days 7 and 8, which was probably caused by accidentally using a glucose-only medium on those days. After that perturbation, the populations resumed their previous trajectories. Negative frequency-dependent selection thus maintains the polymorphism.

This stable coexistence suggests that the Cit⁻ cells are superior to the Cit⁺ cells in competition for glucose, allowing the former to persist as glucose specialists. Indeed, the Cit⁻ cells have a shorter lag phase and higher growth rate on glucose than do the Cit⁺ cells (Fig. 2). These differences were also evident when we monitored the intraday dynamics of mixtures of Cit⁻ and Cit⁺ cells (data not shown).

Discussions and Future Directions

E. coli cells cannot grow on citrate under oxic conditions, and that inability has long been viewed as a defining characteristic of this important, diverse, and widespread species. In a long-term experiment, we propagated 12 populations of E. coli, all founded from the same ancestral strain, in a medium containing glucose, which is the limiting resource, and abundant citrate. For more than 50,000 generations, none of them evolved the capacity to use the citrate, although billions of mutations occurred in each population, such that any typical base pair mutation would have been tested many times in each one. It is clearly very difficult for E. coli to evolve this function. In fact, the mutation rate of the ancestral strain from Cit⁻ to Cit⁺ is immeasurably low; even the upper bound is 3.6 × 10⁻ⁱ⁵ per cell generation, which is three orders of magnitude below the typical base pair mutation rate.

Nevertheless, one population eventually evolved the Cit⁺ function, whereas all of the others remain Cit⁻ after more than 40,000 generations.

We demonstrated that the evolution of this new function was contingent on the history of the population in which it arose. In particular, we showed that one or more earlier mutations potentiated the evolution of this function by increasing the mutation rate to Cit⁺, although even the elevated rate is much lower than a typical mutation rate. The potentiated cells are not generally hypermutable. Rather, their potentiation appears to be specific to the Cit⁺ function, which suggests two possible mechanisms. One mechanism is epistasis, whereby the functional expression of the mutation that finally yielded the Cit⁺ phenotype requires interaction with one or more mutations that evolved earlier. A second possibility is that the physical production of the mutation that produced the Cit⁺ phenotype requires some previous mutation that allows the final sequence to be generated. For example, the insertion of a mobile genetic element creates new sequences at its junctures, and one of these new sequences might then undergo a mutation that generates a final sequence that could not have occurred without the insertion. The E. coli genome has many insertion-sequence elements (53), some of which have been active in the LTEE (54–56). Whatever the mechanism, this potentiation made the Cit⁺ function mutationally accessible, and a weak Cit⁺ variant emerged by 31,500 generations.

The origin of the Cit⁺ function also had profound consequences for the ecology and subsequent evolution of that population. This new capacity was refined over the next 2,000 generations, leading to a massive population expansion as the Cit⁺ cells evolved to exploit more efficiently the abundant citrate in their environment. Although the Cit⁺ cells continued to use glucose, they did not drive the Cit⁻ subpopulation extinct because the Cit⁻ cells were superior competitors for glucose. Thus, the overall diversity increased as one population gave rise evolutionarily to an ecological community with two members, one a resource specialist and the other a generalist.

The evolution of the new Cit⁺ function represents a key innovation that involves multiple steps, and it provides an explicit demonstration of the importance of historical contingency in evolution. It also transcends the phenotypic boundaries of a diverse and well-studied species, and led to an ecological transition from a single population to a two-member community. Our future research on this fascinating case of evolution in action will revolve around four themes: genetics, physiology, ecology, and speciation.

What is the genetic basis of this evolutionary innovation? The emergence of the Cit⁺ phenotype in population Ara-3 indicates at least two important genetic events: the origin of the function in its weak form, and its subsequent refinement for efficient use of citrate. The replay experiments indicate an even more complex pattern that must involve, at a minimum, three important genetic events. At least one mutation in the LTEE was necessary to produce a genetic background with the potential to generate Cit⁺ variants, while the distribution and dynamics of Cit⁺ mutants in fluctuation tests indicate at least two additional mutations are involved. To find the relevant mutations, we will perform whole-genome resequencing, which has become a powerful approach that is well suited to experimental evolution (57–59). We expect to find dozens of mutations relative to the ancestor (22), which will complicate identification of those changes that were important specifically for the origin of the Cit⁺ function. However, some of the key changes should become apparent if we also resequence a Cit⁻ clone from the same population around the time that the Cit⁺ variants first emerged. Once candidate genes and mutations have been identified, we can examine the other 19 Cit⁺ variants from our replay and mutation-rate experiments for parallel changes.

We are especially eager to find the potentiating mutation or mutations. We want to know whether the potentiating mutation interacts epistatically with a later mutation to allow expression of the Cit⁺ function or, alternatively, whether it was physically required for the later mutation to occur. We also want to test whether the potentiating mutation was itself beneficial or, alternatively, a neutral or deleterious change that fortuitously hitchhiked to high frequency. We anticipate that identifying the potentiating mutation will be especially challenging, however, because its only known phenotype is to increase the rate of production of certain mutants that are themselves extremely rare.

Once we have identified all of the relevant mutations, it might be interesting to model the population dynamics that govern the occurrence of this new function. Such a model would require not only all of the relevant mutation rates but also the ecological phenotypes of the mutants, including their growth rates on glucose and citrate as well as their abilities to transition between the two resources. A satisfactory model should also reflect the stochastic origin of mutations, the role of random drift, and the possibility of alternative mutational paths to the phenotype of interest (9, 60).

What physiological mechanism has evolved that allows aerobic growth on citrate? E. coli should be able to use citrate as an energy source after it enters the cell, but it lacks a citrate transporter that functions in an oxygen-rich environment. One possibility is that the Cit⁺ lineage activated a “cryptic” transporter (41), that is, some once-functional gene that has been silenced by mutation accumulation. This explanation seems unlikely to us because the Cit⁻ phenotype is characteristic of the entire species, one that is very diverse and therefore very old. We would expect a cryptic gene to be degraded beyond recovery after millions of years of disuse. A more likely possibility, in our view, is that an existing transporter has been coopted for citrate transport under oxic conditions. This transporter may previously have transported citrate under anoxic conditions (43) or, alternatively, it may have transported another substrate in the presence of oxygen. The evolved changes might involve gene regulation, protein structure, or both (61).
What will be the long-term fates of the coexisting Cit^+ and Cit^- subpopulations? We showed they stably coexist owing to the inferiority of the Cit^- cells in competition for glucose. However, the Cit^- lineage might eventually acquire mutations that compensate for its inferior performance on glucose, thus undermining the coexistence. Compensation for maladaptive side-effects of adaptations, including resistance to phages and antibiotics, has been observed in many other experiments with bacteria (62–66). Moreover, the Cit^- subpopulation is much larger than the Cit^- subpopulation, so it should experience more beneficial mutations even without compensation. On the other hand, coexistence would be strengthened if selection in the Cit^- subpopulation favors specialization on citrate. In the same way that we established multiple populations for retrospective replays in this study, we can establish multiple communities to examine their prospective evolution. We can also vary environmental factors, such as the presence or absence of glucose, and the presence or absence of the Cit^- subpopulation, to investigate how they influence the future evolution of the Cit^- lineage.

Will the Cit^- and Cit^- lineages eventually become distinct species? According to the biological species concept widely used for animals and plants, species are recognized by reproductive isolation between species (67). Although the bacteria in the LTEE are strictly asexual, we can nonetheless imagine testing this criterion by producing recombinant genotypes. In particular, we could move mutations that are substituted in the evolving Cit^- lineage into a Cit^- background to test whether they reduce fitness in their ancestral context. One could also perform the reciprocal experiment, although we anticipate more rapid evolution in the Cit^- lineage because it has acquired a key innovation that substantially changed its ecological niche. Such experiments would require, of course, controls to examine the fitness effects of the same mutations in the lineage where they arose. If the Cit^- lineage is indeed evolving into a new species, then we expect, with time, that more and more of the beneficial mutations substituted in that lineage would be detrimental in the ecological and genetic context of its Cit^- progenitor.

In any case, our study shows that historical contingency can have a profound and lasting impact under the simplest, and thus most stringent, conditions in which initially identical populations evolve in identical environments. Even from so simple a beginning, small happenstances of history may lead populations along different evolutionary paths. A potentiated cell took the one less traveled by, and that has made all the difference.

Materials and Methods

The Long-term Evolution Experiment. The LTEE is described in detail elsewhere (2, 22). Briefly, two ancestral clones of E. coli B were each used to found six populations. The ancestors differ by a single mutation that allows one of them to use arabinose (Ara^-). Ara^- and Ara^- cells make red and white colonies, respectively, on tetrazolium–arabinose (TA) plates, but the mutation is neutral for animals and plants, species are recognized by reproductive isolation between species (67). Although the bacteria in the LTEE are strictly asexual, we can nonetheless imagine testing this criterion by producing recombinant genotypes. In particular, we could move mutations that are substituted in the evolving Cit^- lineage into a Cit^- background to test whether they reduce fitness in their ancestral context. One could also perform the reciprocal experiment, although we anticipate more rapid evolution in the Cit^- lineage because it has acquired a key innovation that substantially changed its ecological niche. Such experiments would require, of course, controls to examine the fitness effects of the same mutations in the lineage where they arose. If the Cit^- lineage is indeed evolving into a new species, then we expect, with time, that more and more of the beneficial mutations substituted in that lineage would be detrimental in the ecological and genetic context of its Cit^- progenitor.

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