

Evolution of high mutation rates in experimental populations of *E. coli*

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Most mutations are likely to be deleterious, and so the spontaneous mutation rate is generally held at a very low value¹. Nonetheless, evolutionary theory predicts that high mutation rates can evolve under certain circumstances^{2–4}. Empirical observations have previously been limited to short-term studies of the fates of mutator strains deliberately introduced into laboratory populations of *Escherichia coli*^{5–7}, and to the effects of intense selective events on mutator frequencies in *E. coli*⁸. Here we report the rise of spontaneously originated mutators in populations of *E. coli* undergoing long-term adaptation to a new environment. Our results corroborate computer simulations of mutator evolution in adapting clonal populations⁴, and may help to explain observations that associate high mutation rates with emerging pathogens⁹ and with certain cancers¹⁰.

As a means of studying evolution directly in the laboratory, 12 replicate experimental populations of a clonal strain of *E. coli* were established¹¹. Because these populations were founded from a single ancestral clone, mutation provided their only source of genetic variation. The glucose-limited environment in which these populations were propagated was essentially novel at the outset of the experiment and it provided considerable scope for adaptive evolution. Substantial evolutionary increases in fitness and changes in certain other phenotypic features in these populations have been documented elsewhere^{11–15}.

We measured mutation rates in the common ancestral strain and in the 12 experimental populations after they had been evolving for 10,000 generations. Most of the populations retained the ancestral mutation rate, but three populations (designated Ara⁻2, Ara⁻4 and Ara⁺3) displayed mutation rates that were between one and two orders of magnitude higher than those in the ancestor (Fig. 1). The times at which mutator phenotypes arose during the evolution of populations Ara⁻2, Ara⁻4 and Ara⁺3, as determined by screening for the mutator phenotype in isolates stored periodically throughout the experiment, are shown in Fig. 2. Once a lineage displayed a mutator phenotype, all subsequent isolates from that lineage also tested as mutators through 10,000 generations. This observation indicates that mutators rose to and remained at high frequencies in these populations.

To ascertain a genetic basis for the observed mutator phenotypes, we transformed 10,000-generation clonal isolates from populations Ara⁻2, Ara⁻4 and Ara⁺3, and an isolate from the ancestral strain REL606 with multicopy plasmids bearing wild-type alleles of seven known general mutator loci. The ancestral mutation rate was fully restored in the Ara⁻2, Ara⁻4 and Ara⁺3 strains only by the presence of wild-type alleles of genes in the methyl-directed mismatch repair pathway¹⁶ (Fig. 3). The mutation rate in the ancestral strain, REL606, was not significantly affected by the plasmids (with a single exception: see Fig. 3). In the Ara⁺3 strain the *mutS*⁺ allele alone restored the ancestral mutation rate. In the Ara⁻2 and Ara⁻4 strains the ancestral rate was restored completely by *uvrD*⁺, and the data also indicated a partial effect of *mutL*⁺. Recent studies have suggested that there is a mechanistic interaction between the *mutL*

and *uvrD* gene products, such that a defect in one may in some cases be complemented by increased production of the other (P. Modrich, personal communication).

Most mutations are deleterious, so mutator alleles are likely on average to have negative effects on fitness. In an evolving clonal population, however, a deleterious mutator can rise to high frequency (hitch-hike) in association with an adaptive mutation, provided that the selective cost of the mutator does not outweigh the selective benefit of the adaptive mutation. Hitch-hiking of mutators with adaptive mutations was demonstrated previously in chemostat populations of *E. coli*⁶, but these results left open the question of whether such events would be likely to occur in natural populations. When a mutator was introduced above a relatively high threshold number relative to the wild type, the mutator population always acquired an adaptive mutation before the wild-type population, and the mutator rose to high frequency. However, when the mutator was introduced in lower numbers, the wild-type population always acquired a beneficial mutation first and displaced the mutator.

In the evolution experiment we have described here, as in natural populations, mutator alleles must have arisen by mutation. This raises the question of how mutators reached high frequencies in 3 of our 12 populations. Computer simulations⁴ and analytical modeling (P.J.G. and P.D.S., unpublished data) suggest that rare mutators may occasionally hitch-hike to high frequencies in finite asexual populations as a consequence of chance associations with adaptive mutations. We think that such chance hitch-hiking events are the likely explanation for our results. Sufficient time may not have elapsed for them to have been observed in earlier studies of

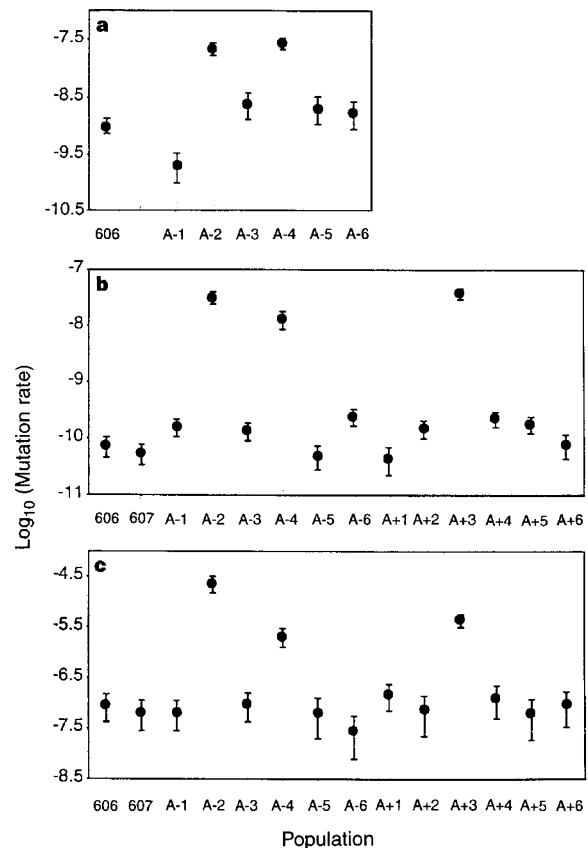


Figure 1 Rates of mutation in 12 Ara⁻ and Ara⁺ experimental populations (A – 1 to A – 6 and A + 1 to A + 6) at 10,000 generations and in their common ancestors REL606 (Ara⁻) and REL607 (Ara⁺). Error bars give approximate 95% confidence intervals. **a**, Reversion to Ara⁺ in isolates from the 7 Ara⁻ populations. **b**, Mutation to nalidixic-acid resistance. **c**, Mutation to bacteriophage T5 resistance.

competition between mutator and wild-type strains⁵⁻⁷, as those studies were carried out for only a few tens or hundreds of generations, as opposed to the 10,000 generations in our study. Although we cannot formally exclude the possibility that certain mutations in mismatch repair might increase fitness directly, it is much more likely that the evolved mutators we observed are deleterious or at best effectively neutral. Indeed, known mutator

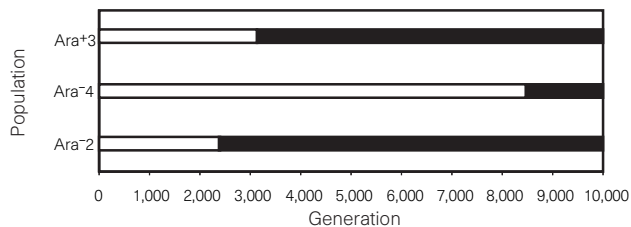


Figure 2 Time of appearance and evolutionary persistence of mutator phenotypes in experimental populations Ara⁻², Ara⁻⁴ and Ara⁺³. White bars, wild type; black bars, mutator.

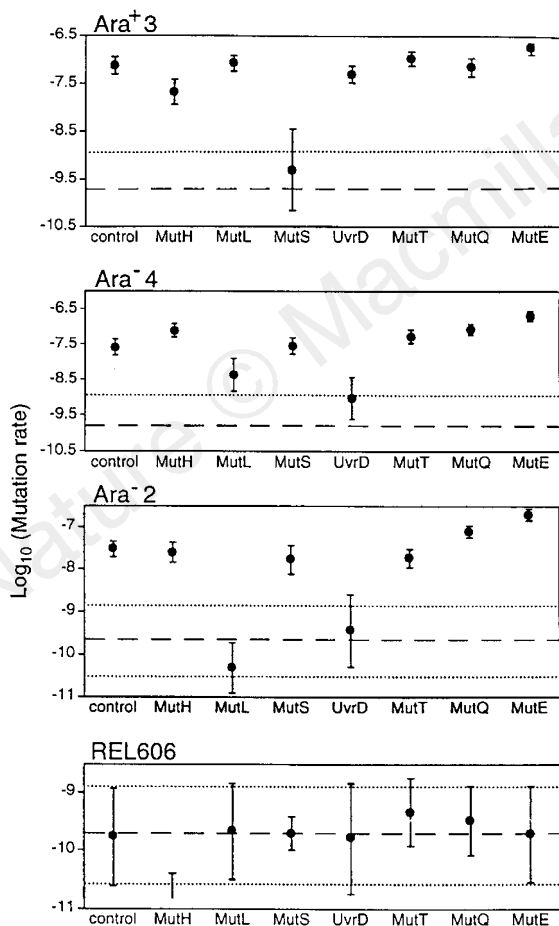


Figure 3 Mutation rates to nalidixic-acid resistance in 10,000-generation isolates from populations Ara⁺³, Ara⁻⁴ and Ara⁻² and ancestor REL606 transformed with plasmids bearing wild-type alleles of seven known general mutator loci. Controls shown are plasmid free. Error bars give approximate 95% confidence limits. Only an upper confidence limit is shown for the case in which no mutants were obtained in any culture. For visual comparison, the dashed horizontal line and dotted horizontal lines in each panel illustrate the mutation rate and approximate 95% confidence limits, respectively, measured in the ancestral control.

alleles of *mutS*, *mutL* and *uvrD* do not increase cell fitness under our experimental conditions (C. Zeyl, unpublished data).

A prediction of the simulations conducted by Taddei *et al.*⁴ is that mutators will hitch-hike in some, but not all, finite asexual populations undergoing adaptation. Our experimental observations are consistent with this prediction. A further prediction is that fitness will be higher on average in mutator populations than in those that retain the wild-type mutation rate. However, this fitness effect is very small and subtle: approximately half of the mutator populations in the simulations of Taddei *et al.* did not show higher fitness, and the observed increases in fitness over contemporaneous wild-type populations were slight (approximately 1%) in the remainder (see Fig. 3b in ref. 4). We tested for a relationship between mutability and fitness in our experimental populations in several ways, none of which gave statistical significance. For example, we tested for a correlation between mutability ranks (as calculated from data obtained in screens for mutator phenotypes; see Methods) and relative fitness ranks averaged across the duration of the experiment in all 12 populations. The results were inconclusive ($r = 0.28$, $n = 12$, one-tailed $P = 0.19$). It is clear from our experiments and the simulations of Taddei *et al.*⁴ that increased mutation rate and not increased fitness is the more striking consequence of the hitch-hiking process.

Our finding that asexual populations can evolve high mutation rates in a relatively benign environment may help to explain observations associating mismatch repair mutators with certain cancers¹⁰ and with pathogenicity in *E. coli* and *Salmonella*⁹. As in our experimental populations, high mutation rates may evolve as a chance byproduct of adaptation in clonal tumour lineages and in populations of asexual pathogens^{17,18}. The potential for faster evolution once high mutation rates have evolved may have important health implications. □

Methods

Experimental system. The common ancestral strain was REL606, an Ara⁻ clone of *E. coli* B obtained by B. R. Levin from S. Lederberg¹⁹. A spontaneous Ara⁺ revertant was selected from REL606 and designated REL607. Six clones each from REL606 and REL607 were used to found the 12 experimental populations, which were propagated at 37 °C by daily 100-fold dilutions into 10 ml of fresh Davis minimal medium²⁰ supplemented with glucose at 25 μg ml⁻¹. Population sizes fluctuated between approximately 5 × 10⁶ cells after dilution and 5 × 10⁸ cells at the stationary phase. Isolates from each population were stored at -80 °C at 100-generation intervals during the first 2,500 generations of the experiment, and at 500-generation intervals thereafter.

Mutation rate measurements. Fluctuation tests²¹ to estimate a given rate of mutation were conducted simultaneously on all strains to provide a controlled comparison. Before testing, strains were revived and regrown for 3 days in the original experimental medium to re-establish the physiological conditions that had prevailed in the evolving populations. In general, 24 or more cultures of a given strain were grown from inocula of approximately 1,000 cells for each fluctuation test. For a given mutation rate measurement, all strains to be compared were grown separately in the same batch of Davis minimal medium. Cultures were grown to stationary phase before selective plating. (An additional 24 h in stationary phase had no discernible effect on the mutation rate in pilot studies.) Final population sizes were estimated by growing and sampling three extra cultures taken at random for each strain and measuring cell densities using a Coulter particle counter; the means were used in mutation rate calculations. Ara⁺ mutants were enumerated on Davis minimal agar medium supplemented with 4 mg ml⁻¹ arabinose; Nal^r mutants were enumerated on LB agar medium²² containing 20 μg ml⁻¹ of nalidixic acid; T5^r mutants were enumerated by plating cultures with excess phage T5 in soft agar.

Time of origin and persistence of mutator phenotypes. Mutator screens were conducted at approximately 500-generation intervals over the 10,000 generations of the experiment. Overnight cultures in 5 ml of Davis minimal medium (supplemented with 1,000 μg ml⁻¹ glucose) were assayed for Nal^r mutant numbers at the stationary phase. Expected distributions of mutants, based on mutator and ancestral mutation rates, were used to generate criteria

by which isolates could be scored as mutator or non-mutator with >95% confidence.

Complementation tests. Strains were transformed with plasmids carrying wild-type alleles for *mutH* (plasmid pGW1899)²³, *mutL* (pGW1842)²³, *mutS* (pGW1811)²³, *uvrD* (pGT26)²⁴, *mutT* (pSK25)²⁵, *dnaQ* (pMM5)²⁶ and *dnaE* (pMK9)²⁷, according to a standard protocol²⁸. Fluctuation tests were conducted as described above, except that all strains were propagated in LB medium (containing 60 µg ml⁻¹ of ampicillin where the strain was plasmid bearing), and five parallel cultures were used per fluctuation test.

Analysis of fluctuation test data. A local computer program using a Luria–Delbrück distribution-generating algorithm²⁹ was used to calculate maximum-likelihood mutation rates from fluctuation test data. Approximate 95% confidence intervals for the mutation rates illustrated in Fig. 1 were calculated from formulae³⁰. Approximate 95% confidence limits for the mutation rates illustrated in Fig. 3 are based on the theoretical variance of the maximum-likelihood estimate of ln(*m*), assuming normality, where *m* is the expected number of mutations per culture.

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Evidence from Turner's syndrome of an imprinted X-linked locus affecting cognitive function

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Turner's syndrome is a sporadic disorder of human females in which all or part of one X chromosome is deleted¹. Intelligence is usually normal² but social adjustment problems are common³. Here we report a study of 80 females with Turner's syndrome and a single X chromosome, in 55 of which the X was maternally derived (45,X^m) and in 25 it was of paternal origin (45,X^p). Members of the 45,X^p group were significantly better adjusted, with superior verbal and higher-order executive function skills, which mediate social interactions⁴. Our observations suggest that there is a genetic locus for social cognition, which is imprinted⁵ and is not expressed from the maternally derived X chromosome. Neuro-psychological and molecular investigations of eight females with partial deletions of the short arm of the X chromosome⁶ indicate that the putative imprinted locus escapes X-inactivation⁷, and probably lies on Xq or close to the centromere on Xp. If expressed only from the X chromosome of paternal origin, the existence of this locus could explain why 46,XY males (whose single X chromosome is maternal) are more vulnerable to developmental disorders of language and social cognition, such as autism, than are 46,XX females⁸.

An increasing number of mammalian genes are known to be subject to genomic imprinting, defined as parental origin-specific differential gene expression⁵. No imprinted gene has yet been described on the X chromosome in humans⁹, although the *Xist* gene has been shown to be imprinted in the mouse¹⁰. We considered that it should be possible to identify the effects of an X-linked imprinted locus by comparing classes of females with Turner's syndrome. In this chromosomal disorder all, or a substantial part, of one X chromosome is missing as a result of non-disjunction (chromosome loss during gametogenesis or early cleavage of the zygote). In 70% of monosomic (45,X) Turner-syndrome females, the single X chromosome is maternal in origin¹; in the remainder it is paternal. Normal females (46,XX) possess both a maternally derived X chromosome (X^m) and a paternally derived X chromosome (X^p), one of which is randomly inactivated in any given somatic cell⁷. In monosomy X, the single chromosome is never inactivated. Differences in physical or behavioural phenotype between 45,X^p and 45,X^m Turner-syndrome subjects might therefore indicate the existence of an imprinted genetic locus.

Impaired social competence and adjustment are frequent in Turner's syndrome³, but a minority have good social skills¹¹. Intelligence is usually normal in monosomic (45,X) cases². We wished to test the hypothesis that 45,X^p females would be distinguishable from 45,X^m females by their social behaviour.

We karyotyped 80 monosomic (45,X) females and eight with deletions of the short arm of one X chromosome (46,XXp-). The