­Stress-Induced Mutagenesis and Complex Adaptation

Yoav Ram and Lilach Hadany

**Email**: YR - [yoavram@post.tau.ac.il](mailto:yoavram@post.tau.ac.il),

LH - [lilach.hadany@gmail.com](mailto:lilach.hadany@gmail.com) (corresponding author)

**Address**: Dept. of Molecular Biology and Ecology of Plants, Tel Aviv University

Tel-Aviv 69978, Israel. Tel. +972.3.640.6886

**Running title:** Complex Adaptation with SIM

**Keywords**: stress-induced variation, mutation rate, mathematical model, simulations, trade-off

# Summary

Because mutations are mostly deleterious, mutation rates should be reduced by natural selection. However, mutations also provide the raw material for adaptation. Therefore, evolutionary theory suggests that the mutation rate must balance between *adaptability* – the ability to adapt – and *adaptedness* – the ability to remain adapted. We model an asexual population crossing a fitness valley and analyze the rate of complex adaptation with and without stress-induced mutagenesis – the increase of mutation rates in response to stress or maladaptation. We show that stress-induced mutagenesis increases the rate of complex adaptation without reducing the population mean fitness, thus breaking the evolutionary trade-off between *adaptability* and *adaptedness*. Our theoretical results support the hypothesis that stress-induced mutagenesis promotes adaptation and provide quantitative predictions of the rate of complex adaptation with different mutational strategies.

# Introduction

There is experimental, clinical and theoretical evidence that high mutation rates increase the rate of adaptation and that during adaptive evolution, constitutive mutators - alleles that constitutively increase the mutation rate - can rise in frequency because of the beneficial mutations they generate [1–3]. However, during evolution in a stable environment, constitutive mutators become associated with poor genetic backgrounds due to increased accumulation of deleterious mutations; this was evidenced both in the lab [4] and in the clinic [5]. Classical models suggest the "reduction principle", which states that natural selection reduces the mutation rate in a stable environment [6,7]. But many adaptations require new beneficial mutations, especially in asexual populations. This tension between the effects of beneficial and deleterious mutations leads to "the rise and fall of the mutator allele" [8], where mutator alleles increase in frequency in a maladapted population, only to be eliminated by natural selection when the population is well-adapted. This dynamic was studied using experimental evolution [9,10], mathematical analysis, and simulations [11–13].

Thus, the mutation rate must balance between two evolutionary traits, as Leigh [14] suggested: *adaptability* – the capacity to adapt to new environmental conditions – and *adaptedness* – the capacity to remain adapted to existing conditions.

Stress-induced mutagenesis (SIM) - the increase of mutation rates in stressed or maladapted individuals - has been demonstrated in several species, including both prokaryotes and eukaryotes [15]. SIM has been observed in lab strains [16,17] and natural populations of *Escherichia coli* [18, but see 19], and in other species of bacteria such as Pseudomonads [20], *Helicobacter pylori* [21], *Vibrio cholera* [22] and *Streptococcus pneumonia* [23]. SIM has also been observed in yeast [24,25], algae [26], nematodes [27], flies [28], and human cancer cells [29]. Several stress responses regulate the mutation rate in bacteria by shifting replication to error-prone DNA polymerases [30] and by inhibiting the mismatch repair system [31]. These stress responses include the SOS DNA-damage response, the RpoS-controlled general or starvation stress response, and the RpoE membrane protein stress response [32].

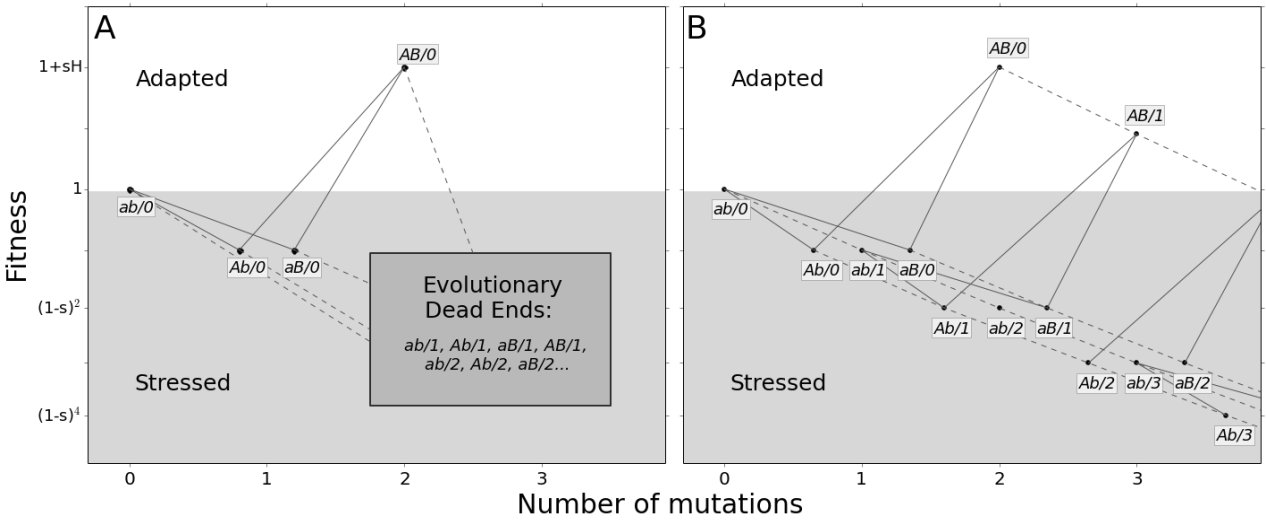
It is still not clear how SIM affects evolution and adaptation. Some authors have proposed that SIM has a significant impact on *adaptability* or *evolvability* [17,33,34], but there is no theoretical treatment of this impact. On the other hand, the effect of SIM on *adaptedness* was studied with deterministic [35] and stochastic [36] models. These articles showed that without beneficial mutations SIM doesn't affect the mean fitness of asexual populations in stable environments, in contrast with constitutive mutagenesis, which decreases the population mean fitness. More recently, we have shown that with rare beneficial mutations, if maladapted individuals increase their mutation rate then the population mean fitness of asexual populations increases [37].

Here, we analyze population genetics models of adaptive evolution to explore the rate of complex adaptation on rugged fitness landscapes, in which adaptations require two separately deleterious mutations [38,39]. We develop analytic approximations and stochastic simulations and compare normal, constitutive, and stress-induced mutagenesis. We show that stress-induced mutagenesis can break the trade-off between *adaptability* and *adaptedness* by increasing the rate of complex adaptation without decreasing the population mean fitness.

# ­Model

We model a population of *N* haploid asexual individuals with a large number of loci in full linkage. The model includes the effects of mutation, selection, and genetic drift. Individuals are characterized by their genotype in two specific bi-allelic loci – *ab*, *Ab*, *aB*, and *AB* – and by the number of deleterious mutations they carry in the rest of the non-specific loci. For example, *aB/3* is the *aB* genotype with additional three deleterious mutations in non-specific loci.

We focus on adaptation to a new rugged fitness landscape. The fitness of the wildtype *ab/0* is 1, the fitness of the single mutants *Ab*/0 and *aB*/0 is 1-*s*, and the double mutant *AB/0* has the highest fitness 1+*sH*, where *s* is the selection coefficient and *H* is the relative advantage of the double mutant. This is the simplest case of a rugged fitness landscape: the single mutants *Ab* and *aB* are fitness valleys between the local and global fitness peaks *ab*/0 and *AB/0* (Figure 1).

****

**Figure 1 – Adaptation on a rugged fitness landscape.** The figure shows the fitness of the possible genotypes, which are represented by the allele combination at the specific loci (*ab*, *Ab*, *aB*, and *AB*) and the number of deleterious alleles across the genome following the forward-slash ('/'). The y-axis represents fitness: the wildtype *ab/0* has fitness 1; the fittest genotype *AB/*0 has fitness 1+*sH*; deleterious alleles, either at the *A/a* and *B/b* loci, or at the non-specific loci, reduce fitness by 1-*s*. The x-axis represents the number of accumulated mutations (genotypes are jittered to increase separation). Solid lines represent mutations at the *a/A* and *b/B* loci, occurring with probability *µ*. Dashed lines represent deleterious mutations in the rest of the genome, occurring with rate *U*. Mutagenesis is induced in stressed genotypes with fitness <1 (gray background). Fit genotypes, with fitness ≥1, do not hypermutate (white background). **(A)** In the analytic model genotypes with deleterious alleles in non-specific loci are considered "Evolutionary Dead Ends" and do not contribute to adaptation. **(B)** In the simulations individuals can accumulate up to 25 deleterious alleles (the figure only shows three). Multiple mutations can occur simultaneously but are not shown for simplicity of the illustration.

Deleterious mutations in the non-specific loci independently (multiplicatively) reduce the fitness of the individual by 1-*s*. Mutations occur in the specific loci with probability *µ*. The number of new mutations per replication in the rest of the genome (the non-specific loci) is Poisson distributed with an average *U*. The model neglects back-mutations and compensatory mutations due of their minor short term effects.

We consider three mutational strategies: normal mutagenesis (NM), where there is no increase in the mutation rate; constitutive mutagenesis (CM), where all individuals always increase their mutation rate by *τ*,the mutation rate fold increase; and stress-induced mutagenesis (SIM), where only stressed or maladapted individuals increase their mutation rate by *τ*. Individuals are considered stressed if their fitness is below a specific threshold, so stress can be caused by a deleterious mutation (either in the specific *A/a* and *B/b* loci or in non-specific loci); there is indeed evidence that in bacteria, starvation [16,40], growth in stationary-phase [41], and several antibiotics [42,43] reduce fitness, induce mutagenesis via regulated stress responses, and are mitigated by beneficial mutations.

The main analysis assumes that the effect of SIM on the mutation rate of an individual with fitness *ω* is

|  |  |
| --- | --- |
| . | (1) |

This equation models a scenario in which an environmental change – *i.e.*, appearance of a new ecological niche or a new carbon source – provides an opportunity for adaptation without affecting the fitness of the wildtype (*ab/0*). We also study a different scenario in which the environmental change reduces the absolute fitness of the wildtype so that it is also stressed – see section 3.5. The threshold relationship between fitness and the mutation rate eq. 1 is relaxed in Appendix E in which we explore continuous relationships between fitness and the mutation rate.

We are interested in calculating the adaptation rate of a population homogenous for each of the above mutational strategies (NM, CM, or SIM). The adaptation process is separated into two ­­­­distinct stages. In the first stage, a double mutant *AB* appears in the population, usually in a single copy. In the second stage, the double mutant either goes to extinction or avoids extinction, increases in frequency, and goes to fixation.

We analyzed this model with two methods. The first is analytic (Figure 1A), in which we assume that: (i) genotypes with deleterious backgrounds (deleterious alleles in the non-specific loci) do not contribute to the adaptation process; and (ii) the number of deleterious alleles per individual before the appearance of a double mutant is at a mutation-selection balance (MSB) and is Poisson distributed with mean *U/s* (Haigh 1978). The former assumption requires that mutation is weaker than selection (); the later assumption only requires that mutation is not much stronger than selection. Specifically, the expected number of mutation-free individuals is at least one: [45].

The second method is a stochastic Wright-Fisher simulation with selection, mutation and genetic drift (Figure 1B), in which: (i) individuals with a deleterious background can contribute to adaptation; (ii) a mutation-free population evolves towards a MSB without assuming a Poisson distribution of the number of deleterious alleles.

## Wright-Fisher simulations

We track the number of individuals in each genotype class: *ab/x*, *Ab/x*, *aB/x*,and *AB/x*, where *x*≥0 is the number of deleterious alleles in non-specific loci. The simulations start with a single-peak smooth fitness landscape (the fitness of *AB/x*  is (1-*s*)2+*x*) and a mutation-free population (all individuals start in the optimal *ab/0* genotype with fitness 1) that accumulates deleterious mutations over the first 5,000 generations of the simulation. With *s*=0.05 and 0.005, 180 and 1,800 generations are enough for the average number of deleterious alleles per individual to reach 99.99% of its MSB value, *U/s* [46].

After 5,000 generations the fitness landscape changes to a rugged one, making *AB* the optimal genotype with fitness 1+*sH* (Figure 1B). The simulation then proceeds until an *AB* genotype appears and either fixates in the population or goes extinct (either all or no individuals are in the *AB* classes, respectively). Therefore, each simulation provides one sample of the waiting time for the appearance of a double mutant and one sample of the probability of fixation of a double mutant. At least 1,000 simulations were performed for each parameter set.

Table 1 summarizes the model parameters with estimated values for *E. coli*.

# Results

## ­­­­Appearance of a double mutant

We are interested in the waiting time for the appearance of a double mutant *AB* either by a double mutation in a wildtype individual *ab*, or via a single mutation in a single mutant *Ab* or *aB* (Figure 1A). Denoting the population size by *N,* we note that (i) if then double mutants are already expected at the MSB and adaptation will not require new mutations; (ii) if then no single mutants are expected at the MSB and double mutants must be generated by a double mutation in a wildtype individual. In this case, increasing the mutation rate of individuals with fitness below 1 will have no effect on the appearance of the double mutant and there is no point in analyzing the effect of SIM.

Combining the two constraints we get this constraint on the population size *N*: . This constraint is reasonable for bacterial populations (see Table 1).

The frequencies of wildtype (*ab*) and single mutants (*aB* and *Ab* combined) that are mutation-free at the MSB are roughly and , respectively. The probability that an offspring of a wildtype or single mutant parent is a double mutant *AB* is and , respectively. The probability that such an offspring is also mutation-free in the rest of its genome (the only mutations that occurred were at the specific loci) is . Therefore, the probability *q* that a random offspring is a double mutant, given there are no double mutants in the current generation, is approximated by

|  |  |
| --- | --- |
| . | (2) |

The first expression assumes that individuals with a deleterious background don't contribute to adaptation and that the MSB distribution of deleterious alleles is Poisson. The second expression also assumes that mutation is much weaker than selection: *U<<*s.

With SIM the mutation rate of single mutants is increased *τ*-fold and the probability that a random offspring is a double mutant is

|  |  |
| --- | --- |
| . | (3) |

These expressions use the same assumptions as in eq. 2. The second expression also assumes that .

Appendix A includes full derivations of the above equations and Figure S1 compares them with simulations results.

## Fixation probability of the double mutant

Assuming an advantage to the double mutant (*H*>1)and a large population size (see the above constraint on *N*), a double mutant has two possible fates after its appearance: fixation or extinction. Following Eshel [47], the fixation probability *ρ* of the double mutant is (see Appendix B)

|  |  |
| --- | --- |
| . | (4) |

That is, the fixation probability of the double mutant is roughly twice its adaptive advantage. This is a classic result of population genetics theory [48,49].

The fixation probability with SIM equals that of NM and CM because the mutation rate of the wildtype *ab* equals that of the double mutant *AB* (but see an exception in section 3.5).

## Adaptation rate

From the probability *q* that a random offspring is a double mutant, we can derive the probability that one or more double mutants appear in the next generation: . This is a good approximation because *Nq* is very small due to the constraint on *N*. Once a double mutant appears it goes to fixation with probability *ρ*.

When fixation is much faster than appearance of the double mutant *AB*, the time for adaptation *T* can be approximated by the waiting time for a double mutant that goes to fixation. This waiting time follows a geometric distribution with rate *Nqρ* and therefore the adaptation rate *ν* (the inverse of the waiting time for adaptation) is approximately

|  |  |
| --- | --- |
| . | (5) |

Plugging eqs. 2-4 in eq. 5, we get these approximations:

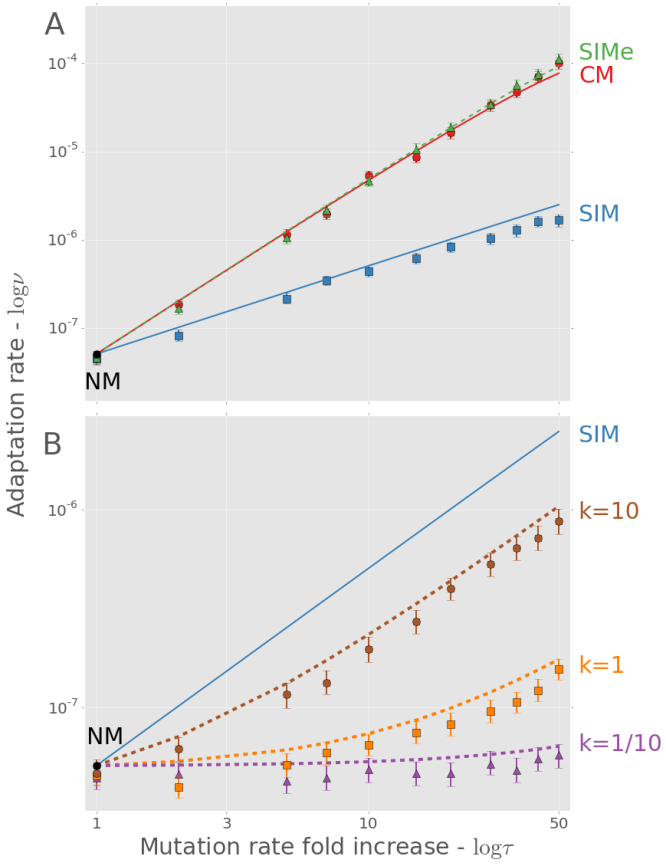
|  |  |
| --- | --- |
|  | (6) |
|  | (7) |
|  | (8) |

NM is normal mutagenesis, CM is constitutive mutagenesis, and SIM is stress-induced mutagenesis. The middle expression in each equation is the full approximation, which assumes a Poison distribution and no contribution of deleterious genotypes to adaptation. The right hand sides are first order approximations that assume mutation is much weaker than selection ( for NM and SIM, for CM) and that 1<τ<1/*U*. See Table 1 for description of model parameters and an article by Weinreich and Chao [50] for a result similar to eq. 6.

The main conclusions from eqs. 6-8: First, adaptation with CM is faster than with NM. Second, adaptation with SIM is also faster than with NM, but not as fast as with CM because the mutation-free wildtype (*ab/0*) does not hypermutate.

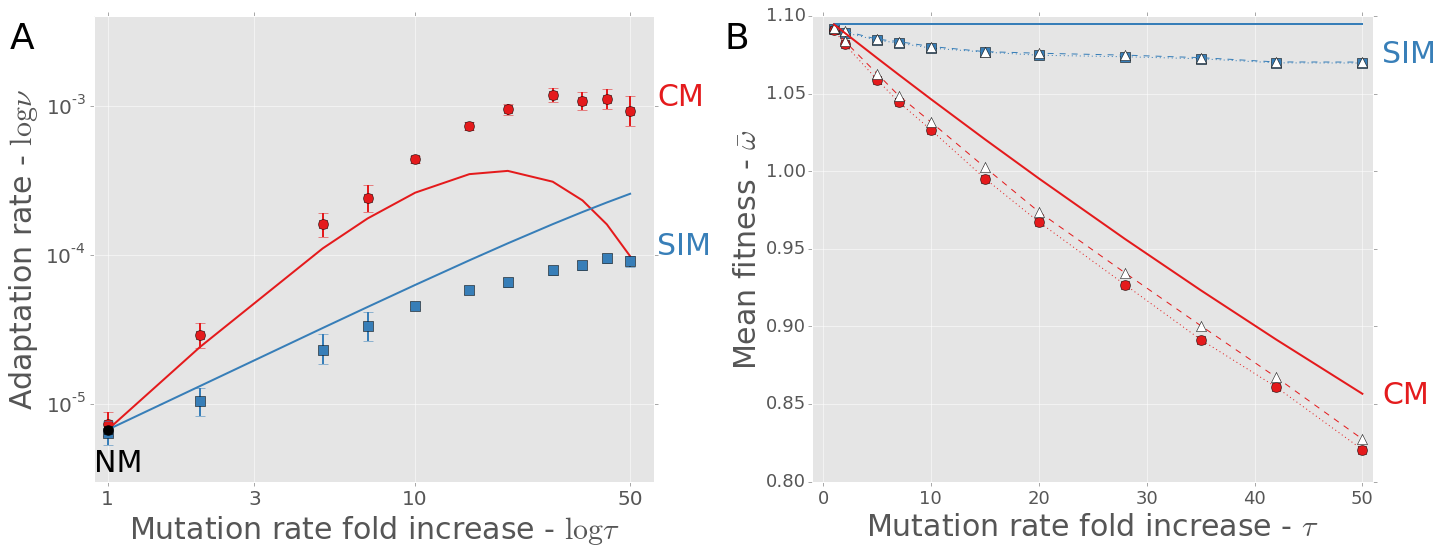
If mutation is weaker than selection () then the adaptation rate with CM increases with *τ2* and the adaptation rate with SIM increases with *τ*. In addition, because the fixation probability is the same for NM, CM and SIM, the differences in the adaptation rate are due to differences in the appearance probability *q* (Figure S1); see section 3.5 for a different scenario in which SIM also increases the fixation probability.

Figure 2A compares the analytic approximations with simulation results for the weak mutation regime (). This regime is relevant for asexual microbes in which the deleterious mutation rate is generally 10-4-10-3 mutations per genome per generation and selection coefficients are estimated to be between 10-1 and 10-2 (see Table 1). When the mutation rate fold increase *τ* is high (>10), the approximations slightly overestimate the adaptation rate because the double mutant *AB* is more likely to appear on a deleterious background (*AB/1* instead of *AB/0*). Because the fitness of *AB/1* is higher than that of the wildtype *ab/0* (this happens because *H*>(1-*s*)-1≈1+*s*), the double mutant can go to fixation even when it appears on a deleterious background, sweeping the deleterious alleles with it to fixation in a process called "genetic hitch-hiking" [51]. However, these sweeps result in a lower fixation probability for the double mutant (Figure S2).

****

**Figure 2 – Complex adaptation with different mutational strategies.** The figure shows the adaptation rate *ν* as a function of the mutation rate increase *τ* (both in log scale). **(A)** A black circle is normal mutagenesis (NM*; τ*=1); solid line with circles is constitutive mutagenesis (CM); solid line with squares is stress-induced mutagenesis (SIM); dashed lines with triangles is stress-induced mutagenesis with environmental stress (SIMe; see section 3.5). Lines are analytic approximations; markers are the means of stochastic simulation results; error bars represent 95% confidence interval of the mean (at least 1,000 replicates per point; computed with bootstrap with 1,000 samples per point). Parameters (see Table 1): *U*=0.0004, *s*=0.05, *β*=0.0002, *H*=2, *N*=106. **(B)** Markers are the results of adaptation with SIM with different continuous relationships between fitness and mutation rate, defined by a curvature parameter *k* and mutation rate fold increase *τ*=10 (see Online Appendix E and Figure E1)**.** Each dashed line is an analytic approximation of the continuous SIM next to it, using a SIM strategy with a threshold relationship and with *τ*=4.61, 1.45, and 1.05, top to bottom. Online version in colour.

The adaptation rate of SIM with continuous relationships between fitness and mutation rate (Online Appendix E) is similar to that of SIM with threshold relationships (Figure 2B). This is because the main factor determining the adaptation rate is the mutation rate increase of the wildtype and the single mutants (*ab*, *aB*, and *Ab*), because individuals with more than a single mutation do not have a significant contribution to adaptation.



**Figure 3 – Adaptation with strong mutation.** When the deleterious mutation rate is high – here *U=s/10* – the adaptation process can lead to hitch-hiking of deleterious alleles with the beneficial double mutant. **(A)** The adaptation rate *ν* as a function of the mutation rate increase *τ* (both in log scale). A black circle for normal mutagenesis (NM*; τ*=1); red solid line and circles for constitutive mutagenesis (CM); blue solid line and squares for stress-induced mutagenesis (SIM). Lines are analytic approximations. Markers are the means of stochastic simulations results. Error bars represent 95% confidence interval of the mean (at least 1,000 simulations per point; computed with bootstrap with 1,000 samples per point). Parameters (see Table 1): *U*=0.005, *s*=0.05, *β*=0.0002, *H*=2, *N*=106. **(B)** The population mean fitness after successful fixation of the beneficial double mutant as a function of the mutation rate increase *τ*. Solid lines are analytic approximations neglecting adaptation from deleterious background *(e-τU*(1+*sH*)); dotted lines with filled squares (SIM) and circles (CM) are the means of stochastic simulation results; dashed lines with white triangles are predictions based on the genotype on which *AB* appeared in the simulations, including MSB but disregarding the effects of drift during the fixation process (which only has a significant effect with CM due to higher mutation rates in the wildtype). Error bars are too small to see. Same parameter values as in panel A. Online version in colour.

## The trade-off between *adaptability* and *adaptedness*

Next, we explore how different mutational strategies (NM, CM and SIM) balance between *adaptability* – the ability to adapt to new conditions– and *adaptedness* – the ability to remain adapted to current conditions. For this purpose we define *adaptedness* asthe population mean fitness in a stable environment and *adaptability* as *ν the* rate of complex adaptation.

We used the above approximations (eqs. 6-8) to calculate the rate of complex adaptation of populations with NM, CM and SIM. We also extended an existing model [37] to calculate the population mean fitness at the mutation-selection balance. This extended model includes rare back- or compensatory mutations (which have a stronger effect on mutation-selection balance dynamics than on adaptive dynamics) and allows more than one mutation to occur in the same individual and generation. The details of this model and the calculation of population mean fitness with various mutational strategies are given in Online Appendix D.

The mutation rate with CM is constant and uniform across the population, and the population mean fitness mainly depends on the fitness and mutation rate of the fittest individuals. Therefore, the population mean fitness decreases when the mutation rate increases; this decrease is due to generation of deleterious mutations in the fittest individuals. The adaptation rate, however, increases with the mutation rate (eq. 7). This trade-off between *adaptability* and *adaptedness* constraints the population: after a long period of environmental stability it can lose the potential for adaptation, and after a long period of environmental change the population can be susceptible to reduced fitness and mutational meltdowns [52].

However, this trade-off between *adaptability* and *adaptedness* can be broken if mutation rates are not uniform across the population. Increased mutation rates in unfit individuals increase the population mean fitness, as long as beneficial (or compensatory) mutations can occur [37]. Figure D1 shows this advantage of SIM over NM in terms of the difference in population mean fitness (). Moreover, increased mutation rates in unfit individuals also increase the adaptation rate (eq. 8; Figure 2). Therefore, SIM breaks the trade-off between *adaptability* and *adaptedness*.



**Figure 4 – The trade-off between *adaptedness* and *adaptability*.** The figure shows the relative *adaptedness* and the relative *adaptability* of different mutational strategies in comparison to normal mutagenesis (NM). *Adaptedness* is defined by the population mean fitness at MSB, (see Online Appendix D). *Adaptability* is defined by the rate of complex adaptation, (eqs. 6-8). Constitutive mutagenesis (CM) increases the mutation rate of all individuals *τCM*-fold; Stress-induced mutagenesis (SIM) increases the mutation rate of stressed individuals *τSIM*-fold; Mixed strategies (dashed line) increase the mutation rate of all individuals *τCM*-fold and of stressed individuals an additional *τSIM*-fold. SIM breaks off the *adaptability-adaptedness* trade-off of CM, increasing the *adaptability* without compromising the *adaptedness* of the population. Parameters (see Table 1): *N*=106, *U*=­0.0004, *β*=0.0002, *s*=0.05, *H*=2, *τ*<<s/*U* (Online version in colour.).

Figure 4 shows the adaptation rate and population mean fitness of CM and SIM compared to NM for different values of *τ*, the mutation rate fold increase.

Any realistic rate of adaptation *ν* can be realized using both CM and SIM. The highest mean fitness will always be attained with SIM, which has a small advantage over NM (that cannot be seen in this figure, but see Figure D1) due to the increased generation of beneficial mutations in individuals with low fitness. If for some rate of adaptation the mutation rate fold increase *τ* required by SIM is too high (*i.e.*, *τU*>*s*), the same adaptation rate can be realized by a mixed strategy (dashed line in Figure 4). For example, a 96-fold increase in adaptation rate can be achieved with CM with *τ*=10, with SIM with *τ*=96, or with a mixed strategy with *τCM*=7and *τSIM*=2 in which all individuals increase their mutation rate 7-fold and stressed individuals further increase their mutation rate 2-fold. However, these increases in adaptation rates have a price: the mutational load will decrease the population mean fitness from 0.9996 with NM to 0.996 with CM and 0.9972 with the mixed strategy. This price in not paid by populations with SIM because the mean fitness mainly depends on the mutation rate of fit individuals.

## Environmental stress

So far, we considered the case where the environmental change creates an opportunity for adaptation without affecting the absolute fitness of the population – for example, a new ecological niche can be favorable without affecting the well-being of the current population. In that scenario, the wildtype *ab* was not stressed and did not hypermutate.

Next, we consider a different scenario in which an environmental change affects the well-being of the entire population: for example, exposure to an antibiotic drug or a host immune response. In this case the environmental change doesn't just create an opportunity for adaptation but also causes stress in the entire population. We use a subscript *e* to denote quantities related with this scenario.

As before the double mutant *AB* is resistant to the stress (*i.e.* the drug or immune response) and therefore has a higher fitness than either the wildtype or the non-resistant single mutant*s*. However, in this scenario the wildtype *ab* is also stressed and therefore hypermutates with SIM (compare with eq. 1):

|  |  |
| --- | --- |
| . | (9) |

This scenario has an important biological relevance, as SIM has been implicated in the evolution of drug resistance in bacteria and yeast [34,53,54] and could be involved in the evolution of pathogen virulence and the evolution of drug resistance and progression in cancer cells [55].

We assume that after the environmental change the SIMe population has reached a new MSB [46] with mutation rate, before the appearance of the double mutant (with s=0.05 and *U*=0.0004, for example, the average number of deleterious mutations is 0.99∙*U/s* after 90 generations, whereas the adaptation time is well over 1,000 generations). Under this assumption, the adaptation rate with SIMe is (see Appendix C for full derivation)

|  |  |
| --- | --- |
| . | (10) |

That is, adaptation with SIMe is faster than with CM (Figure 2A). The fixation probability of double mutants is higher with SIMe than with CM, because the mutation rate of double mutants is lower than that of the rest of the population. This difference in mutation rates confers an additional selective advantage to the double mutants (see Appendix C) which increases their fixation probability:

|  |  |
| --- | --- |
| . | (11) |

This additive advantage increases linearly with *τ* with a slope of and can be significant: for *s*=0.05, *H*=2 and *U*=0.0004, increasing the mutation rate of stressed individuals 10-fold increases the fixation probability by 3.6%. The increased fixation probability was verified by simulations (Figure S2).

# Discussion

We studied the effect of stress-induced mutagenesis (SIM) on both the *adaptability* – the capacity of populations to adapt to new complex conditions– and the *adaptedness* – the ability of populations to stay adapted to existing conditions [14]. We showed that SIM breaks the trade-off between *adaptability* and *adaptedness*, allowing rapid adaptation to complex environmental challenges without compromising the population mean fitness in a stable environment.

In addition to the pure strategies of constitutive mutagenesis (CM) and SIM, our model also considers a mixed mutational strategy. There are two examples of such a mixed strategy. First, if individuals have incomplete information regarding their condition (this is the case in most realistic biological scenarios) then we expect errors in the induction of mutagenesis: induction of mutagenesis without stress and failure to induce mutagenesis under stress. In this case the population would, on average, use a mixed strategy. Second, a mutator allele can increase the mutation rate constitutively and further increase it under stress – for example, a recent study with *Pseudomonas aeruginosa* found that although the *mutS*, *mutY* and *mutM* mutator alleles always increase the mutation rate in comparison with the wildtype, the level of this increase depends on the level of stress the cell experiences [56].

Although our model focuses on rugged landscapes in which single mutants have lower fitness (*s*>0), one can also consider cases in which single mutants have intermediate fitness (0>*s>sH*) [57]. We previously shown that SIM is favored by selection in these cases [37]; however, further work is required to analyze the effect of SIM on adaptation rate such scenarios.

Our model does not assume direct fitness costs for any of the mutational strategies. A "cost of DNA replication fidelity" [58] – the energy and time expended in order to maintain a low mutation rate – could make both CM and SIM more successful. The "cost of fidelity" may require further study, but empirical evidence suggests that it doesn't play an important role in the evolution of the mutation rate [40,59–61]. Another fitness cost might be associated with the regulation of the mutation rate: for individuals to determine if their condition calls for the induction of mutagenesis, they must invest resources and energy in costly sensory mechanisms. However, such mechanisms already exist for various unrelated purposes, such as the maintenance of cell cycle and homeostasis. Therefore, we consider these mechanisms as "free" in terms of fitness costs. Moreover, in *E. coli* mutagenesis is induced by several stress responses that serve other cellular functions [16,32], and this is probably the case in other organisms as well.

Our model focuses on asexual populations, ignoring recombination, segregation, and sexual reproduction. These mechanisms are important for adaptation on a rugged fitness landscape both because they help to cope with deleterious mutations and because they allow different single mutants to produce double mutants without an increased mutation rate. We expect that recombination will reduce the advantage of SIM over NM in terms of population mean fitness [35], direct competitions [62], and adaptation rate (due to the Fisher-Muller effect).

Mean fitness and adaptation rate are both population-level traits. But simply because SIM has the most efficient balance between these traits doesn't mean it will necessarily evolve, because individual-level selection and population-level selection can act in opposing directions. In a previous article we have demonstrated that 2nd order selection can lead to the evolution of SIM [37]: in an asexual population evolving on a smooth fitness landscape, selection favored SIM over both NM and CM. In the current article we show that selection also favors SIM on a rugged fitness landscape (Online Appendix F).

Several authors have suggested that the mutation rate must balance between *adaptability* and *adaptedness*: Kimura found a mutation rate that balances between mutational and substitutional load [6]; Johnson and Barton found an optimal mutation rate that balances the generation of beneficial and deleterious mutations during adaptation [63]; Leigh found an optimal mutation rate that balances the generation of deleterious mutations and maintenance of standing variation in a changing environment [64]; Komarova and Wodartz found an optimal rate of chromosome loss that balances the unmasking of recessive alleles and genetic load during carcinogenesis [65]; Komarova et al. and Agur et al. found a time-dependent mutation rate strategy that optimizes carcinogenesis [66] and adaptive immune response [67], respectively. Compared to these findings, SIM operates at a larger time scale because it allows individuals to switch between rates optimized for stressful and benign conditions according to the environmental and genetic circumstances.

Complex traits, coded by multiple genes, present an open evolutionary problem, first described by Sewall Wright in 1931: if different alleles are separately deleterious but jointly advantageous, how can a population evolve from one co-adapted gene complex to a fitter one, crossing a less fit "valley"? Wright suggested the "shifting-balance theory of evolution" [38,39]. His solution is valid [68–70] but possibly limited to specific parameter ranges [71–74]. As a result, other mechanisms have been proposed: increased phenotypic variance after population bottlenecks [75]; environmental fluctuations [76]; environmental heterogeneity [77]; fitness-associated recombination [78]; stochastic tunneling in large asexual populations [79]; and intermediate recombination rates [80]. Our model of complex adaptation is similar to that of Weinreich and Chao [50], but our model includes various mutational strategies and the effects of stress and deleterious mutations. Our results (Figure 2) suggest that SIM can help resolve the problem of fitness valley crossing by reducing the time required for a population to shift an adaptive peak.

Our results provide theoretical basis to the conjecture that SIM facilitates adaptation. This conjecture can be tested experimentally, for example, with *E. coli*, where it is possible to interfere with the regulation of mutagenesis [34]. The adaptation time with and without SIM can be measured in an experimental population adapting on a two-peak fitness landscape [81]. These measurements can then be compared to our analytic approximations to determine the relative advantage and disadvantage of the different mutational strategies.

## Conclusions

Stress-induced mutagenesis has been implicated as a driver of adaptive evolution for several decades [17,33,82]. We provide theoretical treatment of this concept. Our results show that stress-induced mutagenesis increases the rate of complex adaptation, and that in contrast to constitutive mutagenesis it does not jeopardize the fitness of populations under stable conditions. Because mutation is a fundamental force in every biological system, these results have important implications on many fields in the medical and life sciences, including epidemiology [53], oncology [83,84], ecology [18], and evolutionary biology [85].

# Acknowledgments

We thank U. Obolski for advice on statistical analysis. We are grateful to A. F. Agrawal, T. F. Cooper//the editor// and two anonymous reviewers for helpful comments on an earlier version of the manuscript. This research has been supported in part by the Israeli Science Foundation 1568/13 (LH) and by Marie Curie reintegration grant 2007–224866 (LH).

# Data accessibility

The data used in this study, as well as the code necessary to reproduce the figures, will be made available on Dryad before publication.

# Appendixes

## Appendix A: Appearance of a double mutant

In the following analysis we assume that and – see model overview for details. This also means that and . The probability *q* that a random offspring in the next generation is *AB* given there are no *AB* in the current generation can be approximated by:

.

Using the above assumptions, this resolves to:

.

Taking the derivative with respect to *U* and denoting (*g* can be thought of as the number of non-specific loci in the genome):

.

So *q* increases with *U* because the right hand side is guaranteed to be true under the assumption .

For a population with SIM

.

The last approximation assumes that *.* Rearranging the last result, we find the approximation

|  |  |
| --- | --- |
| . |  |

Taking the derivative with respect to *τ*,

,

because *q*, *U*, and *τ* are all positive. So the condition guarantees that *qSIM*increases with *τ*, and it is also sufficient for (not shown).

## Appendix B: Fixation of a double mutant

Following Eshel [47], the fixation probability *ρ* of the double mutant *AB* is

,

where *α* is the fitness of the double mutant relative to the population mean fitness and assuming that fitness is measured by the average number of progeny which is Poisson distributed:

.

Here, we only consider progeny without new deleterious mutations; their fraction is *e-U*. This factor cannot be ignored because there is variation in mutation rates within the population (see "minor technical point" by Johnson and Barton [63]). At this stage, double mutants are still very rare, so we can use the population mean fitness at the MSB. The population mean fitness can be approximated by (see *supporting information*). Therefore,

Assuming *sH* is small () we can approximate this by

.

## Appendix C: Fixation of a double mutant with SIMe

With SIMe the mutation rate of *ab* is *τU* while that of *AB* is only *U*. We assume the population reached a MSB before the fixation of *AB* because convergence to MSB [46] is much faster than adaptation. Following the derivation in Appendix B, the relative fitness of SIMe is

.

Plugging that in the fixation probability,

.

This can be simplified by a 1st order approximation for :

.

Because , the right hand side is greater than 1. Therefore,

.

Because the appearance with SIMe is the same as with CM, the adaptation rate with SIMe can be written as

.

# References

1. Sniegowski, P. D., Gerrish, P. J., Johnson, T. & Shaver, A. 2000 The evolution of mutation rates: separating causes from consequences. *BioEssays* **22**, 1057–66. (doi:10.1002/1521-1878(200012)22:12<1057::AID-BIES3>3.0.CO;2-W)

2. Denamur, E. & Matic, I. 2006 Evolution of mutation rates in bacteria. *Mol. Microbiol.* **60**, 820–7. (doi:10.1111/j.1365-2958.2006.05150.x)

3. De Visser, J. A. G. M. 2002 The fate of microbial mutators. *Microbiology* **148**, 1247–52.

4. Funchain, P., Yeung, A., Stewart, J. L., Lin, R., Slupska, M. M. & Miller, J. H. 2000 The consequences of growth of a mutator strain of *Escherichia coli* as measured by loss of function among multiple gene targets and loss of fitness. *Genetics* **154**, 959–70.

5. Montanari, S. et al. 2007 Biological cost of hypermutation in *Pseudomonas aeruginosa* strains from patients with cystic fibrosis. *Microbiology* **153**, 1445–54. (doi:10.1099/mic.0.2006/003400-0)

6. Kimura, M. 1967 On the evolutionary adjustment of spontaneous mutation rates. *Genet. Res.* **9**, 23–34. (doi:10.1017/S0016672300010284)

7. Liberman, U. & Feldman, M. W. 1986 Modifiers of mutation rate: a general reduction principle. *Theor. Popul. Biol.* **30**, 125–42.

8. Giraud, A., Radman, M., Matic, I. & Taddei, F. 2001 The rise and fall of mutator bacteria. *Curr. Opin. Microbiol.* **4**, 582–585. (doi:10.1016/S1369-5274(00)00254-X)

9. Sniegowski, P. D., Gerrish, P. J. & Lenski, R. E. 1997 Evolution of high mutation rates in experimental populations of E. coli. *Nature* **387**, 703–5. (doi:10.1038/42701)

10. Wielgoss, S. et al. 2012 Mutation rate dynamics in a bacterial population reflect tension between adaptation and genetic load. *Proc. Natl. Acad. Sci.* **110**, 222–227. (doi:10.1073/pnas.1219574110)

11. Taddei, F., Radman, M., Maynard Smith, J., Toupance, B., Gouyon, P.-H. & Godelle, B. 1997 Role of mutator alleles in adaptive evolution. *Nature* **387**, 700–2. (doi:10.1038/42696)

12. Kessler, D. & Levine, H. 1998 Mutator Dynamics on a Smooth Evolutionary Landscape. *Phys. Rev. Lett.* **80**, 2012–2015. (doi:10.1103/PhysRevLett.80.2012)

13. Tenaillon, O., Toupance, B., Le Nagard, H., Taddei, F. & Godelle, B. 1999 Mutators, population size, adaptive landscape and the adaptation of asexual populations of bacteria. *Genetics* **152**, 485–93.

14. Leigh, E. G. J. 1970 Natural Selection and Mutability. *Am. Nat.* **104**, 301–305.

15. Galhardo, R. S., Hastings, P. J. & Rosenberg, S. M. 2007 Mutation as a stress response and the regulation of evolvability. *Crit. Rev. Biochem. Mol. Biol.* **42**, 399–435. (doi:10.1080/10409230701648502)

16. Foster, P. L. 2007 Stress-induced mutagenesis in bacteria. *Crit. Rev. Biochem. Mol. Biol.* **42**, 373–97. (doi:10.1080/10409230701648494)

17. Rosenberg, S. M., Shee, C., Frisch, R. L. & Hastings, P. J. 2012 Stress-induced mutation via DNA breaks in *Escherichia coli*: A molecular mechanism with implications for evolution and medicine. *BioEssays* , 1–8. (doi:10.1002/bies.201200050)

18. Bjedov, I., Tenaillon, O., Gérard, B., Souza, V., Denamur, E., Radman, M., Taddei, F. & Matic, I. 2003 Stress-induced mutagenesis in bacteria. *Science (80-. ).* **300**, 1404–9. (doi:10.1126/science.1082240)

19. Katz, S. & Hershberg, R. 2013 Elevated mutagenesis does not explain the increased frequency of antibiotic resistant mutants in starved aging colonies. *PLoS Genet.* **9**, e1003968. (doi:10.1371/journal.pgen.1003968)

20. Kivisaar, M. 2010 Mechanisms of stationary-phase mutagenesis in bacteria: mutational processes in pseudomonads. *FEMS Microbiol. Lett.* **312**, 1–14. (doi:10.1111/j.1574-6968.2010.02027.x)

21. Kang, J. M., Iovine, N. M. & Blaser, M. J. 2006 A paradigm for direct stress-induced mutation in prokaryotes. *FASEB J.* **20**, 2476–85. (doi:10.1096/fj.06-6209com)

22. Baharoglu, Z. & Mazel, D. 2011 *Vibrio cholerae* triggers SOS and mutagenesis in response to a wide range of antibiotics: a route towards multiresistance. *Antimicrob. Agents Chemother.* **55**, 2438–41. (doi:10.1128/AAC.01549-10)

23. Henderson-Begg, S. K., Livermore, D. M. & Hall, L. M. C. 2006 Effect of subinhibitory concentrations of antibiotics on mutation frequency in *Streptococcus pneumoniae*. *J. Antimicrob. Chemother.* **57**, 849–54. (doi:10.1093/jac/dkl064)

24. Heidenreich, E. 2007 Adaptive mutation in *Saccharomyces cerevisiae*. *Crit. Rev. Biochem. Mol. Biol.* **42**, 285–311. (doi:10.1080/10409230701507773)

25. Rodriguez, G. P., Romanova, N. V, Bao, G., Rouf, N. C., Kow, Y. W. & Crouse, G. F. 2012 Mismatch repair-dependent mutagenesis in nondividing cells. *Proc. Natl. Acad. Sci.* **109**, 6153–8.

26. Goho, S. & Bell, G. 2000 Mild environmental stress elicits mutations affecting fitness in Chlamydomonas. *Proc. R. Soc. B Biol. Sci.* **267**, 123–9. (doi:10.1098/rspb.2000.0976)

27. Matsuba, C., Ostrow, D. G., Salomon, M. P., Tolani, A. & Baer, C. F. 2012 Temperature, stress and spontaneous mutation in *Caenorhabditis briggsae* and *Caenorhabditis elegans*. *Biol. Lett.* , 8–12. (doi:10.1098/rsbl.2012.0334)

28. Sharp, N. P. & Agrawal, A. F. 2012 Evidence for elevated mutation rates in low-quality genotypes. *Proc. Natl. Acad. Sci.* **109**, 6142–6. (doi:10.1073/pnas.1118918109)

29. Bristow, R. G. & Hill, R. P. 2008 Hypoxia and metabolism: Hypoxia, DNA repair and genetic instability. *Nat. Rev. Cancer* **8**, 180–92. (doi:10.1038/nrc2344)

30. Ponder, R. G., Fonville, N. C. & Rosenberg, S. M. 2005 A switch from high-fidelity to error-prone DNA double-strand break repair underlies stress-induced mutation. *Mol. Cell* **19**, 791–804. (doi:10.1016/j.molcel.2005.07.025)

31. Debora, B. N., Vidales, L. E., Ramírez, R., Ramírez, M., Robleto, E. A., Yasbin, R. E. & Pedraza-Reyes, M. 2010 Mismatch Repair Modulation of MutY Activity Drives *Bacillus subtilis* Stationary-Phase Mutagenesis. *J. Bacteriol.* **193**, 236–45. (doi:10.1128/JB.00940-10)

32. Al Mamun, A. A. M. et al. 2012 Identity and function of a large gene network underlying mutagenic repair of DNA breaks. *Science (80-. ).* **338**, 1344–8. (doi:10.1126/science.1226683)

33. Tenaillon, O., Denamur, E. & Matic, I. 2004 Evolutionary significance of stress-induced mutagenesis in bacteria. *Trends Microbiol.* **12**, 264–70. (doi:10.1016/j.tim.2004.04.002)

34. Cirz, R. T. & Romesberg, F. E. 2007 Controlling mutation: intervening in evolution as a therapeutic strategy. *Crit. Rev. Biochem. Mol. Biol.* **42**, 341–54. (doi:10.1080/10409230701597741)

35. Agrawal, A. F. 2002 Genetic loads under fitness-dependent mutation rates. *J. Evol. Biol.* **15**, 1004–1010. (doi:10.1046/j.1420-9101.2002.00464.x)

36. Shaw, F. H. & Baer, C. F. 2011 Fitness-dependent mutation rates in finite populations. *J. Evol. Biol.* **24**, 1677–84. (doi:10.1111/j.1420-9101.2011.02320.x)

37. Ram, Y. & Hadany, L. 2012 The evolution of stress-induced hypermutation in asexual populations. *Evolution* **66**, 2315–28. (doi:10.1111/j.1558-5646.2012.01576.x)

38. Wright, S. 1931 Evolution in Mendelian Populations. *Genetics* **16**, 97–159.

39. Wright, S. 1988 Surfaces of selective value revisited. *Am. Nat.* **131**, 115–123. (doi:10.1086/284777)

40. Shee, C., Gibson, J. L., Darrow, M. C., Gonzalez, C. & Rosenberg, S. M. 2011 Impact of a stress-inducible switch to mutagenic repair of DNA breaks on mutation in *Escherichia coli*. *Proc. Natl. Acad. Sci.* **108**, 13659–13664. (doi:10.1073/pnas.1104681108)

41. Yeiser, B., Pepper, E. D., Goodman, M. F. & Finkel, S. E. 2002 SOS-induced DNA polymerases enhance long-term survival and evolutionary fitness. *Proc. Natl. Acad. Sci.* **99**, 8737–41. (doi:10.1073/pnas.092269199)

42. Cirz, R. T., Chin, J. K., Andes, D. R., de Crécy-Lagard, V., Craig, W. A. & Romesberg, F. E. 2005 Inhibition of mutation and combating the evolution of antibiotic resistance. *PLoS Biol.* **3**, e176. (doi:10.1371/journal.pbio.0030176)

43. Gutierrez, A. et al. 2013 β-Lactam antibiotics promote bacterial mutagenesis via an RpoS-mediated reduction in replication fidelity. *Nat. Commun.* **4**, 1610. (doi:10.1038/ncomms2607)

44. Haigh, J. 1978 The accumulation of deleterious genes in a population - Muller’s Ratchet. *Theor. Popul. Biol.* **14**, 251–267. (doi:10.1016/0040-5809(78)90027-8)

45. Gessler, D. D. G. 1995 The constraints of finite size in asexual populations and the rate of the ratchet. *Genet. Res.* **66**, 241. (doi:10.1017/S0016672300034686)

46. Gordo, I. & Dionisio, F. 2005 Nonequilibrium model for estimating parameters of deleterious mutations. *Phys. Rev. E* **71**, 18–21. (doi:10.1103/PhysRevE.71.031907)

47. Eshel, I. 1981 On the survival probability of a slightly advantageous mutant gene with a general distribution of progeny size - a branching process model. *J. Math. Biol.* **12**, 355–362. (doi:10.1007/BF00276922)

48. Fisher, R. A. 1930 *The Genetical Theory of Natural Selection*. Oxford: Clarendon Press.

49. Patwa, Z. & Wahl, L. M. 2008 The fixation probability of beneficial mutations. *J. R. Soc. Interface* **5**, 1279–89. (doi:10.1098/rsif.2008.0248)

50. Weinreich, D. M. & Chao, L. 2005 Rapid evolutionary escape by large populations from local fitness peaks is likely in nature. *Evolution* **59**, 1175–82.

51. Maynard Smith, J. & Haigh, J. 1974 The hitch-hiking effect of a favourable gene. *Genet. Res.* **23**, 23–35. (doi:10.1017/S0016672300014634)

52. Lynch, M., Bürger, R., Butcher, D. & Gabriel, W. 1993 The mutational meltdown in asexual populations. *J. Hered.* **84**, 339–44.

53. Obolski, U. & Hadany, L. 2012 Implications of stress-induced genetic variation for minimizing multidrug resistance in bacteria. *BMC Med.* **10**, 1–30. (doi:10.1186/1741-7015-10-89)

54. Shor, E., Fox, C. a. & Broach, J. R. 2013 The yeast environmental stress response regulates mutagenesis induced by proteotoxic stress. *PLoS Genet.* **9**, e1003680. (doi:10.1371/journal.pgen.1003680)

55. Karpinets, T., Greenwood, D., Pogribny, I. & Samatova, N. 2006 Bacterial stationary-state mutagenesis and Mammalian tumorigenesis as stress-induced cellular adaptations and the role of epigenetics. *Curr. Genomics* **7**, 481–96.

56. Torres-Barceló, C., Cabot, G., Oliver, A., Buckling, A. & MacLean, R. C. 2013 A trade-off between oxidative stress resistance and DNA repair plays a role in the evolution of elevated mutation rates in bacteria. *Proc. R. Soc. B Biol. Sci.* **280**, 20130007. (doi:10.1098/rspb.2013.0007)

57. Christiansen, F. B., Otto, S. P., Bergman, A. & Feldman, M. W. 1998 Waiting with and without recombination: the time to production of a double mutant. *Theor. Popul. Biol.* **53**, 199–215. (doi:10.1006/tpbi.1997.1358)

58. Dawson, K. J. 1998 Evolutionarily stable mutation rates. *J. Theor. Biol.* **194**, 143–57. (doi:10.1006/jtbi.1998.0752)

59. Giraud, A., Matic, I., Tenaillon, O., Clara, A., Radman, M., Fons, M. & Taddei, F. 2001 Costs and benefits of high mutation rates: adaptive evolution of bacteria in the mouse gut. *Science (80-. ).* **291**, 2606–8. (doi:10.1126/science.1056421)

60. Loh, E., Salk, J. J. & Loeb, L. A. 2010 Optimization of DNA polymerase mutation rates during bacterial evolution. *Proc. Natl. Acad. Sci.* **107**, 1154–9. (doi:10.1073/pnas.0912451107)

61. Gentile, C. F., Yu, S.-C., Serrano, S. A., Gerrish, P. J. & Sniegowski, P. D. 2011 Competition between high- and higher-mutating strains of *Escherichia coli*. *Biol. Lett.* **7**, 422–4. (doi:10.1098/rsbl.2010.1036)

62. Tenaillon, O., Le Nagard, H., Godelle, B. & Taddei, F. 2000 Mutators and sex in bacteria: conflict between adaptive strategies. *Proc. Natl. Acad. Sci.* **97**, 10465–70. (doi:10.1073/pnas.180063397)

63. Johnson, T. & Barton, N. H. 2002 The effect of deleterious alleles on adaptation in asexual populations. *Genetics* **162**, 395–411.

64. Leigh, E. G. J. 1973 The evolution of mutation rates. *Genetics* **73**, Suppl 73:1–18.

65. Komarova, N. L. & Wodarz, D. 2004 The optimal rate of chromosome loss for the inactivation of tumor suppressor genes in cancer. *Proc. Natl. Acad. Sci. U. S. A.* **101**, 7017–21. (doi:10.1073/pnas.0401943101)

66. Komarova, N. L., Sadovsky, A. V & Wan, F. Y. M. 2008 Selective pressures for and against genetic instability in cancer: a time-dependent problem. *J. R. Soc. Interface* **5**, 105–21. (doi:10.1098/rsif.2007.1054)

67. Agur, Z., Mazor, G. & Meilijson, I. 1991 Maturation of the humoral immune response as an optimization problem. *Proc. R. Soc. B Biol. Sci.* **245**, 147–50. (doi:10.1098/rspb.1991.0101)

68. Crow, J. F., Engels, W. R. & Denniston, C. 1990 Phase Three of Wright’s Shifting-Balance Theory. *Evolution* **44**, 233. (doi:10.2307/2409403)

69. Wade, M. J. & Goodnight, C. J. 1991 Wright’s shifting balance theory: an experimental study. *Science (80-. ).* **253**, 1015–8. (doi:10.1126/science.1887214)

70. Peck, S. L., Ellner, S. P. & Gould, F. 2000 Varying migration and deme size and the feasibility of the shifting balance. *Evolution* **54**, 324–7. (doi:10.1111/j.0014-3820.2000.tb00035.x)

71. Moore, F. B.-G. & Tonsor, S. J. 1994 A Simulation of Wright’s Shifting-Balance Process: Migration and the Three Phases. *Evolution* **48**, 69. (doi:10.2307/2410004)

72. Gavrilets, S. 1996 On phase three of the shifting-balance theory. *Evolution* **50**, 1034–1041. (doi:10.2307/2410644)

73. Coyne, J. A., Barton, N. H. & Turelli, M. 2000 Is Wright’s shifting balance process important in evolution? *Evolution* **54**, 306–317. (doi:10.1111/j.0014-3820.2000.tb00033.x)

74. Whitlock, M. C. & Phillips, P. C. 2000 The exquisite corpse: a shifting view of the shifting balance. *Trends Ecol. Evol.* **15**, 347–348. (doi:10.1016/S0169-5347(00)01930-3)

75. Whitlock, M. C. 1995 Variance-induced peak shifts. *Evolution* **49**, 252. (doi:10.2307/2410335)

76. Whitlock, M. C. 1997 Founder effects and peak shifts without genetic drift: adaptive peak shifts occur easily when environments fluctuate slightly. *Evolution* **51**, 1044. (doi:10.2307/2411033)

77. Hadany, L. 2003 Adaptive peak shifts in a heterogenous environment. *Theor. Popul. Biol.* **63**, 41–51. (doi:10.1016/S0040-5809(02)00011-4)

78. Hadany, L. & Beker, T. 2003 Fitness-associated recombination on rugged adaptive landscapes. *J. Evol. Biol.* **16**, 862–870. (doi:10.1046/j.1420-9101.2003.00586.x)

79. Weissman, D. B., Desai, M. M., Fisher, D. S. & Feldman, M. W. 2009 The rate at which asexual populations cross fitness valleys. *Theor. Popul. Biol.* **75**, 286–300. (doi:10.1016/j.tpb.2009.02.006)

80. Weissman, D. B., Feldman, M. W. & Fisher, D. S. 2010 The rate of fitness-valley crossing in sexual populations. *Genetics* **186**, 1389–410. (doi:10.1534/genetics.110.123240)

81. Schrag, S. J., Perrot, V. & Levin, B. R. 1997 Adaptation to the fitness costs of antibiotic resistance in *Escherichia coli*. *Proc. R. Soc. B Biol. Sci.* **264**, 1287–91. (doi:10.1098/rspb.1997.0178)

82. Cairns, J., Overbaugh, J. & Miller, S. 1988 The origin of mutants. *Nature* **335**, 142–5. (doi:10.1038/335142a0)

83. Tomlinson, I. & Bodmer, W. F. 1999 Selection, the mutation rate and cancer: ensuring that the tail does not wag the dog. *Nat. Med.* **5**, 11–2. (doi:10.1038/4687)

84. Bielas, J. H., Loeb, K. R., Rubin, B. P., True, L. D. & Loeb, L. A. 2006 Human cancers express a mutator phenotype. *Proc. Natl. Acad. Sci. U. S. A.* **103**, 18238–42. (doi:10.1073/pnas.0607057103)

85. Cotton, S. 2009 Condition-dependent mutation rates and sexual selection. *J. Evol. Biol.* **22**, 899–906. (doi:10.1111/j.1420-9101.2008.01683.x)

86. Kibota, T. T. & Lynch, M. 1996 Estimate of the genomic mutation rate deleterious to overall fitness in *E. coli*. *Nature* **381**, 694–6. (doi:10.1038/381694a0)

87. Gordo, I., Perfeito, L. & Sousa, A. 2011 Fitness effects of mutations in bacteria. *J. Mol. Microbiol. Biotechnol.* **21**, 20–35. (doi:10.1159/000332747)

88. Drake, J. W., Charlesworth, B., Charlesworth, D. & Crow, J. F. 1998 Rates of spontaneous mutation. *Genetics* **148**, 1667–86.

89. Wielgoss, S., Barrick, J. E., Tenaillon, O., Cruveiller, S., Chane-Woon-Ming, B., Médigue, C., Lenski, R. E. & Schneider, D. 2011 Mutation rate inferred from synonymous substitutions in a long-term evolution experiment with *Escherichia coli*. *G3* **1**, 183–186. (doi:10.1534/g3.111.000406)

90. Hall, L. M. C. & Henderson-Begg, S. K. 2006 Hypermutable bacteria isolated from humans--a critical analysis. *Microbiology* **152**, 2505–14. (doi:10.1099/mic.0.29079-0)

91. Pupo, G. M. & Richardson, B. J. 1995 Biochemical genetics of a natural population of *Escherichia coli*: seasonal changes in alleles and haplotypes. *Microbiology* **141**, 1037–44. (doi:10.1099/13500872-141-4-1037)

92. Berg, O. G. 1996 Selection intensity for codon bias and the effective population size of *Escherichia coli*. *Genetics* **142**, 1379–82.

# Tables

Table 1 – Model parameters and estimated values for *E. coli*

|  |  |  |  |
| --- | --- | --- | --- |
| Symbol | Name | Estimate | References |
| *s* | Selection coefficient | 0.001-0.03 | [86,87] |
| *H* | Double mutant advantage | 1-10 | [87] |
| *U* | Genomic deleterious mutation rate | 0.0004-0.003 | [88,89] |
| *µ* | Site-specific mutation rate | *U*/5000 | [87] |
| *τ* | Fold-increase in mutation rate | 1-100 | [18,90] |
| *N* | Population size | 105-1010 | [91,92] |

­­­