

# Mutation–Selection Balance: Ancestry, Load, and Maximum Principle

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**We analyze the equilibrium behavior of deterministic haploid mutation–selection models. To this end, both the forward and the time-reversed evolution processes are considered. The stationary state of the latter is called the ancestral distribution, which turns out as a key for the study of mutation–selection balance. We find that the ancestral genotype frequencies determine the sensitivity of the equilibrium mean fitness to changes in the corresponding fitness values and discuss implications for the evolution of mutational robustness. We further show that the difference between the ancestral and the population mean fitness, termed mutational loss, provides a measure for the sensitivity of the equilibrium mean fitness to changes in the mutation rate. The interrelation of the loss and the mutation load is discussed. For a class of models in which the number of mutations in an individual is taken as the trait value, and fitness is a function of the trait, we use the ancestor formulation to derive a simple maximum principle, from which the mean and variance of fitness and the trait may be derived; the results are exact for a number of limiting cases, and otherwise yield approximations which are accurate for a wide range of parameters. These results are applied to threshold phenomena caused by the interplay of selection and mutation (known as error thresholds). They lead to a clarification of concepts, as well as criteria for the existence of error thresholds.**

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**Key Words:** mutation–selection model; clonal reproduction; branching process; backward processes; mutation load; epistasis; mutational robustness; error threshold; statistical physics.

## 1. INTRODUCTION

A lot of research in theoretical population genetics has been directed towards mutation–selection models in multilocus systems and infinite populations. One is

usually interested in the statistical properties of the equilibrium distribution of genotypes, like the means and variances of fitness and trait(s). The standard approach to determine these starts out from the equilibrium condition for the genotype frequencies (which takes the form of an eigenvalue equation if the

population is haploid). On this basis, a wealth of methods and results has been created; for a comprehensive and up-to-date account, see Bürger (2000).

In this article, we present an alternative route, which relies on a time-reversed version of the mutation–selection process and its stationary distribution—to be called the *ancestral distribution*, as opposed to the equilibrium distribution of the forward process. We will apply this approach to tackle a rather general class of models for haploids, or diploids without dominance. It is only assumed that fitness is a function of a trait, and genotypes with equal trait values have equivalent mutation patterns. In fact, this is a standard assumption, and is often implied without special mention. It applies, for example, if (geno)types are identified with a collection of loci with two alleles each (wildtype and mutant), which mutate independently and according to the same process, and the number of (deleterious) mutations plays the role of the trait. The assumption of permutation invariance (with respect to the loci) is certainly a distortion of biological reality, but, even in this simplified setting, general answers have previously been considered impossible (Charlesworth, 1990), and researchers have resorted to more specific choices of the fitness function and the mutation model (e.g., quadratic fitness functions and unidirectional mutation).

With the help of the ancestral distribution, we will be able to tackle general fitness functions (with arbitrary epistasis), as well as general mutation schemes (including arbitrary amounts of back mutation), from the permutation invariant class. The mutation–selection equilibrium will be characterized through a maximum principle which relates the equilibrium population to the ancestral one, and may be evaluated explicitly to yield expressions for the mean fitness and the mean trait value, as well as the variances of these quantities. The results are exact for a number of limiting cases, and otherwise yield approximations which are accurate for a wide range of parameters.

The results will then be used to settle the long-standing issue of characterization and classification of error threshold phenomena in this model class. An error threshold may be generally, but vaguely, circumscribed as a critical mutation rate beyond which mutation can no longer be controlled by selection and leads to genetic degeneration; for review, see Eigen *et al.* (1989). Some, but by no means all, mutation–selection models display such behavior. It turns out that a consistent definition of an error threshold is rather subtle and must be sorted out first. On this basis, we will classify mutation–selection models according to their threshold behavior (if any).

TABLE I

**Glossary of Repeatedly Used Notation**

|                 |  |           |
|-----------------|--|-----------|
| $a$             | ancestor frequencies                       | 2.2       |
| $G, g$          | mutational loss                            | (24)      |
| $g$             | mutational loss function                   | (31)      |
| $\mathbf{H}$    | evolution matrix                           | (4)       |
| $\mathbf{I}$    | identity matrix                            | 2.1       |
| $i, j, k, \ell$ | genotype/class labels                      | 2.1       |
| $L, l$          | mutation load                              | 2.4       |
| $m$             | mutation rates                             | 2.1       |
| $\mathbf{M}$    | mutation matrix                            | 2.1       |
| $N$             | number of mutation classes/sequence length | 2.1       |
| $p$             | population frequencies                     | 2.1       |
| $\mathbf{Q}$    | generator of reversed process              | 2.2       |
| $R, r$          | reproduction rates                         | 2.1       |
| $r$             | fitness function                           | (27)      |
| $\mathbf{R}$    | reproduction matrix                        | 2.1       |
| $s^\pm$         | mutational effects                         | 2.4       |
| $s$             | (binary) sequence                          | 2.1       |
| $\mathbf{T}$    | time evolution matrix                      | 2.1       |
| $t$             | time                                       | 2.1       |
| $U^\pm, u^\pm$  | genomic mutation rates                     | 2.1       |
| $u^\pm$         | mutation functions                         | (27)      |
| $V, v$          | variances                                  | 2.4       |
| $X, x$          | mutational distance                        | 2.4       |
| $\mathcal{X}$   | mutation classes                           | 2.1       |
| $Y, y$          | arbitrary trait                            | 2.7       |
| $z$             | relative reproductive success              | 2.2       |
| $\gamma$        | overall factor for reproduction rates      | 5.3       |
| $\kappa$        | biallelic mutation asymmetry parameter     | 2.1       |
| $\lambda_{max}$ | largest eigenvalue of $\mathbf{H}$         | 2.1       |
| $\mu$           | overall mutation rate                      | 2.1, (56) |

*Note.* Symbols are given together with the section or equation in which they are defined. Symbols whose scope is only a single section are not shown.

Since the article treats quite a number of topics, we start out with a brief reader’s guide to the main results here and give hints on what parts can be skipped at a first reading. Let us also mention that Table I contains a glossary of repeatedly used notation.

The scene is set in Section 2, where we will introduce the model (the continuous-time mutation–selection model) and establish its relationship with a multitype branching process. Two concepts that are central to this paper, the *ancestral distribution* and the *mutation class limit*, are developed in this section. Section 2.2 introduces the ancestral distribution as the stationary distribution of the time-reversed branching process and links the algebraic properties of the model to a probabilistic picture that also helps to shape biological intuition. In order to allow quick progress to the results in the remainder of the article, however, we have summarized the main points in Section 2.3. In Section 2.4, the means and variances of the trait and of fitness

with respect to the equilibrium population and with respect to the ancestors are introduced. In Section 2.5, the difference between the ancestral and the population mean fitness, termed mutational loss, is shown to provide a measure for the sensitivity of the equilibrium mean fitness to changes in the mutation rate. This result is used and expanded in some of the applications in Sections 5 and 6, but can be skipped at first reading. Sections 2.6 and 2.7 are mainly concerned with the mutation class limit, along with the proper scaling of fitness functions and mutation schemes. Like the well-known infinite-sites limit, this limit assumes an infinite number of types, but uses a different scaling. As a consequence, it is valid if the total mutation rate is large relative to typical fitness differences between types. In this paper, the mutation class limit is used to derive analytic expressions for means and variances of fitness and the trait for the general case with back mutations and a non-linear fitness function. It is also crucial for our discussion of threshold behavior in Section 6.

Section 3 is a condensed summary of the main results related to the maximum principle. The mean fitness at mutation–selection balance equals the maximum of the difference between the fitness function and the so-called *mutational loss function*, where the position of the maximum determines the mean ancestral trait. Once these means are known, explicit expressions are available for the mean trait and the variances of fitness and trait. The derivations are postponed to Section 4, which may be skipped at first reading.

The following two sections are devoted to applications. Both are, to a large extent, independent of each other and rely only on the matter introduced in Sections 2 and 3. In Section 5, we first discuss the evolutionary significance of the mutational loss, and then turn to the mutation load. Explicit expressions are derived for small (back) mutation rates; but arbitrary mutation rates are covered by the maximum principle, which may be interpreted as a generalized version of Haldane’s principle. Consequences for the evolution of mutational effects and for mutational robustness are discussed. Finally, a note is added as to the accuracy of the expressions for the means and variances.

In Section 6, we first analyze the definitions available for the error threshold. It will turn out that various notions must be distinguished, which coincide only in special cases. For each of these notions, a criterion for the existence of an error threshold is derived from the maximum principle. Furthermore, the phenomena are illustrated by means of examples and discussed with respect to their biological implications. Section 7 provides a summary and an outlook.

Appendix A describes the connection between our mutation–selection model and a system of quantum-statistical mechanics, which had been used previously (e.g., Baake *et al.*, 1997; Baake and Wagner, 2001) to solve a more restricted model class, and which also served as the source of concepts and methods for the current article. However, the body of the paper does not require any knowledge of physics and remains entirely within the framework of population genetics and classical probability theory. Appendices B and C, finally, contain the proofs from Sections 4 and 6, respectively.

## 2. MODEL SETUP

### 2.1. The Model

We consider the evolution of an infinite population of haploid individuals (or diploids without dominance) under mutation and selection. Disregarding environmental effects, we take individuals to be fully described by their genotypes, which are labeled by the elements of  $\{1, \dots, M\}$ . Let  $p_i(t)$  be the relative frequency of type  $i$  at time  $t$ , so that  $\sum_i p_i(t) = 1$ , and let  $\mathbf{p}(t) = (p_1(t), \dots, p_M(t))^T$  with T denoting transposition. Throughout this article we will use the formalism for overlapping generations, which works in continuous time, and only comment on extensions to the analogous model for discrete generations. The standard system of differential equations which describes the evolution of the vector  $\mathbf{p}(t)$  is (Crow and Kimura, 1970; Hofbauer, 1985; see also Bürger, 2000)

$$\dot{p}_i(t) = [R_i - \bar{R}(t)]p_i(t) + \sum_j [m_{ij}p_j(t) - m_{ji}p_i(t)]. \quad (1)$$

Here,  $R_i$  is the Malthusian fitness of type  $i$ , which is connected to the respective birth and death rates as  $R_i = B_i - D_i$ , and  $\bar{R}(t) = \sum_i R_i p_i(t)$  designates the mean fitness. Further,  $m_{ij}$  is the rate at which a  $j$ -individual mutates to  $i$ , and the dot denotes the time derivative. In this model, mutation and selection are assumed to be independent processes which go on in parallel. However, mutation may also be viewed as occurring during reproduction. In this case, the mutation rate is given by  $m_{ij} := v_{ij}B_j$ , where  $v_{ij}$  is the respective mutation *probability* during a reproduction event. Since, formally, this leads to the same model, it need not be discussed further.

For some of the main results of this article, further assumptions on the mutation scheme are required. To this end, we collect genotypes into classes  $\mathcal{X}_k$  of equal fitness,  $0 \leq k \leq N$ , and assume mutations only to occur

between neighboring classes. Let  $R_k$  denote the fitness of class  $k$  and  $U_k^\pm$  the mutation rate from class  $\mathcal{X}_k$  to  $\mathcal{X}_{k\pm 1}$  (i.e., the total rate for each genotype in  $\mathcal{X}_k$  to mutate to some genotype in class  $\mathcal{X}_{k\pm 1}$ ), with the convention  $U_0^- = U_N^+ = 0$ . Thus, we obtain a variant of the so-called single-step mutation model:

$$\begin{aligned} \dot{p}_k(t) = & [R_k - \bar{R}(t) - U_k^+ - U_k^-]p_k(t) \\ & + U_{k-1}^+p_{k-1}(t) + U_{k+1}^-p_{k+1}(t). \end{aligned} \quad (2)$$

(Here, the convention  $p_{-1}(t) = p_{N+1}(t) = 0$  is used.) We can, for example, think of  $\mathcal{X}_0$  as the *wildtype* class with maximum fitness and fitness only depending on the number of mutations carried by an individual. If, further, mutation is modeled as a continuous point process (or if multiple mutations during reproduction can be ignored), Eq. (1) reduces to Eq. (2), with an appropriate choice of mutation classes. Depending on the realization one has in mind, the  $U_k$  then describe the total mutation rate affecting the whole genome or just some trait or function.

In most of our examples, we will use the Hamming graph as our genotype space. Here, genotypes are represented as binary sequences  $s = s_1s_2\dots s_N \in \{+, -\}^N$ , thus  $M = 2^N$ . The two possible values at each site, + and -, may be understood either in a molecular context as nucleotides (purines and pyrimidines) or, on a coarser level, as wildtype and mutant alleles of a biallelic multilocus model. We will assume equal mutation rates at all sites, but allow for different rates,  $\mu(1 + \kappa)$  and  $\mu(1 - \kappa)$ , for mutations from + to - and for back mutations, respectively, according to the scheme depicted in Fig. 1.

Clearly, the biallelic model reduces to a single-step mutation model (with the same  $N$ ) if the fitness landscape<sup>1</sup> is invariant under permutation of sites. To this end, we distinguish a reference genotype  $s_+ = +\dots+$ , in most cases the wildtype or master sequence, and assume that the fitness  $R_s$  of sequence  $s$  depends only on the Hamming distance  $k = d_H(s, s_+)$  to  $s_+$  (i.e., the number of mutations, or “-” signs in the sequence). The resulting total mutation rates between the Hamming classes  $\mathcal{X}_k$  and  $\mathcal{X}_{k\pm 1}$  read

$$U_k^+ = \mu(1 + \kappa)(N - k) \quad \text{and} \quad U_k^- = \mu(1 - \kappa)k \quad (3)$$

if mutation is assumed to be an independent point process at all sites. We usually have the situation in mind in which fitness decreases with  $k$  and will therefore speak of  $U_k^+$  and  $U_k^-$  as the *deleterious* and *advantageous*

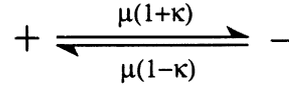


FIG. 1. Rates for mutations and back mutations at each site or locus of a biallelic sequence.

mutation rates. However, monotonic fitness is never assumed, unless this is stated explicitly.

In much of the following, we will treat the general model (1), which builds on single genotypes, and the single-step mutation model (2), in which the units are genotype classes, with the help of a common formalism. To this end, note that both models can be recast into the following general form using matrices of dimension  $M$ , respectively  $N + 1$ :

$$\dot{\mathbf{p}}(t) = (\mathbf{H} - \bar{R}(t)\mathbf{I})\mathbf{p}(t). \quad (4)$$

Here,  $\mathbf{I}$  is the identity. The evolution matrix  $\mathbf{H} = \mathbf{R} + \mathbf{M}$  is composed of a diagonal matrix  $\mathbf{R}$  that holds the Malthusian fitness values, and the mutation matrix  $\mathbf{M} = (M_{ij})$  with either off-diagonal entries reading  $m_{ij}$ , or with  $U_k^\pm$  on the secondary diagonals. The diagonal elements in each case are  $M_{ii} = -\sum_{j \neq i} M_{ji}$ , hence the column sums vanish. Where the more restrictive form of the single-step model is needed, this will be stated explicitly. Unless we talk about unidirectional mutation ( $U_k^- \equiv 0$  for the single-step mutation model), we will always assume that  $\mathbf{M}$  is irreducible (i.e., each entry is non-zero for a suitable power of  $\mathbf{M}$ ).

Let now  $\mathbf{T}(t) := \exp(t\mathbf{H})$ , with matrix elements  $T_{ij}(t)$ . Then, the solution of (4) is given by (see, e.g., Bürger, 2000, Chapter III.1)

$$\mathbf{p}(t) = \frac{\mathbf{T}(t)\mathbf{p}(0)}{\sum_{i,j} T_{ij}(t)p_j(0)} \quad (5)$$

as can easily be established by differentiating and using  $\sum_{i,j} H_{ij}p_j(t) = \sum_i R_i p_i(t) = \bar{R}(t)$ . Due to irreducibility, the population vector converges to a unique, globally stable equilibrium distribution  $\mathbf{p} := \lim_{t \rightarrow \infty} \mathbf{p}(t)$  with  $p_i > 0$  for all  $i$ , which describes mutation–selection balance. By the Perron–Frobenius theorem,  $\mathbf{p}$  is the (right) eigenvector corresponding to the largest eigenvalue,  $\lambda_{max}$ , of  $\mathbf{H}$ .

## 2.2. The Branching Process—Forward and Backward in Time

Our approach will heavily rely on genealogical relationships, which contain a more detailed information than the time course of the relative frequencies (5) alone. Let us, therefore, reconsider the mutation–

<sup>1</sup>We use the notion of a *fitness landscape* (Kauffman and Levin, 1987) as synonymous with *fitness function* for the mapping from genotypes to individual fitness values.

selection model as a branching process. Branching processes have been classical tools in population genetics to approximate the fixation rates of a single mutant type in a finite population. This approach goes back to Haldane (1927) (see also Crow and Kimura, 1970), and has been used in many recent applications as well (e.g., Barton, 1995; Otto and Barton, 1997).

We pursue a different route here by considering the process of mutation, reproduction and death as a (continuous-time) *multitype* branching process, as described previously for the quasi-species model (Demetrius *et al.*, 1985; Hofbauer and Sigmund, 1988, Chapter 11.5). Let us start with a *finite* population of individuals, which reproduce (at rates  $B_i$ ), die (at rates  $D_i$ ), or change type (at rates  $M_{ij}$ ) independently of each other, *without any restriction on population size*. Let  $Y_i(t)$  be the random variable denoting the number of individuals of type  $i$  at time  $t$ , and  $n_i(t)$  the corresponding realization; collect the components into vectors  $\mathbf{Y}$  and  $\mathbf{n}$ , and let  $\mathbf{e}_i$  be the  $i$ th unit vector. The transition probabilities for the joint distribution,  $\Pr(\mathbf{Y}(t) = \mathbf{n}(t) | \mathbf{Y}(0) = \mathbf{n}(0))$ , which we will abbreviate as  $\Pr(\mathbf{n}(t) | \mathbf{n}(0))$  by abuse of notation, are governed by the differential equation<sup>2</sup>

$$\begin{aligned} & \frac{d}{dt} \Pr(\mathbf{n}(t) | \mathbf{n}(0)) \\ &= - \left( \sum_i \left( B_i + D_i + \sum_{j \neq i} M_{ji} \right) n_i(t) \right) \Pr(\mathbf{n}(t) | \mathbf{n}(0)) \\ &+ \sum_i B_i (n_i(t) - 1) \Pr(\mathbf{n}(t) - \mathbf{e}_i | \mathbf{n}(0)) \\ &+ \sum_i D_i (n_i(t) + 1) \Pr(\mathbf{n}(t) + \mathbf{e}_i | \mathbf{n}(0)) \\ &+ \sum_{\substack{i,j \\ i \neq j}} M_{ij} (n_j(t) + 1) \Pr(\mathbf{n}(t) - \mathbf{e}_i + \mathbf{e}_j | \mathbf{n}(0)). \end{aligned} \quad (6)$$

The connection of this stochastic process with the deterministic model described in Section 2.1 is two-fold. Firstly, in the limit of an infinite number of individuals ( $n := \sum_i n_i(0) \rightarrow \infty$ ), the sequence of random variables  $\mathbf{Y}^{(n)}(t)/n$  converges almost surely to the solution  $\mathbf{y}(t)$  of  $\dot{\mathbf{y}} = \mathbf{H}\mathbf{y}$  with initial condition  $\mathbf{y}(0) = \mathbf{n}(0)/n$  (Ethier and Kurtz, 1986, Chapter 11, Theorem 2.1). That is,  $\Pr(\lim_{n \rightarrow \infty} \mathbf{Y}^{(n)}(t)/n = \mathbf{y}(t)) = 1$ , and the superscript  $(n)$  denotes the dependence on the number of individuals. The connection is now clear since  $\mathbf{p}(t) := \mathbf{y}(t) / \sum_i y_i(t)$  solves the mutation–selection equation (1).

<sup>2</sup>Note that differentiability of the transition *probabilities* is guaranteed in a finite-state, continuous-time Markov chain, provided the transition *rates* are finite, cf. Karlin and Taylor (1975, Chapter 4) and Karlin and Taylor (1981, Chapter 14).

Secondly, taking expectations of  $Y_i$  and marginalizing over all other variables, one obtains the differential equation for the conditional expectations

$$\begin{aligned} & \frac{d}{dt} E(Y_i(t) | \mathbf{n}(0)) = (B_i - D_i) E(Y_i(t) | \mathbf{n}(0)) \\ &+ \sum_j [M_{ij} E(Y_j(t) | \mathbf{n}(0)) - M_{ji} E(Y_i(t) | \mathbf{n}(0))]. \end{aligned} \quad (7)$$

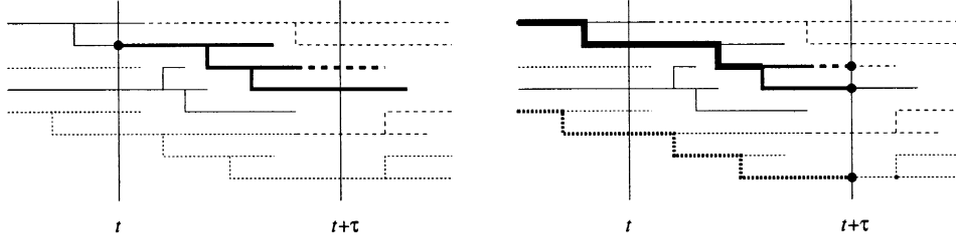
Clearly, our evolution matrix  $\mathbf{H}$  appears as the infinitesimal generator here, and the solution is given by  $\mathbf{T}(t)\mathbf{n}(0)$ , where  $\mathbf{T}(t) := \exp(t\mathbf{H})$  (see also Hofbauer and Sigmund, 1988, Chapter 11.5). In particular, we have  $E(Y_i(t) | \mathbf{e}_j) = T_{ij}(t)$  for the expected number of  $i$ -individuals at time  $t$ , in a population started by a single  $j$ -individual at time 0 (a “ $j$ -clone”). In the same way,  $T_{ij}(\tau)$  is the expected number of descendants of type  $i$  at time  $t + \tau$  in a  $j$ -clone started at an arbitrary time  $t$ , cf. left panel of Fig. 2. (Note that due to the independence of individuals and the Markov property, the progeny distribution depends only on the age of the clone, and on the founder type.) Further, the expected total size of a  $j$ -clone of age  $\tau$ , irrespective of the descendants’ types, is  $\sum_i T_{ij}(\tau)$ .

Initial conditions come into play if we consider the reproductive success of a clone *relative to the whole population*. A population of independent individuals, with initial composition  $\mathbf{p}(t)$ , has an expected mean clone size  $\sum_{i,j} T_{ij}(\tau) p_j(t)$  at time  $t + \tau$  (note that  $t$  always means “absolute” time, whereas  $\tau$  denotes a time increment). The expected size of a single  $j$ -clone at time  $t + \tau$ , relative to the expected mean clone size of the whole population, then is

$$z_j(\tau, t) := \sum_i T_{ij}(\tau) / \sum_{k,\ell} T_{k\ell}(\tau) p_\ell(t). \quad (8)$$

The  $z_j$  express the expected relative success of a type after evolution for a time interval  $\tau$ , in the sense that, if  $z_j(\tau, t) > 1$  ( $< 1$ ), we can expect the clone to flourish more (less) than average (this does in general not mean that type  $j$  is expected to increase (decrease) in abundance relative to the initial population). Clearly, the values of  $z_j$  depend on the fitness of type  $j$ , but also on its mutation rate and the fitness of its (mutated) offspring. (If there is only mutation, but no reproduction or death, one has a Markov chain and  $z_j(\tau, t) \equiv 1$ .)

We now consider *lines of descent*, as in the right panel of Fig. 2. To this end, we randomly pick an individual alive at time  $t + \tau$ , and trace its ancestry back in time; this results in an unbranched line (in contrast to the lineage forward in time). Let  $Z_{t+\tau}(t)$  denote the type



**FIG. 2.** The multitype branching process. Individuals reproduce (branching lines), die (ending lines), or mutate (lines changing type) independently of each other; the various types are indicated by different line styles. Left: The fat lines mark the clone founded by a single individual (bullet) at time  $t$ . Right: The fat lines mark the lines of descent defined by three individuals (bullets) at time  $t + \tau$ . After coalescence of two lines, their ancestor receives twice the “weight”, as indicated by extra fat lines.

found at time  $t \leq t + \tau$ , where we will drop the index for easier readability. We seek its probability distribution  $\Pr(Z(t) = j)$ . Since the (relative) clone size  $z_j(\tau, t)$  also determines the expected (relative) frequency of lines present at time  $t + \tau$  that contain a  $j$ -type ancestor at time  $t$ , we have

$$\Pr(Z(t) = j) = z_j(\tau, t) p_j(t) =: a_j(\tau, t). \quad (9)$$

The  $a_j(\tau, t)$  define a probability distribution ( $\sum_j a_j(\tau, t) \equiv 1$ ), which will be of major importance, and may be interpreted in two ways. Forward in time,  $a_j(\tau, t)$  is the frequency of  $j$ -individuals at time  $t$ , weighted by their relative number of descendants after evolution for some time  $\tau$ . Looking backward in time,  $a_j(\tau, t)$  is the fraction of the ( $\mathbf{p}$ -distributed) population at time  $t + \tau$  whose ancestor at time  $t$  is of type  $j$ . We shall therefore refer to  $\mathbf{a}(\tau, t)$  as the *ancestral distribution* at the earlier time,  $t$ .

Let us, at this point, expand a little further on this backward picture by explicitly constructing the time-reversed process. This is done in the usual way, by writing the joint distribution of parent–offspring pairs (i.e., pairs  $Z(t)$  and  $Z(t + \tau)$ ) in terms of forward and backward transition probabilities. On the one hand,

$$\begin{aligned} \Pr(Z(t + \tau) = i, Z(t) = j) &= \Pr(Z(t + \tau) = i | Z(t) = j) \Pr(Z(t) = j) \\ &= P_{ij}(\tau) a_j(\tau, t). \end{aligned} \quad (10)$$

Here,  $P_{ij}(\tau) := \Pr(Z(t + \tau) = i | Z(t) = j)$  may be obtained by rewriting the (conditional) expectations defining the (forward) branching process as  $T_{ij}(\tau) = P_{ij}(\tau) \sum_k T_{kj}(\tau)$ , which gives

$$P_{ij}(\tau) = T_{ij}(\tau) / \sum_k T_{kj}(\tau). \quad (11)$$

On the other hand,

$$\begin{aligned} \Pr(Z(t + \tau) = i, Z(t) = j) &= \Pr(Z(t) = j | Z(t + \tau) = i) \Pr(Z(t + \tau) = i) \\ &= \tilde{P}_{ji}(\tau, t) p_i(t + \tau), \end{aligned} \quad (12)$$

where  $\tilde{P}_{ji}(\tau, t) := \Pr(Z(t) = j | Z(t + \tau) = i)$  is the transition probability of the *time-reversed* process and is obtained from (10) and (12) as  $\tilde{P}_{ji}(\tau, t) = a_j(\tau, t) P_{ij}(\tau) \times (p_i(t + \tau))^{-1}$ . With Eqs. (8), (9), and (11), one therefore obtains the elements of the backward transition matrix  $\tilde{\mathbf{P}}$  as

$$\tilde{P}_{ji}(\tau, t) = p_j(t) \frac{T_{ij}(\tau)}{\sum_{k,\ell} T_{k\ell}(\tau) p_\ell(t)} (p_i(t + \tau))^{-1}. \quad (13)$$

By differentiating  $\tilde{\mathbf{P}}(\tau, t)$  with respect to  $\tau$  and evaluating it at  $\tau = 0$ , one obtains the matrix  $\mathbf{Q}(t)$  governing the corresponding backward process in continuous time. Its elements read  $Q_{ji}(t) = (d/d\tau) \tilde{P}_{ji}(\tau, t)|_{\tau=0} = p_j(t) (H_{ij} - \delta_{ij} \bar{\mathbf{R}}(t)) (p_i(t))^{-1} - \delta_{ij} \dot{p}_i(t) / p_i(t)$ . Using (4) this simplifies to

$$Q_{ji}(t) = \begin{cases} p_j(t) H_{ij} (p_i(t))^{-1}, & i \neq j, \\ -\sum_{k \neq i} p_k(t) H_{ik} (p_i(t))^{-1}, & i = j. \end{cases} \quad (14)$$

Note that the backward process is, in general, state-dependent (it does not generate a Markov chain). Note also that time reversal works in the same way if sets of types  $\mathcal{X}_k$  instead of single types are considered, as long as mutation and reproduction rates are the same within classes. Furthermore, an analogous treatment is possible both for mutation coupled to reproduction, as well as for discrete generations.

As to the asymptotic behavior of our branching process, it is well known that, for irreducible  $\mathbf{H}$  and  $t \rightarrow \infty$ , the time-evolution matrix  $\exp(t(\mathbf{H} - \lambda_{\max} \mathbf{I}))$  becomes a projector onto the equilibrium distribution  $\mathbf{p}$ , with matrix elements  $p_i z_j$  (e.g., Karlin and Taylor,

1981, the appendix). Here,  $\mathbf{z}$  is the Perron–Frobenius (PF) left eigenvector of  $\mathbf{H}$ , normalized such that  $\sum_i z_i p_i = 1$ . As suggested by our notation, one also has

$$\lim_{\tau, t \rightarrow \infty} \mathbf{z}(\tau, t) = \mathbf{z}, \quad (15)$$

which may be seen from (8).<sup>3</sup> We therefore term  $z_i$  the *relative reproductive success* of type  $i$ .

At stationarity, the matrix governing the backward process simplifies to  $Q_{ji} = p_j(H_{ij} - \delta_{ij}\lambda_{\max})p_i^{-1}$ , which can now be interpreted as a Markov generator. Further, the (asymptotic) ancestor distribution, given by  $a_i = z_i p_i$ , turns out to be the stationary distribution of the backward process, since  $\sum_i Q_{ji} a_i = \sum_i p_j(H_{ij} - \delta_{ij}\lambda_{\max})p_i^{-1} z_i p_i = \sum_i p_j z_i (H_{ij} - \delta_{ij}\lambda_{\max}) = 0$ . Due to ergodicity of the backward process ( $\mathbf{Q}$  is irreducible if  $\mathbf{H}$  is),  $\mathbf{a}$  is, at the same time, the distribution of types along each line of descent (with probability 1).

### 2.3. The Equilibrium Ancestor Distribution

As we saw in the last subsection, there is a simple link between the algebraic properties of  $\mathbf{H}$  and the probabilistic structure of the mutation–selection process at equilibrium, which may be summarized as follows. The PF right eigenvector  $\mathbf{p}$  (with  $\sum_i p_i = 1$ ) determines the composition of the population at mutation–selection balance; the corresponding left eigenvector  $\mathbf{z}$  (normalized so that  $\sum_i z_i p_i = 1$ ) contains the asymptotic offspring expectation (or relative reproductive success) of the various types; and the ancestral distribution, defined by  $a_i = p_i z_i$ , gives the asymptotic distribution of types that are met when lines of descent are followed backward in time (cf. Fig. 2). Figure 3 shows  $\mathbf{p}$ ,  $\mathbf{a}$ , and  $\mathbf{z}$  for a single-step mutation model with a linear fitness function. One sees that  $\mathbf{z}$  decreases exponentially.

For the single-step mutation model, we may directly transform the eigenvalue equation  $\mathbf{H}\mathbf{p} = \lambda_{\max}\mathbf{p}$  into an equation for  $\mathbf{a}$ . To this end, we define a diagonal transformation matrix  $\mathbf{S}$  with non-zero elements  $S_{kk} = \prod_{\ell=1}^k \sqrt{U_{\ell}^- / U_{\ell-1}^+}$  and obtain a symmetric matrix by  $\tilde{\mathbf{H}} := \mathbf{S}\mathbf{H}\mathbf{S}^{-1}$ . The corresponding PF right and left eigenvectors are given by  $\tilde{\mathbf{p}} = \mathbf{S}\mathbf{p}$  and  $\tilde{\mathbf{z}} = \mathbf{S}^{-1}\mathbf{z}$ . But now,

<sup>3</sup> Both  $\mathbf{z}$  and  $\mathbf{p}$  also admit a more stochastic interpretation. If the population does not go to extinction, one has  $\lim_{t \rightarrow \infty} Y_i / (\sum_j Y_j(t)) = p_i$  almost surely, i.e., the stochasticity is in the population size, not in the relative frequencies (Kesten and Stigum, 1966; Hofbauer and Sigmund, 1988, Chapter 11.5). Further, for the *critical process* generated by  $\mathbf{H} - \lambda_{\max}\mathbf{I}$ , one has  $\lim_{t \rightarrow \infty} t \Pr(\mathbf{Y}(t) \neq 0 | \mathbf{Y}(0) = \mathbf{e}_j) = z_j / C$  and  $\lim_{t \rightarrow \infty} (1/t) E(Y_i(t) | \mathbf{Y}(0) = \mathbf{e}_j, \mathbf{Y}(t) \neq 0) = C p_i$ , where  $C$  is a constant; this is the continuous-time analog of a result by Jagers (1975, p. 94). Note that, in the long run, the expected offspring depend on the founder type only through the probability of non-extinction of its progeny.

as  $\tilde{\mathbf{H}}$  is symmetric, we have  $\tilde{\mathbf{z}} \sim \tilde{\mathbf{p}}$  (where  $\sim$  means proportional to). Hence, due to  $a_k = z_k p_k = \tilde{z}_k \tilde{p}_k \sim \tilde{p}_k^2$ , one has  $\tilde{p}_k \sim \sqrt{a_k}$ . Thus, we obtain the following explicit form of the eigenvalue equation for  $\tilde{\mathbf{H}}$ , which will be crucial for the derivation of our main results:

$$[R_k - U_k^+ - U_k^-] \sqrt{a_k} + \sqrt{U_{k-1}^+ U_k^-} \sqrt{a_{k-1}} + \sqrt{U_k^+ U_{k+1}^-} \sqrt{a_{k+1}} = \lambda_{\max} \sqrt{a_k}. \quad (16)$$

Note that Eq. (16) relates the mean fitness of the *equilibrium* population ( $\bar{R} = \lambda_{\max}$ ) to the *ancestor* frequencies  $a_k$ .

### 2.4. Observables and Averages

In this section we define the observables, i.e., measurable quantities, that are used to describe the population on its evolutionary course. Besides the usual population mean, we shall also introduce the mean with respect to the ancestor distribution (see Section 2.3).

We shall consider means and variances of two observables in the following. To this end, we characterize each type (or class)  $i$  by its *fitness value*  $R_i$  and its *mutational distance*  $X_i$  from the reference genotype (or the class  $\mathcal{X}_0$ ). For the biallelic model in particular, mutational distance corresponds to the Hamming distance to  $s_+$ . If, in addition, this is the fittest type,  $X_i$  just gives the number of deleterious mutations. But in general it can also be used to describe the value of any additive trait with equal contributions of sites or loci. Similarly, for single-step mutation, we define  $X_k$  to be the distance from the class  $\mathcal{X}_0$ , thus  $X_k = k$  for class  $\mathcal{X}_k$ . Again,  $X_k$  may be viewed as (the genetic contribution to) any character with discrete values that depends linearly on the mutation classes.

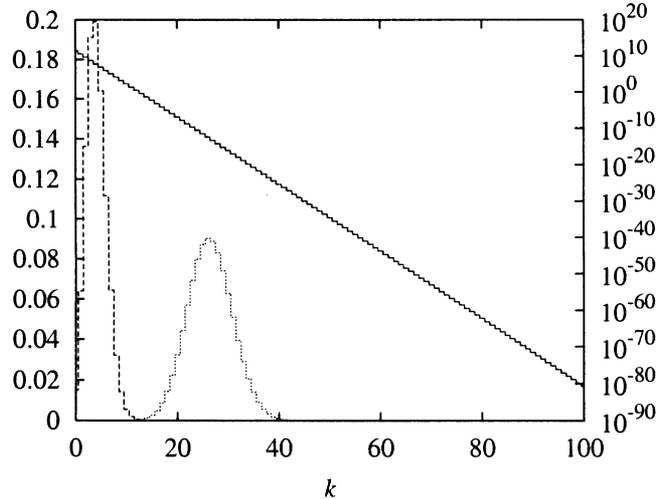
*Population average:* Representing an arbitrary observable as  $(O_i)$ , such as  $(R_i)$  or  $(X_i)$ , we will denote its *population average* as

$$\bar{O}(t) := \sum_i O_i p_i(t). \quad (17)$$

By omission of the time dependence we will indicate the corresponding equilibrium average.

As to mean fitness,  $\bar{R}(t)$  determines the mutation load,  $L(t) := R_{\max} - \bar{R}(t)$ . Here,  $R_{\max} = \max_i R_i$  is the fitness of the fittest genotype, in line with the usual convention (see, e.g., Ewens, 1979; Bürger, 2000). It is well known that the equilibrium value  $\bar{R} := \lim_{t \rightarrow \infty} \bar{R}(t)$  is given by the largest eigenvalue,  $\lambda_{\max}$ , of the evolution matrix  $\mathbf{H}$ .

For the variance of fitness,  $V_R(t) = \sum_i (R_i - \bar{R}(t))^2 \times p_i(t)$ , we differentiate  $\bar{R}(t)$  according to (1), i.e.,



**FIG. 3.** Equilibrium values of population frequencies  $p_k$  (dotted line), ancestor frequencies  $a_k$  (dashed line), and relative reproductive success  $z_k$  (solid line) for the biallelic model with additive fitness  $R_k = \gamma(N - k)$  (where  $\gamma$  is the loss in reproduction rate due to a single mutation), point mutation rate  $\mu = 0.2\gamma$ , mutation asymmetry parameter  $\kappa = \frac{1}{2}$ , and sequence length  $N = 100$ . The logarithmic right axis refers to  $z_k$  only.

$(d/dt)\bar{R}(t) = \sum_i R_i \dot{p}_i(t) = V_R(t) + \sum_{i,j} R_i M_{ij} p_j(t)$ , and hence

$$\begin{aligned} V_R(t) &= \frac{d}{dt} \bar{R}(t) - \sum_{i,j} R_i M_{ij} p_j(t) \\ &= \frac{d}{dt} \bar{R}(t) + \sum_j \left( \sum_i (R_j - R_i) M_{ij} \right) p_j(t). \end{aligned} \quad (18)$$

The interpretation of this completely general formula is as follows: In the absence of mutation, Eq. (18) just reproduces Fisher’s fundamental theorem, i.e., the variance in fitness equals the change in mean fitness, as long as there is no dominance (see, e.g., Ewens, 1979). If mutation is present, however, a second component emerges, which is given by the population mean of the mutational effects on fitness, weighted by the corresponding rates. It may be understood as the *rate of change in mean fitness due to mutation alone*. At mutation–selection balance, this second term is obviously the only contribution to variance in fitness.

For the single-step mutation model in particular, we can define *deleterious* and *advantageous mutational effects* separately as  $s_k^+ = R_k - R_{k+1}$  and  $s_k^- = R_{k-1} - R_k$ , respectively. For decreasing fitness values (which is the usual case, but not strictly presupposed here) these are positive. This way we obtain

$$\begin{aligned} V_R &= \overline{s^+ U^+ - s^- U^-} = \overline{s^+ U^+} - \overline{s^- U^-} \\ &\quad + \text{Cov}(s^+, U^+) - \text{Cov}(s^-, U^-) \end{aligned} \quad (19)$$

for the equilibrium variance, a result we will rely on in the following.

Just as for the fitness distribution, we define the population mean,  $\bar{X}(t) = \sum_{i=0}^N X_i p_i(t)$ , and variance,  $V_X(t) = \sum_i (X_i - \bar{X}(t))^2 p_i(t)$ , of the mutational distance.

*Ancestral average:* We will also need the ancestral average of our observables, that is, the average with respect to the ancestral distribution defined in Eq. (9):  $\hat{O}(\tau, t) := \sum_i O_i a_i(\tau, t) = \sum_i z_i(\tau, t) O_i p_i(t)$ . In the following, we will only be concerned with the ancestral distribution in equilibrium, i.e., with both  $t$  and  $\tau$  going to infinity. We obtain the ancestral average of any observable ( $O_i$ ) in this limit as

$$\hat{O} := \sum_i O_i a_i = \sum_i z_i O_i p_i. \quad (20)$$

These averages may be read forward in time (corresponding to a weighting of the current population with expected offspring numbers), and backward in time (corresponding to an averaging w.r.t. the distribution of the ancestors). A third interpretation is available if the mutation matrix is irreducible, which entails that the equilibrium backward process defined by  $\mathbf{Q}$  is ergodic. Then, with probability 1, the equilibrium ancestral average also coincides with the average of the observable over a lineage backwards in time. Note that the information so obtained is not contained in the population average, which is merely a “time-slice” average. The ancestral mean adds a time component to the averaging procedure, which provides extra information on the evolutionary dynamics. In Appendix A, we shall show that our ancestral averaging coincides

with the way observables are evaluated in a system of quantum statistical mechanics.

### 2.5. Linear Response and Mutational Loss

We now come to another interpretation of the equilibrium ancestor frequencies introduced in Section 2.2. Consider the derivative of the equilibrium mean fitness with respect to the  $i$ th fitness value in a general system of parallel mutation and selection (1):

$$\begin{aligned} \frac{\partial \bar{R}}{\partial R_i} &= \frac{\partial}{\partial R_i} \left[ \sum_{j,k} z_j H_{jk} p_k \right] \\ &= a_i + \bar{R} \frac{\partial}{\partial R_i} \left[ \sum_j z_j p_j \right] = a_i, \end{aligned} \quad (21)$$

where we made use of the normalization condition  $\sum_j z_j p_j = \sum_j a_j \equiv 1$ . The ancestor frequency  $a_i$  therefore measures the *linear response* (or *sensitivity*) of the equilibrium mean fitness to changes in the  $i$ th fitness value.<sup>4</sup> A similar calculation for the response to changes in the mutation rates results in

$$\frac{\partial \bar{R}}{\partial M_{ij}} = (z_i - z_j) p_j. \quad (22)$$

Using (21) and (22), we can express the equilibrium mean fitness as follows:

$$\bar{R} = \hat{R} + \sum_{i,j} z_i M_{ij} p_j = \sum_i R_i \frac{\partial \bar{R}}{\partial R_i} + \sum_{i,j} M_{ij} \frac{\partial \bar{R}}{\partial M_{ij}}. \quad (23)$$

Let us give a variational interpretation for the ancestor mean fitness as well. To this end, we define the *mutational loss*  $G$  of the system as the difference between ancestor and population mean fitness in equilibrium. Assume now that we change all mutation rates  $M_{ij}$  by variations in a common factor  $\mu$ . From (23) and (21) we then find that the mutational loss relates to the linear response of the equilibrium mean fitness to changes in the mutation rates as

$$G := \hat{R} - \bar{R} = -\mu \frac{\partial \bar{R}}{\partial \mu}. \quad (24)$$

Actually, this relation holds for arbitrary (clonal) mutation–selection systems, in particular also if mutation and reproduction are coupled (in which case the mutation *rates* are replaced by mutation *probabilities*).

Equations (21) and (24) may also be used to determine the change in mean fitness if  $\mathbf{H}$  changes to  $\mathbf{H} + \Delta\mathbf{H}$ , to linear order in  $\Delta\mathbf{H}$ . (Small changes in the

fitness values, or mutation rates, may be due to environmental changes, or changes in the genetic background.) Clearly,  $\mathbf{H} + \Delta\mathbf{H}$  has  $\bar{R} + \Delta\bar{R}$  as the largest eigenvalue, with  $\Delta\bar{R} \simeq \sum_i \Delta R_i (\partial \bar{R} / \partial R_i) + \sum_{i,j} \Delta M_{ij} \times (\partial \bar{R} / \partial M_{ij})$  to linear order in  $\Delta R_i$  and  $\Delta M_{ij}$ . If only fitness values are affected, (21) yields

$$\Delta\bar{R} \simeq \sum_i \Delta R_i a_i, \quad (25)$$

where the  $a_i$  belong to the original system. If only the mutation rates change by variations in a common factor  $\mu$ , (24) leads to

$$\Delta\bar{R} \simeq -\frac{\Delta\mu}{\mu} G. \quad (26)$$

We will come to further interpretation and discussion of the mutational loss and the response relations in Section 5.1.

### 2.6. Fitness Functions and Mutation Models

For many of the results and all of our examples, we will restrict our treatment to the case of the single-step mutation model as described by Eq. (2). Although most of our results do not depend on this particular choice we will, for simplicity, concentrate on this scheme here, and only briefly discuss the possible extensions. We will start out with a discussion of fitness functions and mutation schemes in this context. Depending on whether the phenotype or the genotype is considered the primary quantity for the model, the inherent approximation mainly concerns the mutation or the fitness part, respectively.

If  $X_k$  ( $0 \leq k \leq N$ ) are the values of a quantitative trait on which selection acts, fitness may be taken as an arbitrary function of it. The essential assumption, in this case, is that genotypes with equal trait values have equivalent mutation patterns, with mutation in single steps as an additional simplification. This is the original view in which this assumption first appeared, with  $X_k$  as the electric charge of proteins (Ohta and Kimura, 1973). The numerous papers to follow have been reviewed by Bürger (1998, 2000).

If, on the other hand,  $X_k$  is the number of mutations with respect to the wildtype (i.e.,  $X_k = k$  as in the biallelic model), single-step mutation is a natural approximation and directly emerges if mutation and reproduction are modeled as independent processes. The essential simplification, in this case, consists in the choice of the genotype fitness values, which depend only on  $k$ . This way, only the average epistatic effect is included in the model, whereas any variance among epistatically interacting mutations is disregarded.

<sup>4</sup>If mutation is coupled to reproduction, the linear response to variations in the death rate  $D_i$  is given by  $-a_i$ .

Fitness functions of this kind, although undoubtedly lacking much of the biological complexity, have been used as standard landscapes throughout population genetics literature. While the principal reason for this seems to lie in the large simplifications due to permutation invariance, they already take full account of the limited information on fitness provided by mutation accumulation experiments (e.g., Crow and Simmons (1983), but see also the discussion by Phillips *et al.*, 2000). Further, they include a broad range of examples with vastly diverging properties, ranging from simple additive fitness over *quadratic*—or otherwise polynomial or exponential—landscapes with smoothly varying fitness values (e.g. Charlesworth, 1990) to *truncation selection* (e.g., Kondrashov, 1988) and Eigen’s *sharply peaked* landscape (Eigen *et al.*, 1989).

For a consistent treatment of our model in the mutation class limit  $N \rightarrow \infty$  (to be defined in the next subsection), it will be advantageous to think of the fitness values and mutation rates as being determined by the mutational distance *per class* (or site),  $x_k := X_k/N \in [0, 1]$ :

$$R_k = Nr_k = Nr(x_k), \quad U_k^\pm = Nu_k^\pm = Nu^\pm(x_k). \quad (27)$$

Here,  $r_k$  and  $u_k^\pm$  are also introduced as fitness and total mutation rates per class. They can now be thought of as being defined, without loss of generality, by three functions  $r$  and  $u^\pm$  on the compact interval  $[0,1]$ . We will refer to  $r$  as the *fitness function*, and to  $u^+$  and  $u^-$  as the (deleterious and advantageous) *mutation functions* of the model. Both  $u^+$  and  $u^-$  are assumed to be continuous and positive, with boundary conditions  $u^-(0) = u^+(1) = 0$ , and  $r$  to have at most finitely many discontinuities, being either left or right continuous at each discontinuity in  $]0,1[$ . This should include all biologically relevant examples. For the biallelic model, the mutation functions are simple linear functions of  $x$ :

$$u^+(x) = \mu(1 + \kappa)(1 - x), \quad u^-(x) = \mu(1 - \kappa)x. \quad (28)$$

Note that the classical stepwise mutation model (Ohta and Kimura, 1973) is not covered by this framework, since its genotype space  $\mathbb{Z}$  is inherently non-compact.

## 2.7. Three Limiting Cases

Our primary aim in the following sections is to establish simple relations for the equilibrium means and variances of mutational distance and fitness that lend themselves to biological interpretation. Whereas these relations are approximations in the general case, they rest on three limiting cases as pillars, for which they hold as exact identities. All three are biologically meaningful

by themselves, two of them are well studied, and we will show that the formulas reduce to well-known results there.

The first case is the *limit of vanishing back mutations*, defined by  $U_k^- \equiv 0$  in our model. The second one is a *limit of linearity*, in which fitness and mutation rates depend linearly on some trait  $Y_k = Ny_k = Ny(x_k)$  with  $Y_0 = 0$  and  $Y_N = N$ , such as

$$\begin{aligned} r(x) &= r_0 - \alpha y(x), & u^+(x) &= \beta^+(1 - y(x)), \\ u^-(x) &= \beta^- y(x). \end{aligned} \quad (29)$$

Note that, if  $Y_k$  is proportional to the mutational distance  $X_k = k$ , the fitness function is linear whereas the mutation functions  $u^\pm$  reproduce the mutation scheme of the biallelic model if  $\beta^\pm = \mu(1 \pm \kappa)$ . This limit can be understood as the limit of vanishing epistasis, in which the system is known as the Fujiyama model in the sequence space literature (cf. Kauffman, 1993).

The third case is the limit of an infinite number of mutation classes,  $N \rightarrow \infty$ , which we will call *mutation class limit* for short. In the case of the biallelic multilocus model, this limit has been used and discussed in a recent publication (Baake and Wagner, 2001). It addresses the situation of weak or almost neutral mutations, where the average mutational effect (over the mutation classes) is small compared to the mean total mutation rate,  $U \gg s$ . The limit further assumes that differences in mutation rate between neighboring (pairs of) classes are small compared to the mean rate itself. In this case, genetic change by mutation proceeds in many steps of small average effect and the model is a genuine multiclass model in the sense that typically a large number of classes are relevant in mutation–selection equilibrium. Note that only the *average* mutational effect must be small; this includes the possibility of single steps with much larger effect (such as in truncation selection).

Technically, the limit  $N \rightarrow \infty$  is performed such that the mutational effects  $s^\pm$  and the fitness values and mutation rates *per class*,  $r$  and  $u^\pm$ , remain constant. If fitness values and mutation rates are defined by the three functions  $r$  and  $u^\pm$  as described above (27), increasing  $N$  simply leads to finer “sampling” of the functions.

With this kind of scaling, the means and variances *per class* of the observables defined in Section 2.4 approach well-defined limits, which then serve as approximations for the original model with finite  $N$ . We will denote them by the corresponding lower case letters, i.e.,  $\hat{r} := \hat{R}/N$ ,  $v_X := V_X/N$ , etc.; an additional subscript will indicate the limit value, e.g.,  $\bar{x}_\infty := \lim_{N \rightarrow \infty} \bar{x}$ . Note that it is, in general, the *variance per class* of a given quantity that is meaningful in this limit, not the variance of the *quantity per class* (e.g.,  $\text{Var}(X/N)$ ), which tends to zero

(cf. Section 4.4). The described limit is the biological analog of the *thermodynamic limit* in statistical physics. We will further discuss this issue for physically interested readers in Appendix A.

Let us finally compare the mutation class limit with the more familiar *infinite-sites limit*, which, when applied to the biallelic model, also leads to a stepwise mutation model with an infinite number of classes (as found, e.g., in Bürger, 2000). Both limits, however, approximate an original situation with a large, but finite number of types in quite different ways. In the infinite-sites limit, the original model is *extrapolated* to an infinite one by adding new states at the boundaries, where the population distribution is (assumed to be) small. In contrast, the present approach arrives at the limit by *interpolation* of the types of the finite model. Mathematically, this leads to a non-compact state space (such as  $\mathbb{Z}$ ) in the infinite-sites limit, whereas the state space in the mutation class limit is a compact interval (bounded by the extreme types of the original model). To approximate biological observables of the finite model in the limit, the approaches use a different scaling. In the infinite-sites case, the range of Malthusian fitness parameters  $R$  usually diverges (depending on how the extrapolation is done), while the total (“genomic”) mutation rate  $U$  is kept constant. In the mutation class limit, both  $R$  and  $U$  diverge with  $N$ , but the ratio  $U/R$  is kept constant. These differences in scaling result in different ranges of validity of the two limits. The mutation class limit assumes  $U \gg s$ , it is accurate if the total mutation rate is large or fitness differences are small, and allows a sizable fraction of sites to be mutated ( $\bar{x} = \bar{X}/N$  may approach a non-zero limit). In this article, we are mainly interested in this regime, in particular in Section 6, where we discuss error thresholds. Infinite-sites models, on the other hand, typically assume  $U \ll s$ . Then back mutations can be neglected, and the bulk of the population is concentrated on just a few classes with a finite number of mutations.

### 3. RESULTS FOR OBSERVABLE MEANS AND VARIANCES

In this section, we want to give a short summary of our main findings for the single-step mutation model. Derivations and a more extended discussion are postponed to Sections 4 and 5.

A key result of this article is the following estimate of the equilibrium mean fitness, which states a maximum principle and holds as an exact identity in the three limiting cases described in the preceding section:

$$\bar{r} \simeq \bar{r}_\infty = \sup_{x \in [0,1]} (r(x) - g(x)). \quad (30)$$

Here, the function  $g$  is defined as twice the difference between the arithmetic and geometric mean of the mutation functions

$$g(x) = u^+(x) + u^-(x) - 2\sqrt{u^+(x)u^-(x)}. \quad (31)$$

For reasons that will become clear in Section 5.1, we will call it *mutational loss function*. For the biallelic model, it reads explicitly

$$g(x) = \mu(1 + \kappa - 2\kappa x - 2\sqrt{(1 - \kappa^2)x(1 - x)}). \quad (32)$$

In general,  $\bar{r}_\infty$  describes the equilibrium mean fitness  $\bar{r}$  to leading order in  $1/N$  and to next to leading order in  $u^-$ . The approximation is indeed rather accurate already for moderately large  $N$  and/or weak back mutation rates, cf. Section 5.3 and the examples in Section 6.

The maximum principle (30) is closely linked to the ancestor distribution. In particular, if the maximum is attained at a unique value, this is precisely the ancestor mean  $\hat{x}_\infty$ :

$$\bar{r}_\infty = r(\hat{x}_\infty) - g(\hat{x}_\infty) = \hat{r}_\infty - g(\hat{x}_\infty), \quad (33)$$

where the relation  $r(\hat{x}_\infty) = \hat{r}_\infty$  can be proved for all three limiting cases. A corresponding relation for the population mean  $\bar{x}_\infty$ ,

$$\bar{r}_\infty = r(\bar{x}_\infty), \quad (34)$$

holds in the mutation class limit and the linear case, if this equation has a unique solution (e.g., for strictly monotonic  $r$ ).

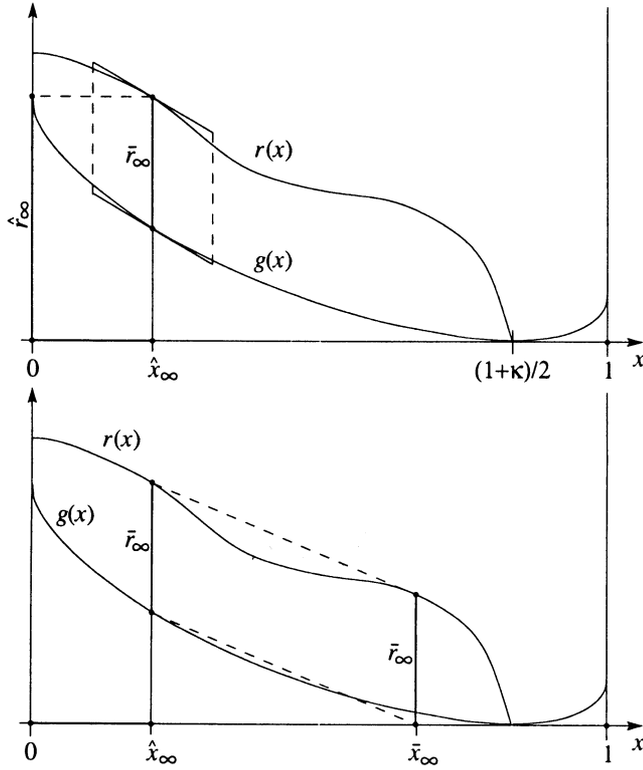
The variances per site of fitness and of distance from wildtype are then given by

$$v_{R,\infty} = -r'(\bar{x}_\infty)(u^+(\bar{x}_\infty) - u^-(\bar{x}_\infty)) \quad \text{and} \\ v_{X,\infty} = \frac{v_{R,\infty}}{(r'(\bar{x}_\infty))^2}, \quad (35)$$

provided  $r$  is differentiable, in which case  $-r'(\bar{x}_\infty)$  is the population mean of the mutational effects. For the biallelic model this is explicitly

$$v_{R,\infty} = -r'(\bar{x}_\infty)\mu(1 + \kappa - 2\bar{x}_\infty) \quad \text{and} \\ v_{X,\infty} = -\frac{\mu(1 + \kappa - 2\bar{x}_\infty)}{r'(\bar{x}_\infty)}. \quad (36)$$

If  $r$  has a jump discontinuity at  $x_{jump}$  from  $r^+$  to  $r^-$  and we have  $r^+ \leq \bar{r}_\infty \leq r^-$ , then  $\bar{x}_\infty = x_{jump}$  and  $v_{R,\infty}$  diverges. In this case,  $V_{r,\infty} = \lim_{N \rightarrow \infty} V_R/N^2$  is finite (cf. the



**FIG. 4.** Graphical constructions for the observable means following the results in Section 3. Upper part:  $\bar{r}_\infty$  is the maximal distance  $r(x) - g(x)$ , cf. (30). This is attained at  $x = \hat{x}_\infty$ , cf. (33), where  $r'(\hat{x}_\infty) = g'(\hat{x}_\infty)$ . Lower part:  $\bar{x}_\infty$  is the solution of  $\bar{r}_\infty = r(\bar{x}_\infty)$ , cf. (34).

example of truncation selection in Fig. 13 and the one in Fig. 11):

$$V_{r,\infty} = (r^+ - \bar{r}_\infty)(\bar{r}_\infty - r^-). \quad (37)$$

The results presented here lead to simple graphical constructions of the means as shown in Fig. 4. This allows for an intuitive overview over the dependence of these quantities on (the shape of) the fitness function and mutation rates, without the need for explicit calculations.

## 4. DERIVATIONS

We now come to the proofs and some first interpretation of the results presented in the previous section. Our starting point is the mutation–selection equilibrium of the single-step model (2) for finite  $N$ , i.e., the

eigenvalue equation

$$[r(x_k) - u^+(x_k) - u^-(x_k)]p_k + u^+(x_{k-1})p_{k-1} + u^-(x_{k+1})p_{k+1} = \bar{r}p_k. \quad (38)$$

For most of our calculations, we will use the equivalent equation for the ancestor distribution, cf. (16),

$$[r(x_k) - u^+(x_k) - u^-(x_k)]\sqrt{a_k} + \sqrt{u^+(x_{k-1})u^-(x_k)}\sqrt{a_{k-1}} + \sqrt{u^+(x_k)u^-(x_{k+1})}\sqrt{a_{k+1}} = \bar{r}\sqrt{a_k}, \quad (39)$$

which is the eigenvalue equation for the largest eigenvalue of the symmetric matrix  $\tilde{\mathbf{H}}$ . For the latter, Rayleigh's principle is applicable, which is a general maximum principle involving the full  $(N+1)$ -dimensional space:  $\bar{r} = \sup_{\mathbf{y}} \sum_{k,\ell} y_k \tilde{\mathbf{H}}_{k\ell} y_\ell / \sum_k y_k^2$ , with non-zero  $\mathbf{y}$ . In the following subsections we will show, for each of the three limiting cases (cf. Section 2.7) separately, how it boils down to the simple scalar maximum principle (30) and relation (33), and give a biological interpretation. We will then come to the derivation of the other identities.

### 4.1. Unidirectional Mutation

We start with the limit of unidirectional mutation, since exclusion of back mutations leads to a considerably simpler situation, and we can show how our findings connect to well-known results. To be specific, we assume  $u_k^- \equiv 0$  and  $u_k^+ > 0$  for  $k < N$ . All results then follow fairly directly from the equilibrium condition (38).

Owing to  $u_k^- \equiv 0$ , the equilibrium distribution  $\mathbf{p}$  in general depends on initial conditions. But  $u_k^+ > 0$  implies that for any such  $\mathbf{p}$ , there exists a particular label  $\hat{k}$ ,  $0 \leq \hat{k} \leq N$ , which divides all classes of genotypes into two parts according to

$$p_k = 0, \quad k < \hat{k}, \quad p_k > 0, \quad k \geq \hat{k}. \quad (40)$$

Equivalently, we obtain for the corresponding left eigenvector  $\mathbf{z}$ :

$$z_k = 0, \quad k > \hat{k}, \quad z_k > 0, \quad k \leq \hat{k}. \quad (41)$$

Since  $a_k = p_k z_k$ , this shows that the only non-zero element of the ancestral distribution is  $a_{\hat{k}} = 1$ , and that  $\hat{k}$  is the equilibrium ancestor mean  $\hat{X}$  of the mutational distance from the reference class  $\mathcal{X}_0$ . In line with this, the mutational distance of every line of ancestors in equilibrium dynamics converges to  $\hat{k}$  (with probability 1). For the classes with non-vanishing frequency, the fitness and total mutation rate are thus related

according to

$$\begin{aligned}\hat{r} - \bar{r} &= r(\hat{x}) - \bar{r} = u^+(\hat{x}), \\ r(x_k) - \bar{r} &< u^+(x_k), \quad k > \hat{k},\end{aligned}\quad (42)$$

the first part of which corresponds to (33).

Although the equilibrium distribution is not unique, (42) implies that the one with maximal mean fitness (which is the only stable one and is automatically obtained in the limit of vanishing back mutations,  $u^- \rightarrow 0$ , or by starting with a population with  $p_0(t=0) > 0$ ) is characterized by

$$\bar{r} = r(\hat{x}) - u^+(\hat{x}) = \max_k (r(x_k) - u^+(x_k)) \quad (43)$$

for arbitrary choices of  $r(x_k)$  and  $u^+(x_k)$ . Obviously, (43) is the discrete version of the maximum principle given in Eq. (30).

If the sequence  $r(x_k)$  or the sequence  $u^+(x_k)$  is monotonically decreasing (as in the biallelic model),  $\hat{k}$  is also the fittest class present in the equilibrium population:

$$\hat{r} = r(\hat{x}) = \max_k \{r(x_k) | p_k \neq 0\}. \quad (44)$$

If additionally  $\hat{k}$  coincides with the class of maximal fitness, i.e.,  $\hat{r} = r_{max}$ , then (42) is a special case of Haldane’s principle, which relates the mutation load  $l$  to the deleterious mutation rate of the fittest class (Kimura and Maruyama, 1966; Bürger, 1998):

$$l = r_{max} - \bar{r} = u^+(\hat{x}). \quad (45)$$

In derivations of (variants of) Eq. (45), it is often tacitly assumed that the equilibrium frequency of the fittest class is non-zero. This, however, is in general not the case and must be made explicit here since we are also interested in the change of the equilibrium distribution with varying mutation rates. This can lead to a shift in  $\hat{k}$  and hence in  $\hat{r}$ .

## 4.2. The Linear Case

If fitness values and mutation rates depend linearly on some trait  $Y$ , as described in (29), the maximum principle holds as an exact identity. This may be derived from (39) by a short direct calculation, which we present in Appendix B.1.

For an interpretation of this result, first consider a trait proportional to the mutational distance  $X$  from the reference class, in which case the system coincides with the Fujiyama model. Since this is a model without epistasis, the means and variances are easily obtained (O’Brien, 1985; Baake and Wagner, 2001). In particular, they are independent of the number of classes. What is more, our derivation shows that they only rely on a

linear dependence of fitness and mutation functions on some trait, as well as the boundary conditions for the mutation functions. This means that they remain unchanged if mutation classes are permuted, or even subjoined or removed.

## 4.3. Mutation Class Limit

Since the proof of the maximum principle (30) and relation (33) in the limit  $N \rightarrow \infty$  is somewhat technical we will just give a sketch here and defer the details to Appendix B.2. The main idea is to look at the system locally, i.e., at some interval of mutation classes in (38) and (39). This will provide us with upper and lower bounds for the mean fitness of a system with finite  $N$  (denoted by  $\bar{r}_N$ ). In the limit  $N \rightarrow \infty$ , they can then be shown to converge to the same value  $\bar{r}_\infty = \lim_{N \rightarrow \infty} \bar{r}_N$ .

For a lower bound, let us consider submatrices of the evolution matrix  $\mathbf{H}$  that, for any class  $\mathcal{X}_k$ , consist of the rows (and columns) corresponding to  $\mathcal{X}_{k-m}$  through  $\mathcal{X}_{k+n}$ . Each of them describes the evolution process on a certain interval of mutation classes at whose boundaries there is mutational flow out, but none in. Thus, each largest eigenvalue,  $\bar{r}_{k,m,n}$ , corresponding to the local growth rate, is a lower bound for  $\bar{r}_N$ . In order to estimate  $\bar{r}_{k,m,n}$ , it is advantageous to use the formulation in ancestor form—with the same local growth rates as largest eigenvalues of the corresponding symmetric submatrices of  $\tilde{\mathbf{H}}$ . Here, lower bounds can be found due to Rayleigh’s principle, and follow from evaluating the corresponding quadratic form for the vector  $(1, 1, \dots, 1)^T$ :

$$\begin{aligned}\bar{r}_N \geq \bar{r}_{k,m,n} \geq \frac{1}{n+m+1} \left[ \sum_{\ell=k-m}^{k+n} (r_\ell - g_{N,\ell}) \right. \\ \left. - \sqrt{u_{k-m-1}^+ u_{k-m}^-} - \sqrt{u_{k+n}^+ u_{k+n+1}^-} \right],\end{aligned}\quad (46)$$

where  $g_{N,\ell} = u_\ell^+ + u_\ell^- - \sqrt{u_{\ell-1}^+ u_\ell^-} - \sqrt{u_\ell^+ u_{\ell+1}^-}$ .

For an upper bound, consider a local maximum of the ancestor distribution, i.e., a  $k^+$  such that  $a_{k^+} \geq a_{k^+ \pm 1}$  (with the convention  $a_{N+1} = a_{-1} = 0$  such a maximum always exists). Evaluating (39) for this  $k^+$  then yields the inequality

$$\bar{r}_N \leq r_{k^+} - g_{N,k^+} \leq \sup_k (r_k - g_{N,k}). \quad (47)$$

Let now  $r_k = r(x_k)$  and  $u_k^\pm = u^\pm(x_k)$  be given by continuous functions as described in Eq. (27), and analogously  $g_{N,k} = g_N(x_k)$ . (The more general case with a finite number of steps in  $r$  is treated in Appendix B.2.)

For an increasing number of mutation classes, fitness values and mutation rates of neighboring classes will then become more and more similar on the scale of the total range of values. More generally, we can use that  $x_k - x_{k \pm i} = \pm i/N \rightarrow 0$  for any finite  $i$  as  $N \rightarrow \infty$ . Defining, for each  $x \in [0, 1]$ , an appropriate sequence  $(k_N) = (k_N(x))$ , such that  $x_{k_N} \rightarrow x$ , we therefore obtain  $r(x_{k_N \pm i}) - g_N(x_{k_N \pm i}) \rightarrow r(x) - g(x)$ , with  $g(x)$  as defined in Eq. (31). Evaluating  $\bar{r}_{k_N, m, n}$  for increasing submatrix dimension  $n + m \rightarrow \infty$  in that limit, we have  $\lim_{n+m \rightarrow \infty} \lim_{N \rightarrow \infty} \bar{r}_{k_N, m, n} = r(x) - g(x)$  for each  $x$ . Combining this with the upper bound (47), in which  $\sup_k (r_k - g_{N, k}) \leq \sup_{x \in [0, 1]} (r(x) - g_N(x)) \rightarrow \sup_{x \in [0, 1]} (r(x) - g(x))$  due to the uniform convergence  $g_N \rightarrow g$  (see Appendix B.2), gives

$$\sup_{x \in [0, 1]} (r(x) - g(x)) \leq \bar{r}_\infty \leq \sup_{x \in [0, 1]} (r(x) - g(x)), \quad (48)$$

which implies the maximum principle (30). As shown at the end of Appendix B.2, the ancestral distribution is sharply peaked around those  $x$  at which  $r(x) - g(x)$  is maximal. Thus, whenever the supremum is unique (which is the generic case), Eq. (33) follows.

#### 4.4. Mean Mutational Distance and the Variances

In this subsection, we derive and discuss the results for the mean mutational distance and the variances, which hold in the linear case and for  $N \rightarrow \infty$ .

If fitness is linear in an arbitrary trait  $y_k = y(x_k)$ , the relation  $\bar{r} = r(\bar{y})$  is immediate. For the variance formulas, we must additionally assume that fitness is linear in the mutational distance,  $r(x) = r_{max} - \alpha x$ , or, equivalently, that all mutational effects are equal. Thus, the covariances in the general formula (19) vanish, and  $v_R = \alpha(\bar{u}^+ - \bar{u}^-)$ . Due to linearity, this also determines the variance in mutational distance as  $v_X = (\bar{u}^+ - \bar{u}^-)/\alpha$ . These relations do not require that  $u^\pm(x)$  are linear in  $x$ ; they reduce to (35) if this is the case.

In the mutation class limit, let us first assume  $r$  to be continuously differentiable on  $[0, 1]$  with derivative  $r'$ . Expressing  $v_{R, \infty}$  as the limit variance for increasing system size  $N$ , and using (19) for the variance of each finite system,  $v_{R, N}$ , we obtain

$$\begin{aligned} v_{R, \infty} &= \lim_{N \rightarrow \infty} \sum_{k=0}^N \left( \frac{r_k - r_{k+1}}{N-1} u_k^+ - \frac{r_{k-1} - r_k}{N-1} u_k^- \right) p_k \\ &= -\overline{r'(u^+ - u^-)}_\infty. \end{aligned} \quad (49)$$

Here, we made use of the fact that the mutational effects converge to the corresponding values of  $-r'$ , i.e., the negative slope of the fitness function.

Since  $r'$  is bounded, (49) in particular shows that  $v_{R, \infty}$  is finite, and hence

$$\begin{aligned} V_{r, \infty} &= \lim_{N \rightarrow \infty} \left[ \sum_{k=0}^N r_k^2 p_k - \left( \sum_{k=0}^N r_k p_k \right)^2 \right] \\ &= \lim_{N \rightarrow \infty} N^{-1} v_{R, N} = 0. \end{aligned} \quad (50)$$

For increasing  $N$ , the distribution of fitness values *per class* therefore concentrates around  $\bar{r}$ . In the limit, if  $r$  is invertible at  $\bar{r}_\infty$ , this fixes the mean mutational distance at  $\bar{x}_\infty = r^{-1}(\bar{r}_\infty)$ , cf. (34), which approximates the mean distance  $\bar{x}_N = \bar{X}_N/N$  of a finite system to leading order in  $N^{-1}$ .

With this, we have  $v_{R, \infty} = -r'(\bar{x}_\infty)(u^+(\bar{x}_\infty) - u^-(\bar{x}_\infty))$ , cf. (35), which approximates  $v_{R, N} = V_{R, N}/N$ . Note that the leading order term w.r.t.  $N^{-1}$  is proportional to  $-r'(\bar{x}_\infty)$ , which is the population mean of the mutational effects in the limit:  $\overline{s_N^\pm} \rightarrow \overline{s_\infty^\pm} = \overline{s_\infty} = -r'(\bar{x}_\infty)$ . (The local curvature of  $r$  only gives rise to higher-order corrections.) Obviously, the leading order depends only on the effective deleterious mutation rate,  $u^+(\bar{x}_\infty) - u^-(\bar{x}_\infty)$ , if this does not vanish. Otherwise, the dominant term is of higher order in  $N^{-1}$ .

The variance in  $x$  can be obtained via the linear approximation  $r(x) \simeq r(\bar{x}_\infty) + r'(\bar{x}_\infty)(x - \bar{x}_\infty)$  as  $v_{X, \infty} = v_{R, \infty}/(r'(\bar{x}_\infty))^2$ , cf. (35). In contrast to  $v_R$ ,  $v_X$  decreases with increasing mutational effects at  $\bar{x}$ . Interestingly,  $\sqrt{v_R/v_X}$  can serve as an estimate for the mean mutational effect (at least in our simple setup)—a quantity which is difficult to determine experimentally. For our numerical examples in Sections 5.3 and 6, this works reasonably well (not shown).

Comparing the results with those for the linear case above, we see that, given  $\bar{r}$ , the infinite mutation class limit can be interpreted as a local linear approximation. This does not mean, however, that non-linearities (i.e., epistasis) are ignored. They enter indirectly through the mean fitness as determined by the maximum principle.

For fitness functions with kinks, the derivation is analogous, as long as the left- and right-hand sided limits of  $r'$ , and thus the mutational effects in the limit  $N \rightarrow \infty$ , remain bounded. If  $r'$  diverges at  $\bar{x}_\infty$ , or if there is even a jump in the fitness function,  $v_R$  diverges according to the above relation. In the latter case,  $V_{r, \infty}$  is finite and determined by the fraction of the population below and above the jump, which yields (37).

## 5. INTERLUDE: APPLICATIONS AND DISCUSSION

### 5.1. Mutational Loss

A central role in this article is played by the mutational loss  $G$ , which was defined in Eq. (24) as the difference between the ancestor and population mean fitness in equilibrium. Let us now add some further interpretation to this quantity. Recapitulating relations (22)–(24), we obtain for  $g := G/N$  in the framework of the general mutation–selection model (1):

$$\begin{aligned} g = \hat{r} - \bar{r} &= -\frac{1}{N} \sum_{i,j} z_i M_{ij} p_j \\ &= \frac{1}{N} \sum_{i,j} M_{ij} (z_j - z_i) p_j = -\mu \frac{\partial \bar{r}}{\partial \mu}. \end{aligned} \quad (51)$$

It is instructive to compare  $g$  with the mutation load  $l = r_{max} - \bar{r}$ . Both quantities describe the effect of mutation on the equilibrium mean fitness. But whereas the mutation load compares the biological system with a fictitious system free of mutations, the loss is essentially a response quantity: In analogy with (26), we have  $\delta \bar{r} \simeq -(\delta \mu / \mu) g$ . Since mutation rates are usually not switched on or off in nature, but may be subject to gradual change, the mutational loss seems to be the quantity of more direct relevance for questions connected with the evolution of mutation rates.

From the above, we see that the loss can be understood as the linear component of the load. In particular, loss and load will coincide if the latter is linear in  $\mu$ . This holds for unidirectional mutation as long as the wildtype has non-vanishing equilibrium frequency (and, more generally, below a wildtype threshold, see Section 6.2.2), where  $r_{max} = \hat{r}$ . In general, however, non-linear terms in  $\mu$  will also contribute to the load and we find  $l > g$ .

The genetic load concept has often been criticized, since the reference genotype (usually the one with maximum fitness) is often extremely unlikely to be found in the population at all. This argument is made precise by Ewens (1979, Chapter 9.2) and Gillespie (1991, Chapter 6.2) for the substitution and the segregation load in finite populations. An analogous point may be made against the mutation load, even in infinite populations: The equilibrium frequency of the fittest class is often close to zero (or may even vanish for unidirectional mutation). Therefore, measurements of  $r_{max}$  in real populations are difficult, if not impossible,

and the evolutionary significance of the reference type seems questionable.

This problem is circumvented in the definition of the mutational loss. As a response quantity,  $g$  is well defined as long as it is meaningful to think of a system as in equilibrium. Measurements of  $g$  could make use of marker techniques in (bacterial or viral) clones in order to determine clone sizes (and thus  $z$ ) and ancestor frequencies, or determine directly the response of  $\bar{r}$  to changes in mutation rates, e.g., by comparing strains with different mutation repair efficiencies.

Up to this point, we have entirely concentrated on the mutational loss as a response quantity. There is, however, a second line of interpretation, which clarifies the role of  $g$  in the equilibrium dynamics and also sheds some light on the maximum principle. If an individual mutates from  $j$  to  $i$ , its offspring expectation changes by  $z_j - z_i$ , where the sign determines whether a loss (+) or gain (–) is implied. Since the mutational flow from  $j$  to  $i$  in equilibrium is  $M_{ij} p_j$ , the entire system loses offspring at rate  $\sum_{i,j} (z_j - z_i) M_{ij} p_j$ , which is the same as  $G$  (compare with Eq. (23) or (51)).

The mutational loss does not include any information about the destination of the “lost” offspring. This, however, may easily be found by recalling that, asymptotically, every ancestor of type  $i$  leaves  $z_i p_i$  descendants of type  $j$  in the equilibrium population. Further,  $p_i(z_i - 1) = a_i - p_i$  is the excess offspring produced by an  $i$ -individual. We thus come to a picture of a constant flow of mutants from the ancestor to the equilibrium population.

Let us now turn to the mutational loss function  $g(x)$ . Recall that, in the derivation of the maximum principle in the mutation class limit, we obtained  $r(x) - g(x)$  as the leading eigenvalue of a local open subsystem around  $x$ ; if  $\bar{r}_\infty$  is the death rate due to population regulation in the entire system,  $r(x) - \bar{r}_\infty - g(x)$  is the net growth rate of the subsystem at  $x$ . Hence,  $g(x)$  must describe the rate of mutational loss due to the flow out of the local system. This can be made more precise within the framework of large-deviation theory, which will be presented in a future publication. If  $r(x) - g(x)$  has a unique maximum, in which case the ancestor distribution has a single peak, the maximum principle (30) along with  $\bar{r} \rightarrow \bar{r}_\infty$  and  $\hat{r} \rightarrow r(\hat{x}_\infty)$  as  $N \rightarrow \infty$  implies that the mutational loss  $g = G/N$  converges to  $g(\hat{x}_\infty)$ . Thus,  $g(\hat{x})$  can be taken as an approximation to the actual mutational loss  $g$  in this case.

Let us finally add a remark concerning the influence of epistasis on the mutational loss (in the sense of a response quantity) in the single-step model. Following the suggestion of Phillips *et al.* (2000), we speak of

negative (positive) epistasis w.r.t. some class  $k$  if  $R_{k+1} - R_k < (>) R_k - R_{k-1}$ . This entails *synergistic* (*antagonistic*) interaction of deleterious mutations. This way, negative and positive epistasis are connected to concavity and convexity of the fitness function, and thus to its second derivative (if well-defined) being negative, respectively, positive.

Let us now keep the mutation rates fixed and compare fitness functions with different degrees of epistasis. Let  $g$  be a decreasing loss function, and  $r$  and  $r_s$  two decreasing fitness functions which are either convex or concave, and only differ in an open subinterval of  $[0, 1]$  that includes  $\hat{x}$  (the ancestral mean trait under  $r$ ). Assume that  $r_s - r$  is a concave function in our subinterval. Then  $r_s$  describes more negative, or less positive, epistasis than  $r$ . Under the above assumption,  $r_s - r$  has a unique maximum whose position we denote by  $x_0$ . As is most easily seen from the graphical representation of the maximum principle (Fig. 4), one then finds  $\hat{x}_s > \hat{x}$  whenever  $x_0 > \hat{x}$  (and vice versa), where  $\hat{x}_s$  is the ancestral mean trait under the modified fitness function. Since  $g(x)$  is decreasing, it follows that  $g(\hat{x}_s) < g(\hat{x})$  if  $x_0 > \hat{x}$ . If  $\hat{x}$  is small (as may be considered typical of realistic examples), increased negative epistasis will reduce the loss. The opposite may be said of decreased negative or increased positive epistasis, in line with the fact that the loss is maximal for the sharply peaked landscape, which displays extreme positive epistasis.

## 5.2. Haldane's Principle and Evolution of Mutational Effects

As we have seen in the discussion of the unidirectional case, the maximum principle reduces to a well-known form of the Haldane–Muller principle in that limit. Using the concept of the ancestor distribution, we will now re-analyze this principle in the broader context of models with back mutations. We will also discuss consequences for the evolution of mutational effects and mutational robustness.

For models without back mutations to the fittest genotype with non-zero equilibrium frequency, Haldane's principle says that the difference in fitness between this type,  $\hat{i}$ , and the population mean is equal to the total mutation rate for  $\hat{i}$ . For our general model this reads  $L = R_{max} - R_{\hat{i}} + \sum_{j \neq \hat{i}} M_{j\hat{i}}$ , where  $M_{j\hat{i}}$  is the mutation rate from the fittest type (or class)  $\hat{i}$  to some other type (or class)  $j \neq \hat{i}$ . Note in particular that the load is independent of the mutant fitness values if the wildtype itself has non-zero frequency in equilibrium, i.e.,  $\hat{i} = 0$ . We will assume  $R_0 = R_{max}$  for simplicity in this section.

If back mutations to the fittest class are present, but mutation rates, denoted by  $u$ , are small compared to the fitness advantage,  $u \ll s$ , the relation for the load is modified by a correction term of order  $u^2/s$  (Bürger and Hofbauer, 1994). In the following, we will reproduce this result in our setting by deriving an explicit expression of the correction term for the single-step model. We will also show that this leading order contribution of the back mutations is exactly contained in the estimate of  $\bar{r}$  as derived from the maximum principle.

Let us assume, for notational simplicity, that the wildtype is also the fittest type present in the equilibrium population, and remains so if back mutations are switched off. Suppose that the back mutation rates  $u_k^-$  are small compared to the fitness effects, but not necessarily the deleterious mutation rates  $u_k^+$ . We then obtain, to linear order in  $u_1^-$ ,

$$l \simeq u_0^+ - \frac{\partial \bar{r}}{\partial u_1^-} \Big|_{u_1^- = 0} u_1^- = u_0^+ - \frac{p_1}{p_0} \Big|_{u_1^- = 0} u_1^-, \quad (52)$$

where we have used Eq. (22) for the derivative of  $\bar{r}$  with respect to the mutation rates, and  $z_0 = 1/p_0$ ,  $z_1 = 0$  for  $u_1^- = 0$ . Calculating  $p_1/p_0$  from the equilibrium condition for the mutation–selection equation, we find

$$l = u_0^+ - \frac{u_0^+ u_1^-}{s_0^+ - u_0^+ + u_1^+} + O([u^-]^2). \quad (53)$$

This is in accordance with the result of Bürger and Hofbauer (1994) if also  $u_0^+, u_1^+ \ll s_0^+$ .

On the other hand, starting with a linear interpolation of the fitness and mutation functions of the form  $r(x) = r_0 + Nx(r_1 - r_0)$ ,  $u^+(x) = u_0^+ + Nx(u_1^+ - u_0^+)$ , and  $u^-(x) = Nx u_1^-$  for  $0 \leq x \leq 1/N$ , we find the load by using  $\bar{r}$  from the maximum principle. To linear order in  $u^-$ , a lengthy but elementary calculation yields that  $r(x) - g(x)$  is maximized at  $Nx = u_0^+ u_1^- / (s_0^+ - u_0^+ + u_1^+)^2$ , and we again obtain Eq. (53) for the load. We can therefore conclude that the maximum principle, when applied to finite  $N$ , still gives results that are correct to linear order in the back mutation rates (cf. Section 5.3).

In the preceding paragraphs, back mutations have merely played the role of a small perturbation of the system with unidirectional mutation. Our main interest in this article, however, lies in the case of sufficiently large mutation rates—or sufficiently small fitness effects of mutations (as in a nearly neutral landscape) such that the equilibrium distribution is no longer dominated by one or a few wildtype states, but is dispersed over many classes. This is exactly the situation in which one would assume back mutations to become important, with effects beyond a second-order correction term. At the same time, this is the domain of validity of the mutation

class limit, in which the maximum principle is also exact. We then obtain

$$l = r_{max} - r(\hat{x}) + g(\hat{x}) \quad [\leq g(0) = u^+(0)] \quad (54)$$

as an estimate for the mutation load. Clearly, the load is no longer independent of the fitness function as soon as the ancestral mean fitness  $\hat{r} = r(\hat{x})$  differs from the wildtype fitness. Note, however, that the only quantity that matters is the deviation of the ancestor mean fitness from the wildtype fitness.

It is instructive to compare the load for different fitness functions. Let  $r_s$  and  $r$  be fitness functions with  $r(0) = r_s(0) = r_{max}$ , and  $r_s(x) \geq r(x)$  for all  $x \in [0, 1]$ . By the maximum principle, the load with  $r_s$  cannot be larger than that with  $r$ . If  $r_s(x) > r(x)$  at  $x = \hat{x}$ , the ancestral genotype under  $r$ , the load with  $r_s$  is strictly smaller than with  $r$ . In this sense, higher mutant fitness tends to decrease the mutation load (and vice versa).

Let us now extend these thoughts to the evolution of mutational effects. To this end, we consider a general mutation–selection model (i.e., not restricted to permutation invariant fitness or single step). Assume there is an additional modifier locus, which is tightly linked to the other loci and changes the fitness of one or several of the original types or classes. (In the biallelic model, this may, for example, happen through epistatic interactions outside our permutation invariant fitness scheme.)

Let now a modifier be introduced into the equilibrium population at low frequency at time  $t = 0$  (by mutation or migration), and consider its fate for  $t \rightarrow \infty$ . If there is no further mutation at the modifier locus, the modifier will asymptotically fix (or get lost) in terms of relative frequencies,  $p(t) = y(t)/(\sum_i y_i(t))$ , if the modified system has a larger (smaller) leading eigenvalue than the original one, in which case we write  $\delta\bar{r} > 0$  ( $\delta\bar{r} < 0$ ). If  $\delta\bar{r} = 0$ , the modifier will equilibrate at an intermediate frequency, the exact value of which depends on the initial conditions.

The above argument is analogous to the clonal competition mechanism as described for mutation rate modifiers in asexual populations (for review, see Sniegowski *et al.*, 2000). It requires slight modification if mutation at the original loci is unidirectional. Here, the fate of the modifier also depends on the genetic background it is introduced into at  $t = 0$ . If the fitness modifications are so small that the fittest type present remains the same in equilibria with and without modifier, the modifier will always get lost if it does not already occur in individuals of that type at  $t = 0$ . This follows since all other types asymptotically expect no offspring.

Note that the competition mechanism just described works *within* the population, the separation of genotypes with and without modifier being due to tight linkage of the modifier to the primary loci. In particular, no group selection is implied.

What consequences, now, does this have for the possibility of mutational effects to evolve? Again, the answer involves the ancestor distribution. We have seen in (25) that changing the fitness values  $r_i$  to  $r_i + \delta_i$  will change the equilibrium mean fitness by

$$\delta\bar{r} \simeq \sum_i \delta_i a_i \quad (55)$$

to first order in the  $\delta_i$ . From this, we now obtain the following intuitive picture: In order for a modification to prevail in an equilibrium population, it has to invade the ancestors; otherwise, it will be “washed away”.

Let us discuss this in some more detail. According to our above discussion, the fate of the modifier is entirely determined by  $\delta\bar{r}$  if we have back mutations. Now, the right-hand side of Eq. (55) may be interpreted as the selection coefficient of the modifier with respect to the ancestor distribution—assuming that the modifier is statistically independent of the other loci. In order to understand why this quantity governs the leading order of  $\delta\bar{r}$ , consider infinitesimally small fitness changes  $\delta_i$ , in which case Eq. (55) becomes exact. Here, mutation will indeed drive the modifier distribution towards statistical independence in an initial period of time. In order to eventually spread to fixation, the modifier now has to compete successfully against those types whose descendants make up the equilibrium population at an even later time. These, however, follow the ancestor distribution. In this sense,  $\delta\bar{r}$  may be understood as measuring the modifier’s growth within the ancestor population. In the same vein, the vector of the ancestor frequencies can be seen as the gradient of the mean fitness, pointing into the direction of the indirect (i.e., second order) selection pressure exerted on the fitness values  $r_i$ .

This long-term picture is in sharp contrast to the initial growth of the modifier in the population, which is determined by its selection coefficient with respect to the *equilibrium population* and of course depends on the distribution of the modifier over the types at  $t = 0$ . If  $\delta\bar{r}$  is positive (negative), the modifier will asymptotically fix (vanish) even if its initial selection coefficient is negative (positive). Note, however, that this process may be very slow if  $\delta\bar{r}$  is small.

If there is no back mutation to the fittest class (or type)  $\hat{i}$  present, this is the absorbing state of the backward process, in which all lineages end, and the ancestor distribution is entirely concentrated there ( $a_i =$

1,  $a_j = 0, j \neq \hat{i}$ ). So Eq. (55) leads back to the prediction of Haldane's principle that the mean fitness is independent of the mutant fitness values in this case. In order to "invade the ancestors", a modifier must be introduced into the fittest type in the first place, and increase its fitness.

Assume now that the wildtype fitness is kept fixed but mutational effects at the wildtype are modified by variations of the mutant fitness values. Such modifiers are *canalizing* (or modifiers for mutational robustness) if they increase the mutant fitnesses, and *decanalizing* (modifiers for antirobustness) if they decrease the mutant fitnesses (cf. Wagner *et al.*, 1997). It is now clear from Eq. (55) that only an *increase* of mutant fitness values may lead to an evolutionary advantage. Independent of the fitness landscape or of mutation patterns, we thus never find a potential for the evolution of antirobustness in mutation–selection models; however, mutational robustness may, indeed, evolve. Here, modifiers increasing the fitness of mutant classes with large ancestor frequencies will be under particularly large (positive) selection pressure. If modifiers have deleterious side effects, these may even be the only ones that persist and go to fixation.<sup>5</sup>

Let us, for further analysis, consider two limiting cases of the mutation scheme now. If mutation is unidirectional, neither modifications for robustness nor for antirobustness will change the mean fitness (at least under the usual assumption that the wildtype is present in the original equilibrium). We may conclude that there is no selection pressure on the mutant fitness values at all in this simple setting, and hence no potential for these to evolve either.<sup>6</sup> On the other hand, if the mutation matrix is symmetric,  $M_{ij} = M_{ji}$ , the ancestor frequencies are proportional to the square of the population frequencies,  $a_i \sim p_i^2$ . Thus, the landscape is evolvable exactly in those regions in which the equilibrium frequency of the population distribution is high.

Note that we may come to different results here depending on whether genotype classes or single genotypes are the relevant entities. If mutation between genotypes is symmetric (as in our biallelic model with  $\kappa = 0$ ), modifiers of single genotypes will be particularly

<sup>5</sup>Note, however, that no predictions are made here concerning invadability of modifier mutations, or fixation probabilities, if random drift becomes a weighty factor.

<sup>6</sup>For very large, but finite populations (where Muller's ratchet does not operate but there is drift among classes of equal fitness) the fixation probability of clones with and without the modifier is ultimately determined only by the initial sizes of the respective wildtype classes (Gabriel and Bürger, 2000). Any modifier which enters the wildtype class at low frequency will therefore get lost from this class and, consequently, from the population with high probability.

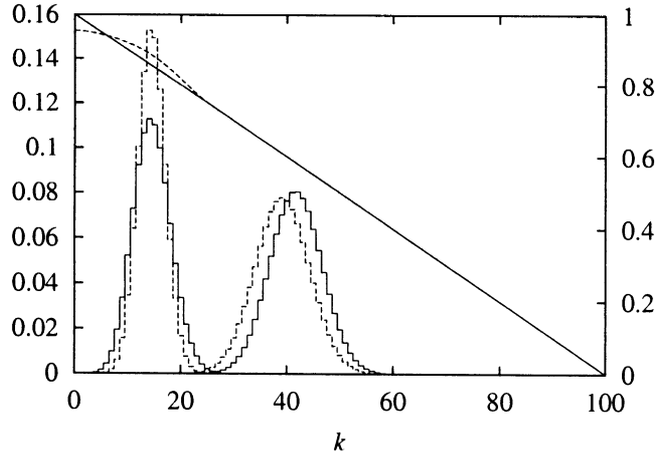
important if the corresponding equilibrium frequency is high. For modifiers of whole genotype classes, however, the asymmetric mutation scheme with respect to the classes is relevant, and the maximum of the ancestor distribution will in general deviate from the maximum of the population distribution.

In order to see what happens between these limiting cases, let us restrict our discussion again to the single-step mutation model. Here, the ancestor distribution becomes sharply concentrated around  $\hat{x}$  with an increasing number of mutation classes (cf. Appendix B.2). Similar to the case of unidirectional mutation, only a very minor part of the fitness function will thus experience appreciable selection pressure. Note that this part need neither extend to the types which contain the bulk of the equilibrium distribution (concentrated around  $\bar{r} < \hat{r}$ ), nor the largest fitness values at  $r_{max} > \hat{r}$ . If robustness modifiers have deleterious side effects, only those which lead to buffering in the ancestor region will prevail at all. Therefore, if robustness evolves by the mechanism described, the strongly *differential* selection pressure might lead to the emergence of synergistic epistasis at the same time. This is illustrated in Fig. 5, where modification of the fitness function leads to a flattening near its "summit" at  $x < \hat{x}$  relative to the "slope" at  $x > \hat{x}$ . The example also shows that an increase in fitness around  $\hat{r}$  may compensate for a deleterious side effect of the modifier mutation which decreases the wildtype fitness.

### 5.3. Accuracy of the Approximation

In this subsection we wish to illustrate the accuracy of the analytical expressions for means and variances given in Section 3. To pay respect to the invariance of the equilibrium distributions under scaling of both reproduction and mutation rates with the same factor, we introduce  $\gamma$  as an overall constant for the reproduction rates. It should be chosen to represent roughly the average effect of a single mutation on the reproduction rate in a mutant genotype (with the maximum number of mutations considered) as compared to the wildtype. This does not exclude the possibility that effects of single mutations may be quite large. In the figures, both reproduction and mutation rates are given in units of this constant, i.e., as  $r/\gamma$ , respectively  $\mu/\gamma$ .

Figure 6 displays an example of a biallelic model that deviates from all three exact limiting cases described in Section 2.7, and, for comparison, three modifications that are closer to one of the exact limits each. All numerical values, also in the rest of this article and in Figs. 3 and 5, are virtually exact and, if not noted



**FIG. 5.** Comparison of population frequencies  $p_k$  (near  $k = 40$ ) and ancestor frequencies  $a_k$  (near  $k = 15$ ) for the biallelic model with  $\mu = 0.365\gamma$  (where  $\gamma$  is the loss in reproduction rate due to a single mutation as in Fig. 3),  $\kappa = \frac{1}{2}$ , and  $N = 100$ . The right axis refers to the fitness functions used: additive fitness  $r(x)/\gamma = 1 - x$  (solid lines), and a modified version (dashed lines) that is favored with respect to the additive one. The modified fitness is increased in regions of high ancestor frequencies. In this particular example, it is slightly decreased at the wildtype and unchanged in other regions of vanishing ancestor frequencies, but note that the success of a modification is independent of the fitness values there.

otherwise, obtained by the power method (Wilkinson (1965, Chapter 9) also known as von Mises iteration) with the evolution matrix  $\mathbf{H}$ . For continuous fitness functions, the approximate expressions for the observable means agree with the exact ones up to corrections of order  $N^{-1}$  (as indicated by numerical comparison, not shown) or of order  $(u^-)^2$  (cf. Section 5.2). For fitness functions with jumps, the error seems to be at most of order  $N^{-1/2}$  (cf. Fig. 13); for a jump at  $x = 0$  such as in the sharply peaked landscape; however, the corrections to  $\bar{r}$  appear to be still of order  $N^{-1}$  for the biallelic model (cf. Fig. 7).

Further examples, exhibiting more conspicuous features, are shown in Section 6. For most of them, one will also find good agreement of numerical and analytical values for the means for sequences of length  $N = 100$ ; for the variances, however, one sometimes needs longer ones, like  $N = 1000$ . In the biallelic model, we generally find stronger deviations for higher mutation rates, as in this regime back mutations become more and more important, whereas for small mutation rates, deviations are of linear order in  $\mu$ .

## 6. MORE APPLICATIONS: THRESHOLD PHENOMENA

In this section, we will take a closer look at how the equilibrium behavior of a mutation–selection system

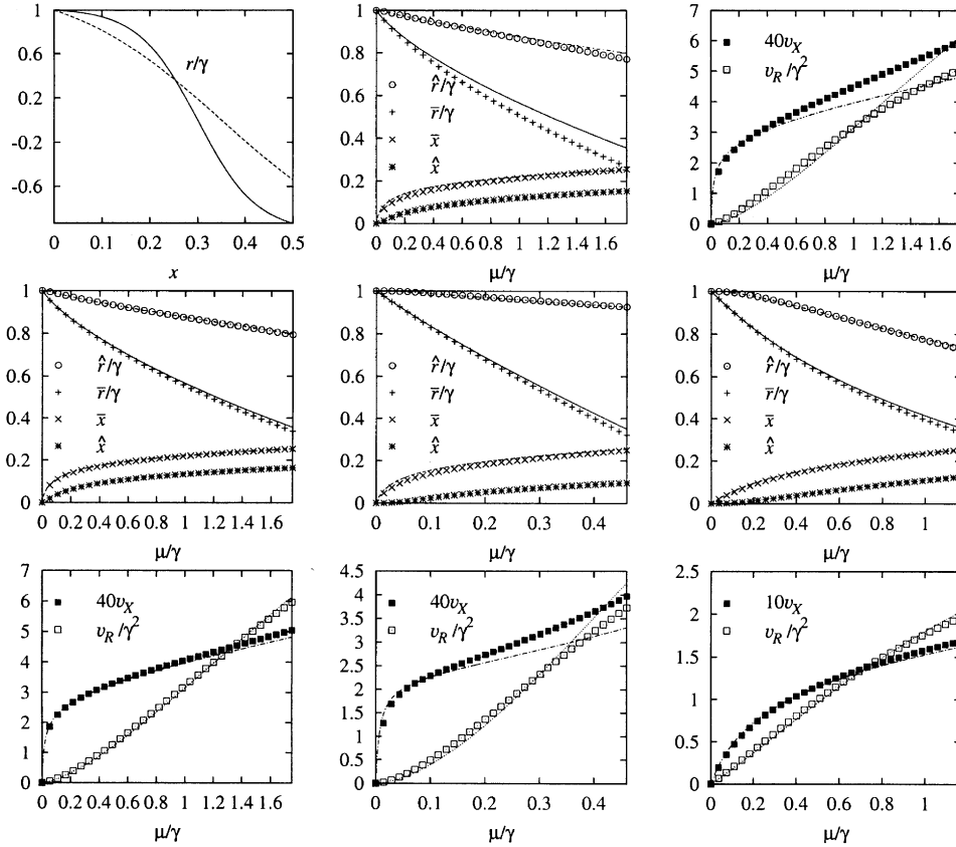
changes if the mutation rates are allowed to vary relative to the corresponding mutational effects. In order to keep the overall shapes of the fitness and mutation functions constant, we vary all mutation rates by a common scalar factor  $\mu \geq 0$ . Concentrating on the single-step mutation model in this section, we choose  $\mu$  as the mean mutation rate over all classes,

$$\mu = (2N)^{-1} \sum_{k=0}^N (u_k^+ + u_k^-) \quad (56)$$

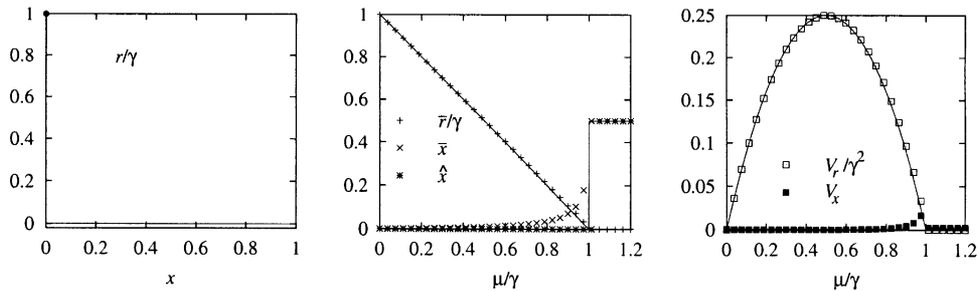
(recall that  $u_0^- = u_N^+ = 0$ ). This is consistent with the definition of  $\mu$  as the mean point mutation rate for the biallelic model, cf. Eq. (28) and Fig. 1. By slight abuse of notation, we define the shape of the mutational loss function as  $g(1, x) = \mu^{-1}g(x)$  (which does not depend on  $\mu$ ), and introduce  $\mu$  as a variable parameter via  $g(\mu, x) = \mu g(1, x)$ .

### 6.1. Mutation Thresholds

Consider a population in mutation–selection balance. Usually, if mutation rates change slightly, the population will move on to a new equilibrium with the observables, like means and variances of traits and fitness, close to the old ones. At certain *critical mutation rates*, however, threshold phenomena may occur, associated with much larger effects on traits or fitness. The prototype of this kind of behavior is the so-called *error threshold*, first observed in a model of prebiotic evolution many years ago (Eigen, 1971) and discussed in



**FIG. 6.** The top row refers to a biallelic model that deviates from all three exact limiting cases described in Section 2.7 in having a strongly non-additive fitness function  $r/\gamma$  (left, solid line), symmetric site mutation ( $\kappa = 0$ ), and small sequence length ( $N = 20$ ). The mean values of the observables (middle) and corresponding variances (right) are shown as a function of the mutation rate  $\mu/\gamma$ , both for the model itself (symbols) and according to the expressions given in Section 3 (lines, sometimes hidden by symbols). Even here, we find reasonable agreement. Deviations, however, are visible for larger mutation rates. As can be seen from the last two rows, going towards any of the three exact limits, i.e., increasing the number of mutation classes (left,  $N = 100$ ), going to more asymmetric mutation (middle,  $\kappa = 0.8$ ), or using a different fitness function with less curvature (right,  $r/\gamma$ : top left, dashed line) these deviations vanish quickly. In the case of increasingly asymmetric mutation, however, this is not true for the variances, since the approximation becomes only exact here in either of the other two limits (cf. Section 4.4).



**FIG. 7.** The error threshold of the *sharply peaked landscape* (left) with  $r(0) = \gamma$  (bullet) and  $r(x) = 0$  for  $x > 0$  (line), for the biallelic model with symmetric mutation ( $\kappa = 0$ ). The observable means are shown in the middle, the variances on the right. Symbols correspond to  $N = 100$ , lines to the expressions in Section 3. The ancestral fitness  $\hat{r}(\mu)$  (not shown) jumps from  $\gamma$  to 0 at  $\mu = \gamma$ . Note that  $V_r$  follows the scaling described by (49) and is given by (37) for  $N \rightarrow \infty$ .

numerous variants ever since (for review, see Eigen *et al.*, 1989; Baake and Gabriel, 2000).

In the following, we will discuss and classify “error threshold like” behavior in our model class. We shall, however, avoid the term error threshold as the collective name for all threshold effects that may be observed, but rather, and more generally, speak of *mutation thresholds*. This is because the definition of the error threshold is closely linked to the model in which it had been observed originally, namely the quasi-species model with the sharply peaked fitness landscape. While many effects of the original error threshold will turn out to generalize easily to the much larger class of models considered here, the criterion of the *loss of the wildtype*, which has frequently been taken as the defining property of the error threshold, seems to be applicable only in special cases.

We want to be as general as possible as far as the fitness model and mutation schemes are concerned, but specific about the responsible evolutionary forces. Error thresholds have also been described as driven by the joint action of mutation and segregation (Higgs, 1994) or recombination (Boerlijst *et al.*, 1996). We will not consider these phenomena.

Let us now define the notion mutation threshold. Ideally, a characterization should give a precise mathematical definition in the modeling framework which, at the same time, captures biologically significant behavior. As may be seen from the varying and sometimes incompatible definitions that have previously been suggested for the error threshold (see, e.g., the discussion in Baake and Gabriel, 2000), this can be a complex problem. Let us therefore start with a verbal description:

*A mutation threshold* for a particular trait or fitness is the pronounced change of the equilibrium distribution of the trait or fitness values within a narrow range of mutation rates. Here, the threshold phenomenon is purely due to the interplay of mutation and selection.

Note that we only consider effects on distributions, not on absolute numbers. This demarcates mutation thresholds from mutational meltdown effects (cf. Gabriel *et al.*, 1993).

In order to come to a stringent mathematical definition, a two-fold limit must be considered for any general mutation–selection model (1). These are the infinite population limit, which we assumed right from the beginning, and the limit of an infinite number of mutation classes.

Application of the *infinite population limit* is a direct consequence of the last condition in the verbal definition above. As mutation thresholds result from mutation and

selection alone, they must persist in the absence of genetic drift. Hence, unlike drift effects (like Muller’s ratchet), these phenomena cannot be avoided by increasing population size. For the purposes of analysis and classification, therefore, deterministic models provide the right framework. Of course, aspects of thresholds should also persist in (large) finite populations, if the phenomena are biologically relevant. For some models this has been confirmed in numerical studies (Nowak and Schuster, 1989; Bonhoeffer and Stadler, 1993): While certain properties of the threshold (such as the critical mutation rate) may be altered by finite population size, the threshold effect as such is not eliminated by drift.

The *infinite mutation class limit*, on the other hand, is needed to give the vague notion of a “pronounced change” a more precise meaning in mathematical terms. Our intention is to specify this notion as a *discontinuous* change of a biological observable (or, at least, of one of its derivatives) as a function of  $\mu$ . In any finite system with back mutations, however, this clearly conflicts with the fact that the population frequencies are analytic functions of the mutation rates.<sup>7</sup> The same problem also arises for the definition of phase transitions in physics. Phase transitions, therefore, are defined as non-analyticity points of the free energy in the *thermodynamic limit* (i.e., for infinitely large systems). Since the infinite mutation class limit is just the counterpart of the thermodynamic limit in our models (cf. Appendix A), we take this concept of theoretical physics as our guideline and characterize different types of mutation thresholds by discontinuities or kinks in the equilibrium mean and/or variance of some trait or of fitness as a function of  $\mu$  in the limit  $N \rightarrow \infty$ . (Therefore, we will omit the subscript  $\infty$  throughout this section.)

Let us add a few comments concerning this strategy:

1. Firstly, and most importantly, the proposed procedure is in accordance with the original definition of the error threshold: In the quasi-species model, a kink in the wildtype frequency (and thus the mean fitness) as a function of the total mutation rate was first established by an approximate formula for finite sequence length by Eigen (1971), which was later found to be exact in the limit  $N \rightarrow \infty$  (Swetina and Schuster, 1982). The finite system is thus effectively approximated by an infinite one. In order to capture

<sup>7</sup>This follows from the Perron–Frobenius theorem and the fact that the PF eigenvalue and eigenvector depend analytically on the matrix entries. Since the PF eigenvalue is real and unique under the above conditions, it never crosses with the second largest eigenvalue as a function of any model parameter, such as mutation rates.

- the behavior of the finite system in the limit, the total mutation rate and the selective advantage of the wildtype must scale with the number of classes  $N$  (thus leaving the *mean* mutational effect per class constant; cf. Franz and Peliti, 1997). The equivalence of this phenomenon with a magnetic phase transition has first been established by Leuthäusser (1987), and was later used by Tarazona (1992) and many others.
2. Whereas we have introduced the mutation class limit mainly as an approximation for real systems with a finite number of classes, its use in the present context rather has a conceptual reason. Analogous to phase transitions in physics, the threshold should be considered as a property of the limit that manifests itself (as a “pronounced change”) in finite systems as well (cf. the numerical examples in Figs. 7, 9, and 11–15).
  3. Discontinuities in the biological observables can also arise in finite systems if the evolution matrix  $\mathbf{H}$  is reducible (as for unidirectional mutation). Then mutation thresholds can be directly defined for finite  $N$ . This has previously been done by Wiehe (1997) and will be discussed in Section 6.3 below.

## 6.2. Description of Threshold Types

Following the lines of the above reasoning, we now come to a description of different types of mutation thresholds. In our list we will not include *any* discontinuous change that might occur, but rather concentrate on pronounced changes of potential evolutionary significance. To this end, we will take the original error threshold of the sharply peaked landscape as our reference and analyze four of its characteristic properties, namely (cf. Fig. 7):

- A kink in the population mean fitness,
- the loss of the wildtype from the population,
- complete mutational degradation, and
- a jump in the population mean of the mutational distance (or some additive trait).

For these threshold effects, we will check whether and how they extend to the permutation-invariant class of mutation–selection models. We will discuss their origin, analyze how they are related, and formulate criteria for the fitness function to exhibit each threshold effect, or *type of threshold*, separately.

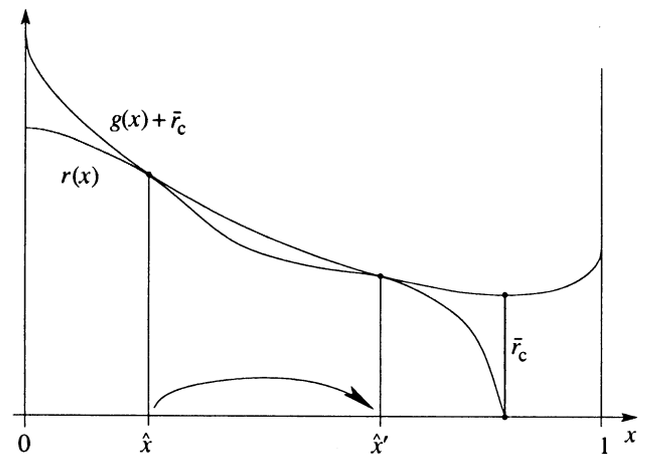
**6.2.1. Fitness Thresholds.** As we will see below, the kink in the population mean fitness is, in many respects,

the most fundamental aspect to classify mutation thresholds. We therefore discuss it first.

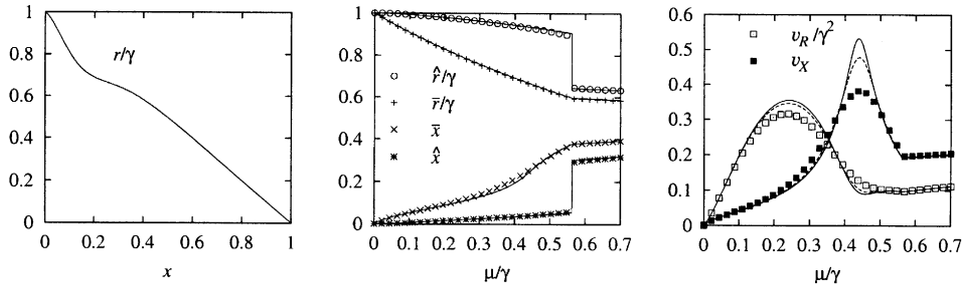
*Phenomenon:* The most pronounced change that may happen to the *fitness* distribution at some critical mutation rate  $\mu_c$  is characterized by a kink in the mean fitness  $\bar{r}$  as a function of  $\mu$  (i.e., a jump in its derivative). We will refer to this phenomenon as a mutation threshold in fitness, or *fitness threshold* for short. Using Eq. (51) and the maximum principle, we see that an alternative definition can be given in terms of the ancestor distribution. Here, a fitness threshold is defined by a jump in the mutational loss (as a function of  $\mu$ ),  $g = g(\hat{x}) = -\mu \partial \bar{r} / \partial \mu$ , corresponding to jumps in  $\hat{x}$  and the ancestor mean fitness  $\hat{r} = r(\hat{x})$ . As a consequence of the kink in  $\bar{r}$ , the mean mutational distance  $\bar{x}$ , and the variances  $v_R$  and  $v_X$ , will typically show a kink as well.

*Interpretation and graphical representation:* The origin of a fitness threshold is easily understood from the maximum principle. For a generic choice of  $\mu$ , the function  $r(x) - g(\mu, x)$  is maximized for a unique  $x = \hat{x}$ . For some fitness functions, however, there are particular values of  $\mu$  that lead to multiple solutions. It is precisely this phenomenon of two distinct ancestor distributions becoming degenerate with respect to the maximum principle which marks the threshold. This may be illustrated graphically as shown in Fig. 8.

Let us add a remark concerning the transferability of these notions to the original “biological” model with fixed, finite  $N$ . In defining fitness thresholds in the mutation class limit, we have tacitly assumed that the



**FIG. 8.** Graphical construction of the fitness threshold, following Fig. 4. At the critical mutation rate  $\mu_c$ , the maximum of  $r(x) - g(x)$  is not unique. Thus, with  $\mu$  being increased across  $\mu_c$ , the mean of the ancestor distribution jumps from a position of relatively high fitness and high mutational loss,  $\hat{x}$ , to lower fitness genotypes with less mutational loss at  $\hat{x}'$ . The figure also shows how the population mean fitness is constructed at the threshold.



**FIG. 9.** Means (middle) and variances (right) for a biallelic model with asymmetric mutation ( $\kappa = 0.4$ ), and a fitness function  $r/\gamma$  (left) that displays strong positive epistasis near  $x = 0.15$ . One therefore observes a *fitness threshold* ( $\mu_c/\gamma \simeq 0.562$ ). Symbols correspond to  $N = 100$ , dashed lines to  $N = 500$ , and solid lines to the expressions in Section 3.

fitness function  $r$  reasonably interpolates the discrete fitness values of the original model. In order to avoid “pseudo-thresholds” driven by purely local features of the fitness function on a scale smaller than  $1/N$ , the effects should be stable under different interpolations. Note that one way to assure this is to apply the maximum principle only to the discrete point set  $\{x_k\} = \{k/N\}$  and ask for a jump in  $\hat{x}$  over more than one mutation class. In any case, the example in Fig. 9 and those in Figs. 11–15 show that the threshold effects are usually clearly visible also for finite  $N$ .

*Criterion:* To derive a criterion for the existence of a fitness threshold for a given fitness function  $r$ , we use the following argument. According to the above definition, a fitness threshold is signaled by a jump in  $\hat{x}$ . Thus, in any fitness landscape *without* a threshold,  $\hat{x}(\mu)$  varies continuously from the wildtype position  $x_{\min} := \lim_{\mu \rightarrow 0} \hat{x}(\mu)$  to the position of the mutation equilibrium,  $x_{\max} := \lim_{\mu \rightarrow \infty} \hat{x}(\mu)$ , where  $g(x_{\max}) = 0$  (for the biallelic model,  $x_{\max} = (1 + \kappa)/2$ ). Therefore, at each  $x$  in the half-open interval  $[x_{\min}, x_{\max}[$  the maximum in (30) is attained for some finite  $\mu$ . If  $r$  and  $u^\pm$  are twice continuously differentiable in the closed interval  $[x_{\min}, x_{\max}]$ , then  $g$  is twice continuously differentiable in  $]x_{\min}, x_{\max}[$  and we arrive at the following sufficient condition for the non-existence of a fitness threshold:

$$\forall x \in ]x_{\min}, x_{\max}[ \exists \mu > 0: \quad r'(x) = g'(\mu, x) \quad \text{and} \quad r''(x) < g''(\mu, x). \quad (57)$$

Expressing  $\mu = \mu(x)$  through the derivatives of  $r$  and  $g$ , we can state an existence condition in the following general form, cf. Appendix C.1.

*There is a fitness threshold in the mutation–selection equilibrium at some critical mutation rate  $\mu_c$  if and only if*

$$\sup_{x \in [x_{\min}, x_{\max}]} \left( r''(x) - \frac{r'(x)g''(x)}{g'(x)} \right) \geq 0. \quad (58)$$

For the biallelic model, this reads

$$\sup_{x \in [x_{\min}, x_{\max}]} \left( r''(x) - \frac{-r'(x)}{2x(1-x)(1-2x+2\kappa\sqrt{x(1-x)/(1-\kappa^2)})} \right) \geq 0. \quad (59)$$

In the special case that the supremum in (58) is zero, but is assumed only in a single point  $x_0$ , there is actually no jump in  $\hat{x}$ . Here, we obtain limiting cases of a threshold, in the sense that a jump in  $\hat{x}$  may be obtained by arbitrarily small changes in the slope or curvature of  $r$  or  $g$ . Typically, this limiting behavior is indicated by an infinite derivative of the function  $\hat{x}(\mu)$  at  $\hat{x} = x_0$  (cf. Appendix C.1).<sup>8</sup>

Discontinuities in the fitness function or its derivatives can formally be included in (58) by considering left- and right-hand sided limits separately. For a kink in  $r$ , we formally set  $r'' = \infty$  or  $r'' = -\infty$ , respectively, if  $r'$  increases or decreases at this point (which makes (58) true in the former, but not in the latter case). Finally, a jump in  $r$  always results in a fitness threshold.

Note that the criteria presented here do not indicate whether there are one or multiple thresholds for a given combination of  $r$  and  $u^\pm$ . Neither do they provide direct information about the value of  $\bar{r}$  at the threshold, or about  $\mu_c$ . In fact, (58) and (59) are independent of the scalar factor  $\mu$ , but only depend on the shapes of the mutation and fitness function. Answers to these questions, however, are easily derived from the maximum principle for any specific  $r$  and  $u^\pm$ , and may also be

<sup>8</sup> In physics, this kind of behavior corresponds to the important class of continuous phase transitions, cf. Appendix A. In the biological models, however, these non-generic limiting cases do not seem to justify a category of their own.

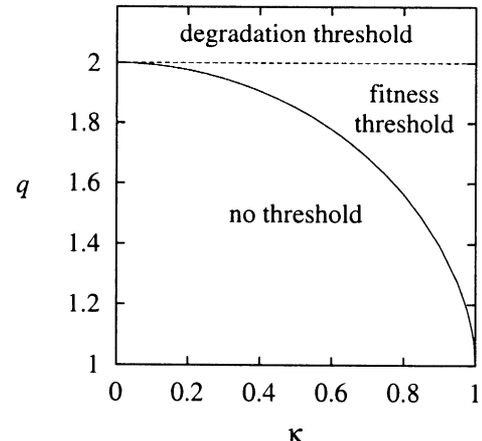
obtained from the graphical construction, cf. the discussion in the preceding paragraph.

*Discussion:* Under what conditions should we expect a fitness threshold to exist in a mutation–selection system? The above criterion (58) compares  $r''$ , which measures the epistasis of the fitness function, with  $g''$  weighted by a factor  $r'/g'$ . Under the reasonable assumption that the rate of back mutations  $u^-$  increases with the distance to the wildtype, whereas the rate of deleterious mutations  $u^+$  decreases, we have  $g' < 0$  and  $r'/g' > 0$  for decreasing fitness functions. Typically, if the curvature of  $u^+$  and  $u^-$  is not too large, we also find  $g'' > 0$ . The criterion then shows that a finite minimum strength of positive epistasis ( $r'' > 0$ , cf. the end of Section 5.1) is required for a fitness threshold. For the biallelic model with  $r(x) = (x_{max} - x)^q$ , this is shown in Fig. 10. Vanishing curvature or even concavity of the mutational loss function,  $g'' \leq 0$ , on the other hand, may even lead to thresholds for fitness functions with negative epistasis.

As will become apparent in Appendix A, the fitness threshold as defined above is the biological counterpart of a first-order phase transition in physics. Since  $\hat{x}$ , which translates into the magnetization, plays the role of the order parameter, the phase transition is generically first order, and continuous only in the limiting case mentioned above. Note that positive epistasis with quadratic exponent  $q = 2$  in a biallelic model with symmetric site mutation ( $\kappa = 0$ ), as has been discussed by Baake and Wagner (2001), is just such a limiting case. The physical analogy shows that a fitness threshold is indeed a true collective phenomenon on the level of the sites or loci. The essential self-enhancing effect simply is that in regions of positive epistasis the selection pressure decreases with any new deleterious mutation.

**6.2.2. Wildtype Thresholds.** The *loss of the wildtype* is the classic criterion for the original error threshold as defined by Eigen (1971): For the sharply peaked landscape, the frequency  $p_0$  of the wildtype (or master sequence) remains finite for small mutation rates even for  $N \rightarrow \infty$ , but vanishes above the critical mutation rate. The same effect may be observed for any fitness function with a jump at the wildtype position  $x_{min}$ .<sup>9</sup> Note that this does not depend on whether we assume the wildtype class to contain only a single or a large number of genotypes (the latter case has sometimes been called the *phenotypic error threshold*, cf. Huynen et al., 1996).

<sup>9</sup>As the mean fitness varies continuously, the wildtype frequency in the limit decreases linearly with the mutation rate, until the mean fitness reaches the lower value at the jump. For larger mutation rates, the wildtype frequency in the limit is zero due to the sharpness of the population distribution for  $N \rightarrow \infty$  (cf. Section 4.4).



**FIG. 10.** The figure shows, as a solid line, the minimum exponent  $q$ , parametrizing epistasis of the fitness function  $r(x) = (x_{max} - x)^q$ , that is needed to obtain a fitness threshold in the biallelic model as a function of the asymmetry parameter  $\kappa$  of the site mutations rate. The exponent varies continuously from quadratic (for symmetric site mutation,  $\kappa = 0$ ) to linear (for unidirectional mutation,  $\kappa = 1$ ). For  $q > 2$  (dashed line), the fitness threshold is also a degradation threshold (see Section 6.2.3). For this combination of fitness and mutation functions, a wildtype threshold only occurs for unidirectional mutation ( $\kappa = 1$ ).

If  $r$  is continuous at  $x_{min}$ , however, the population distribution spreads over a large number of mutation classes with similar fitness for any finite mutation rate. While for finite  $N$  the frequency in any class remains positive for arbitrary  $\mu$  (as long as there are back mutations), the frequency of any single mutation class (including the wildtype class) vanishes for  $N \rightarrow \infty$ . According to the original definition, error thresholds therefore depend on strongly decanalized wildtypes in the sense that deleterious mutations with small mutational effects are virtually absent. While such a model was found to be adequate in certain cases, such as the evolution of coliphage Q $\beta$  and certain viruses (cf. Eigen and Biebricher, 1988), and could be favored by pleiotropy (Waxman and Peck, 1998), slightly deleterious mutations are generally assumed to occur in most biologically relevant situations (Kimura, 1983, Chapter 8.7; Ohta, 1998).

Still, one may ask for some related phenomenon that goes together with the loss of the wildtype in all models in which this effect is observed,<sup>10</sup> but defines a threshold also in a broader model class. The fitness threshold as defined above does not meet this requirement, since

<sup>10</sup>I.e., basically for fitness functions with a jump at the wildtype, and for certain models with unidirectional mutation, see the discussion in Section 6.3.

fitness functions with a jump at the wildtype may well have multiple fitness thresholds, but only lose their wildtype once. Instead, we will give a definition which is based on the ancestor distribution.

*Phenomenon:* We define the wildtype threshold as the largest mutation rate  $\mu_c^- > 0$  below which the ancestral mean fitness coincides with the fitness of the wildtype:

$$\hat{r}(\mu) = \hat{r}(0) = r_{max}, \quad \mu < \mu_c^-. \quad (60)$$

The threshold may equivalently be defined as the largest  $\mu_c^-$  below which  $\hat{x}(\mu) = x_{min}$ . As a consequence, the population mean fitness  $\bar{r}$  responds linearly to an increase of the wildtype fitness if  $\mu < \mu_c^-$ , but becomes independent of (sufficiently small) changes in the wildtype fitness above the threshold.

Note that for unidirectional mutation, the ancestral average  $\hat{x}$  (in general) also denotes the fittest class with non-vanishing equilibrium frequency for any finite  $N$ , cf. Eq. (44). In this special case, the wildtype thus indeed vanishes from the population at  $\mu_c^-$ . Threshold criteria in models with special unidirectional mutation schemes have been derived previously, see the discussion below in Section 6.3.

*Criterion:* For a wildtype threshold to occur,  $r(x) - g(\mu, x)$  must be maximized at  $x = x_{min}$  for some  $\mu > 0$ . Assuming  $r$  and  $g$  to be continuously differentiable for  $x > x_{min}$ , we arrive at the criterion

$$\lim_{x \searrow x_{min}} \frac{g(1, x) - g(1, x_{min})}{r(x) - r(x_{min})} = \lim_{x \searrow x_{min}} \frac{g'(1, x)}{r'(x)} < \infty, \quad (61)$$

see Appendix C.2 for a proof. Fitness functions with a jump at the wildtype position lead to a threshold for any continuous  $g$ .

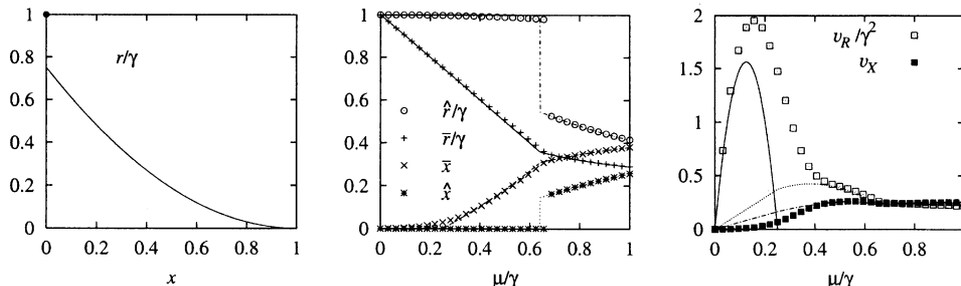
*Discussion:* Note first that a wildtype threshold will always lead to non-analytic behavior of  $\hat{x}(\mu)$  and  $\bar{r}(\mu)$  in  $\mu_c^-$  and is therefore closely related to a fitness threshold.

In general, however, it need not show up as a prominent feature with a jump in means or variances as functions of the mutation rate. If we have a fitness threshold with a jump in  $\hat{x}(\mu)$  at  $\hat{x} = x_{min}$ , however, this will also be a wildtype threshold. In a system with a series of thresholds, the wildtype threshold (if it exists) is always the one with the smallest  $\mu_c$ .

The existence of a wildtype threshold, and also the “loss of the wildtype” where applicable, depends on the strength of the deleterious mutational effect at the wildtype, measured by  $r'(x_{min})$ . The degree to which the wildtype requires a fitness advantage to avoid the threshold depends on the mutational loss function. If  $g$  has a finite derivative at  $x_{min}$ , we always obtain a threshold if the mutational effects do not tend to zero. In many important situations, like the biallelic model and  $x_{min} = 0$ , however, a wildtype threshold requires fitness functions with a rather sharp peak, like  $r(x) \sim -x^p$  with  $p \leq 1/2$  or the one used in the example in Fig. 11. Note that this result depends on back mutations, which make the slope of  $g$  diverge at  $x = 0$ . For  $u^- \equiv 0$ , however, the situation changes drastically, and we obtain a threshold if only  $r'(0) < 0$ , as described above.

Since  $g(0) = u^+(0)$ , we see from Eqs. (30) and (33) that  $\bar{r}$  and  $\bar{x}$  (but not necessarily the variances) are unaffected by back mutations for mutation rates below the wildtype threshold. Further, the mutation load coincides with the mutational loss,  $l = r_{max} - \bar{r} = \hat{r} - \bar{r} = u^+(0)$ , and therefore provides a meaningful measure for changes in  $\bar{r}$  if the mutation rate is varied. In this sense  $\mu_c^-$  may be seen as a point up to which back mutations can be safely ignored.

**6.2.3. Degradation Thresholds. Phenomenon:** A far reaching effect of the error threshold is that selection



**FIG. 11.** Means (middle) and variances (right) for a model with symmetric mutation ( $\kappa = 0$ ),  $N = 100$  (symbols), and the fitness function  $r(x) = \frac{3}{4}\gamma(1-x)^2$  with an additional single peak of height  $\gamma$  at  $x = 0$  (left). Due to the latter, one finds a *wildtype threshold* ( $\mu_c^-/\gamma \simeq 0.641$ ), which is also a fitness threshold. Lines correspond to the expressions in Section 3. For  $1 \leq \bar{r}/\gamma < \frac{3}{4}$ , i.e.,  $0 \leq \mu/\gamma < \frac{1}{4}$ , the variance in fitness no longer follows Eq. (36), but scales differently and is given by (37) for  $N \rightarrow \infty$  (see the discussion in Section 4.4). For finite  $N$ , we can approximate  $v_R$  by a combination of both relations, where (37) and (36) dominate for small and large  $\mu$ , respectively. Note that  $\bar{r}$  is analytic at  $\mu/\gamma = \frac{1}{4}$ ; we thus have no fitness threshold at this point.

altogether ceases to operate. We define a degradation threshold as the smallest mutation rate  $\mu_c^+$  above which the population mean fitness is insensitive to any further increase of the mutation rate:

$$\frac{\partial \bar{r}}{\partial \mu} = -g(1, \hat{x}(\mu)) = 0, \quad \mu > \mu_c^+. \quad (62)$$

This is equivalent to the condition  $\hat{x}(\mu) = x_{max}$  for  $\mu > \mu_c^+$ . Also, the other means and variances then coincide with their values in mutation equilibrium, and the population is degenerate.

*Criterion:* Selection ceases to operate according to the above definition if and only if  $r(x) - g(\mu, x)$  is maximal at  $x = x_{max}$  (where  $g(\mu, x_{max}) = 0$ ) for any finite  $\mu > \mu_c^+$ . Since  $g$  is continuous and strictly positive for  $x < x_{max}$  and  $\mu > 0$ , it is sufficient to compare the asymptotic behavior of  $r$  and  $g$  in the neighborhood of  $x_{max}$ , cf. Appendix C.3:

$$\lim_{x \nearrow x_{max}} \frac{r(x) - r(x_{max})}{g(1, x)} = \lim_{x \nearrow x_{max}} \frac{r'(x)}{g'(1, x)} < \infty. \quad (63)$$

*Discussion:* The degradation threshold is related to the fitness threshold in an analogous way as the wildtype threshold above. In particular, we always find non-analytic behavior of  $\hat{x}(\mu)$  and  $\bar{r}(\mu)$  at  $\mu_c^+$ , but not necessarily a jump or a kink. However, a fitness threshold with a jump of  $\hat{x}(\mu)$  onto  $x_{max}$  is necessarily a degradation threshold. If there is a series of thresholds connected with a system fulfilling (63), the degradation threshold obviously is the last one as  $\mu$  increases.

Criterion (63) implies an important necessary condition for a degradation threshold, namely  $r(x_{max}) > -\infty$ , i.e., genotypes should not be lethal at this point. This parallels a well-known sufficient condition for the existence of a normalizable limit distribution for arbitrary mutation rates in models with non-compact state space (Moran, 1977; Bürger, 2000, p. 128). From a biological point of view, a finite value of  $r(x_{max})$  means

that not the whole genome, but only the part relevant for a specific function or phenotypic property is included in the model, and the genetic background is under sufficiently strong selection to be stable under the mutation rates considered and guarantees survival of the population. We may then obtain mutational degradation w.r.t. *the function under consideration* if this function is less robust under mutation than the background and fitness thus levels out at a finite value. Essentially, this is the threshold criterion previously given by Wagner and Krall (1993) in their treatment of single-step models with unidirectional mutation (see the discussion in Section 6.3).

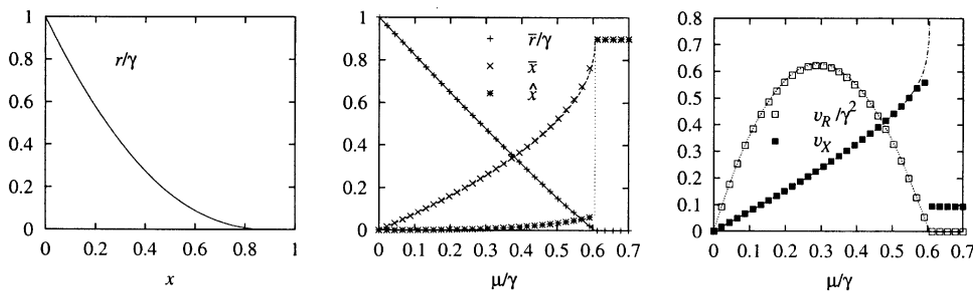
For the more general model with back mutations, we see that  $r(x)$  must approach the fitness level at  $r(x_{max})$  sufficiently fast in order to fulfill (63). For the biallelic model, it is easy to show that we need positive epistasis with at least a quadratic exponent, i.e.  $r \sim r(x_{max}) + \alpha(x_{max} - x)^2$ . Clearly, we always obtain mutational degradation if  $r(x) = r(x_{max})$  already for  $x < x_{max}$ , corresponding to the reasonable assumption that a minimum of non-random coding region is needed for the gene or function considered to show a fitness effect at all. An example for a degradation threshold is given in Fig. 12.

Note finally that we obtain a degradation threshold that at the same time is a wildtype threshold (and a fitness threshold with a jump of  $\hat{x}$  from  $x_{min}$  to  $x_{max}$ ) if and only if

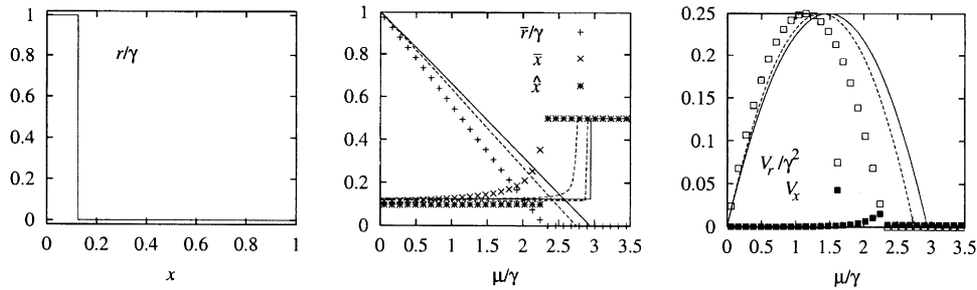
$$\sup_{x \in [x_{min}, x_{max}]} \left( r(x) - r(x_{max}) - g(x) \frac{r(x_{min}) - r(x_{max})}{g(x_{min})} \right) \leq 0 \quad (64)$$

as is most easily seen with the help of the graphical representation, cf. Fig. 8. Clearly, Eq. (64) is fulfilled for the sharply peaked landscape used in Fig. 7, but also for truncation selection, see Fig. 13.

**6.2.4. Trait Thresholds. Phenomenon:** As stated above, there is usually a kink in the population mean



**FIG. 12.** Means (middle) and variances (right) for a model with asymmetric mutation ( $\kappa = 0.8$ ),  $N = 100$  (symbols), and the fitness function  $r(x) = \gamma(x_{max} - x)^q / (x_{max})^q$  with  $x_{max} = (1 + \kappa)/2 = 0.9$  and  $q = 2.2$  (left). As  $q > 2$ , one finds a *degradation threshold* ( $\mu_c^+/\gamma \simeq 0.606$ ), which is also a fitness threshold, cf. Fig. 10. As  $\hat{r}$  behaves just like  $r(\hat{x})$  with a similar accuracy of the approximation, it is not shown here.



**FIG. 13.** Means (middle) and variances (right) for a model with symmetric mutation ( $\kappa = 0$ ) and *truncation selection*, i.e.,  $r(x) = \gamma$  for  $x \leq \frac{1}{8}$  and  $r(x) = 0$  otherwise (left). As in the sharply peaked landscape, cf. Fig. 7, one finds a combined fitness, wildtype, degradation, and trait threshold ( $\mu_c/\gamma \simeq 2.94$ ). Also, the variance in fitness follows a different kind of scaling as described by (49) and is given by (37) for  $N \rightarrow \infty$ . Symbols correspond to  $N = 100$ , dashed lines to  $N = 1000$ , and solid lines to the expressions in Section 3. As  $\bar{r}$  behaves just like  $r(\bar{x})$  with similar accuracy, it is not shown here. Note that the deviations of the approximate expressions are somewhat stronger (of order  $N^{-1/2}$ ) for fitness functions with jumps, cf. Section 5.3.

of the mutational distance  $\bar{x}(\mu)$  (or some other trait) at a fitness threshold. The most pronounced change in the equilibrium distribution of  $x$ , however, is a jump of  $\bar{x}$  at some mutation rate  $\mu_c^x$ , referred to as a *trait threshold*. Since a discontinuous change in  $\bar{x}$  is usually accompanied by a jump in the *local* mutation rates  $u^\pm(\bar{x})$  as well as  $r'(\bar{x})$ , it typically also leads to jumps in  $v_X$  and  $v_R$ . The mean fitness, however, is not at all affected at such points (if they do not coincide with a fitness threshold as defined above).

*Criterion:* Since the equilibrium mean fitness  $\bar{r}(\mu)$  as a function of the mutation rate is always continuous, we easily conclude from  $\bar{r} = r(\bar{x})$  that a jump in  $\bar{x}$  occurs if and only if the fitness function is *not* strictly decreasing from  $x_{min}$  to  $x_{max}$ .

*Discussion:* Obviously, any fitness landscape with a trait threshold also fulfills (58) and thus also has a fitness threshold, but not vice versa. We have  $\mu_c \geq \mu_c^x$  (i.e., the jump in  $\bar{x}$  in general precedes the fitness transition with the jump in  $\hat{x}$ ); see the example in Fig. 14. This shows, in particular, that with varying mutation rate there may be large changes in the phenotype that may be accompanied by changes in the fitness variance, but have virtually no effect on mean fitness. Trait and fitness thresholds should, therefore, be clearly distinguished. In contrast to the fitness threshold or a phase transition in physics, the trait threshold is not driven by collective (self-enhancing) action, but simply mirrors a local feature of the fitness function.

### 6.3. Unidirectional Mutation

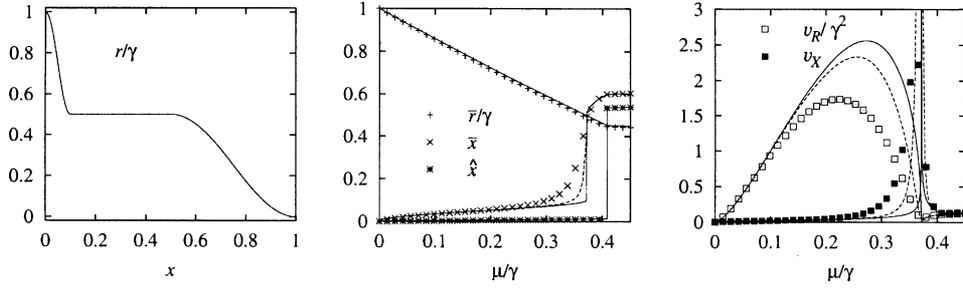
In this section, we briefly discuss how the definitions of mutation thresholds specialize for unidirectional

mutation ( $\kappa = 1$  for the biallelic model). An example is given in Fig. 15. We shall also take the chance to make contact with previous results on threshold criteria by Wagner and Krall (1993) and by Wiehe (1997), where related models were studied.

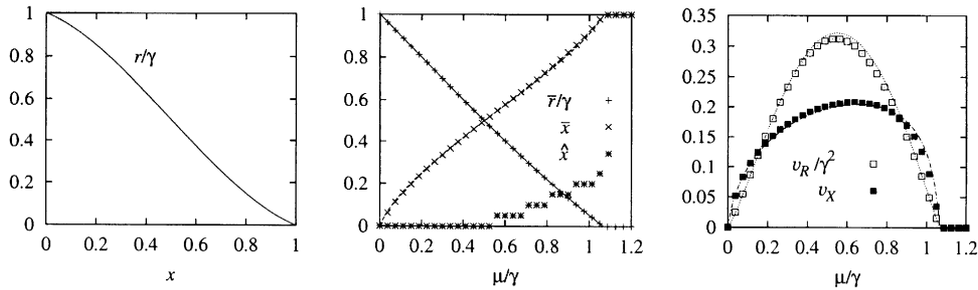
For the above definition of thresholds, the maximum principle in the mutation class limit has played a central role. Since, for vanishing back mutations, it reduces to Haldane's principle and is also exact for finite  $N$ , many of the notions above can be formulated directly, avoiding the limit. As has been described in Section 4.1, the ancestral mean of the mutational distance (or trait) agrees with the minimal  $x$  in the equilibrium population. Since this minimum can only assume discrete values for finite  $N$ , jumps in  $\hat{x}$  will necessarily occur for some  $\mu$ . For a system with a large number of mutation classes (which we consider here), this should, however, not be regarded relevant. In line with the above reasoning on the applicability of the fitness threshold definition for finite systems and previous definitions of the error threshold for unidirectional mutation, it seems reasonable to restrict the term threshold to the first and the last jump, i.e., the loss of the wildtype (Wagner and Krall, 1993; Wiehe, 1997) and the point of complete mutational degradation (Wiehe, 1997), and to jumps of  $\hat{x}$  over more than one class.

Threshold criteria are easily found as analogs of the above relations (note that  $g$  reduces to  $u^+$  if back mutations vanish). As condition for the existence of a fitness threshold with a jump over more than one class, for example, we obtain for monotonic  $u_k^+$ :

$$\max_k \left[ \frac{s_k^+}{s_{k+1}^+} - \frac{u_k^+ - u_{k+1}^+}{u_{k+1}^+ - u_{k+2}^+} \right] \geq 0. \quad (65)$$



**FIG. 14.** Means (middle) and variances (right) for a model with asymmetric mutation ( $\kappa = 0.5$ ) and a fitness function  $r/\gamma$  (left) with an ambiguity for  $r(x)/\gamma = 0.5$ . Thus, one finds a *trait threshold* ( $\mu_c^x/\gamma \simeq 0.372$ ) which precedes a *fitness threshold* ( $\mu_c/\gamma \simeq 0.408$ ), cf. Section 6.2.4. Symbols correspond to  $N = 100$ , dashed lines to  $N = 500$ , and solid lines to the expressions in Section 3. As  $\hat{r}$  behaves just like  $r(\hat{x})$  with similar accuracy, it is not shown here.



**FIG. 15.** Means (middle) and variances (right) for a model with *unidirectional mutation* ( $\kappa = 1$ ),  $N = 20$  (symbols), and the fitness function  $r/\gamma$  shown on the left. The means  $\bar{r}$  and  $\hat{x}$  were calculated via the discrete maximum principle (43). For  $\bar{x}$  and the variances the population distribution was calculated explicitly using the recursion following from (2) for  $U_k^- \equiv 0$ ; solid lines refer to the expressions from Section 3. One observes both a wildtype and a degradation threshold. As  $\hat{r}$  is exactly  $r(\hat{x})$ , it is not shown here.

We *always* find a wildtype threshold with loss of the fittest class if the total range of fitness values is *finite*. If there are lethal genotypes ( $r_k = -\infty$ ), we obtain *no* such threshold if (and only if) the mutation rate at all non-lethal types is larger or equal to the mutation rate at the wildtype. For the special case of a constant mutation rate ( $u_k^+ = \text{const}$ ,  $k < N$ ,  $u_N^+ = 0$ ), this reproduces the criterion by Wagner and Krall (1993). A degradation threshold is found if and only if there are no lethals.

For the special case of the biallelic model with linearly decreasing mutation rates, Eq. (65) reduces to  $\max_k s_k^+ / s_{k+1}^+ \geq 1$  and we obtain a threshold for any degree of positive epistasis, but also for linear parts of the fitness function (with any three fitness values on a straight line). For fitness functions of the form  $r(x) = r_0(1-x)^\alpha$ , finally, we can confirm the result by Wiehe (1997) that wildtype and degradation thresholds coincide if and only if  $\alpha \leq 0$  (no or negative epistasis). Note that the result by Wiehe (1997) was derived for a different mutation scheme (with mutations coupled to reproduction).

#### 6.4. Variation of Fitness Values and Sequence Lengths

Up to this point, we have discussed mutation thresholds as effects that may occur as the mutation rate varies, while the fitness function and the number of mutation classes are kept fixed (note that the mutation class limit is always understood as an approximation to a given finite system). Here are two alternative points of view.

Firstly, we can consider threshold effects as the *fitness values* vary, while the mutation rates remain constant. As already mentioned in the discussion of Haldane's principle (Section 5.2), mean fitness is largely independent of local variations in the fitness function, but only depends on the shape of  $r$  in regions with substantial weight in the ancestral distribution. For most values of the mutation rate, this has a unique peak, and therefore only the neighborhood of the mean ancestral mutational distance,  $\hat{x}$ , matters. At fitness thresholds, however, we find, in general, two peaks at which variations in  $r$  can change the mean fitness.

Secondly, in line with the original work of Eigen (1971), we can increase *sequence length* while leaving the mutation rate *per site* fixed, thus altering the *total* mutation rate. The question of interest then is: Given a certain (fixed) fitness advantage of some function, and fixed mutation rates per site in a molecular model, how long can the coding region for the function become and still be maintained intact by selection? In this case, with  $u^\pm/\gamma \sim N$  (where  $\gamma$  denotes the range of fitness values under consideration), we obtain thresholds which are inversely related to sequence length,  $\mu_c \sim 1/N$ , in all situations above. Note that this is in accordance with the original findings for the sharply peaked landscape (cf. Eigen and Biebricher, 1988), but at variance with results by Wiehe (1997). The latter are artificial effects caused by the use of a different scaling of the fitness functions, in which  $\gamma$  is not kept fixed, but increases with  $N$  in just those cases where conflicting results have been found.

### 6.5. Implications of Mutation Thresholds

At mutation thresholds, mutation–selection balance is unstable with respect to small changes in the model parameters. There is no real lower limit on the mutation rates at which these phenomena may happen, but for fitness and degradation thresholds mutation rates must be comparable to the average mutational effect  $\gamma$  to obtain effects of significant magnitude (cf. our examples in Figs. 9 and 11–15). In this case, the *average* effect of the mutations considered will be very slightly deleterious (or almost neutral) for realistic values of the mutation rate. The model then pays respect to the rationale that these mutations are the relevant ones for the discontinuous behavior. Since they may be numerous (cf., e.g., Kondrashov, 1995), their collective effect may nevertheless be quite large. Mutations with much stronger effects, on the other hand, will only occur at very low frequency in the population and contribute smooth changes to the system observables if the mutation rate is varied. They may therefore be excluded from these considerations.

An important consequence of the original error threshold of the sharply peaked landscape (and, more generally, of any degradation threshold in our typology), which has been stressed in particular by Eigen (1971) as well as by Maynard Smith and Szathmáry (1995, Chapter 4.3), is its potential importance for the evolution of mutation rates. Since the total mutation rate increases with the sequence length (see the previous subsection), site mutation rates must evolve below the threshold value to allow functions to prevail that need a certain minimum length of the coding region as their

genetic basis. This might have been a severe problem for early replicators since the mutational repair mechanisms required to reduce the mutation rate depend on enzymes with relatively large coding regions. Since we find degradation thresholds for a rather broad class of fitness functions, this is also a plausible hypothesis with respect to our more general model class.<sup>11</sup>

A closer look at the effect of thresholds on the mutational loss reveals yet another mechanism by which degradation thresholds, and fitness thresholds as well, may be important for the evolution of mutation rates, even if mutational repair itself is not the function endangered. Assume that the mutation rate may be reduced by modifications of the replication accuracy. Recall further that the mutational loss  $g = \hat{r} - \bar{r}$  provides a measure for the indirect fitness advantage  $\delta\bar{r}$  gained by the decrease of  $\mu$ . Therefore, a system beyond a degradation threshold (where  $g = 0$ ) will never experience any selection pressure for decreasing mutation rates, and thus cannot evolve in this direction. But even a fitness threshold (with a jump in  $g$ , but  $g > 0$  for  $\mu > \mu_c$ ) may have a similar effect. This is because modifiers for reduced mutation usually have deleterious physiological side effects, dubbed the “cost of fidelity” (see Sniegowski *et al.*, 2000, for a recent review). Clearly, for the modifier to prevail, the indirect fitness advantage  $\delta\bar{r}$  gained by the decrease of  $\mu$  must be at least as high. Therefore, a jump in  $g$  separates two different evolutionary regimes: for  $\mu < \mu_c$ , much larger costs can be counteracted than for  $\mu > \mu_c$ .

In a second line of interpretation, the critical mutation rate of an error threshold has often been argued to provide a strict upper limit that must be avoided in all real organisms. Certain kinds of viruses are perceived as thriving just below that value as to maximize their adaptability in a changing environment (Eigen and Biebricher, 1988). While it is certainly true that wildtype sequences or certain functions can get lost at threshold points, it is, however, much more difficult to argue why evolution should care about them. After all,  $g$  drops at the threshold, thus making a further increase in adaptability less costly. Further, the equilibrium mean fitness changes continuously with the mutation rate in arbitrary deterministic mutation–selection systems, even at threshold points. Mutation thresholds, therefore, cannot be seen as strict limits constraining the evolution of mutation rates. This may be different if further evolutionary forces are relevant, most importantly drift.

<sup>11</sup>Note that the fitness effect of mutational repair is always an indirect one caused by an increase in the copying accuracy in parts of the sequence that are directly related to fitness.

Indeed, numerical studies show that the mean fitness (averaged over time) may drop discontinuously at critical mutation rates in a finite population (Nowak and Schuster, 1989). A jump in the mean fitness has also been found for sexually reproducing populations with dominance (Higgs, 1994). This, however, is outside our model class and, according to our definition, no longer a property of a *mutation* threshold but essentially a drift (or segregation) phenomenon.

Let us finally turn to yet another effect that has previously been described as characteristic of the error threshold (e.g., Bonhoeffer and Stadler, 1993). Assume that mutation classes increase in size with the distance from the wildtype (as in the case for the biallelic model for  $k$  up to  $N/2$ ), which is reflected by asymmetric mutation rates between neighboring classes. Then, a jump in  $\bar{x}$ , as at the critical mutation rate  $\mu_c^*$  of a trait threshold, entails a *delocalization effect*. It should be stressed, however, that this effect has *no* direct consequences for the evolution of mutation rates, which are entirely connected to the population mean fitness and thus only to fitness, wildtype, and degradation thresholds.

## 7. SUMMARY AND OUTLOOK

The findings of this article, and the future directions they might lead to, fall into three parts, which we would like to discuss in turn.

*Ancestors:* As a crucial concept for the study of (asexual) mutation–selection models, we have identified the ancestor distribution of genotypes, or genotype classes, which, in mutation–selection balance, is the equilibrium distribution of the time-reversed evolution process. The ancestor frequency of the  $i$ th genotype (or class) is given as  $a_i = z_i p_i$ , where  $z_i$ , the relative reproductive success, and  $p_i$ , the equilibrium frequency, are the  $i$ th entries of the left and right leading (PF) eigenvectors of the evolution matrix. In the biology–physics analogy laid down in Appendix A, the ancestor distribution corresponds to the distribution of the bulk magnetization in spin models. Biologically, measurements of ancestor frequencies in real population should in principle be possible by marker techniques. In the equilibrium dynamics, the ancestors permanently feed the swarm of mutants that is observed at any instant of time. Significant evolutionary change is indicated by modification of this ancestor population. We have shown this in a couple of instances.

If the fitness values  $R_i$  are subject to change, the  $a_i$  measure the sensitivity of the equilibrium mean fitness

$\bar{R}$  to these changes. The net total change in  $\bar{R}$  is given (to linear order) by Eq. (25). If the fitness changes are due to a modifier mutation, Eq. (55) can be read as the selection coefficient of this modification with respect to the ancestors. Such a modifier will asymptotically fix if and only if it increases the fitness of the ancestors. The vector of ancestor frequencies can therefore be seen as the gradient which points into the direction of the effective selection pressure on the fitness function and determines the course of evolution—given that the appropriate modifier mutations are available.

Since the  $a_i$  are non-negative, selection will always favor an increase of fitness values. In the case of modifiers that change the mutant fitness values, we thus find a tendency for the evolution of robustness, or canalization, in systems with back mutations, whereas antirobustness, or decanalization, cannot result in this simple setup. As the selection pressure is strongly differential, one can even speculate that this mechanism is a cause for negative (synergistic) epistasis (as in the example in Fig. 3), which is considered a rather general phenomenon by many (Crow and Simmons, 1983; Phillips et al., 2000, and references therein). As always with indirect selection, however, selective forces are weak and probably of relevance only in large populations and for rather high mutation rates. In the limiting case of unidirectional mutation, the ancestor distribution is concentrated at the wildtype (if present in the equilibrium population). Then, only modifiers that increase the wildtype fitness will go to fixation, whereas modifications of mutant fitness values have no effect on the equilibrium mean fitness—in line with the predictions of the Haldane–Muller principle.

We have defined the *mutational loss*  $G$  as the difference between ancestor and population mean fitness, which equals the long-term loss in progeny that the equilibrium system suffers due to mutation. Eq. (24) shows that the loss determines the change in the equilibrium mean fitness if the mutation rate is subject to change. Again, it is thus the ancestor distribution that provides the link between external variations of model parameters and the equilibrium response. We always have  $G \leq L$ , with  $L$  the mutation load, and equality only in systems without back mutation to the fittest type. Measurements of the mutational loss should be possible by fitness measurements in mutator strains or by direct determination of the ancestor fitness distribution using genetic markers.

The ancestor concept, as introduced in this article, is independent of modeling assumptions on fitness landscapes and mutation schemes. We have derived a few basic results that hold for this general case, and extend

the Haldane–Muller principle. Under additional assumptions much stronger results may be obtained, as we have seen for the single-step mutation model. We expect that, of the many successful approximation methods that are routinely applied to the population distribution, some could also be applicable for the ancestor distribution and yield further interesting results. However, in order to apply this approach to more general situations, namely including genetic drift, the concept will have to be extended. The question is whether it is possible to characterize the distribution of genotypes on a single lineage backward in time, and to relate this to the mutation–selection–drift equilibrium.

*The maximum principle:* The reformulation of the equilibrium condition in terms of ancestor variables leads to a maximum principle for the equilibrium mean fitness, which we have exploited for the single-step mutation model. In this model, fitness is an arbitrary function of the number of mutations (or some other additive trait). Mutation proceeds stepwise on the mutation classes, but mutation rates (as well as back mutation rates) may vary from class to class. Here, the maximum principle may be recast into a particularly simple form, which yields the mean fitness as the maximum of the difference between the fitness function and the mutational loss function (see Eqs. (30) and (31)). The position of the maximum determines the mean ancestral genotype and the corresponding value of the mutational loss function yields the mutational loss  $G$  (Eq. (33)). The simplicity of the maximum principle results from the fact that maximization is over one single scalar variable only, and may be performed explicitly, or with the help of a simple graphical construction (Fig. 4). A different maximum principle has been suggested previously for mutation–selection models (Demetrius, 1983). It relies on general variational principles in the framework of ergodic theory, in which maximization is over all possible genealogies, and therefore not constructive.

Our maximum principle is exact in three independent limiting cases, namely unidirectional mutation, models with a linear dependence of both mutation rates and fitness on an underlying trait (including multilocus wildtype-mutant models without epistasis), and in the limit of an infinite number of mutation classes. For small back mutation rates,  $u^- \ll (u^+ + s)$ , the resulting estimate for the equilibrium mean fitness is exact to linear order in  $u^-$ . In general, the maximum principle holds as an approximation that leads to quantitatively reasonable results for a wide range of parameters and quickly becomes accurate if one of the exact limits is approached.

Starting from the mean fitness, we have explicitly calculated the fitness variance and the mean and variance of the trait. All formulas are collected in Section 3. The fitness variance is both proportional to the mean mutational effect and the mean difference of deleterious and back mutation rates; the trait variance has the same dependence on the mutation rates, but is inversely proportional to the mean mutational effect (Eq. (35)). These formulas give the amount of genetic variability that is maintained by the balance between mutation and selection.

Extensions of the maximum principle to a larger model class is possible in various ways. Following the lines of this paper, it is relatively straightforward to include double or multiple mutations in the theory. Poisson-distributed mutations (which emerge naturally in the biallelic model if mutation is coupled to reproduction) can also be treated. A necessary ingredient is that the evolution matrix can still be symmetrized by transformation to the ancestor frequencies.

The models discussed here all assume fitness to depend only on the distance to a reference class (the Hamming distance to the reference type in the biallelic case). Especially in a molecular context, this is, of course, a severe oversimplification. But also in classical population genetics, the importance of variance of additive and epistatic effects has often been highlighted (see, e.g., Bürger and Gimelfarb, 1999; Phillips *et al.*, 2000). Progress in this direction can be made by applying methods of inhomogeneous mean-field theory from statistical physics to the biallelic model. Here, it is possible to derive a simple maximum principle for models in which groups of sites or loci have different weights assigned that scale their respective direct and epistatic fitness effects (H.W., unpublished results). With similar techniques, fitness landscapes with more than one trait, such as the multiple quantitative trait model (Taylor and Higgs, 2000), can also be treated. Here, the equilibrium mean fitness is derived from a maximum principle over an  $n$ -dimensional space, if  $n$  is the number of traits. Finally, multilocus models with more than two alleles per locus (or states per site) may be considered. In the molecular context, an explicit treatment of the four-letter case with Kimura 3ST mutation scheme (cf. Swofford *et al.*, 1995) has already been given by Hermisson *et al.* (2001).

*Mutation thresholds:* Inspired by the definition of phase transitions in statistical physics, we have used the concept of the mutation class limit to define threshold behavior in mutation–selection models as the discontinuous change of statistical observables (such as the mean

fitness or the mean number of mutations) with the mutation rate  $\mu$ . Four different types of thresholds have been singled out, which all coincide in Eigen's original *error threshold* for the model with the sharply peaked landscape, but should be distinguished for general fitness functions. With the help of the maximum principle, criteria have been given to characterize the fitness functions and mutation schemes that lead to each type of threshold.

*Fitness thresholds* are characterized by a kink in the population mean fitness and a jump in the mutational loss  $G$ . They precisely occur at mutation rates for which the equilibrium ancestor distribution that solves the maximum principle is non-unique in the mutation class limit. The evolutionary significance of a fitness threshold lies in its potential impact on the evolution of mutation rates. Since the mutational loss jumps and may take much smaller values for  $\mu$  exceeding the critical mutation rate, the gain in mean fitness by reduction of  $\mu$  may be very small until the threshold is reached. If this gain in fitness must (over)compensate costs connected with mutational repair, evolution for lower mutation rates might be slowed down in the presence of a threshold. For the existence of fitness thresholds, positive (antagonistic) epistasis is required for many mutation schemes. Small convex parts of the fitness function, however, may already be sufficient. Fitness thresholds are collective phenomena and correspond to phase transitions in physics.

Whereas the loss of the wildtype from the population is not a well-defined notion for most of the models treated here, we consider the ancestor mean in the mutation class limit instead. A *wildtype threshold* is then characterized by a critical mutation rate  $\mu_c^- > 0$  below which the ancestor mean fitness coincides with the wildtype fitness, and the ancestor distribution is concentrated at the wildtype. Below a wildtype threshold, the system behaves, in many respects, as a system with unidirectional mutation. For the biallelic model, wildtype thresholds occur only for fitness functions with very sharp peaks at the wildtype position.

A *degradation threshold* is characterized by the fact that selection altogether ceases to operate and the mean fitness does not change any further for mutation rates exceeding a critical value  $\mu_c^+$ . A necessary condition for a degradation threshold is that the fitness function does not diverge to minus infinity. This is reminiscent of a threshold criterion derived for a model with unidirectional mutation by Wagner and Krall (1993). Degradation thresholds have similar implications for the evolution of mutation rates as fitness thresholds.

A *trait threshold*, finally, is characterized by a jump in the trait or mean number of mutations  $\bar{X}$ . In the sequence space picture, a trait threshold is connected with (partial) delocalization of the equilibrium population in genotype space. It is important to note that a trait threshold is not a collective phenomenon but is simply caused by non-monotonic parts of the fitness function. The delocalization effect is not connected with any significant change in the mean fitness (unless the trait threshold goes together with a fitness threshold), and thus has no direct impact on the selection pressure on the mutation rate.

The types of thresholds found here should also be observable in mutation–selection models with more general fitness landscapes and mutation schemes. Explicit threshold criteria can be obtained at least in some cases, such as the four-state model treated by Hermisson *et al.* (2001) (J.H., unpublished result).

## APPENDIX A: THE CONNECTION TO PHYSICS

For a number of models from statistical physics, a relation to mutation–selection models has been demonstrated, see Baake and Gabriel (2000) for an overview. In the present investigation, too, concepts and techniques from theoretical physics have served as a guideline for the analysis. Most importantly, the maximum principle (30) derives from the physical principle by which a system seeks to minimize its free energy. In our definitions of mutation thresholds, we also exploited the correspondence between thresholds and physical phase transitions, which has been first pointed out by Leuthäusser (1987).

Whereas such correspondences can be very fruitful, they require a detour through the physical world, which remains unsatisfactory from the biological point of view. Therefore, our intention in the body of the article has been to develop and discuss concepts entirely within the biological framework. Nevertheless, for readers with a physical background, as well as for biologists who are familiar with the interface to statistical mechanics, we will briefly sketch the relationship between both approaches. This may, on the one hand, facilitate further transfer of methods; on the other hand, limitations of certain “imported” concepts in the biological context become obvious. Last but not least, it is exactly this connection which resolves a few issues that had remained enigmatic so far.

Concentrating on the biallelic model with types  $s$  in this appendix, we can rely on a connection to a model of quantum statistical mechanics that was previously established by Baake *et al.* (1997) (see also Wagner *et al.*, 1998). More precisely, the evolution operator of the biallelic model with symmetric mutation was shown to be exactly equivalent to the Hamilton operator of an Ising quantum chain (up to a minus sign). Generalizing this slightly to include asymmetric mutation rates, and assuming a suitable ordering of genotypes, we may represent the quantum chain Hamiltonian as

$$\mathbf{H} = \mu \left[ \sum_n (\sigma_n^x - \mathbf{I}) - \kappa \sum_n (i\sigma_n^y + \sigma_n^z) \right] + \sum_I \eta_I \prod_{n \in I} \sigma_n^z = \mathbf{M} + \mathbf{R}. \quad (66)$$

Here,

$$\sigma_n^a := \underbrace{\mathbf{I} \otimes \cdots \otimes \mathbf{I}}_{n-1 \text{ copies}} \otimes \sigma_n^a \otimes \underbrace{\mathbf{I} \otimes \cdots \otimes \mathbf{I}}_{N-n \text{ copies}} \quad (67)$$

where  $a$  equals  $x$ ,  $y$  or  $z$ , and  $\sigma^{x,y,z}$  are *Pauli's matrices*, i.e.,

$$\sigma^x := \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad \sigma^y := \begin{pmatrix} 0 & -i \\ i & 0 \end{pmatrix} \quad \text{and} \\ \sigma^z := \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}. \quad (68)$$

The last sum in (66) runs over all index sets  $I \subset \{1, \dots, N\}$ , and  $\eta_I$  are the interaction coefficients of the spins within the chain, or with a longitudinal field. The collection of the  $\eta_I$  determines the fitness function. Further, there are transversal fields in  $x$  and  $y$  directions that account for mutation. Note that the Hamiltonian is non-Hermitian for asymmetric mutation.

This equivalence was used by Wagner *et al.* (1998) and Baake and Wagner (2001) to solve the model for a couple of fitness functions and symmetric mutation ( $\kappa = 0$ ) with the help of methods from quantum statistical mechanics. In the current investigation, we have chosen an equivalent formulation, which remains entirely within classical probability, to analyze more general mutation and fitness schemes. In order to briefly sketch the connection between the approaches, we first symmetrize  $\mathbf{H}$  by means of a similarity transform, i.e.,  $\tilde{\mathbf{H}} = \mathbf{S}\mathbf{H}\mathbf{S}^{-1}$  with  $\mathbf{S} := \prod_{n=1}^N (\cosh(\theta/2)\mathbf{I} + \sinh(\theta/2)\sigma_n^z)$  and  $\theta = \text{artanh}(\kappa)$ . (Note that this transformation relies on the sequence space representation of  $\mathbf{H}$  (66), in contrast to the symmetrization in Section 2.2, which starts out from a mutation class representation.)

The central concept now required is the *quantum mechanical expectation*  $\langle \mathbf{O} \rangle$  of an operator  $\mathbf{O}$ , defined by

$$\langle \mathbf{O} \rangle(t) := \text{tr}(\mathbf{O}\boldsymbol{\rho}(t)), \quad (69)$$

where  $\boldsymbol{\rho}(t)$  is the so-called *density operator*

$$\boldsymbol{\rho}(t) := \exp(t\tilde{\mathbf{H}}) / \text{tr}(\exp(t\tilde{\mathbf{H}})) \quad (70)$$

and  $t$  corresponds to the inverse temperature. For the choice  $\mathbf{O} := (1/N) \sum_{n=1}^N \sigma_n^z$ , one obtains the *quantum mechanical magnetization*, which is the crucial *order parameter* for the quantum chain.

We will concentrate on the limit  $t \rightarrow \infty$  (the ground state), where  $\boldsymbol{\rho}(t)$  becomes identical with the time evolution operator of the critical branching process we have met in Section 2.2. That is,  $\boldsymbol{\rho} = \lim_{t \rightarrow \infty} \boldsymbol{\rho}(t) = \lim_{t \rightarrow \infty} \exp(t(\tilde{\mathbf{H}} - \lambda_{\max}\mathbf{I})) = \tilde{\mathbf{p}}\tilde{\mathbf{p}}^T / \langle \tilde{\mathbf{p}}, \tilde{\mathbf{p}} \rangle$ , where  $\langle \cdot, \cdot \rangle$  denotes the scalar product and  $T$  means transposition. In this limit, the quantum mechanical expectation of any diagonal operator  $\mathbf{O}$  (defined by the elements  $O_{ss}$ ) therefore turns out to coincide with the corresponding *ancestral average* (cf. Section 2.4):

$$\begin{aligned} \langle \mathbf{O} \rangle &= \text{tr}(\mathbf{O}\boldsymbol{\rho}) = \frac{\langle \tilde{\mathbf{p}}, \mathbf{O}\tilde{\mathbf{p}} \rangle}{\langle \tilde{\mathbf{p}}, \tilde{\mathbf{p}} \rangle} = \sum_s O_{ss} \frac{\tilde{p}_s^2}{\sum_{s'} \tilde{p}_{s'}^2} \\ &= \sum_s O_{ss} a_s = \hat{\mathbf{O}}, \end{aligned} \quad (71)$$

where we have used that  $a_s = \tilde{p}_s^2 / \sum_{s'} \tilde{p}_{s'}^2$ , for symmetric  $\tilde{\mathbf{H}}$ , in line with Section 2.2. In particular, the quantum mechanical magnetization (given by  $O_{ss} = (N - 2d_{\text{H}}(s, s^+)/N)$ ), which has, so far, appeared as a crucial but technical quantity unrelated to any biological observable, now emerges as the mean ancestral genotype  $\hat{x}$  (up to a factor and an additive constant). In contrast, the classical magnetization  $\sum_s O_{ss} p_s$ , which we had termed surplus previously, translates into the population average  $\bar{x}$ . Let us note in passing that the change in normalization performed in Baake *et al.* (1998, Eqs. (7), (11)) and Baake and Wagner (2001, Eqs. (55), (56)) to formulate Rayleigh's principle for the PF eigenvalue (i.e.,  $\lambda_{\max} = \sup \langle \mathbf{x}, \tilde{\mathbf{H}}\mathbf{x} \rangle$  where the supremum is taken over all  $\mathbf{x}$  with  $\langle \mathbf{x}, \mathbf{x} \rangle = \|\mathbf{x}\|_2 = 1$ ) is equivalent to our ancestral transformation in (16). This way, we may take advantage of  $L^2$  theory although the original problem is inherently in the realm of  $L^1$ . Finally, the expectation of the *non-diagonal* operator  $\mathbf{M}$  is  $\langle \mathbf{M} \rangle = \sum_{s,s'} \tilde{p}_s \tilde{\mathbf{M}}_{s,s'} \tilde{p}_{s'} = \sum_{s,s'} z_s \tilde{\mathbf{M}}_{s,s'} p_{s'}$  (with  $\tilde{\mathbf{M}} := \mathbf{S}\mathbf{M}\mathbf{S}^{-1}$ ), which we have identified with the loss  $G$  in offspring due to mutation (cf. Section 5).

The concept of ancestral distributions is very general and does not rely on our special dynamical system. It also applies to discrete dynamical systems, as long as

they are linear (or may be transformed to a linear system) and admit a unique stable stationary state. This is true if a system is defined by an iteration matrix  $\mathbf{T}$  for which the Perron–Frobenius theorem holds (hints in this direction may be found in Demetrius, 1983). In particular, this applies to a discrete-time version of the quasi-species model defined by  $T_{ss'} = v_{ss'} w_{s'}$ , where  $v_{ss'}$  is the mutation probability from  $s'$  to  $s$ , and  $w_{s'}$  is Wrightian fitness of  $s'$ . As observed by Leuthäusser (1986, 1987) and reviewed by Baake and Wagner (2001, Appendix II), this model is equivalent to a classical two-dimensional Ising model with row transfer matrix  $\mathbf{T}$ , whereas the rows correspond to genotypes, and the columns to generations. Hence, every 2D configuration corresponds to one line of descent, *conditional on non-extinction at present*.

Here, considerable confusion has arisen in the literature as to the distinction and meaning of *surface* and *bulk* magnetizations (Leuthäusser, 1987; Tarazona, 1992; Franz and Peliti, 1997). *Surface* quantities correspond to the last row (in the time direction) of a configuration with open boundary conditions, i.e., the current population; therefore, surface averages are population averages. In contrast, *bulk* quantities are averages over the entire 2D configuration. In the limit of an infinite number of rows, they become identical with averages over a single row “in the middle” of the configuration (i.e., at infinite distance from both the first and the last row), as given by  $\lim_{n \rightarrow \infty} \text{tr}(\mathbf{T}^n \mathbf{O} \mathbf{T}^n) / \text{tr}(\mathbf{T}^{2n})$ . Therefore, the bulk average is, again, the ancestral average (also compare with (69) and (71)).

Everything we have said so far holds for arbitrary, finite  $N$ . Clearly, the infinite mutation class limit  $N \rightarrow \infty$  is the thermodynamic limit of the statistical mechanics system with its extensive scaling of energy and magnetic field terms. Technical aspects related to this scaling in the biological system are covered by Baake and Wagner (2001). While the thermodynamic limit may be taken as a matter of course in most classical situations in solid-state physics, the adequacy of the corresponding limit as an approximation in biological applications must be thoroughly considered. In particular, the mutation class limit should be clearly distinguished from the infinite-sites limit, which is widely used in theoretical population genetics; see the discussion in Section 2.7, and Baake and Wagner (2001).

Clearly, the fitness thresholds described in Section 6 correspond to the phase transitions of the physical system, in the sense of a non-analytic point of the free energy of the classical Ising system or the ground state energy of the quantum chain (the mean fitness in the biological model). Most importantly, the idea to use the

thermodynamic limit for the mathematical definition of the concept is taken from physics. As we have pointed out (Section 6.1), this is in accordance with the original definition of the error threshold for the sharply peaked landscape. It should be noted, however, that the fitness functions of the biological system typically lack the symmetries inherent in physics. As a consequence, the usual classification of phase transitions in physics according to orders of the non-analyticity as well as the consideration of critical exponents does not seem to be particularly meaningful in the biological context. Fitness thresholds are typically first order and exhibit a jump in the ancestral mean  $\hat{x}$ , which parallels the physical magnetization. Note at this point that neither the population mean  $\bar{x}$  (as suggested by Tarazona, 1992; Franz and Peliti, 1997) nor the mean fitness itself (as implicitly in Higgs, 1994) should be mistaken as an order parameter, in the sense that jumps in these quantities do not characterize first-order phase transitions.

## APPENDIX B: PROOFS FROM SECTION 4

### B.1. The Additive Case

Let us prove here that, if fitness and mutation rates depend linearly on some trait  $y_k = y(x_k)$  as described in (29), the system (39) reduces to just two equations, one corresponding to the necessary extremum condition following from (30), the other being the defining Eq. (33) for the ancestral mean  $\hat{y}$  (for  $y(x) = x$ ). For the sake of simplicity, we write  $x_k$  instead of  $y_k$  here. Taking the difference of two arbitrary equations of the linear system (39), say for  $k$  and  $\ell$ , divided by  $\sqrt{a_k}$  and  $\sqrt{a_\ell}$ , respectively, we get

$$\begin{aligned} & (\beta^+ - \beta^- - \alpha)(x_k - x_\ell) \\ & + \sqrt{\beta^+ \beta^-} \left( \sqrt{x_k(1-x_{k-1})} \sqrt{\frac{a_{k-1}}{a_k}} \right. \\ & - \sqrt{x_\ell(1-x_{\ell-1})} \sqrt{\frac{a_{\ell-1}}{a_\ell}} + \sqrt{x_{k+1}(1-x_k)} \sqrt{\frac{a_{k+1}}{a_k}} \\ & \left. - \sqrt{x_{\ell+1}(1-x_\ell)} \sqrt{\frac{a_{\ell+1}}{a_\ell}} \right) = 0. \end{aligned} \quad (72)$$

With the *ansatz*

$$\frac{a_{k-1}}{a_k} = C \frac{x_k}{1-x_{k-1}} \Leftrightarrow \frac{a_{k+1}}{a_k} = C^{-1} \frac{1-x_k}{x_{k+1}} \quad (73)$$

Equation (72) can be divided by  $(x_k - x_\ell)$  and becomes independent of  $k$  and  $\ell$ . Note that (73) also takes care of the boundary conditions  $a_{-1} = a_{N+1} = 0$  if  $x_0 = 0$  and  $x_N = 1$ . Summing both sides of  $(1 - x_{k-1}) \times a_{k-1} = Cx_k a_k$  over  $k$ , we obtain  $C = (1 - \hat{x})/\hat{x}$  and thus from (72)

$$\beta^+ - \beta^- - \alpha + \sqrt{\beta^+ \beta^-} \frac{1 - 2\hat{x}}{\sqrt{\hat{x}(1 - \hat{x})}} = 0, \quad (74)$$

which is exactly the extremum condition  $r'(\hat{x}) = g'(\hat{x})$  following from (30). Together with the negative second derivative this implies the maximum principle.

On the other hand, we can use (73) to eliminate  $a_{k\pm 1}$  from (39). After multiplication by  $\sqrt{a_k}$  this reads

$$\left[ r_0 - \alpha x_k - \bar{r} - \beta^+(1 - x_k) - \beta^- x_k + \sqrt{\beta^+ \beta^-} \left( x_k \sqrt{\frac{1 - \hat{x}}{\hat{x}}} + (1 - x_k) \sqrt{\frac{\hat{x}}{1 - \hat{x}}} \right) \right] a_k = 0 \quad (75)$$

and we obtain, by summation over  $k$ ,

$$\begin{aligned} \bar{r} &= r_0 - \alpha \hat{x} - \beta^+(1 - \hat{x}) - \beta^- \hat{x} + 2\sqrt{\beta^+ \beta^-} \hat{x}(1 - \hat{x}) \\ &= r(\hat{x}) - g(\hat{x}), \end{aligned} \quad (76)$$

which corresponds to (33). Since fitness is assumed linear in the trait, the mean values with respect to the population distribution are also related via  $\bar{r} = r(\bar{x})$ .

## B.2. The Case $N \rightarrow \infty$

Let  $u^\pm : [0, 1] \rightarrow \mathbb{R}_{\geq 0}$  be continuous and positive, but fulfill  $u^+(1) = u^-(0) = 0$ . Let  $r : [0, 1] \rightarrow \mathbb{R}$  have at most finitely many discontinuities, being either left or right continuous at each discontinuity in  $]0, 1[$ . Then, with the scaling described at the end of Section 2.6, the maximum principle (30) holds in the limit  $N \rightarrow \infty$ .

For a proof, we follow the arguments and notation introduced in Section 4.3. First note that the lower bound for  $\bar{r}_N$  in (46) is itself greater than or equal to

$$\begin{aligned} \rho_{k,m,n} &:= \inf_{y \in I_{k,m,n}} (r(y) - g(y)) - \sup_{y \in I_{k,m,n}} |g(y) - g_N(y)| \\ &= \frac{\sqrt{u_{k-m-1}^+ u_{k-m}^-} + \sqrt{u_{k+n}^+ u_{k+n+1}^-}}{m+n+1}, \end{aligned} \quad (77)$$

where  $I_{k,m,n} = [(k-m)/N, (k+n)/N]$  and the rules for inf/sup have been applied. We will now construct a sequence  $\rho_N(x) := \rho_{k_N(x), m_N(x), n_N(x)}$  for each  $x \in [0, 1]$ , using suitable sequences for the indices, such that

$$\rho_N(x) \rightarrow r(x) - g(x). \quad (78)$$

Since, by definition,  $\lim_{N \rightarrow \infty} \bar{r}_N = \bar{r}_\infty$ , Eqs. (46), (77), and (78) will then establish  $\bar{r}_\infty \geq \sup_{x \in [0,1]} (r(x) - g(x))$ ,

from which, together with the upper bound in (48), the claim will follow.

Note first that, for  $x = 0$  or  $x = 1$ ,  $\rho_N(x) = \rho_{xN,0,0} = r(x) - g(x)$  holds for arbitrary  $N$ . Now, fix  $x \in ]0, 1[$ . If  $r$  is continuous in  $[x-d, x]$  for a suitable  $d > 0$ , let  $k_N(x) := \lfloor xN \rfloor$ ,  $m_N(x) = \lfloor d\sqrt{N} \rfloor$ , and  $n_N(x) \equiv 0$ . Otherwise  $r$  is continuous in  $[x, x+d]$  for some  $d > 0$ , and we define  $k_N(x) := \lfloor xN \rfloor$ ,  $m_N(x) \equiv 0$ , and  $n_N(x) = \lfloor d\sqrt{N} \rfloor$ . With these choices, the last term in (77) vanishes for  $N \rightarrow \infty$  since  $m_N(x) + n_N(x) \rightarrow \infty$ , and the enumerator is bounded. So does the supremum term because of the uniform convergence  $g_N \rightarrow g$ :  $\sup_{y \in I_{k_N, m_N, n_N}} |g(x) - g_N(x)| \leq \sup_{y \in [0,1]} |g(x) - g_N(x)| \rightarrow 0$ . The latter follows from the uniform continuity of  $\sqrt{u^\pm}$  since, in

$$\begin{aligned} &|g(x) - g_N(x)| \\ &= \left| \left( \sqrt{u^+ \left( x - \frac{1}{N} \right)} - \sqrt{u^+(x)} \right) \sqrt{u^-(x)} \right. \\ &\quad \left. + \sqrt{u^+(x)} \left( \sqrt{u^- \left( x + \frac{1}{N} \right)} - \sqrt{u^-(x)} \right) \right|, \end{aligned} \quad (79)$$

the terms in parentheses vanish uniformly in  $x$  as  $N \rightarrow \infty$  and  $\sqrt{u^\pm(x)}$  is bounded. Finally, the infimum term in (77), and thus  $\rho_N(x)$ , converges to  $r(x) - g(x)$  since  $x_{k_N(x)} \rightarrow x$ ,  $r$  is continuous in all  $I_N \ni x$ , and  $|I_N| = (m_N(x) + n_N(x))/N \rightarrow 0$ . This was to be shown.

Now, let us prove that the ancestor distribution is concentrated around those  $x$  for which  $r(x) - g(x)$  is maximal, from which Eq. (33) follows if the maximum is unique. Multiplying the evolution equation in ancestor form (39) by  $\sqrt{a_k}$ , we obtain, after summation over  $k$ :

$$\begin{aligned} \bar{r}_N &= \sum_{k=0}^N [(r(x_k) - u^+(x_k) - u^-(x_k))a_k \\ &\quad + \sqrt{u^+(x_{k-1})u^-(x_k)}\sqrt{a_k a_{k-1}} \\ &\quad + \sqrt{u^+(x_k)u^-(x_{k+1})}\sqrt{a_{k+1} a_k}]. \end{aligned} \quad (80)$$

Using  $\sqrt{a_k a_{k\pm 1}} \leq \frac{1}{2}(a_k + a_{k\pm 1})$ , we get

$$\bar{r}_N \leq \sum_{k=0}^N [r(x_k) - g_N(x_k)]a_k = \hat{r}_N - (\widehat{g}_N)_N \quad (81)$$

with  $g_N(x_k)$  as defined in Section 4.3. Since  $\bar{r}_N \rightarrow \bar{r}$  and  $g_N(x) \rightarrow g(x)$  uniformly in  $x \in [0, 1]$ , we can find for any given  $\varepsilon > 0$  some  $N_\varepsilon$ , such that for all  $N > N_\varepsilon$ :

$$\sum_{k=0}^N [r(x_k) - g(x_k)]a_k > \bar{r} - \varepsilon^2. \quad (82)$$

We now divide this sum into two parts,  $\sum_k := \sum_{k_\leq} + \sum_{k_\geq}$ . The first part,  $\sum_{k_\leq}$ , collects all  $k$  with  $r(x_k) - g(x_k) > \bar{r} - \varepsilon$ , the second part contains the rest.

We then obtain

$$\begin{aligned} \bar{r} - \varepsilon^2 &< \sum_{k=0}^N [r(x_k) - g(x_k)]a_k \\ &\leq \bar{r} \sum_{k>} a_k + (\bar{r} - \varepsilon) \sum_{k\leq} a_k = \bar{r} - \varepsilon \sum_{k\leq} a_k \end{aligned} \quad (83)$$

and thus  $\sum_{k\leq} a_k < \varepsilon$ . We conclude that for  $N$  sufficiently large, the ancestor distribution is concentrated in those mutation classes for which  $r(x) - g(x)$  is arbitrarily close to its maximum,  $\bar{r}$ .

## APPENDIX C: PROOFS FROM SECTION 6

### C.1. Proof of (58)

We first prove that the negation of (58),

$$r''(x) - \frac{r'(x)g''(x)}{g'(x)} < 0 \quad \forall x \in [x_{min}, x_{max}] \quad (84)$$

implies (57) and is therefore a sufficient condition for the absence of a fitness threshold. We start by showing that both  $r$  and  $g$  are strictly decreasing in  $]x_{min}, x_{max}[$ . To see this, suppose there exists an  $x > x_{min}$  with  $r'(x) = 0$ , and let  $x_r$  be the smallest such  $x$ . Then either  $g'(1, x_r) = 0$  and  $\lim_{x \rightarrow x_r} (r''(x) - \frac{r'(x)g''(x)}{g'(x)}) = r''(x_r) - r''(x_r) = 0$  in contradiction to (84), or  $g'(1, x_r) \neq 0$ , in which case we obtain  $r''(x_r) < 0$  in contradiction to  $r'(x) < 0$  for  $x \in ]x_{min}, x_r[$ . On the other hand, imagine  $g'(x) = 0$  for some  $x \in ]x_{min}, x_{max}[$ , and let  $x_g$  be the largest such  $x$ . Then, since  $g'(x) < 0$  for  $x \in ]x_g, x_{max}[$ , we have  $g''(x_g) \leq 0$  and thus  $\lim_{x \rightarrow x_g} g''(x)/g'(x) = +\infty$  for the right-sided limit, which again contradicts (84) since  $r'(x_g) < 0$ . Therefore,  $\mu(x) := r'(x)/g'(1, x)$  is well defined everywhere in  $]x_{min}, x_{max}[$ , it guarantees  $r'(x) = g'(\mu(x), x)$ , and (84) yields  $r''(x) < g''(\mu(x), x)$ , which completes the first part of the proof.

We now prove that (58) implies a threshold. Assume first that the supremum in (58) is larger than zero. Due to the continuity of  $r$ ,  $g$ , and their derivatives, we then find an  $x_0$  in  $]x_{min}, x_{max}[$  with  $r''(x_0) - r'(x_0)g''(x_0)/g'(x_0) > 0$ . This, however, implies  $r''(x_0) - g''(x_0) > 0$  whenever  $r'(x_0) - g'(x_0) = 0$ . Therefore, the maximum of  $r(x) - g(x)$  is never attained at  $x_0$  and we must have a jump in  $\hat{x}(\mu)$ . If the supremum in (58) is exactly zero, we argue as follows. For the absence of a threshold, we need a continuous function  $\hat{x}(\mu)$  whose inverse, by the maximum principle, is  $\mu(\hat{x}) = r'(\hat{x})/g'(1, \hat{x}) > 0$  for  $\hat{x} \in ]x_{min}, x_{max}[$ . For the derivative of  $\mu(\hat{x})$ , we find

$\mu'(\hat{x}) = [r''(\hat{x}) - r'(\hat{x})g''(\hat{x})/g'(\hat{x})]/g'(1, \hat{x})$ , which must be non-negative in the absence of a threshold. Consider now those  $\hat{x}$  at which the supremum in (58) is attained. For  $g'(1, \hat{x}) \neq 0$ , we have  $\mu'(\hat{x}) = 0$ . Since  $\hat{x}'(\mu) = 1/\mu'(\hat{x})$ ,  $\hat{x}(\mu)$  has a diverging derivative at these points, and a jump if the supremum is attained (and thus  $\mu'(\hat{x}) = 0$ ) on a whole interval. Finally, we also obtain a jump if the supremum is attained on an interval where also  $g'(1, x) = 0$  as then the whole interval is degenerate as a maximum. We exclude the special case that  $g'(1, x) = 0$  at an isolated  $x$  to avoid lengthy technicalities.

### C.2. Proof of (61)

Note first that existence of a wildtype threshold obviously implies a lower bound of  $1/\mu_c^-$  on the left-hand side of (61). Assume, on the other hand, that there are sequences  $x_i$  in  $[x_{min}, x_{max}]$  and  $\mu_i > 0$  with  $\mu_i \rightarrow 0$  and  $r(x_i) - \mu_i g(1, x_i) > r(x_{min}) - \mu_i g(1, x_{min})$  for all  $i$ . Let then  $x_j \rightarrow x_\infty$  be a convergent subsequence. Since  $r$  and  $g$  are assumed to be continuous, we have  $r(x_\infty) \geq r(x_{min})$  and hence  $x_\infty = x_{min}$ , since  $r(x_{min})$  is the unique maximum of  $r$  in  $[x_{min}, x_{max}]$ . Thus, we find

$$\frac{g(1, x_j) - g(1, x_{min})}{r(x_j) - r(x_{min})} > \frac{1}{\mu_j} \rightarrow \infty, \quad (85)$$

contradicting (61) and proving the criterion.

### C.3. Proof of (63)

The proof is analogous to the case of the wildtype threshold. On the one hand, existence of the threshold implies the criterion with a bound  $\mu_c^+$ . On the other hand, if we have sequences  $x_i$  in  $[x_{min}, x_{max}]$  and  $\mu_i$  with  $r(x_i) - \mu_i g(1, x_i) > r(x_{max})$  for  $\mu_i \rightarrow \infty$ , we can again choose a convergent subsequence  $x_j \rightarrow x_\infty$ . Since  $g(1, x)$  is continuous and  $x_{max}$  is the only zero of  $g$  in  $[x_{min}, x_{max}]$ , we have  $x_\infty = x_{max}$ . As in the wildtype case above, this contradicts (63) and proves the criterion.

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