

Evolvability Suppression to Stabilize Far-Sighted Adaptations

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Abstract The opportunistic character of adaptation through natural selection can lead to *evolutionary pathologies*—situations in which traits evolve that promote the extinction of the population. Such pathologies include imprudent predation and other forms of habitat overexploitation, or the tragedy of the commons, adaptation to temporally unreliable resources, cheating and other antisocial behavior, infectious pathogen carrier states, parthenogenesis, and cancer, an intraorganismal evolutionary pathology. It is known that hierarchical population dynamics can protect a population from invasion by pathological genes. Can it also alter the genotype so as to prevent the generation of such genes in the first place, that is, suppress the evolvability of evolutionary pathologies? A model is constructed in which one locus controls the expression of the pathological trait, and a series of modifier loci exist that can prevent the expression of this trait. It is found that multiple *evolvability checkpoint* genes can evolve to prevent the generation of variants that cause evolutionary pathologies. The consequences of this finding are discussed.

Keywords

Evolvability, metapopulation, hierarchies, cheats, far-sighted

I Introduction

Adaptation through natural selection is an opportunistic process, in that it is driven by the selective forces of the immediate moment upon the individual organism. Yet traits that provide immediate advantage to the individual may be detrimental to the population or species, or detrimental over longer time scales. This has been understood since Darwin. Conversely, traits may impose an immediate disadvantage to the individual, yet be advantageous to the population or species, or advantageous over longer time scales. Whether or how such *far-sighted* traits can evolve has been a challenge for evolutionary mechanics.

Far-sighted traits include such phenomena as altruistic behavior—behavior that benefits not its carrier, but other individuals. Other phenomena include cooperation, social organization, prudent predation, population regulation, and multicellular organization.

The opposite of far-sighted traits could be called “short-sighted,” “cheating” [35], “selfish,” or “greedy.” Interestingly, in computer science, the term “greedy” is used to describe algorithms that are short-sighted in that they respond only to immediate conditions without regard to long-term consequences. It is well known that for certain problems, greedy algorithms can lead to outcomes that are ultimately suboptimal.

Examples of short-sighted adaptations include:

- cheating, defection, and other antisocial behavior,
- meiotic drive [33],

- parthenogenesis [19],
- overpopulation [67],
- imprudent predation [51] and other forms of habitat overexploitation—the *tragedy of the commons* [24],
- cannibalism [22],
- cancer (the organism being the population) [45, 54],
- adaptation to temporally unreliable resources [26],
- viable but infectious pathogen carrier states [29],
- evolution of endosymbionts to the detriment of the host [63].

An interesting example of how a straightforward trait such as longevity can become a short-sighted trait is explored by Kirchner and Roy [29] in the situation of chronic infection. They begin with the observation that longer life spans can result in larger reservoirs of persistently infected hosts, and these reservoirs can in turn reduce population size. While longer life spans provide a reproductive advantage to the individual, in this situation they impose a viability disadvantage to the population. Thus, under certain ecological circumstances, even longevity can be a short-sighted trait.

1.1 Dynamical Hierarchies

Far-sighted traits by their very definition pose mechanistic difficulties for their evolution, since they may impose a disadvantage on their carriers while offering long-term advantages to a population. The primary mechanistic solution to allow the evolution of far-sighted traits is a hierarchical structuring of the population—which includes group selection [67], kin selection [20], structured demes [66], reciprocal altruism [57], viscous populations [21, 58, 40, 41], multilevel selection [15], lineage selection [46], and metapopulation dynamics [30, 31, 37].

Hierarchical structures that allow the evolution of far-sighted traits include a broad spectrum of possibilities. They may be discrete, such as a colony, isolated deme, or multicellular organism; they may be ephemeral, as in *structured demes* [64]; or they may be continuous, such as viscous populations [21, 40, 41]. The requisite feature for all, however, is that the compositions of populations in different places be able to diverge from one another, so as to allow different fates to befall the different compositions.

1.2 Evolutionary Stability through Evolvability Suppression

The key theoretical concept to understanding when hierarchical dynamics can produce far-sighted traits is *evolutionary stability* [36, 12]. Evolutionary stability poses two complementary questions:

1. Will genetic variation that enhances a far-sighted trait survive and flourish when introduced into a population?
2. Will genetic variation for a short-sighted trait (e.g., cheater mutations) be driven to extinction when introduced into a population?

The existing theoretical solutions to the problem of far-sighted traits mainly take the evolutionary stability approach: conditions are found that prevent the invasion of short-sighted variants when they are introduced into a population, or prevent the extermination of far-sighted traits by short-sighted traits.

In the case when selection, genetic, and population parameters are in the right range, both conditions for the stability of far-sighted traits can be met. In the example above of longevity and parasite load, Kirchner and Roy [29] find parameters in a metapopulation model that satisfy each of

these evolutionary stability criteria, and prevent the evolution of longer life spans, reduce the parasite burden, and improve the population viability. Whether, or how often, these parameter conditions are met in nature continues to be a controversial issue [16].

The evolutionary stability approach to the evolution of far-sighted traits contains within it a hidden presupposition: namely, that mutations that enhance or break down far-sighted traits—that is, perturbations to the equilibria—occur with enough frequency to destabilize an unstable equilibrium. For example, the theory for the evolution of altruism assumes that cheater mutations will arise with enough frequency to make the stability of altruistic traits an issue. The ability of a genome to generate variants in the direction of a certain trait is referred to as that trait's *evolvability*. Therefore, the evolutionary stability approach to understanding the evolution of far-sighted traits assumes the evolvability of these traits.

The possibility has been raised; however, the evolutionary stability of far-sighted traits against invasion by short-sighted traits need not always be a problem, because the evolvability of phenotypes that break down far-sighted traits may not necessarily exist. Let us take this possibility to the next level: Suppose that evolution could find a way to *prevent the generation* of the short-sighted traits in the first place. This would provide another mechanism to stabilize far-sighted traits.

A number of studies in recent years have examined the question of how the evolvability of traits may itself evolve as a systematic outcome of evolutionary forces [3, 7, 44, 4–6, 61, 62, 8, 34, 14, 47, 56]. The focus of population genetics theory has traditionally been on the fate of genetic variation within populations, rather than on the processes that originate it. Levinton [32, p. 494] states this succinctly:

Evolutionary biologists have been mainly concerned with the *fate of variability* in populations, not the *generation of variability*. . . . This could stem from the dominance of population genetic thinking, or it may be due to a general ignorance of the mechanistic connections between the genes and the phenotype. Whatever the reason, the time has come to reemphasize the study of the origin of variation.

Suppose that the organism could be mutated so that it could no longer even generate the short-sighted traits. Would such mutations have a selective advantage? In other words, could genotypes with this suppressed evolvability of the short-sighted traits come to predominate in the population? Would it require hierarchical population dynamics? If so, this would be a novel, higher-order evolutionary phenomenon made possible by hierarchical population dynamics. The suppression of evolvability would constitute a second-order form of evolutionary stability.

Wynne-Edwards [68] was perhaps the first person to propose that the suppression of evolvability was a means to stabilize far-sighted traits. However, he did not propose any mechanism by which this could happen [46]. Such a mechanism can be conceived as follows [2]:

Suppose that there was some trait which would evolve by individual selection if variation for it existed, but which would increase the chance of the deme to go extinct (for example, the size variation in the smaller of the two competing species in Roughgarden's models [52]). Now, suppose the variation between populations was for the *rate of introduction* of this phenotypic variant. This could be due to different pleiotropy, different population structure, different genetic systems, or mutation rates. These populations would last longer. Also, this could lock out imprudent predation, and overuse of resources.

The idea is that far-sighted traits could be stabilized by genetic changes that prevent cheats from even being generated. The possibility that these evolvability-suppression mutations could be maintained by metapopulation dynamics [1, reprinted in Appendix] was prompted by a model presented in [48, 49] that showed that extinction and recolonization dynamics could maintain altruism. *Metapopulation dynamics* [30, 31] refers to fragmented populations in which frequent local extinction and recolonization of demes occurs.

Nunney [44] proposes the same mechanism to account for the maintenance of sex (a far-sighted trait) against invasion by asexual mutants (short-sighted traits). He analyzes a metapopulation model where the rate at which asexual mutations arise is subject to variation. In his model, the rate of mutation to viable asexual phenotypes evolves to low values. Nunney [45] also analyzes a model of evolvability suppression against carcinogenesis, and finds that multiple loci could evolve to suppress the evolution of cancers.

In this article, I wish to examine a concrete model for the evolution of evolvability suppressors that will allow for the evolution of multiple suppressor loci, modeling the mutation processes between all alleles, and including a spatial population structure to provide the hierarchical dynamics.

2 The Model

There has been debate about the definition of “far-sighted” and “short-sighted” traits, in particular, “weak” and “strong” altruism [18, 65]. Here I consider traits that are incontrovertibly short-sighted: traits that have a selective advantage but that nevertheless systematically bring about their own long-run extinction and the extinction of the population that carries them [23]. These would be the most extreme outcomes among the short-sighted traits listed above. When the process of survival of the fittest results instead in extinction, I wish to call such a paradoxical outcome an *evolutionary pathology*. The common property of evolutionary pathologies is that the trait under consideration gives its carriers a viability or reproductive advantage, and genetic variation for the trait tends to increase in frequency in the population; yet, once it becomes common, the trait has ecological or group properties that increase the rate of extinction of the population.

The basic question I want to answer here is, can the genome evolve so that it prevents pathological phenotypes from even being produced by mutation? There are two mechanisms by which mutations to the pathological phenotypes could be prevented. One is simply a lowered mutation rate for production of pathological alleles. The other is a requirement that multiple loci be mutated in order to express the pathological phenotype. There are many more possibilities that would involve the latter mechanism, so it is what I examine here. To investigate whether the evolvability of evolutionary pathologies can evolve, the following model is constructed. What the model wishes to test is whether the hierarchical population dynamics can cause the genotype to evolve to be multiple mutational steps away from the pathological trait.

The genotype of individuals consists of a primary locus and numerous modifier loci. In the primary locus, mutation of the wild type gives rise at low frequency to the pathological trait. Specifically, the case will be considered in which all but one of the alleles at the primary locus are wild type, and one is pathological, with uniform mutation between all alleles. A range of allele numbers are investigated, from 2 to 128, to see how the evolvability of the pathology allele puts pressure on the modifier loci to evolve checkpoint alleles.

At the modifier loci, the wild-type alleles are permissive of expression of the pathological trait at the primary locus. A rare gain-of-function mutation of a modifier allele will produce an allele that blocks the expression of the pathological trait. The presence of the gain-of-function allele at any modifier locus blocks the expression of the pathological trait.

Because of their analogy to the checkpoint genes in metazoan organisms that protect cells against evolution into cancer cells [42, 38, 25], I refer to these gain-of-function alleles as *checkpoint-positive* or CHKPT^+ , and to the wild type as *checkpoint-negative* or CHKPT^- . Specifically, the case will be considered in which 127 out of 128 alleles at the primary locus are wild-type CHKPT^- , and the other one has the gain-of-function CHKPT^+ , with uniform mutation between all alleles. Thus under mutational pressure alone, only a small number of these checkpoint alleles would ever be present in the population. If these alleles increase in frequency, under hierarchical population dynamics, to levels far from mutational equilibrium, it will demonstrate that checkpoints can evolve to block the introduction of evolutionary pathologies.

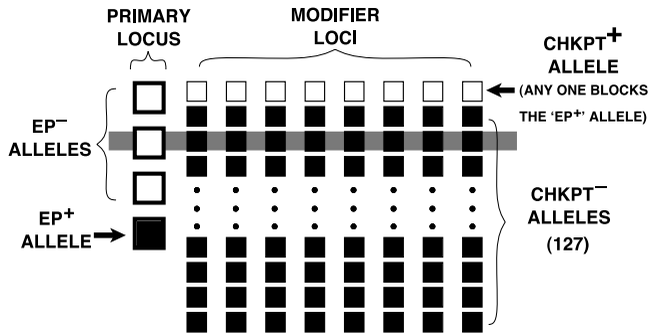


Figure 1. The genome in the evolvability suppression model. EP^+ is the gain-of-function *evolutionary pathology* allele, EP^- is the wild type, $CHKPT^+$ is the gain-of-function modifier allele that blocks expression of the pathological phenotype, and $CHKPT^-$ is the permissive, wild-type modifier allele.

The population is divided up into a lattice of demes. Migration rates between demes are assumed to be lower than the frequency of extinctions of demes, so that migration does not result in a mixing of demes, but in recolonization.

The genetic structure is illustrated in Figure 1. All genotypes are selectively neutral except the one that expresses the pathological trait, which has a selective advantage within the deme. The population structure consists of a 16-by-16 array of demes (with periodic boundaries).

The demes are filled initially with genotypes that have a nonpathology allele at the primary locus, and permissive $CHKPT^-$ alleles at the modifier loci. Other parameters of the model are given in Table 1.

The population reproduces in discrete generations. Organisms are asexual. The Wright-Fisher model of multinomial sampling is used for reproduction: 2000 offspring for each deme are sampled i.i.d. (independently, identically distributed) from the offspring genotype distribution of the parent population of the deme. The offspring genotype distribution is generated under the assumption that:

- mutation rates are equal for all loci;
- mutation rates are symmetric between all alleles; and
- each locus mutates independently of the others.

Table 1. Parameters of the model.

<ul style="list-style-type: none"> • 1 primary locus, 8 modifier loci • 2 to 128 alleles at the primary locus: 1 EP^+, 1 to 127 EP^- • 128 alleles per modifier locus: 1 $CHKPT^+$, 127 $CHKPT^-$ • Mutation rate 0.01 per replication per locus • 16×16 demes in the metapopulation • 2000 individuals per deme • Initial population: all individuals have the wild-type EP^- primary gene, and all modifiers are $CHKPT^-$

These 2000 offspring replace the 2000 parents in the deme.

The deme size remains constant unless the pathological trait appears. I assume that the pathological trait grows in frequency within the deme fast enough to be considered instantaneous relative to the other time scales in the model. Therefore, when a pathological trait appears in a deme, the deme goes extinct before the next iteration of migration, whereupon it is recolonized by a single individual from a random neighboring deme.

This model is designed to be as simple as possible, yet demonstrate the evolution of evolvability in the case of evolutionary pathologies. Clearly there are several avenues in which greater biological realism could be captured, with accompanying increases in the number of free parameters and dynamical variables. These elaborations include intra-deme dynamics of the pathological trait, specific ecological bases for the pathology, sexual reproduction, recombination, migration, and variation in the size of propagules during recolonization. Also, the model could be extended to include modifier variation that is not neutral. These elaborations are deferred to treatments more extensive than the present work.

2.1 Model Details

For concreteness it will be helpful to describe the model mathematically. Let

N be the population size of each deme,

A_p be the number of alleles at the primary locus (only one of which is the pathological allele, EP^+),

A_M be the number of alleles at each modifier locus (only one of which is the $CHKPT^+$ allele),

L be the number of modifier loci, and

μ be the mutation rate per locus per generation.

I assume a uniform mutation model, in which all transitions between alleles are equally likely. Hence this is a *house-of-cards* model [27; 28, p. 15]. This symmetry allows us to lump all the nonpathological alleles EP^- into a single class, indexed with 1, while the pathological allele EP^+ is indexed with 0. Then the transition probabilities under mutation for alleles at the primary locus are

$$\text{Prob}[0 \leftarrow 1] = p_{01} = \mu \left(\frac{1}{A_p} \right),$$

$$\text{Prob}[1 \leftarrow 0] = p_{10} = \mu \left(\frac{A_p - 1}{A_p} \right) = \mu \left(1 - \frac{1}{A_p} \right).$$

The symmetry also allows us to lump all the $CHKPT^-$ alleles into a single class, indexed with 0, while the $CHKPT^+$ allele is indexed with 1 (this choice is made so that the pathological phenotype is expressed in the genotype of all 0s). The transition probabilities under mutation for alleles at each modifier locus are

$$\text{Prob}[0 \leftarrow 1] = m_{01} = \mu \left(\frac{A_M - 1}{A_M} \right) = \mu \left(1 - \frac{1}{A_M} \right),$$

$$\text{Prob}[1 \leftarrow 0] = m_{10} = \mu \left(\frac{1}{A_M} \right).$$

These probabilities give the following transition matrices for mutation at the primary and modifier loci:

$$\mathbf{P} = (1 - \mu)\mathbf{I} + \mu \begin{bmatrix} \frac{1}{A_p} & \frac{1}{A_p} \\ 1 - \frac{1}{A_p} & 1 - \frac{1}{A_p} \end{bmatrix}$$

and

$$\mathbf{M} = (1 - \mu)\mathbf{I} + \mu \begin{bmatrix} 1 - \frac{1}{A_M} & \frac{1}{A_M} \\ \frac{1}{A_M} & 1 - \frac{1}{A_M} \end{bmatrix}.$$

With the assumption that mutation occurs independently at each locus, the transition probability matrix for the entire genotype is

$$\mathbf{T} = \mathbf{P} \otimes \mathbf{M}^{\otimes L}, \tag{1}$$

where $\otimes L$ represents L tensor products of \mathbf{M} .

An alternative representation of the state space for the modifier loci is to lump all the modifier loci together, and simply count the number of CHKPT^+ alleles among them. To get a transition from j to i CHKPT^+ alleles, mutation must produce a net change of $i - j$ in the CHKPT^+ count. We calculate this by summing over all possible ways to get this net change, that is, all the values of k , where k CHKPT^+ alleles mutate to CHKPT^- , and $i - j + k$ CHKPT^- alleles mutate to CHKPT^+ :

$$\begin{aligned} \tilde{m}_{ij} = & \sum_{k=0}^{\min(i,j,L-i,L-j)} \binom{j}{k + \phi(j,i)} m_{01}^{k + \phi(j,i)} (1 - m_{01})^{j - k - \phi(j,i)} \\ & \times \binom{L - j}{k + \phi(i,j)} m_{10}^{k + \phi(i,j)} (1 - m_{10})^{L - j - k - \phi(i,j)}, \end{aligned} \tag{2}$$

where $\phi(i, j) = \max(i - j, 0)$. Clearly Equation 2 defines a Markov chain, but its terms are not easily digested. Its form is useful nevertheless for a numerical implementation of the model, because the values \tilde{m}_{ij} need be calculated only once, and then a single call to a random number generator is sufficient to determine the transition. Direct implementation of Equation 1 would require L random number calls per transition, which is computationally more costly with large population sizes and numbers of generations.

For small mutation rates, Equation 2 is approximated as

$$\tilde{m}_{ij} = \begin{cases} (L - j) m_{10} & \text{if } i = j + 1, \\ 1 - j m_{01} - (L - j) m_{10} & \text{if } i = j, \\ j m_{01} & \text{if } i = j - 1. \end{cases} \tag{3}$$

To complete the description of the evolution within a deme we include Wright-Fisher multinomial sampling. The elements of the Wright-Fisher model are as follows:

\mathbf{x} is the vector of frequencies of each genotype i in the population, corresponding to Nx_i individuals of genotype i ;

$\mathbf{y} = \mathbf{T}\mathbf{x}$ is the vector representing the probability distribution for sampling a genotype among the offspring produced by population \mathbf{x} ;

\mathbf{x}' is the vector of the frequencies of each genotype i in the population in the next generation, corresponding to Nx'_i individuals of genotype i , produced by taking N independent samples from the distribution $\mathbf{y}(\mathbf{x})$;

$n = 2^{L+1}$ is the number of possible genotypes.

Since the population consists of discrete individuals, the frequency vectors are now restricted to a lattice of discrete points on the simplex Δ_n , namely,

$$\Delta_n(N) = \left\{ \mathbf{x} : N x_i \in \{0, 1, \dots, N\}, \sum_{i=1}^n x_i = 1 \right\}.$$

The Wright-Fisher model forms a Markov chain whose transition matrix on frequency vectors is

$$\mathbf{G} = [G_{\mathbf{x}', \mathbf{x}}]_{\mathbf{x}, \mathbf{x}' \in \Delta_n(N)}$$

with entries

$$G_{\mathbf{x}', \mathbf{x}} = N! \prod_{i=1}^n \frac{y_i^{Nx'_i}}{(Nx'_i)!} = \frac{N!}{\prod_{i=1}^n (Nx'_i)!} \prod_{i=1}^n (\mathbf{e}_i^T \mathbf{T}\mathbf{x})^{Nx'_i}, \tag{4}$$

where $\mathbf{e}_i^T = [0 \ 0 \ \dots \ 1 \ \dots \ 0 \ 0]$ has the 1 in the i th position.

The probability distribution on population compositions as a function of the generation time t is thus

$$\boldsymbol{\xi}(t) = \mathbf{G}^t \mathbf{e}_k, \tag{5}$$

where $\boldsymbol{\xi}(t)$ is the vector of probabilities for different population compositions at time t , and k is the genotype that recolonized the deme after its last extinction.

A full analysis of Equation 5 is not necessary to see what is happening in the model. In this model, a deme goes extinct as soon as it generates its first pathological phenotype. Since demes are founded by a single individual, all the genotypes are initially the same, and nonpathological. The model is designed so that mutation pressure alone within a population will steadily reduce the number of CHKP1^+ alleles. When mutation pressure has knocked out all the CHKP1^+ alleles, then the appearance of the pathology allele at the primary locus will produce the pathological

phenotype. As time goes on, the probability that one of the individuals will have mutated to the pathological phenotype increases monotonically. Demes with a greater number of CHKPT^+ alleles will have a greater longevity, and thus be more likely to be the source of propagules to recolonize extinct demes. Through this differential viability of demes, increased numbers of CHKPT^+ alleles in the genome can evolve.

3 Results

The trajectories of a typical run of this model are shown in Figure 2 for the case of one EP^+ and three EP^- alleles at the primary locus. It shows the distribution of the mean number of CHKPT^+ alleles per individual in each deme as evolution progresses.

The distribution of the metapopulation is initialized at generation 0 to fall along on the x axis, that is, all demes have an average of zero CHKPT^+ alleles per individual. Under mutation pressure alone, the population would equilibrate to the dotted line at $0.0625 = 1/16 = (8 \text{ loci})/(128 \text{ alleles per locus})$. After 100 generations of extinction and recolonization dynamics, however, almost all demes have an average of one CHKPT^+ allele per individual. By generation 200, most demes average 2 CHKPT^+ alleles per individual, with a few demes having more or fewer.

As the generations continue, demes averaging just under 3 CHKPT^+ alleles increase to around half of the total number. A fraction of the demes average as many as 4 CHKPT^+ alleles per individual. The consequence of having multiple CHKPT^+ loci is that there is little likelihood that a deme will generate an individual with the pathological trait. Hence, the evolvability of the pathological trait has been suppressed.

It is notable that the distributions are concentrated around integral numbers of CHKPT^+ alleles. This occurs because each deme is recolonized by a single individual with a discrete number of CHKPT^+ alleles.

As mutations accumulate, the distribution shifts downward toward the equilibrium line at 0.0625. Once a deme is recolonized, it “ages” with the mutational loss of CHKPT^+ alleles. There is no dynamic to increase the number of CHKPT^+ alleles above the mutational equilibrium frequency of 0.0625 in an existing deme.

As demes age and more individuals lose their CHKPT^+ alleles, the chance of generating the pathological trait increases and, along with that, deme extinction. Hence, deme extinctions are drawn predominantly from older demes, while deme recolonizations are a uniform sample of individuals in the deme neighborhood. The difference in the distributions of extinctions and recolonizations leads to the increase in the number of CHKPT^+ alleles per individual over the metapopulation.

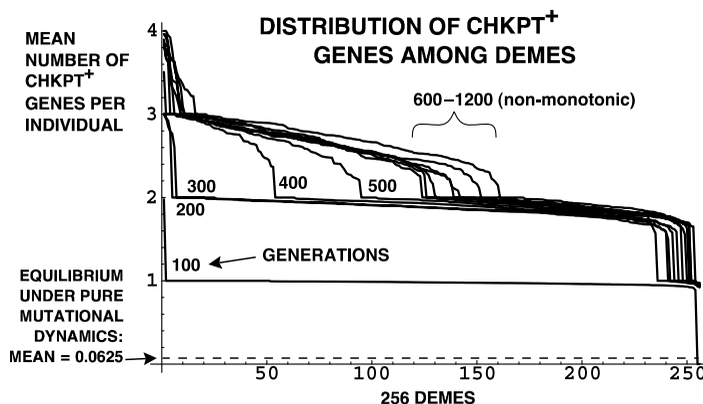


Figure 2. Evolution of evolvability suppression. The mean number of CHKPT^+ alleles per individual in each deme is calculated and plotted against the demes sorted by this value along the X axis. In this run there are one EP^+ and three EP^- alleles at the primary locus.

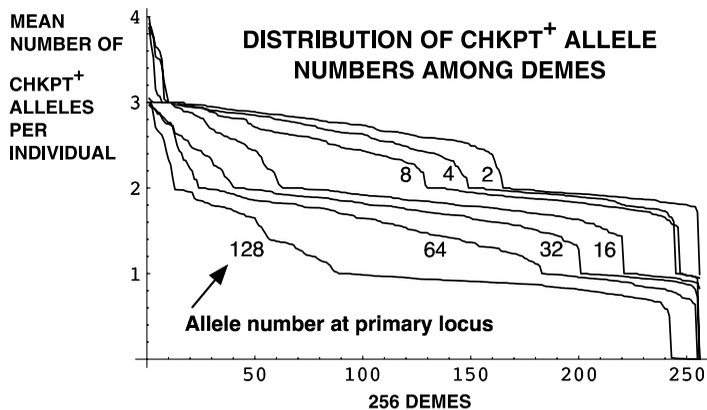


Figure 3. Evolvability suppression as a function of mutation pressure. Distributions in the mean number of CHKPT^+ alleles per individual in each deme are shown as a function of the number of alleles at the primary locus, ranging from 2 to 128. At the primary locus there is always 1 EP^+ allele, and a range of between 1 and 127 EP^- alleles. Census is at generation 1000. The greater the mutation pressure on the EP^+ allele (the fewer the EP^- alleles), the greater the number of CHKPT^+ alleles that evolve to suppress the pathology phenotype.

The effect of allele number at the primary locus, A_p , can be seen in Figure 3. Under the house-of-cards mutation model, the rate of mutation to the EP^+ allele is $p_{01} = \mu/A_p$. Hence, p_{01} is a measure of the baseline evolvability of the pathological trait in the absence of any CHKPT^+ evolvability suppressors. We would expect that the higher the baseline evolvability of the pathological trait is, the greater the pressure for its suppression should be. This is confirmed in Figure 3.

In this figure, the population is always censused at generation 1000, at which time the steady state has been attained. The different distributions represent the outcome for a range of allele numbers A_p at the primary locus. For the minimum number $A_p = 2$, this means that half of all mutations at the primary locus will be to the pathological EP^+ allele. For the maximum number of $A_p = 128$ alleles, only $1/128$ of mutations will produce the pathological EP^+ allele. The system can respond by tuning the number of CHKPT^+ alleles to the evolvability of EP^+ .

This is what is observed. For $A_p = 2, 4, \text{ and } 8$, some fraction of demes is found with mean numbers of CHKPT^+ alleles greater than 3. For $A_p = 16$ and higher, some demes are always found with mean numbers of CHKPT^+ alleles greater than 2. Only when $A_p = 128$ can any demes be found that have no CHKPT^+ alleles.

Thus, the higher the baseline evolvability is for the pathological trait, the greater is the number of evolvability suppressors that will be maintained by extinction and recolonization dynamics.

4 Discussion

The possibility that hierarchically structured populations can affect the evolvability of short-sighted traits (evolutionary pathologies) has been verified in this simple concrete model. Previous work [43] has shown that hierarchically structured populations can lead to the survival of far-sighted traits and to the suppression of short-sighted traits, when genes for them are introduced into a population. Here, we find that even the process that *introduces* short-sighted traits can itself come under selection in hierarchically structured populations, and be caused to suppress the very generation of evolutionary pathologies by mutation.

4.1 Evolvability Modifiers or Simply Epistatic Loci?

It might be asked whether the checkpoint genes modeled here (Figure 1) are really evolvability modifiers, or rather simply genes that epistatically interact with the primary locus to control the pathology phenotype. Indeed, the only distinction between the primary locus and the modifier loci is

in the asymmetry between the number of alleles that allow the pathological trait to be expressed: in the primary locus, 1 out of A_p alleles ($A_p \in \{2, 4, 8, 16, 32, 64, 128\}$) allow expression of the pathological trait; in the modifier loci, 127 out of 128 alleles allow the expression. Loss of the pathological trait is the same whether due to a mutation at the primary locus or to a mutation at one modifier locus to CHKPT^+ .

Thus, one might argue that the results seen here do not constitute an evolution of evolvability, but are simply metapopulation selection against the pathological trait itself, as seen in [29]. This hypothesis, however, would explain the evolution of at most one CHKPT^+ locus. The evolution of additional CHKPT^+ loci has no effect on the phenotype, but rather affects only the rate at which pathological variants arise by mutation—that is, the evolvability of the pathological trait [6]. We observe in the simulation that the metapopulation evolves to the point where most demes average from 2 up to 4 CHKPT^+ loci per individual for small values of A_p .

Nunney [45] investigates a related model of multistep carcinogenesis and finds conditions that would support the evolution of multiple evolvability suppressor loci. The model differs in a number of ways from the present study: instead of a Wright-Fisher model of finite populations, it models cells in a tissue that divide until a certain number is reached. This model is readily analyzed, and when the cell number and mutation rate are high enough, multiple tumor suppressor loci are required to maintain at least one of them in all cells.

The present model is oriented toward multicellular organisms that reproduce in a spatial structure, with extinction and recolonization dynamics. While the model makes for a more protracted analysis, the numerical simulation of the dynamics reveals details of the steady-state distribution of the number of evolvability suppressor alleles. The overall result parallels the results of [45]: strong mutation pressure toward CHKPT^- alleles is overcome by the metapopulation dynamics. The primary consequence is that evolvability of the evolutionary pathology is suppressed, and the rate of deme extinction is reduced because of evolution at the modifier loci.

4.2 Evolvability Suppression and Canalization

The dynamics of this model is closely related to that of neutral network models of canalization. The concept of canalization was proposed in [53, pp. 94–95] and [60]. Canalization is defined as the accumulation of mutations that stabilize a phenotype against either genetic or environmental perturbations. The CHKPT^+ loci stabilize the nonpathological trait against mutations.

The usual condition that promotes canalization is stabilizing selection, when departures from the phenotype are deleterious (canalization thus lowers the mutational load). In the case of evolutionary pathologies, however, instead of stabilizing selection against the pathology phenotype, there is directional selection in favor of it. Only the presence of metapopulation dynamics prevents the immediately advantageous pathology phenotype from fixing in the population. Nevertheless, the metapopulation dynamics has the same long-term effect that stabilizing selection would have if acting against the pathological trait.

4.3 Neutral Network Dynamics in the Model

Since all the different genotypes containing CHKPT^+ alleles or EP^- alleles have equal fitness, they constitute a neutral network under stabilizing selection [13]. The population is forced by metapopulation dynamics to remain on this neutral network, since as soon as a genotype mutates off of the network to the pathology phenotype, its deme goes extinct.

In their study of evolution on neutral networks, van Nimwegen et al. [59] emphasize that “a population seeks out the most connected areas of the neutral network.” By “the most connected areas,” they mean genotypes that produce a large number of neutral mutations, since neutral mutations are what count as connections in the neutral network graph.

However, we see in this model that the feature that the population seeks is not “the most connected areas of the network,” but rather genotypes that are further from the edge of the network. The connectednesses of genotypes are all approximately the same when they have two, three, four,

and more $CHKPT^+$ alleles, in that all of their single-mutant neighbors are still on the network (the double and higher mutations are so rare for small mutation rates that they have little influence). The difference between them is in their distance from the edge of the network, the edge being the genotypes with no $CHKPT^+$ alleles. The number of $CHKPT^+$ loci equals the mutational distance from the edge of the neutral network. Populations are initialized on the edge of the network, and evolve by moving away from the edge. The stronger the baseline evolvability of the pathological trait, the further away from the edge they evolve.

Thus, this model demonstrates an important principle for understanding evolution on neutral networks under stabilizing selection: it is not the local connectedness of regions in the network that makes them attractors, but rather their distance from the boundaries of the network, which is a global feature of network structure.

4.4 The Necessity for Single-Organism Propagules

A crucial requirement of the model for $CHKPT^+$ loci to evolve is that demes be recolonized through extreme population bottlenecks. If recolonized demes were simple copies of a neighboring deme in their genotypes distribution, there would be no way to reverse the mutational pressure toward lower values of $CHKPT^+$ loci. Very strong genetic drift is required to produce occasionally higher numbers of $CHKPT^+$ loci, and the model here maximizes the force of drift by recolonizing with a single individual. Clearly, the magnitude of drift effects will be decreased by propagules with larger numbers of individuals, producing fewer demes with extreme numbers of $CHKPT^+$ loci. Consequently, large propagules will hinder the evolution of $CHKPT^+$ loci.

It is notable that multicellular organisms almost universally start as single individual cells. Michod and Roze [39] point out that this prevents the spread of mutants that defect from multicellular cooperation. When the multicellular organism is taken as the deme, and its death as the extinction of the deme, then the dynamics can be seen to be homologous.

4.5 Plastid Gene Migration to the Nucleus: Driven by Evolvability Suppression?

Evolutionary pathologies can be prophylactically prevented through evolvability suppression, which is the elimination of heritable variation for the pathology at the lower level of the population hierarchy. The model of modifier genes considered here is but one means by which the generation of heritable variation may be suppressed. In this modifier gene model, the mechanism of control is distributed among the lower-level individuals in the population hierarchy, and acts locally within the individuals. Replication and mutation of each lower-level individual occurs independently. The only structures that create dependence between lower-level individuals are their common descent from the single recolonizer of their deme, the cloistering of reproduction within the demes, and the extinction of demes.

If the independence of replication could be taken away from the lower-level individuals, then heritable variation for the pathological trait and natural selection between them could be prevented. If all individuals at the lower level were compelled to share the same phenotype, then no selective differences could arise. A means to achieve this would be to remove the pathology gene from the lower-level individuals, and move its function to their common environment. This constitutes an alternative to modifier genes for evolvability suppression.

One example of this particular phenomenon comes to mind: the transfer of genes from mitochondria and chloroplasts to the nucleus. Mitochondria and chloroplasts correspond to lower-level, endosymbiotic individuals replicating within a deme, the eukaryotic cell. The cell has its own deme-level inheritance mechanism, the nucleus.

The dynamics underlying the migration of genes from endosymbionts to the host genome—a systematically repeated phenomenon—are not resolved. Berg and Kurland [9] investigate whether differential mutation rates can produce the gene migration. The possibility I wish to propose here is that the migration of plastid genes to the nucleus is favored because it is a form of evolvability suppression.

A plastid gene that mutates to increase plastid proliferation, at the expense of cell function, is an evolutionary pathology. If such a gene has migrated to the nucleus, all the plastids will share its phenotype, so heritable variation and differential replication of plastids cannot occur. Hence, only mutations that are beneficial to the cell—the higher level in the hierarchy—will be selected. Thus, plastid genes that migrate into the nucleus are no longer vulnerable to evolutionary pathologies. Furthermore, cellular mechanisms promoting the transfer of plastid genes to the nucleus would be secondarily selected. Thus, evolvability suppression is a hypothesis that may account for the migration of plastid genes to the nucleus.

4.6 The Emergence of Evolvability Suppressors

The model considered here makes a clear distinction between the primary locus and the evolvability modifier loci. In real organisms, however, as well as in the most interesting artificial life systems, genes have emergent functions that are not pre-labeled as primary or modifier loci. Similarly, the phenotypes are not pre-labeled as far-sighted, short-sighted, or pathological, but their effects on the short- or long-term survival of their carriers are emergent properties. These properties may not be inferable from a reductionist description of them. Hence, real organisms and artificial life systems may exhibit evolutionary pathologies as emergent phenomena that can only be observed retrospectively.

Mechanisms that can suppress evolvability may also be emergent. One adaptation may close off the pathway to another adaptation. For example, the evolution of a large mouth could preclude capturing prey in crevices. Evolutionary tradeoffs and developmental constraints are ubiquitous possibilities.

What the present study shows is that a trait that has the happenstance ability to foreclose pathways to evolutionary pathologies may evolve for that reason, and not for what may appear to be its obvious functions.

Mitotic checkpoint genes, which may have originally evolved to ensure viable mitosis, may have been exapted [17] to ensure the *inviability* of defective mitosis in higher organisms.

The suppression of evolvability for evolutionary pathologies becomes a possibility in evolutionary systems that contain structured populations. Structured populations abound in biological systems, and have been investigated in evolutionary computation and artificial life as well [10, 50].

Because of the complexity of organismal phenotypes, the effect of one trait on another trait's evolvability may not be obvious, or even in principle predictable. Hence, structured populations may contain adaptations to suppress evolutionary pathologies where there is no way for the observer to know that this is their purpose. However, if nature, or the experimenter, changes the population structure from multiple demes to a single interacting population, the suppression of evolutionary pathologies may become known by observing its breakdown: genetic variability for pathological traits that had been suppressed under a structured population could be expected to emerge in a newly panmictic population. In biological and artificial life systems where phenotypes and evolutionary dynamics are emergent, the best methodology to explore the suppression of the evolvability of evolutionary pathologies may be through experimental alterations in population structure.

4.7 Conclusion

The model examined here demonstrates another circumstance in which evolvability can evolve. What evolves here is not an organismal phenotype, but the genome's propensity to generate phenotypic variants. In the present case, when the trait confers short-term individual advantage but long-term population disadvantage, under a hierarchically structured population, evolvability evolves to be suppressed. The net effect of these hierarchical evolutionary dynamics is, once again, to increase the survival of the organism.

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Appendix: Group Selection Model for Limitation of Group-Disadvantageous Mutation

Research Notebook, August 4, 1984:

There is a gene which evolves via individual selection but decreases group fitness. There is another gene controlling the rate of *introduction* via mutation of such “selfish” genes. Given that the selfish gene is introduced, we don’t want it to spread over the whole population.

Try Joel Peck's model [48, 49] of group descent: an infinite number of sites. Groups leave offspring groups with probability = f (#of selfish). Each generation, selection acts to increase the % of selfish. The second gene controls the mutation rate to the selfish type.

This model applies not only to behavior but to any trait which may have a population-level negative consequence, or ... a long-term negative consequence.

For the evolution of sex, I imagine that phenotypes causing increased difficulty of producing parthenogens will fix in a sexual population.

Possible difficulty: rates of evolution or selection go to zero as mutation rate goes to 0. So that always some residual chance of the negative mutation remains.

Also, one could have a model where a modifier causes the selfish allele to have a selective disadvantage, or from pleiotropy. [1]

