Synonymous codon usage and selection on proteins

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October 13, 2004

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Selection pressures on proteins are usually measured by comparing homologous nucleotide sequences [57]. Recently we introduced a novel method, termed ‘volatility’, to estimate selection pressures on protein sequences from their synonymous codon usage [38, 39]. Here we provide a theoretical foundation for this approach. We derive the expected frequencies of synonymous codons as a function of the strength of selection, the mutation rate, and the effective population size. We analyze the conditions under which we can expect to draw inferences from biased codon usage, and we estimate the time scales required to establish and maintain such a signal. Our results indicate that, over a broad range of parameters, synonymous codon usage can reliably distinguish between negative selection, positive selection, and neutrality. While the power of volatility to detect negative selection depends on the population size, there is no such dependence for the detection of positive selection. Furthermore, we show that phenomena such as transient hyper-mutators in microbes can improve the power of volatility to detect negative selection, even when the typical observed neutral site heterozygosity is low.

1 Introduction

Nucleotide coding sequences of many organisms exhibit significant codon bias – that is, unequal usage of synonymous codons. Codon bias has been attributed both to neutral processes, such as asymmetric mutation rates, as well as to selection acting on the synonymous codons themselves. The most common selective explanation of codon bias posits that synonymous codons differ in their fitness according to the relative abundances of iso-accepting tRNAs; a codon corresponding to a more abundant tRNA would be used preferentially so as to increase translational efficiency [23, 11, 45]. To a large extent, this hypothesis has successfully explained interspecific variation in genome-wide codon usage for organisms ranging from Escherichia coli to Drosophila melanogaster [1].

Recently, however, we have noted that codon bias in a protein sequence can also result from selection at the amino acid level, even in the absence of direct selection on synonymous codons themselves [38, 39]. Codon bias arises from selection at the amino acid level because of asymmetries in the structure of the standard genetic code. Proteins that experience different selective regimes should exhibit different synonymous codon usage. Following from this observation, we have introduced methods to screen a single genome sequence for estimates of the selection pressures acting on its proteins by comparing their synonymous codon usage [39].

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In this paper, we provide a theoretical discussion of codon usage biases that result from selection at the amino acid level. Our analysis helps to provide a theoretical grounding for techniques of estimating selection pressures on proteins using signals gathered from their synonymous codon usage [38, 39]. Throughout most of this paper, we will ignore any source of direct selection on synonymous codons, and focus on the codon biases that result purely from selection at the amino acid level. To the extent that any other sources of codon bias apply equally across the genome, we have devised a bootstrap method to control for these external sources of codon bias when estimating selection pressures on proteins [38, 39]. In the discussion, however, we describe a range of confounding factors that may vary across the genome in some organisms and limit the applicability of codon-based methods to detect selection.

2 Codon volatility

Codon usage biases can arise from the familiar process of selection on proteins because synonymous codons may differ in their volatility – defined, loosely, as the proportion of a codon’s point mutations that result in an amino acid substitution [38]. Although there are several possible definitions of volatility, which can all be informative, we have recently used the following formal definition [39].

We index the 61 sense codons in an arbitrary order $i = 1 \ldots 61$. We use the notation $aa(i)$ to denote the amino acid encoded by codon $i$. For each codon $i$, let $B(i)$ denote the set of sense codons that differ from codon $i$ by a single point mutation. We define the volatility of codon $i$ by:

$$
\nu(i) = \frac{1}{\#B(i)} \sum_{j \in B(i)} D[aa(i), aa(j)]
$$

where $D$ denotes the Hamming metric, which is zero if two amino acids are identical, and one otherwise. The definition in Eq. 1 applies when all nucleotide mutations occur at the same rate. When differential nucleotide mutation rates are known (e.g. a transition/transversion bias [49]), these rates can be incorporated into the definition of codon volatility by appropriately weighting the ancestor codons [39].

Minor variants of Eq. 1 yield related definitions of codon volatility. For some applications, one may want to allow termination codons in the definition of $B(i)$. It is also natural to consider alternatives to the Hamming metric, $D$, that weight substitutions between amino acids depending upon the differences in their stereochemical properties [33, 38]. A variety of other metrics [47, 53] that reflect the effects of different amino acid substitutions on protein
structure may likewise be incorporated into the definition of codon volatility. In this paper, however, we will focus on the most basic definition of codon volatility (Eq. 1 using the Hamming metric), because variant definitions are based on the same underlying principle and produce similar results in practice [38].

Under the most basic definition of volatility, there are four amino acids (Glycine, Leucine, Arginine, and Serine) whose codons differ in their volatility. As a result, when controlling for amino acid content, we obtain a volatility signal from only those sites that contain one of these four amino acids – which amounts to about 30% of the sites in a typical gene. (If one uses stereochemical metrics [33, 38] for $D$ in the definition of volatility, then $\sim 75\%$ of the sites in a gene contain a volatility signal). Although 30% may seem like a small proportion of sites from which to obtain a signal of selective pressures, it is larger than the proportion of sites often used to detect selection via sequence comparison of recently diverged species [14, 8]. (For example, fewer than 4% of neutral sites exhibit substitutions when comparing human and chimpanzee sequences [8].)

In the following sections we analyze the consequences of selection on proteins for codon usage in general, as well as for the volatility measure in particular. We demonstrate that the expected codon usage at a site, as well as its temporal dynamics, depend upon the strength of positive or negative selection on the amino acid sequence. In Sections 3 through 5 we examine negative selection in infinite and finite populations. In Section 6 we discuss positive selection. Our analysis is initially confined to the patterns of codon usage at a single site under selection at the amino acid level. Proceeding from this analysis, we also discuss codon usage over many sites within a gene or genome, and analyze how many sites are required in principle to detect a reliable signal of selection by inspecting synonymous codon usage.

### 3 Negative Selection and Codon Bias in an Infinite Population

Most nonsynonymous mutations in a protein coding sequence presumably reduce the fitness of an organism. For a large proportion of sites, therefore, natural selection opposes any change in the amino acid. We refer to this type of selection as “negative selection.”

For the purposes of exploring the effect of negative selection on codon usage, we assume that selection cannot discriminate between the synonymous codons for the favored amino acid at a site. However, mutations are more likely to be nonsynonymous, and hence deleterious, if the codon at that site has high volatility. As we will show, this fact results in an effective
preference for the less volatile codons, among those codons that code for the favored amino acid at the site. We emphasize that this preference for a codon of low volatility at a site under negative selection is not caused by a direct fitness difference between synonyms. Rather, more volatile codons will occur less frequently as a second-order consequence of negative selection at the amino acid level, and the structure of the genetic code.

Proteins with a larger number of sites under negative selection will exhibit a statistical bias towards less volatile codons, after controlling for their amino acid content. Here we calculate the expected magnitude of the codon bias as a function of the mutation rate, the strength of negative selection, and, in Section 4, the population size. We also analyze the conditions under which we can expect to detect and draw inferences from this bias, and we estimate the time scales needed to establish and maintain such a signal.

3.1 A simplified genetic code

In an infinite population, we can describe the dynamics of codon usage at an individual site by using the standard multi-allele model first introduced by Haldane [19] and used throughout the literature (e.g. ref. [34] Eq. 2.25 or ref. [21]). This model describes a single site which can assume any of $K$ states. In order to investigate codon usage, we consider $K = 64$ states, corresponding to each of the 64 possible codons. In continuous time, the frequency $x_i$ of individuals with codon $i$ evolves according to

$$\frac{dx_i}{dt} = \sum_{j=1}^{K} x_j(t)w_{ji}M_{ij} - x_iW(t)$$  \hspace{1cm} (2)$$

where $w_j$ is the Malthusian fitness of codon $j$, $W(t) \equiv \sum_j w_jx_j(t)$ is the mean fitness of the population, and $M_{ij}$ is the instantaneous rate of mutation from codon $j$ to codon $i$, with $\sum_j M_{ij} = 0$. Although Eq. (2) is non-linear, the equilibrium frequencies of the “alleles” $i = 1, 2, \ldots K$ are given by the leading eigenvector of the matrix $w_jM_{ij}$ [48]. These frequencies determine the expected equilibrium codon usage at a site. For the purposes of this paper, alternative formulations of the $K$-allele model that treat the processes of selection and mutation separately (e.g. ref. [10] Eq. 6.4.1) yield the exact same results.

The equilibrium solution to Eq. (2) for the full genetic code does not lend itself to intuitive understanding. Transient dynamics are also difficult to calculate in this high-dimensional system. Therefore, in order to highlight the essential points of our analysis, we first consider a “toy” genetic code that retains those features of the true genetic code relevant to the study of synonymous codon usage under negative selection. As we will demonstrate, the solution
for the simplified genetic code yields a complete understanding for the full genetic code as well.

We imagine a simplified genetic system with only three possible codons, \( a_1, a_2, \) and \( b \). Codons \( a_1 \) and \( a_2 \) code for amino acid \( A \), which is favored, and codon \( b \) encodes amino acid \( B \), which has selective disadvantage \( \sigma \). We assume that mutations occur at rate \( u \) between these codons according to the structure

\[
a_1 \rightleftharpoons a_2 \rightleftharpoons b,
\]

so that of the two synonymous codons, \( a_2 \) is more volatile.

According to the standard multi-allele model (Eq. 2), the relative frequencies of codons \( a_1, a_2, \) and \( b \) are described by the equation

\[
\frac{d}{dt} \begin{pmatrix} a_1(t) \\ a_2(t) \\ b(t) \end{pmatrix} = \begin{pmatrix} 1 - u & u & 0 \\ u & 1 - 2u & u(1 - \sigma) \\ 0 & u & (1 - u)(1 - \sigma) \end{pmatrix} \begin{pmatrix} a_1(t) \\ a_2(t) \\ b(t) \end{pmatrix} - W(t) \begin{pmatrix} a_1(t) \\ a_2(t) \\ b(t) \end{pmatrix}, \tag{3}
\]

where \( W(t) = a_1(t) + a_2(t) + (1 - \sigma)b(t) \).

The equilibrium frequencies of codons are given by the leading eigenvector of the matrix in Eq. 3. A simple perturbation analysis of this eigenvector shows that the equilibrium frequency of \( a_1 \) depends monotonically on \( \sigma \), and it exhibits a sharp transition between two regimes: the weak selection regime \( \sigma \ll u \) and the strong selection regime \( \sigma \gg u \). In the weak selection regime, the equilibrium relative frequencies of synonyms are given by the expansion

\[
\frac{\hat{a}_1}{\hat{a}_1 + \hat{a}_2} = \frac{1}{2} + \frac{1}{12} \frac{\sigma}{u} + O \left( \frac{\sigma^2}{u^2} \right). \tag{4}
\]

And in the strong selection regime, the equilibrium relative frequencies are given by

\[
\frac{\hat{a}_1}{\hat{a}_1 + \hat{a}_2} = \frac{\sqrt{5} - 1}{2} - \frac{(5 - 2\sqrt{5})(1 - \sigma)}{5} \frac{u}{\sigma} + O \left( \frac{u^2}{\sigma^2} \right). \tag{5}
\]

In the absence of selection \( (\sigma = 0) \) all three codons occur with equal frequency, as we would expect. In particular, the relative frequency of the two synonymous codons \( a_1 \) and \( a_2 \) equals \( \frac{1}{2} \), regardless of the mutation rate. For weak selection \( (\sigma \ll u) \), this result is still approximately true, according to the perturbation expansion above. In the case of strong negative selection \( (\sigma \gg u) \), the relative frequency of the two synonymous codons is given approximately by the inverse of the golden mean, \( \frac{\sqrt{5} - 1}{2} \approx 0.62 \).

The sharp transition between the weak and strong selection regimes defines \( \sigma = u \) as a critical value for negative selection. For \( \sigma \ll u \) negative selection is ineffective at favoring
the less volatile codon, and the site is effectively neutral. But when $\sigma \gg u$, negative selection favors the less volatile codon, and the magnitude of this effect depends only weakly on the value of $\sigma$. This is an essential point. In the strong selection regime, the magnitude of negative selection is relatively unimportant; volatile codons are disfavored at all sites where $\sigma \gg u$. The transition between the weak and strong selection regimes is shown in Fig. 1.

![Equilibrium frequency vs. Strength of selection](image)

Figure 1: The relationship between selection at the amino acid level and resulting synonymous codon usage. The graph shows relative equilibrium frequency of synonymous codons, $\hat{a}_1/(\hat{a}_1 + \hat{a}_2)$, as a function of the strength of negative selection, $\sigma$. The relative frequency of codon $a_1$ is approximately $1/2$ in the weak selection regime ($\sigma \ll u$), and approximately $\sqrt{5} - 1/2$ in the strong selection regime ($\sigma \gg u$). In this figure $u = 10^{-5}$.

### 3.2 The effective disadvantage of a volatile codon

The critical value of $\sigma$ discussed above can be understood intuitively by considering the “effective selective disadvantage” of the more volatile codon $a_2$ that results indirectly from its volatility. We will use the notion of an “effective selective disadvantage” to aid in our analysis of codon usage at a site under negative selection. But we emphasize that our model (Eq. 2) does not assume any direct fitness difference between synonymous codons.

When the disfavored amino acid $B$ is lethal to the organism, then the effective selective disadvantage of codon $a_2$ is particularly simple to understand. In this case, individuals with
codon $a_2$ are removed from the population at rate $u$ because they mutate to the lethal codon $b$, but receive no back-mutations. Hence the effective selective disadvantage, denoted $s$, of codon $a_2$ versus codon $a_1$ is given by $s = u$. The effective selective disadvantage of $a_2$ does not arise from a fitness difference between synonyms, but rather from selection at the level of amino acids and the structure of the genetic code.

When amino acid $B$ is not lethal the situation is slightly more complicated. Nevertheless, for $\sigma \gg u$, mutations from $a_2$ to $b$ typically die due to negative selection before they mutate back from $b$ to $a_2$. As a result, the effective selective disadvantage will still be $s = u$ in the regime of strong selection. We can make this argument concrete by considering the mutation-selection balance between codon $b$ and codon $a_2$. According to the standard mutation-selection balance, the equilibrium frequency of codon $b$ relative to codon $a_2$ equals $\frac{u}{\sigma}$ in the regime $\sigma \gg u$. Thus for each mutant from $a_2$ to $b$, there are at most of order $\frac{u}{\sigma}$ mutations from $b$ to $a_2$. The net mutation rate from $a_2$ to $b$ is therefore $u \left(1 - \frac{u}{\sigma}\right)$. This is the rate at which individuals of type $a_2$ are lost from the population due to the fact that $a_2$ is more volatile than $a_1$. Thus the effective selective disadvantage of $a_2$ relative to $a_1$ is given by $s = u \left(1 - \frac{u}{\sigma}\right)$. By definition, in the strong selection regime we neglect $\frac{u}{\sigma}$ compared to 1, and the effective selective disadvantage of codon $a_2$ is simply $s = u$.

A similar argument holds for the real genetic code. In this case, the favored amino acid may correspond to several synonymous codons, each with a potentially different volatility. However, the effective selective disadvantage, $s$, of a more volatile codon relative to a less volatile synonym is simply the difference in the number of mutations leading to a disfavored codon ($\sigma \gg u$) times $\frac{u}{3}$, where $u$ is the nucleotide mutation rate. (Note that $\frac{u}{3}$ is the rate of mutation between any two particular nucleotides.) For example, when considering the relative frequencies of codons AGA and CGG at a site under negative selection for Arginine, AGA has selective disadvantage $s = \frac{2}{3}u$ compared to CGG, since AGA has two more disfavored neighbors than CGG. By using the value of the effective selective disadvantage, $s$, we can calculate the equilibrium relative frequency of any pair of synonymous codons in mutation-selection balance, and thereby deduce the relative frequencies of all synonyms. Therefore, we can predict synonymous codon usage in the genetic code without resorting to the full solution of Eq. 2.

An analogous argument can be used to calculate the effective selective disadvantage of codon $a_2$ in the regime of weak selection ($\sigma \ll u$). In this regime, the relative equilibrium frequency of codon $b$ versus codon $a_2$ equals $1 - \frac{u}{2u}$. Thus, the effective selective disadvantage of $a_2$ versus $a_1$ is approximately $s = 0$, plus a small correction of order $\sigma$. In other words, when $\sigma \ll u$ selection between $a_1$ and $a_2$ is effectively neutral; it cannot generate codon bias.
We therefore refer to the regime $\sigma \ll u$ as the “almost neutral regime.” This result holds both for the simplified three-codon model and for the real genetic code.

It is also important to calculate the amount of time required to reach equilibrium codon usage in the presence of strong negative selection. Explicit solution of Eq. 3, assuming $\sigma \gg u$, indicates that the e-fold relaxation time is of order $1/u$ (the selection coefficient is $s \sim u$, and so the time scale for population sizes to change under selection is of order $1/s \sim 1/u$).

In other words, starting from any initial frequencies $a_1(0)$ and $a_2(0)$, these frequencies will become e-fold closer to their equilibrium values after a duration of order $1/u$ generations. The same time scale holds for almost neutral sites ($\sigma \ll u$) and for the real genetic code†. In practice, $u$ will be quite small, and equilibrium volatility is approached very slowly. We will revisit this point when we discuss finite populations, and again when we discuss positive selection.

### 3.3 A specific example: selection for Arginine

In this section we consider a simple example that demonstrates how our analysis applies to the real genetic code. We use Eq. 2 to model the dynamics of $K = 64$ alleles corresponding to the 64 codons, indexed in an arbitrary order. For our example, we consider a single site under negative selection for an Arginine codon. In this case we define

$$M_{ij} = \begin{cases} 
1 - 3u, & \text{if } i = j \\
u/3, & \text{if } i \text{ and } j \text{ differ by a point mutation} \\
0, & \text{otherwise}
\end{cases} \quad (6)$$

where $u$ is the nucleotide mutation rate. We define

$$w_i = \begin{cases} 
1, & \text{if } i \text{ encodes Arginine} \\
1 - \sigma, & \text{if } i \text{ encodes a non-Arginine amino acid} \\
1 - \gamma, & \text{if } i \text{ encodes stop}
\end{cases} \quad (7)$$

so that a codon encoding an amino acid other than Arginine has fitness $1 - \sigma$, and a termination codon has fitness $1 - \gamma$. We analyze this model numerically by calculating the leading eigenvector of the matrix $w_j M_{ij}$, which yields the equilibrium frequencies of all 64 codons.

†For $\sigma \ll u$, the process is almost neutral and the time scale calculation of Section 4.2 applies. The real genetic code has the same dynamics because we still have $s \sim u$ for $\sigma \gg u$ and neutral behavior for $\sigma \ll u$. 
In the case of no selection ($\sigma = \gamma = 0$), we find that all codons occur with the same equilibrium frequency, independent of mutation rate, as we would expect. For almost neutral selection ($\sigma \sim \gamma \ll u$), codon usage is still approximately uniform. In the opposite case when Arginine is favored and all other amino acids (or termination codons) are strongly disfavored (\emph{i.e.} $\sigma \sim \gamma \gg u$), the Arginine codons CGA, CGG, CGC, CGT, AGA, and AGG occur with equilibrium relative frequencies $\approx 0.214 : 0.214 : 0.191 : 0.191 : 0.095 : 0.095$. As expected, under negative selection the more volatile Arginine codons occur with lower relative frequency in equilibrium.

The equilibrium frequencies of Arginine codons determine the expected volatility at a single Arginine site under negative selection. Assuming free recombination \cite{12}, an individual gene consists of many such sites randomly assembled; the mean and standard deviation in the volatility (per site) of a randomly sampled gene are shown in Fig. 2 as a function of the strength of negative selection $\sigma$. Note that the stronger the negative selection, the lower the expected equilibrium volatility. The expected volatility exhibits a sharp transition from high to low values when the strength of negative selection $\sigma$ reaches the mutation rate $u$, as discussed above. On either side of this transition, the volatility is insensitive to $\sigma$. The standard deviations plotted in Fig. 2 correspond to a gene comprised of $L = 200$ such sites, each modeled independently by the multi-allele equation.

According to Fig. 2, $L = 200$ independent sites that each experience neutrality ($\sigma \ll u$) can be distinguished on the basis of their volatility from $L = 200$ sites that experience negative selection ($\sigma \gg u$). The difference in the expected volatility between these two regimes is greater than four standard deviations of the volatility within either regime.

In reality, the selective constraint $\sigma$ will vary greatly across the sites of a given protein. In this case, disregarding the possibility of positive selection, the volatility of a gene (after controlling for its amino acid sequence) essentially reflects the relative number of informative sites that experience negative selection versus neutrality. For example, the volatility of gene $X$ that contains $L = 200$ informative sites under negative selection and an equal number of neutral sites will be significantly greater (with a $Z$-score of about three) than the volatility of gene $Y$ that consists of $2L$ informative sites all under negative selection. A more thorough discussion of variable selection pressures across genes is described in Section 4.1, below.

Table 1 shows the equilibrium relative frequencies of synonymous codons for each of the informative amino acids (G, L, R, and S) under neutrality versus various selective regimes. In Table 1 we assume, as we do throughout this manuscript, that volatility is measured using the Hamming metric and that there is no transition/transversion bias. Corresponding values for different metrics or including a mutational bias may be calculated using the same approach.
Figure 2: The relationship between selection and volatility for a gene comprised of $L = 200$ freely recombining sites under selection for Arginine. The graph shows expected volatility per site in the gene ($\pm 1$ standard deviation, dashed) as a function of the strength of negative selection, $\sigma$. The nucleotide mutation rate is $u = 10^{-5}$. The expected volatility is significantly depressed in the regime of strong negative selection, $\sigma \gg u$. (For this figure we assume $\gamma = 1$; virtually identical results hold for $\gamma = \sigma$.)

As seen in Table 1, the difference in the expected volatility between selective regimes is least extreme (indeed, barely informative) for Glycine sites. The volatility difference is most extreme for serine sites: the highly volatile codons AGT and AGC are not expected to occur at a site under negative selection, but they preferentially occur at a site under positive selection. This extreme case results from the fact that codons AGT and AGC are not connected by synonymous point mutations to the other serine codons. This situation does not imply that codons AGT and AGC should be treated separately from the other serine codons. In fact, when treated as an entire group, the serine codons are particularly informative for positive selection (Table 1).

4 Negative Selection in a Finite Population

The models presented in Section 3 describe the processes of mutation and negative selection in an infinite population. In finite populations, however, genetic drift also affects allelic
Table 1: Equilibrium codon usage under neutrality versus selective regimes. In each selective regime, we report the equilibrium relative abundance of codons, and the resulting mean and standard deviation in volatility per site. The first column corresponds to neutrality ($\sigma = \gamma \ll u$); the second column corresponds to neutrality but with disfavored termination codons ($\sigma \ll u$, $\gamma = 1$); the third column corresponds to strong negative selection in an infinite population ($\sigma \gg u$, $\gamma \gg u$); the fourth column corresponds to the expected frequencies after a positively selected sweep (see Section 6). The final column gives the volatility of each codon, assuming no transition/transversion bias [39].
frequencies. In this section, we study the combined effects of mutation, negative selection, and drift, which we analyze using diffusion equations. These equations can be very complex. A full treatment of even the simplified three-codon genetic code requires a two-dimensional diffusion process, and the real genetic code involves a 63-dimensional process. To make this problem tractable, we use the notion of the “effective selective disadvantage” of more volatile codons, as discussed above. This allows us to consider the dynamics only at the favored codons, thereby reducing the dimensionality of the diffusion process.

The neutral ($\sigma = 0$) or almost neutral ($\sigma \ll u$) regimes are straightforward: here all synonymous codons for the favored amino acid have the same effective fitness. In this regime, each synonymous codon occurs with the same probability in steady state, independent of population size.

For the remainder of this section, we analyze the case of strong negative selection ($\sigma \gg u$) at a single site. We consider a diffusion approximation to the process of mutation, selection, and drift operating only on the synonymous codons, to each of which we assign an effective selective coefficient. For the simplified three-codon genetic system, the more volatile codon $a_2$ has an effective selective disadvantage of $s = u$ compared to codon $a_1$. For the real genetic code, more volatile codons will have a selective disadvantage of this order, but the precise value of $s$ will depend on the specific amino acid in question. In the following analysis, we consider the case of the simplified three-codon system. However, we do not explicitly make the substitution $s = u$, so that our results can also be applied (with a slightly different value of $s$) to the real genetic code.

The time-dependent frequency $f(x, t)$ of allele $a_1$ relative to allele $a_2$ can be described by the Komolgorov forward equation [24]

$$\frac{\partial f(x, t)}{\partial t} = -\frac{\partial}{\partial x} \{a(x)f(x, t)\} + \frac{1}{2} \frac{\partial^2}{\partial x^2} \{b(x)f(x, t)\}$$

where the instantaneous mean and variance in the change of allelic frequency are given by

$$a(x) = sx(1 - x) - ux + u(1 - x)$$
$$b(x) = x(1 - x)/N.$$  

The stationary distribution of allele frequencies $\hat{f}(x)$ satisfies the equation

$$\frac{d}{dx} \{b(x)\hat{f}(x)\} = 2a(x)\hat{f}(x)$$

which has the solution [52]

$$\hat{f}(x) = Cx^{\theta-1}(1 - x)^{\theta-1} e^{Sx}$$

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where $\theta = 2Nu$, $S = 2Ns$, and $C$ is chosen so that $\int_0^1 \hat{f}(x)dx = 1$. Since $s \sim u$ (and thus $S \sim \theta$), the shape of the stationary the distribution $\hat{f}(x)$ falls into two categories: a bell-shaped distribution in the regime $\theta > 1$, and a U-shaped distribution in the regime $\theta < 1$.

In other words, for $\theta > 1$ the steady-state population is typically polymorphic at the locus, much like the infinite population mutation-selection balance. Whereas for $\theta < 1$ the steady-state population is usually near-monomorphic at the locus, occasionally switching between alleles $a_1$ and $a_2$, with a bias (whose strength is determined by $S$) towards allele $a_1$.

In stationary state, the expected frequency of allele $a_1$ is given by

$$M(\theta, S) = \int_0^1 x \hat{f}(x)dx = \frac{1}{2} + \frac{B(\theta + 1/2, S/2)}{2B(\theta - 1/2, S/2)}$$  \hspace{1cm} (11)$$

where $B(x, y)$ is the modified Bessel function of the first kind. Similarly, the variance in the frequency of allele $a_1$ is given by

$$V(\theta, S) = \int_0^1 x^2 \hat{f}(x)dx - M(\theta)^2$$

$$= \frac{1}{4 + 8\theta} + \frac{2\theta B(\theta - 1/2, S/2)B(\theta + 3/2, S/2) - (1 + 2\theta)B(\theta + 1/2, S/2)^2}{(4 + 8\theta)B(\theta - 1/2, S/2)^2}$$  \hspace{1cm} (12-13)$$

We use the standard Taylor series expansion of $B(x, y)$,

$$B(x, y) = \sum_{m=0}^{\infty} \frac{(y/2)^{x+2m}}{m!\Gamma(x+m+1)},$$  \hspace{1cm} (14)$$

to obtain a simple approximation for the mean stationary frequency of allele $a_1$:

$$M(\theta, S) = \frac{1}{2} + \frac{S}{4} + O(\theta^2),$$  \hspace{1cm} (15)$$

valid for $\theta \sim S \ll 1$. This approximation indicates that the difference in expected volatility at a site under neutral versus negative selection is of order $S$, when $\theta \ll 1$.

When $\theta = S = 1$, the mean stationary frequency of allele $a_1$ assumes the value $\frac{1}{e-1} \approx 0.58$. For $\theta \sim S \gg 1$, the mean frequency quickly approaches the asymptotic value $\lim_{\theta \to \infty} M(\theta, \theta) = \frac{\sqrt{5} - 1}{2}$, in agreement with our earlier result for an infinite population.

The results in this section generalize our analysis of an infinite population. For an infinite population, we found that the expected relative frequency of codon $a_1$ equals $\frac{1}{2}$ in the almost neutral regime, and it equals $\frac{\sqrt{5} - 1}{2}$ in the strong selection regime. In a finite population with $\theta \gg 1$, the same results hold. In a finite population with $\theta \ll 1$, the expected relative frequency of the more volatile codon equals $\frac{1}{2}$ in the neutral regime, and it equals $\frac{1}{2} + \frac{Ns}{2}$. 
in the strong selection regime. For any population size, the relative frequency of codon \( a_1 \) depends monotonically on the strength of selection at the amino acid level, \( \sigma \), and it exhibits a sharp transition at the critical value \( \sigma = u \).

It is worth noting that our exact expression (Eq. 11) for the mean stationary frequency of allele \( a_1 \) generalizes earlier work by Bulmer [5] on the relative frequency of two synonymous codons that experience a direct fitness difference. In the limit of small \( \theta \), we find that

\[
\lim_{\theta \to 0} M(\theta, S) = \frac{1}{2} + \frac{B(1/2, S/2)}{2B(-1/2, S/2)} = \frac{1}{1 + e^{-S}},
\]

which agrees with Bulmer’s result (his Equation 6). In other words, Bulmer’s approximation applies only for vanishing small mutation rates (or population sizes).

We can again use the standard Taylor expansion of the Bessel function to obtain a simple expression for the variance in the stationary frequency of allele \( a_1 \),

\[
V(\theta, S) \approx \frac{(3 + 2\theta)(4 + 8\theta) - 3S^2}{16(3 + 2\theta)(1 + 2\theta)^2},
\]

which is a highly accurate approximation for all \( \theta \), provided as usual that \( S \) is of order \( \theta \) or smaller. Note that when \( \theta \ll 1 \) the variance is approximated by \( \frac{1}{4} - \frac{\theta}{2} \), and when \( \theta \gg 1 \) the variance is of order \( \frac{1}{\theta} \).

### 4.1 Inferring Negative Selection in a Finite Population

Our exact (Eq. 11) or approximate (Eq. 15) expressions for the stationary mean frequency of codon \( a_1 \) allow us to determine the minimum number of sites required for codon volatility to distinguish reliably between neutral versus negative selection. When sites are modeled independently (equivalent to the assumption of linkage equilibrium [42]), under neutrality (\( \sigma \ll u; \; s = 0 \)) the relative frequency of codon \( a_1 \) versus codon \( a_2 \) across a gene of length \( L \) is binomially distributed with mean \( \frac{1}{2} \) and variance \( \frac{1}{4L} \). If, on the other hand, the gene experiences negative selection (\( \sigma \gg u; \; s = u \)), then the relative frequency of codon \( a_1 \) is binomially distributed with mean \( M(\theta, S) \) and variance \( M(\theta, S)[1 - M(\theta, S)]/L \). Therefore, in order to reliability reject neutrality at about the 95% confidence level, we require

\[
M(\theta, S) - \frac{1}{2} > 2\sqrt{\frac{1}{4L}}
\]

Using this equation, Fig. 3 shows the minimum number of sites required to reliably distinguish negative selection from neutrality on the basis of codon volatility, under our simplified 'genetic code' consisting of three codons.
Figure 3: The relationship between the scaled population size, $\theta = 2Nu$, and the minimum number of sites required to distinguish negative selection from neutrality, at the 95% confidence level. Sites are assumed to be unlinked. It is important to note that the appropriate effective population size that determines the value of $\theta$ in practice does not necessarily equal the average neutral site heterozygosity (see Section 5).

Eq. 18 applies when comparing a collection of neutral sites against a collection of sites under negative selection. In most situations, however, the selective constraint $\sigma$ will vary across the sites of a protein. For example, consider gene $X$ with $L + J$ sites under negative selection, compared to gene $Y$ with $L$ neutral sites and $J$ sites under negative selection. In this case, the expected frequency of codon $a_1$ in gene $Y$ is $(L/2 + JM(\theta, S))/(L + J)$. Therefore, in order to reliably infer that gene $X$ experiences more negative selection than gene $Y$, at the 95% confidence level we require

$$M(\theta, S) - \frac{L/2 + JM(\theta, S)}{(L + J)} > 2 \sqrt{\frac{L/4 + JM(\theta, S)[1 - M(\theta, S)]}{(L + J)^2}}$$

As Eq. 19 indicates, the power to discriminate between two genes is decreased when both genes contain many sites, $J$, under negative selection and only a few sites, $L$, under different selective regimes. Nevertheless, provided $J \sim L$, the power to discriminate between genes $X$ and $Y$ is decreased by $\sim$20% at most (compared to $J = 0$), and so the minimum number of sites required to detect negative selection (Fig. 3) remains mostly unchanged.

Although the results in this section were derived for a simplified genetic code, the scaling
behavior of these solutions holds for the full genetic code as well – i.e. when comparing neutrality to negative selection, for $\theta \ll 1$ the expected difference in volatility per site will be of order $\theta$; and for $\theta \gg 1$ the expected difference in volatility can be calculated from the infinite population model (Eq. 2 and Table 1).

4.2 Relaxation towards steady state

Although Eq. 11 predicts the steady-state relative frequencies of codons $a_1$ and $a_2$ in the selected regime ($\sigma \gg u$), we have not yet discussed how long it takes, on average, to reach this steady state. In the case of a very large population, $\theta \gg 1$, we know from the infinite population model (Section 2) that the $e$-fold relaxation time to equilibrium is of order $\frac{1}{u}$ generations. In this section, we demonstrate that the same result applies to the time scale of relaxation towards steady state in the regime $\theta \ll 1$.

As usual, we consider a single site under negative selection. In the regime $\theta \ll 1$, we have seen that the steady-state population will spend most of the time in a nearly monomorphic state, with a preference (of order $\theta$) for the less volatile codon, $a_1$. Therefore, in order to calculate the time scale of relaxation towards steady state, we may simply calculate the amount of time required such that, starting with a population fixed for allele $a_2$, the probability of the population remaining fixed for allele $a_2$ has been reduced $e$-fold.

Given a population initially fixed for codon $a_2$, there are $Nu$ mutations to codon $a_1$ generated per generation. Each of these mutations has an effective selective advantage $s = u$ over allele $a_2$, and will therefore fix with probability $2s/(1 - e^{-2Ns})$ \[10\]. Hence the rate of production of a mutation that will eventually fix is given by

$$P_{fix} = \frac{2Nu}{1 - e^{-2Ns}} \approx u,$$

assuming $\theta \ll 1$. According to this calculation, the mean time until fixation of codon $a_1$ is of order $\frac{1}{u}$ generations, which gives the time scale of relaxation to the steady-state codon usage in a finite population under negative selection.

5 About Population Sizes

As discussed above, the strength of the signal of negative selection depends upon the parameter $\theta = 2Nu$. What is the appropriate value of $\theta$ in practice?

Unfortunately, this question is far easier asked than answered. Population geneticists have long struggled to reconcile estimates of $\theta$ deduced from polymorphism data with direct
measurements of $N$ and $u$ across broad taxonomic ranges. The effective population sizes of micro-organisms in particular are topics of active debate. Estimates of $\theta$ are usually obtained by comparing SNP data at neutral (or presumably neutral) sites against the expected site diversity or the expected number of segregating sites under a neutral model \cite{13}. In a recent survey \cite{30} authors have reported an average value of $\theta \approx 0.15$ among the prokaryotes studied. But estimates of $\theta$ for a microbial species can vary by four orders of magnitude, and they depend strongly on assumptions about population structure \cite{3}. To complicate matters further, heterogeneity in mutation rates leads to substantial underestimates of $\theta$ \cite{16}.

Aside from uncertainty in its estimation, the value of $\theta$ deduced from neutral SNP data \cite{30} may not be relevant to questions of selection and volatility. Monomorphism observed at neutral sites may result from non-neutral processes, such as background selection \cite{7} or hitchhiking on periodically sweeping sites \cite{31}. As a result, the variance effective population size estimated from SNP data may not be relevant to other aspects of evolution, such as substitutions at linked weakly selected sites \cite{16}.

One particularly striking example of a discrepancy in the appropriate effective population sizes arises from the consideration of mutator phenotypes. Populations of microbial species periodically experience a transient increase in the mutation rate, often $10^2 - 10^3$ times greater than that of a non-mutator strain \cite{17}. Between 2-20\% of bacterial populations isolated in the wild at any given time exhibit a mutator phenotype \cite{17,36,28}. The mutator phase can be induced in several ways. A defective DNA repair gene may arise and sweep to fixation by hitchhiking on a positively selected mutation \cite{35}. The entire population then experiences an elevated mutation rate until a non-mutator allele sweeps and replaces the mutator \cite{35,12}. A second, perhaps more common mechanism is stress-induced mutagenesis; natural isolates of *E. coli* often experience an increase in their mutation rate in response to stress \cite{4}. As a result of these and other observations, researchers have argued that bacterial populations evolve primarily by periodic acquisition of mutator phenotypes followed by adaptive sweeps and subsequent loss of the mutator \cite{17,12,35}. As we shall see, the effect of this process on synonymous codon usage is dramatic: the expected site diversity is driven by the value of $\theta$ in the wildtype regime ($\theta_w = 2Nu_w$), but the pattern of synonymous codon usage at a site under negative selection is driven by the value of $\theta$ in the mutator regime ($\theta_m = 2Nu_m \gg \theta_w$).

As a simple example of this phenomenon, we have simulated a Fisher-Wright model of a single locus in a population of constant size $N = 1000$. The simulated site is subject to recurrent mutation between “alleles” $a_1$ and $a_2$ at wildtype rate $u_w = 10^{-5}$. As in Section \cite{4} the alleles $a_1$ and $a_2$ differ in fitness by $s$, where $s$ equals the mutation rate. Periodically, we model the fixation of a mutator allele (or, equivalently, the stress-induced mutagenesis
across the entire population) by exogenously increasing the mutation rate to \( u_m = 10^3 \times u_w \) for 100 generations; thereafter we (artificially) enforce a selective sweep at the site, followed by reversion to the wildtype mutation rate. Overall, the population experiences the mutator regime for 5% of the time, consistent with observed frequencies of mutator phenotypes in the wild \[17, 36, 28\]. According to our simulations, the average site diversity, \( 2x(1 - x) \), at a randomly chosen time equals 0.028, which is close to its expected value assuming that \( \theta \) is given by \( \theta_w \): \( \mathbb{E}[2x(1 - x)] = \theta_w = 0.02 \). But the average frequency of allele \( a_1 \) equals 0.611, which is close to its expectation assuming that \( \theta \) is given by \( \theta_m \): \( \mathbb{E}[x] = M(\theta_m, \theta_m) = 0.616 \) (Eq. 11). In other words, the average frequency of the less volatile codon \( a_1 \) is dominated by the mutator periods, but the average site heterozygosity (and any estimate of \( \theta \) based on it) is dominated by the non-mutator periods.

There is a simple, intuitive explanation for this result. The average heterozygosity at the site is low at virtually all times (except during the brief mutator periods) because selective sweeps cause monomorphism, followed by long periods of low \( \theta \). Therefore, the effective \( \theta \) for SNP diversity is small, i.e. close to \( Nu_w \). But the site converges quickly towards the less volatile codon during the mutator periods, since the rate of convergence is determined by \( s = u_m \). And the site is essentially frozen during the non-mutator periods, since the decay rate of volatility is only \( u_w \). Therefore the expected frequency of \( a_1 \) at a random time is primarily determined by the frequency reached during the mutator regime. As is clear from this explanation, the expected frequency of codon \( a_1 \) will, in general, depend upon the stochastic scheduling of mutator periods. For example, the site will converge towards \( M(\theta_m, \theta_m) \) provided the population experiences at least one mutator phase of duration of order \( 1/u_m \) generations, within every \( 1/u_w \) generations. In fact, even if the mutator phases are very brief and infrequent, the average frequency of allele \( a_1 \) can greatly exceed the value predicted by \( \theta \) estimated from the average site heterozygosity.

Although the simple model used in this section does not describe any but the most phenomenological features of mutator alleles, it does reveal an important general observation: the value of \( \theta \) estimated from neutral SNP data does not in general equal the effective value of \( \theta \) that determines synonymous codon usage at a site under negative selection. This result is of utmost importance to any discussion of the relationship between \( \theta \) and the power of volatility to detect negative selection.
6 Positive selection

In the sections above, we have considered selection that opposes a change to the amino acid at a site. This type of negative selection induces a bias towards the less volatile codons for the favored amino acid at a site. However, selection sometimes favors a change in the amino acid at a particular site. In such situations, as we will demonstrate, a site is more likely to be occupied by a codon of greater than average volatility.

A variety of mechanisms are known to cause positive selection. Frequency dependence often induces diversifying selection at a site, whereas an exogenous change in the environment can induce directional selection for a new, specific amino acid. We do not here model all of the various types of positive selection, but rather focus on the essential aspect shared by these mechanisms. We analyze the dynamics at a site that has, for a period of time, experienced negative selection for amino acid A, and that subsequently experiences negative selection for different amino acid, B (for whatever reason). We refer to the change in the selective regime as a positive selection event.

Prior to the onset of positive selection, amino acid A is assigned fitness 1 and all other amino acids fitness $1 - \sigma$; subsequently, amino acid B is assigned fitness 1 and all others fitness $1 - \sigma$. We assume that $N\sigma \gg 1$ (otherwise, the site is effectively neutral at the amino acid level) and that $\sigma \gg u$ (otherwise, the expected codon frequencies are uniform). Once the population shifts to the new amino acid B, it is clear that the site will more likely contain a codon that is more volatile than the average B-codon, because it has just arisen through a nonsynonymous mutation. Since B is now favored, negative selection subsequently operates to reduce the volatility at the site. However, this process takes time. Thus, for some time after the positive selection event, there is a bias toward elevated volatility at the site, which gradually decays. In this section, we analyze this process.

Analogously to previous sections, we initially consider a simplified genetic code consisting of four codons, $a_1$, $a_2$, $b_1$, and $b_2$, the first two of which encode amino acid A, and the latter two amino acid B. Mutations can only occur between codons $a_1$ and $a_2$, $a_2$ and $b_2$, and $b_2$ and $b_1$, creating the mutation structure

$$a_1 \rightleftharpoons a_2 \rightleftharpoons b_2 \rightleftharpoons b_1.$$ 

In this simplified genetic code, codons $a_2$ and $b_2$ are the more volatile codons for their respective amino acids.

After the change in selection from amino acid A to B, a mutation to codon $b_2$ that survives stochastic drift will eventually arise. Thus, at least initially, the more volatile
codon $b_2$ is more prevalent than the less volatile codon $b_1$. During this period, we can detect the signature of the positively selected sweep because of the elevated volatility at the site. However, negative selection for amino acid $B$ will eventually favor codon $b_1$. Therefore, the volatility signature of the positive selection event will be present provided that the time scale of decay toward codon $b_1$ is longer than the interval since the positive selection event.

Fortunately, the time scale of decay towards $b_1$ is quite long. For $\theta \gg 1$, we can use the infinite population model to find this time scale. As discussed above, the time required to reduce the volatility $e$-fold is of order $\frac{1}{u}$. For $\theta \ll 1$, we must use a finite population size calculation. In this regime, the population is nearly monomorphic at almost all times. Following the selective sweep, the site will be monomorphic for $b_2$ with almost unit probability. We are interested in the duration of time required such that probability of being monomorphic for $b_2$ (as opposed to $b_1$) has been reduced $e$-fold. The probability of switching between $b_2$ and $b_1$, however, is of order $u$ per unit time (even before $b_2$ has finished outcompeting $a_2$), according to Eq. 20. Thus, the time scale of decay in a finite population is also $\frac{1}{u}$.

According to this analysis, a selective sweep will result in the presence of a more volatile codon for of order $\frac{1}{u}$ generations – a very long time indeed. (In the case of E. coli, for example, $\frac{1}{u}$ generations is nearly 100,000 years, given $u \approx 5 \times 10^{-10}$ and generation time $\approx 20$ minutes. The generation length and resulting time scale for E. coli in the wild may be much longer yet.) Equivalently, repeated sweeps for amino acid changes at a site will result in the presence of more volatile codons at almost all times, provided that new sweeps occur more often than every $\frac{1}{u}$ generations.

### 6.1 Inferring Positive Selection

The above analysis for a simplified genetic system generalizes in an obvious way to the real genetic code. After a positive selection event at a site, the population switches from a codon for amino acid $A$ to a codon for amino acid $B$. The expected volatility of the new codon is greater than the average volatility of $B$-codons, because the new codon has just arisen through a nonsynonymous mutation. To be more precise, if the population is monomorphic for a random non-$B$ codon before the selective sweep, then after the sweep occurs the expected relative frequencies of the $B$-codons are given, approximately, by their relative volatilities. Subsequent to the selective sweep, the increased volatility at the site will decay on a time scale of order of $\frac{1}{u}$ generations.

There is a critical distinction between the volatility signature of positive selection versus that of negative selection. The depressed volatility at a site under negative selection is
caused by a mutation-selection-drift balance. When the effective population size is small, a large number of sites are required to distinguish negative selection from neutrality. By contrast, the volatility signature of positive selection is not an equilibrium property, and it is not sensitive to population size. Regardless of \( \theta \), the probability of sampling a more volatile codon is significantly elevated immediately after a selective sweep at a site, and this probability decays only at rate \( u \).

As we have seen, a gene that contains many sites under positive selection will exhibit a greater volatility (controlling for its amino acid composition) than a gene under neutral or, especially, negative selection. How many positively selected sites are required in order to detect a reliable signal? In the case of Leucine, for example, the expected volatility of a site that has recently experienced a positively selected sweep is approximately \( 0.660 \pm 0.072 \) (one standard deviation), whereas a neutral Leucine site has expected volatility \( 0.646 \pm 0.073 \), and a Leucine site under negative selection has expected volatility \( 0.632 \pm 0.0070 \) (see Table 1). Therefore, the volatility of about 100 Leucine sites under positive selection will be significantly greater (at the 95% confidence level) than that of 100 neutral sites. Similarly, the volatility of about 25 positively selected Leucine sites will be significantly greater than that of 25 negatively selected sites. Similar results hold for Serine and Arginine; Glycine is less informative.

It is worth noting that the elevated volatility for a positively selected Serine site will decay even more slowly than for other amino acids, because the highly volatile codons ACC and AGT are not connected by synonymous mutations to other serine codons.

7 Discussion

7.1 Codon volatility versus comparative sequence analysis

Selection pressures on proteins are usually estimated by comparing homologous nucleotide sequences [57]. Orthologous genes are identified in different organisms and sequenced; their sequences are then aligned, and the changes that have accumulated since divergence are used to infer the selection pressures that have been acting [18]. When available, sequence variation sampled from individuals within a species can be compared with variation across species to produce an elegant test for adaptive evolution at a locus [32, 42]. In addition, there are a variety of statistical tests designed to detect a departure from neutrality in the site frequency spectrum sampled within a single species (see ref. [27] and references therein). In many cases, the complete distribution of these statistics under the neutral null model are
difficult to derive, but they have been studied through computer simulation [44].

Techniques for estimating selective constraints via sequence comparison are typically applied, independently, to one or several genes at a time. When extensive intra- or inter-specific sequence data are available at a locus of interest, such techniques have proven enormously useful for measuring selection, and it is unlikely that they will be significantly improved by incorporating information about synonymous codon usage. But the accurate estimation of selective constraints requires a large number (approximately six or more [2]) of orthologous sequences for each gene of interest. At the genome-wide scale, comparative data (i.e. orthologous gene sequences) will not be available for all genes, and methods to estimate selective constraints based on sequence comparison will often be inapplicable. Furthermore, the genes under positive selection are often of particular interest, but such genes are even less likely to have identifiable orthologs in related species due to their rapid sequence divergence [39]. Even in the lineage of the Saccharomyces genus, which is currently the best-case scenario for comparative genomics, the genomes of four species have been fully sequenced and only two-thirds of the genes in S. cerevisiae have unambiguously identifiable orthologs in related species [40]. Unlike comparative techniques, the analysis of synonymous codon usage offers a computational tool to screen for selection pressures on all genes in a sequenced genome. Genome-wide screens based on analyzing synonymous codon usage may prove useful in identifying important classes of genes under strong selection, such as the antigens of pathogens [39].

Unlike most comparative statistics that test for a departure from neutrality, estimates of selection based on bootstrapped volatility scores [39] are not ‘estimators’ in a rigorous statistical sense — i.e. statistics whose sampling properties can be derived from a null model, and which can be used in likelihood ratio tests of a null hypothesis [55, 8]. Given the expected relative frequencies of codons that we have derived for each of the three regimes (neutral, negative, and positive selection; Table 1), it may yet be possible to design maximum-likelihood methods that estimate the number of sites of a gene in each regime. This approach will be complicated, however, by other sources of codon bias; see below.

Aside from the different situations in which they are applicable, and differences in the rigor of their derivation, estimates of selection based on codon volatility differ in a fundamental way from most estimates based on sequence comparison. Homologous sequence comparison between species is often used to assess, either by maximum likelihood [18] or maximum parsimony [29], the rates of synonymous and non-synonymous substitutions in a coding sequence. The ratio of these rates, dN/dS, is used as a measure of the selective constraints that have been acting on a protein since the divergence of the species being
compared. An alternative approach, based on a Poisson Random Field (PRF) model of mutation frequencies, uses the site frequency spectrum at a locus sampled from individuals within a species to deduce the average selective pressure for or against amino acid changes in a gene [42]. (Poisson Random Field models can also be used to construct likelihood ratio tests of departure from neutrality [9].) Like most comparative methods, however, both of these models typically assume that all the sites within a gene experience the same selective pressure against amino acid changes (but see the site-by-site likelihood tests of Yang et al. [55]). Under the PRF theory, for example, authors have estimated a very small “average” selection pressure against amino acid changes in *E. coli* genes: $\sigma \sim 10^{-8}$ [20]. This value does not represent the arithmetic average of the true $\sigma$ values across sites, but rather the best-fit constant value of $\sigma$ that would make the PRF model consistent with observed sequence variation at polymorphic sites.

When evolutionary rates are estimated at individual residues [54, 55], however, we find great variation across sites. Moreover, direct experimental measurements of the fitness consequences of amino acid substitutions in micro-organisms reveals huge variation in selection pressures across the residues of an individual protein: a substantial proportion of substitutions are lethal, and a substantial proportion have undetectable effect [50, 51, 58, 41]. Therefore, it is not entirely clear how best to interpret the value of $\sigma \sim 10^{-8}$ estimated for *E. coli* genes using the PRF model, which assumes constant pressure at each residue.

Compared to dN/dS or $\sigma$ estimated by the PRF model, codon volatility quantifies selection pressures in a very different, coarser manner. As discussed above, volatility essentially measures the number of sites in a gene that experience negative ($\sigma \gg u$) versus neutral ($\sigma \ll u$) versus positive selection. Given that, in reality, many amino acid changes to a protein sequence are lethal while other changes have no effect whatsoever, it is reasonable and meaningful to estimate the number of sites in the selected versus neutral regimes. But volatility is not sensitive to variation in selective pressures within either of these regimes. Hence, the volatility measure is in some ways a coarser description of selective pressure than PRF or dN/dS. One should not necessarily expect that volatility will correlate very strongly with dN/dS or PRF estimates, because the latter measures represent some sort of average $\sigma$ over the entire gene, and are thus presumably sensitive to the full range of variation in $\sigma$. A measure based on codon volatility is therefore different from and complementary to dN/dS or PRF estimates of the selective constraints on a genes.

As an aside, it is important to note that the most common model used to estimate dN/dS from divergent nucleotide sequences [18] does not itself reflect the relationship between selection and volatility. dN/dS is often estimated by fitting maximum likelihood parameters
to a simplified Markov-chain model of sequence evolution that ignores population variability [18]. Models that ignore population variability are perfectly reasonable approximations when comparing the sequences of relatively divergent lineages [18]; but such models fail to detect the effect of amino-acid selection on synonymous codon usage. Such models consider only a single sequence that is assumed to represent the dominant genotype in the population at any time. Mutation and selection are modeled simultaneously by adjusting the transition rates between codon states in the sequence [18]. As a result, in equilibrium, the number of transitions into a state per unit time must equal the number of transitions out of that state; and so equilibrium synonymous codon usage does not depend upon the strength of selection in these simplified models [18]. (In fact, under the standard assumption of time-reversibility, such models require as parameters the specification of the equilibrium codon usage [18], and therefore they clearly cannot be used to predict equilibrium codon usage.) Simulations of sequence evolution based on these simplified models (such as the non-frequency-dependent simulations of Zhang [56]) will thus fail to detect the relationship between dN/dS and volatility, whereas more detailed simulations that account for population variability (such as the frequency-dependent simulations of Zhang [56], as well as the non-frequency-dependent simulations in this work) will properly reflect the relationship between selection and volatility, as predicted by Fisher-Wright models of a replicating population.

### 7.2 Other sources of codon bias

Although it came as a surprise to early neutral theorists [25], it is now clear that there are several processes that result in unequal usage of synonymous codons. Many processes that cause codon bias in microorganisms, such as biased nucleotide content or mutation rates, can apply roughly equally to all the genes in a genome. To the extent that other sources of codon bias apply equally across a genome, it is straightforward to control for these biases when comparing the volatilities of genes within a genome to estimate selection pressures on proteins [39].

To the extent that other sources of codon bias differ from gene to gene within a genome, they may (if not properly controlled for) introduce errors into estimates of the relative selection pressures on proteins inferred from codon volatility [39]. Similarly, selection on synonymous codons – particularly selection that varies from gene to gene – will likewise introduce errors into estimates of selection on protein sequences obtained by comparative techniques such as dN/dS [43, 22].

As we have argued, some of the variation in synonymous codon usage across a genome is
caused by the variation in selection pressures on protein sequences. Throughout our analysis, we have specifically ignored any other source of codon biases so as to derive the effects of selection at the amino acid level on codon usage. But in many organisms other processes that vary between genes are certainly operating as well. For instance, it is known that the transition/transversion mutation bias can vary across a genome. Results on *S. cerevisiae*, whose genome exhibits marked variation in the tr/tv bias [40], suggest that this source of variable codon bias will not distort estimates of selection based on volatility: whether or not one accounts for the variation in the tr/tv bias across the genome of *S. cerevisiae* one obtains virtually the same rankings of gene volatilities ($r > 0.99$) [40].

Aside from mutational biases, there are other sources of codon bias that vary from gene to gene in some organisms. In the yeast *S. cerevisiae*, researchers have observed that synonymous codon usage, measured by the Codon Adaptation Index (CAI) [43], is correlated with a gene’s expression level in laboratory conditions [9]. This correlation is thought to be caused by selection for translational efficiency and/or accuracy: a codon corresponding to a more abundant tRNA is expected to be translated more quickly (due to the higher probability per unit time that the appropriate tRNA will “find” the codon) and more accurately (since the correct tRNA will likely have the greatest chance of pairing if it is the most abundant).

Considering this alternative source of biased codon usage, two questions should be asked: do other sources of codon bias distort estimates of selection based on volatility, and how can we control for these confounding factors? Unfortunately we do not have a truly satisfactory answer for either of these questions, but the discussion below may shed some light on the issues involved.

With regards to the first question, we note that the degree to which other sources of codon bias may distort volatility-based estimates of selection will strongly depend on the organism being studied. Some species (such as humans) exhibit a much weaker correspondence between codon frequencies and tRNA abundances than others species; so clearly other sources of codon bias will affect volatility values differently in different species. In a species with a strong correspondence between codon usage and tRNA abundances, the extent to which variation in this source of codon bias across the genome affects volatility will depend on whether volatile codons are (un)preferred: if there is no correlation between volatility and tRNA abundances, then the other sources of codon bias will only introduce random error into volatility estimates, making them less powerful but still reliable. If instead the preferred codons tend to have either high or low volatility, then this effect could introduce systematic errors into volatility estimates. In the latter case, in order to quantify how much codon usage bias is caused by volatility as opposed to other factors, one would require a method to predict
for individual genes the amount of codon bias due to these other factors. Unfortunately we are far from having the necessary level of predictive power for other sources of codon bias in any organism. Although gene expression level is somewhat predictive of codon bias, expression levels do not explain most of the variation in codon bias in any genome studied thus far [19]. Until the various sources of biased codon usage can be reliably disentangled, we cannot reliably quantify the effects of these biases on volatility-based estimates of selection.

The second question, how to control for other sources of biased codon usage, is also difficult to answer at present. As discussed above, an appropriate method to control for other sources of bias would require disentangling the various sources of codon bias in a predictive manner for each gene. While this degree of precision is not currently possible, one approach is to assume that the codon bias measured by CAI is entirely independent of volatility, and then control for CAI using partial correlations. For several reasons, we expect this approach to be conservative, as we illustrate using the yeast *S. cerevisiae* (we use this species as an example because it shows a strong preference for codons that match abundant tRNAs, and because we have reliable dN/dS values for almost two-thirds of its genes, calculated from multiple alignments of closely related species [22]). First, we note that dN/dS is itself strongly correlated with both CAI and gene expression levels [37], and it is therefore impossible to construct any measure of selective constraint that agrees with dN/dS and is not itself strongly correlated with CAI and expression levels in yeast. Second, it is possible that the codon bias measured by CAI is in part caused by volatility (*i.e.* highly expressed genes tend to experience stronger purifying selection and therefore exhibit unusual codon usage biased towards lower volatility), and so controlling for CAI would be inappropriate. Despite several biological hypotheses, there is no accepted mechanistic explanation for the correlation between CAI and dN/dS in yeast [37, 11], and so it is unclear whether controlling for CAI is appropriate. Nevertheless, we have tested the correlation between volatility and dN/dS while controlling for CAI using a partial correlation. We find that even when controlling for CAI (or expression levels), there remains a highly significant correlation between volatility and dN/dS in yeast ($p < 10^{-34}$ [40]). Therefore, even under this conservative test, estimates of selection obtained by volatility are still consistent with estimates obtained by homologous sequence comparison. We interpret this result as evidence that volatility is measuring selective constraints above and beyond any signal inherent in CAI.

Indeed, there is a great deal of empirical evidence indicating that the volatility of a gene is correlated with the selective constraint it experiences. Aside from highly significant correlations between volatility and dN/dS in bacterial species and yeast [39], volatility also reflects a range of other features known to correlate with selection on proteins. In *S. cerevisiae,*
for example, volatility is strongly correlated with the essentiality of genes, the number of their protein-protein interactions, and the degree to which they are preserved throughout the eukaryotic kingdom [40]. Furthermore, volatility is significantly elevated among the known antigens and surface proteins (which experience positive selection) in the pathogens *Mycobacterium tuberculosis*, *Plasmodium falciparum*, and Influenza A virus [38][39]. And volatility is significantly depressed in the genes essential for growth of *M. tuberculosis*, as well as in the genes conserved between related *Mycobacterium* species [39]. Therefore, despite potential confounding sources of codon bias that cannot at present be controlled for with appropriate accuracy, in practice volatility-based methods produce estimates of selection pressures that are consistent with our understanding of protein evolution over a diverse range of taxa.

Finally, we note that there may be direct selection on synonymous codons in order to evade mistranslation [26]. Since mistranslation is far more likely to occur between a codon and an anticodon that differ by a single nucleotide, the definition of volatility (Eq. 1) is appropriate for measuring the selective pressure for or against mistranslation. The strength of this type of selection on synonymous codons would depend upon the mis-incorporation rate of tRNA (which is far higher than the mutation rate) and the detriment of mistranslation (which is likely far lower than that of most mis-sense mutations). It is difficult at present to measure the molecular parameters of tRNA mis-incorporation and its fitness effects; so it is unclear how much of a volatility signal arises from mistranslation avoidance versus standard selection on mis-sense mutations. However strong this signal, though, the volatility of a gene would still reflect the degree to which there is selection to conserve, or not to conserve, the (translated) protein sequence.

**Acknowledgments**

We thank Daniel Fisher, Andrew Murray, and Michael Turelli for their input during the preparation of this manuscript. J.B.P. acknowledges support from the Harvard Society of Fellows. M.M.D. acknowledges support from a Merck Award for Genome-Related Research.

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