# The relationship between dN/dS and scaled selection coefficients

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Computational methods assessing the mode and strength of natural selection have become a cornerstone of comparative sequence analysis. Numerous such methods exist, yet how distinct approaches relate to one another and whether inferences made with competing methods yield comparable or disparate results remains entirely unknown. Here, we elucidate the relationship between two widely-used phylogenetic modeling frameworks: dN/dS models on the one hand and mutation-selection-balance (MutSel) models, which estimate amino acid and/or codon scaled selection coefficients, on the other. We derive a formal mathematical relationship between dN/dS and selection coefficients and use this relationship to gain unprecedented insight into the behaviors, limitations, and applicabilities of these competing models. In particular, we prove that MutSel models correspond strictly to  $dN/dS \leq 1$ , as long as all synonymous changes are neutral. However, if there are fitness differences among synonymous codons, then dN/dS can take on arbitrarily high values even if all selection is purifying. These results broadly demonstrate that Mut-Sel models cannot accomodate positive, diversifying selection and that the dN/dS framework cannot distinguish purifying selection on synonymous codons from positive, diversifying selection on amino acids. Moreover, on the basis of our mathematical relationship between dN/dS and scaled selection coefficients, we to propose a new benchmarking strategy for dN/dS models. We employ this strategy to demonstrate that some of the most widely-used dN/dS models, Goldman-Yang-style models, yields substantially biased dN/dS estimates on realistic sequence data, while the less frequently used Muse-Gaut-style models display much less bias. Strikingly, the models yielding the least-biased and most-precise dN/dS estimates are never the ones displaying the best model fit as assessed by AIC. Thus, selecting models solely according to goodness-of-fit criteria can be counterproductive and yield misleading estimates when the models considered do not exactly correspond to the underlying mechanism that generated the data. In conclusion, establishing mathematical links among modeling frameworks represents a novel, powerful strategy to pinpoint previously unrecognized model limitations and strengths.

## Significance Statement

Phylogenetic models that quantify selection strength in protein-coding sequences are among the most rigorous and sophisticated tools in evolutionary studies. While various such frameworks exist, it is virtually unknown how independent approaches relate. Here, we elucidate the relationship between two competing frameworks, dN/dS and selection coefficient models, to discern previously unrecognized model properties and limitations. We prove that MutSel models correspond strictly to regimes of purifying selection  $(dN/dS \leq 1)$ , although selection on synonymous changes can produce spurious signals of positive selection. We additionally find that most dN/dS models infer systematically biased parameter estimates. Moreover, model fit metrics (AIC) cannot necessarily identify which model produces the most accurate inferences, and thus AIC scores may mislead in model selection.

dN/dS | mutation-selection-balance models | scaled selection coefficients | Markov models of sequence evolution

#### Introduction

The oldest and most-widely used method to infer selection pressure in protein-coding genes calculates the evolutionary rate ratio dN/dS, which represents the ratio of nonsynonymous to synonymous substitution rates. This metric indicates how quickly a protein's constituent amino acids change, relative to synonymous changes, and it is commonly used to identify protein sites that experience purifying selection (dN/dS < 1), evolve neutrally  $(dN/dS \approx 1)$ , or experience positive, diversifying selection (dN/dS > 1) [1, 2, 3, 4]. In phylogenetic contexts, dN/dS is typically calculated using a maximum likelihood (ML) approach [5, 6, 1, 7]. ML methods assume a continuous time Markov model of sequence evolution and have become a staple of comparative sequence analysis since their introduction in the 1990s (see ref [8] for a comprehensive review). Throughout this paper, we will refer to these models as dN/dS-based models.

A second class of Markov models, known as mutationselection-balance (MutSel) models, are increasingly being viewed as a viable alternative to the dN/dS framework. While dN/dS-based models describe how quickly a protein's constituent amino acids change, MutSel models assess the strength of natural selection acting on specific mutations. Couched firmly in population-genetic theory, the MutSel framework estimates site-specific scaled selection coefficients  $S = 2N_e s$ , which indicate the extent to which natural selection favors, or disfavors, particular codon and/or amino acid changes [9, 10, 11, 12]. Although first introduced over 15 years ago [9], MutSel models have seen little use due to their high computational expense. Recently, however, several computationally tractable model implementations have emerged [13, 14], allowing for the first time the potential for widespread adoption.

Over the course of twenty years developement, dN/dS-based models have advanced to a high level of sophistication. These models can accommodate a variety of evolutionary scenarios, including synonymous rate variation [6, 15] and episodic [16, 17] and/or lineage-specific selection [18, 19, 20], and they can also incorporate information regarding protein structure and epistatic interactions [21, 22, 23, 24, 25]. This flexibility, along with accessible software implementations [26, 27, 28], makes dN/dS-based models an attractive analysis choice. On the other hand, some have argued that MutSel

#### **Reserved for Publication Footnotes**

models, given their explicit basis in population-genetics theory and attention to site-specific amino-acid fitness differences, offer a more mechanistically realistic approach to studying coding-sequence evolution [9, 11, 12, 29]. Moreover, a growing body of literature has demonstrated that dN/dS estimates are particularly sensitive to violations in model assumptions, calling into question the general utility of dN/dS-based models [30, 31, 32].

Although both MutSel and dN/dS-based models describe the same fundamental process of coding-sequence evolution along a phylogeny, it is unknown how these two modeling frameworks relate to one another. In particular, as these inference methods have been developed independently, it remains an open question whether or not parameter estimates from one model are comparable to those of the other model. As a consequence, although certain rhetorical arguments may be made in favor of using one method over another, there is currently no formalized, concrete rationale to guide researchers in their methodological choices. Elucidating the relationship between these competing modeling frameworks will more precisely reveal under which circumstances the use of these models is justified and has great potential to reveal previously unrecognized model behaviors, limitations, and capabilities.

Here, we formalize the relationship between these two modeling frameworks by examining the extent to which their respective focal parameters, dN/dS and scaled selection coefficients, yield overlapping information about the evolutionary process. To this end, we derive a mathematical relationship between dN/dS and scaled selection coefficients. We find that dN/dS values can be precisely calculated from scaled selection coefficients, and that dN/dS accurately captures the selective pressures indicated by a given distribution of scaled selection coefficients. Furthermore, we prove that, when synonymous mutations are neutral, it is only possible to recover  $dN/dS \leq 1$ from selection coefficients, demonstrating that MutSel models are inherently only able to model purifying selection. Therefore, these models would be an inappropriate analysis method if positive selection is expected. However, we also find that, when synonymous codons have different fitnesses and hence purifying selection acts on synonymous changes, it is possible to recover dN/dS values above 1, even though classical positive, diversifying selection is not occurring.

Finally, this relationship provides a uniquely rigorous platform to examine the performance of dN/dS-based models. Typically, researchers evaluate performance of a given inference framework through simulations that adhere to the underlying model's assumptions (with a notable exception of ref. [33]). Indeed, simulated data is usually generated according to the same model as the inference framework, allowing for a direct comparison between the true and estimated parameter values. While this strategy is critical for testing whether a model implementation behaves as expected, it cannot assess model performance when the data are generated under a different process than the one modeled in the inference framework. However, in real-world sequence analysis, the inference framework will never exactly match the data-generation process. Therefore, a more sensitive test of model performance would examine how a given method performs when data are simulated under different mechanistic processes, and how sensitive the method is to violations of its assumptions. Unfortunately, such an approach is typically infeasible, because the relationships between parameters of interest among distinct model classes are generally not known.

The relationship we establish here between dN/dS and selection coefficients allows us to overcome this limitation, as we can determine the true dN/dS value directly from Mut-

Sel model parameters. Thus, we can assess performance of dN/dS-based inference frameworks by simulating data with a MutSel model and then comparing inferred dN/dS ML estimates (MLEs) to dN/dS values computed from selection coefficients. Using this strategy, we find, for sequences evolved under a symmetric mutation model, that dN/dS values inferred in an ML framework agreed precisely with those calculated from scaled selection coefficients. However, as mutational asymmetry increases, dN/dS MLEs become increasingly biased away from their true values, under a variety of ML model parameterizations. Surprisingly, the ML model parameterization which produced the most accurate dN/dSestimates was never the model which exhibited the best fit to the data (measured by AIC), ultimately revealing that relying on model fit as a litmus-test for model performance can be an ineffective and misleading strategy.

### **Results and Discussion**

Theoretical model. This section contains a rederivation of results presented in refs. [9, 10], reproduced here to introduce notation and to place the remainder of our work into context. We model sequence evolution using the Halpern-Bruno MutSel modeling framework under the assumptions of a fixed effective population size  $N_e$  and constant selection pressure over time [9, 10, 12, 29]. This continuous-time reversible Markov process is governed by the  $61 \times 61$  transition matrix  $T(t) = e^{Qt}$ , where the matrix  $Q = q_{ij}$  gives the instantaneous substitution probabilities between all 61 sense codons, and diagonal elements of Q satisfy  $q_{ii} = -\sum_{i \neq j} q_{ij}$ . We assume that only single-nucleotide substitutions occur instantaneously.

Let  $f_i^{\text{codon}}$  be the fitness of codon i, and let the selection coefficient acting on a mutation from codon i to codon j be  $s_{ij} = f_j^{\text{codon}} - f_i^{\text{codon}}$  [34, 10]. The fixation probability for this mutation is [35, 9, 10]

$$u_{ij} \approx \frac{2s_{ij}}{1 - e^{-2N_e s_{ij}}} = \frac{1}{N_e} \frac{2N_e s_{ij}}{1 - e^{-2N_e s_{ij}}}.$$
 [1]

We further define  $S_{ij} = 2N_e s_{ij}$  as the scaled selection coefficient for this change [10]. The probability of a substitution from codon i to j is therefore

$$q_{ij} = N_e m_{ij} u_{ij} = m_{ij} \frac{S_{ij}}{1 - e^{-S_{ij}}},$$
 [2]

where  $m_{ij}$  is the codon mutation rate, which represents the rate at which codon i transitions to codon j [9, 34]. If we assume that  $m_{ij}$  only has non-zero entries for single-nucleotide changes, we can write it as  $m_{ij} = \mu_{s_i t_j}$ , where  $\mu_{kl}$  is the per-nucleotide mutation rate,  $s_i$  is the source (i.e., before mutation) nucleotide in codon i, and  $t_j$  is the target (i.e., after mutation) nucleotide in codon j.

We now show how  $S_{ij}$  can be written in terms of mutation rates and stationary (equilibrium) codon frequencies  $P_i$ . As this system statisfies detailed balance (reversibility) [9], we have

$$q_{ij}P_i = q_{ji}P_j. ag{3}$$

From equations [2] and [3], we can write the ratio of substitution probabilities as

$$\frac{P_i}{P_j} = \frac{m_{ji}S_{ji}(1 - e^{-S_{ij}})}{m_{ij}S_{ij}(1 - e^{-S_{ji}})}.$$
 [4]

Using  $S_{ij} = -S_{ji}$ , we find that

$$S_{ij} = \ln\left(\frac{P_j m_{ji}}{P_i m_{ij}}\right).$$
 [5]

This equation, previously derived in ref. [9], establishes a relationship between scaled selection coefficients and the stationary codon frequencies of the Markov chain. Moreover, in the specific case of symmetric mutation rates  $m_{ij} = m_{ji}$ , we have  $S_{ij} = \ln (P_j/P_i)$  [34].

Predicting dN/dS from scaled selection coefficients. We now derive respective expressions for nonsynonymous and synonymous evolutionary rates, which we can divide to obtain the evolutionary rate ratio dN/dS. We write the nonsynonymous rate  $K_{\rm N}$  as

$$K_{\rm N} = \sum_{i} \sum_{j \in \mathcal{N}_i} P_i q_{ij} \,, \tag{6}$$

where  $\mathcal{N}_i$  is the set of codons that are nonsynonymous to codon i and differ from it by one nucleotide, and the substitution probability  $q_{ij}$  is defined in equation [2]. To normalize  $K_N$ , we divide it by the number of nonsynonymous sites, which we calculate according to the mutational opportunity definition of a site [5, 7] as

$$L_{\rm N} = \sum_{i} \sum_{j \in \mathcal{N}_i} P_i m_{ij} \,. \tag{7}$$

Thus, we find that

$$dN = \frac{K_{\rm N}}{L_{\rm N}} = \frac{\sum_{i} \sum_{j \in \mathcal{N}_i} P_i q_{ij}}{\sum_{i} \sum_{j \in \mathcal{N}_i} P_i m_{ij}}.$$
 [8]

Similarly, for dS, the synonymous evolutionary rate  $K_S$  per synonymous site  $L_S$ , we find

$$dS = \frac{K_{S}}{L_{S}} = \frac{\sum_{i} \sum_{j \in \mathcal{S}_{i}} P_{i} q_{ij}}{\sum_{i} \sum_{j \in \mathcal{S}_{i}} P_{i} m_{ij}}, \qquad [9]$$

where  $S_i$  is the set of codons that are synonymous to codon i and differ from it by one nucleotide substitution. The quantities  $K_S$  and  $L_S$  are defined as in Eqs. [6] and [7] but sum over  $j \in S_i$  instead of  $j \in \mathcal{N}_i$ . Moreover, if we assume that mutation rates are symmetric and that all synonymous codons have equal fitness (i.e. synonymous mutations are neutral), the synonymous fixation rate satisfies  $u_{ij}|_{j \in S_i} = 1/N_e$  [36], and hence the substitution probability becomes  $q_{ij} = m_{ij}$ . In this circumstance, the value for dS reduces to 1.

#### MutSel models strictly describe purifying selection.

We examined the relationship between dN/dS and scaled selection coefficients by simulating 200 distributions of amino acid scaled fitness values,  $F_a^{\rm aa} = 2Nf_a^{\rm aa}$ , from a normal distribution  $\mathcal{N}(0,\sigma^2)$ . We drew a unique  $\sigma^2$  for each fitness distribution from a uniform distribution  $\mathcal{U}(0,4)$ . Higher values for  $\sigma^2$  correspond to larger fitness differences among amino acids, causing selection to act more strongly against nonsynonymous changes. Thus, high  $\sigma^2$  values indicate strong purifying selection, low values indicate weak purifying selection, and  $\sigma^2 = 0$  indicates that all amino acids are equally fit. We note that these  $F_a^{\rm aa}$  quantities correspond exactly to the aminoacid propensity parameters estimated by currently available site-specific MutSel inference methods [13, 14].

We then converted each amino-acid fitness distribution to a corresponding set of codon fitnesses, as described in *Methods*. Briefly, for 100 of the distributions, we assumed that all synonymous codons had the same fitness, but for the other 100 distributions we allowed synonymous codons to have different fitnesses. In other words, the former 100 distributions do not incorporate purifying selection on synonymous changes

whereas the latter 100 distributions do. Using equations [6] - [9], we computed dN/dS for each distribution of codon fitnesses. For these calculations, we assumed the symmetric mutation model HKY85 [37], which is specified by the parameters  $\mu$ , the nucleotide mutation rate, and  $\kappa$ , the ratio of transititions to transversions. Specifically, transititions occur at a rate  $\mu\kappa$  and transversions at a rate  $\mu$ . We used  $\mu = 10^{-6}$  for all simulations, while we selected a unique value for  $\kappa$  for each simulation from  $\mathcal{U}(1,6)$ .

Under neutral evolution, we expect that dN/dS=1, and as purifying selection increases in strength, dN/dS should correspondingly decrease. Therefore, we expect that dN/dS will decline with the variance  $(\sigma^2)$  of the distribution of amino acid fitness values. Indeed, we observed a strong, negative correlation between these quantities (Figure 1). The larger the fitness differences among amino acids (higher  $\sigma^2$ ), the lower dN/dS, properly reflecting increased purifying selection. This correlation was much stronger for fitness distributions without synonymous selection (Figure 1A) than for those with synonymous selection (Figure 1B). This difference emerged because fitness differences among synonymous codons obscured underlying amino-acid fitness differences. Even so, selection on synonymous codons did not negate the significant correlation between dN/dS and overall selection strength.

Importantly, Figure 1A demonstrates that, in the limiting case when  $\sigma^2$  approaches 0, and thus all codons have virtually the same fitness, dN/dS converges to 1. In other words, when the protein-coding sequence evolved neutrally, selection coefficients correctly yielded a  $dN/dS \approx 1$ . Furthermore, we never recovered dN/dS > 1 when synonymous changes were neutral, revealing a key property of MutSel models: they inherently cannot describe positive, diversifying selection. Indeed, in Appendix 1, we prove that scaled selection coefficients strictly yield  $dN/dS \leq 1$ , under the assumptions that synonymous changes are neutral and mutation is symmetric. This proof formalizes the MutSel model's underlying assumption that selection pressure is constant over the phylogeny and that the protein evolves under equilibrium conditions. Although this proof assumes symmetric mutation rates, we have found numerically that dN/dS remains bounded from above by 1 even when mutations rates are asymmetric (Figure S1).

Purifying selection on synonymous changes can produce dN/dS > 1. The restriction  $dN/dS \le 1$  does not hold when synonymous changes are not neutral, as seen in Figure 1B. In other words, even though the Halpern-Bruno model explicitly assumes that the system is at equilibrium [9, 29], dN/dS can readily be greater than 1. In fact, it is theoretically possible to achieve arbitrarily high dN/dS values when synonymous codon substitutions carry fitness changes. In the most extreme case of synonymous selection, where only a single codon per amino acid is selectively tolerated, the number of synonymous changes becomes  $K_{\rm S}=0$ , and thus the value for dN/dS approaches infinity.

Given that the MutSel model framework assumes an overarching regime of purifying selection, this finding might seem paradoxical. However, the logical argument that dN/dS>1 represents positive, diversifying selection assumes that the rate of synonymous change may be used as a neutral benchmark, an assumption clearly violated when selection acts on synonymous changes. Thus, the traditional signal of positive, diversifying selection, a dN/dS value in excess of one, can result simply from strong synonymous fitness differences.

That sequences evolving under purifying selection can spuriously bear the hallmark of positive, diversifying selection highlights the pitfalls of naively interpreting dN/dS values. Indeed, evolutionary constraints which induce synony-

mous selection are pervasive and affect virtually all domains of life [38], from viruses [39, 40] to plants [41] to Metazoa [42, 43, 44, 45, 46]. Recent work has shown that synonymous rate variation is common across myriad proteins and contributes to evolutionary rate heterogeneity in up to 42% of known protein families [47]. For example, exonic splicing enhancers [48, 49, 50], regions contributing to mRNA secondary structure such as translation-initiation sites [51, 48, 38, 39, 40], and DNA- and RNA-binding sites [49] all experience moderate to strong synonymous selection. It has additionally been suggested that up to 18% of mutational fitness effects in RNA viruses, whose genomes frequently feature sites with dN/dS > 1 [52, 53, 54, 55, 25], are caused by selection acting on synonymous changes [39]. Finally, both selection against protein misfolding and for translation efficiency tend to induce synonymous selection in a gene-specific manner [56, 57], most notably in highly expressed genes [58, 46]. Therefore, while synonymous selection may not dominate genomes in organisms with relatively small effective population sizes [43, 45], it certainly acts strongly at specific sites; indeed,

and/or small, local regions

Recent work also suggests that up to 42% of sites in a given protein, from all domains of life, may experience substantial synonymous selection [47]. As dN/dS ratios are typically measured on a per-site basis, we expect that some sites with dN/dS > 1 may in fact be false positives in the detection of positive, diversifying selection. We offer several approaches to ease this concern in Conclusions.

Relationship between dN/dS and scaled selection coefficients provides a novel benchmarking approach. The relationship we have established between dN/dS and scaled selection coefficients offers a unique opportunity to assess the robustness of dN/dS-based inference methods. It is conventional practice in model development to benchmark models against data simulated according to the model itself. While crucial for testing whether a given model has been correctly implemented, this strategy inherently cannot discern how the model behaves when data arose from a different mechanistic process. To overcome this limitation, we applied a novel benchmarking approach which used the theoretical relationship among modeling frameworks to assess the accuracy and specific utility of those models. This approach, outlined in Figure 2A, entails comparing dN/dS values calculated from selection coefficients to those inferred by an dN/dSbased model. In addition, because MutSel models are based explicitly on population genetics theory, these simulated alignments are likely far more similar to real sequence data than are alignments simulated under a dN/dS-based model, and therefore provide excellent benchmarking data.

Using the selection coefficients and symmetric mutation rates from the previous two subsections, we simulated alignments using standard methods [7] according to the Halpern-Bruno MutSel model [9]. We then inferred a dN/dS value for each alignment using the GY94 matrix [5, 1], which estimates dN/dS with the parameter  $\omega$ . Throughout the remaining text, we refer to dN/dS inferred using ML as  $\omega$  or  $\omega$  maximum likelihood estimate (MLE), and to dN/dS computed using equations [6] - [9] simply as dN/dS. We found that dN/dS values agree nearly perfectly with

We found that dN/dS values agree nearly perfectly with  $\omega$  MLEs (Figure 2B), and indeed this relationship was robust to both synonymous selection and uneven nucleotide composition (simulated alignments featured GC contents ranging from 0.21-0.89). Additionally, Figure 2C demonstrates that  $\omega$  converged to the true dN/dS value as the size of the data set (i.e., simulated alignment length) increased. These

results unequivocally show that, when nucleotide mutation is symmetric, dN/dS-based model-inference methods behave exactly as expected, yielding precisely accurate dN/dS estimates. This finding has important implications for modeling choices; although the MutSel framework might model the sequence evolution in a way that more mechanistically matches the evolutionary process, dN/dS-based models may suffice to model selective forces in phylogenetic data.

Biased dN/dS estimates under asymmetric mutation **models.** We next sought to test the accuracy of dN/dSbased models using more realistic parameter values. this end, we determined codon fitness distributions from 498 unique distributions of experimentally-derived, site-specific amino-acid fitnesses for H3N2 influenza nucleoprotein (NP) [59]. We combined each of these fitness distributions with three sets of experimentally-determined mutation rates, either for NP [59], yeast [60], or polio virus [61], to determine  $498 \times 3 = 1494$  distinct distributions of steady-state codon frequencies (see *Methods* for details). While all three mutation matrices were asymmetric, each featured a differing degree of mutational bias; specifically, the mean ratios  $\mu_{ij}/\mu_{ji}$  for NP, yeast, and polio mutation rates are 1.03, 1.69, and 5.25, respectively. For each resulting set of stationary codon frequencies, in combination with its respective set of mutation rates, we calculated dN/dS and simulated alignments from which we inferred  $\omega$ . Note that we assumed no selection on synonymous codons for these calculations.

dN/dS-based model matrices account for nucleotide mutational bias by incorporating either target codon [5] or target nucleotide [6] frequencies; these frameworks are known, respectively, as GY-style and MG-style models [62]. For example, the instantaneous rate matrix element giving the substitution probability from codon AAA to AAG would contain the target codon frequency  $P_{\rm AAG}$  in GY-style models but the target nucleotide frequency  $\pi_{\rm G}$  in MG-style models. Although the GY-style and MG-style matrices appear to follow different forms, it is possible to write MG-style matrices such that they conform to the reversible GY-style framework, as we show in Appendix 2.

Previous works have suggested that MG-style and GY-style models yield different  $\omega$  estimates [15, 63], so we inferred  $\omega$  according to both GY- and MG-style frameworks. For GY-style models, we used the frequency estimators F61 [5], F3x4 [5], CF3x4 [62], and F1x4 [6]. For MG-style models, we considered both a parameterization in which four global nucleotide frequency parameters were used and a parameterization which employed twelve nucleotide frequency parameters to allow for different frequencies at each codon position. We term the former framework MG1 and the latter MG3. Note that our MG1 corresponds to the original MG-style model [6], whereas our MG3 corresponds to the so-called MG94×HKY84 model [15].

Figure 3 shows the resulting relationships between dN/dS and  $\omega$  MLEs for each set of mutation rates (NP, yeast, and polio), across model frequency parameterizations. Figure 3A displays the estimator bias, defined as the average difference between the true dN/dS value and the  $\omega$  MLEs. Figure 3B displays the precision in this relationship, measured by the squared correlation coefficient  $r^2$  between dN/dS and  $\omega$ . The exact bias and  $r^2$  values are given in Tables S1 and S2, respectively, and full regression plots for  $\omega$  vs. dN/dS are shown in Figure S1.

Two distinct trends emerge from Figure 3. First, asymmetry in the mutational process consistently induced significant bias in  $\omega$  estimates. Most often, the model underestimated  $\omega$  relative to the true dN/dS value. Based on simulations without any selection (dN/dS=1), ref. [63] had previously sug-

gested that GY-style models produce negatively biased  $\omega$  estimates. Our results generalize this finding and show that this bias is pervasive, remains approximately constant through a wide range of dN/dS values, and is not limited to the GY-style framework (Figure 3A, Table S1, Figure S1). Furthermore, we show that this bias systematically increased in magnitude as the underlying mutational process became more asymmetric. Indeed, for all frequency parameterizations,  $\omega$  MLEs were most accurate under NP mutation rates, and both accuracy and precision tended to decrease as mutational bias progressed from yeast to polio mutation rates. Second, frequency parameterizations which more closely matched the mechanistic process that generated the data (MG1 and MG3) generally outperformed all other frequency estimators. In particular, MG1 clearly performed the best of all frequency estimators considered, featuring by far the least amount of estimator bias for the highly asymmetric polio mutation rates. Thus, going forward, we highly recommend that researchers employ MGstyle matrix frameworks in their dN/dS inferences to minimize bias.

Strikingly, when we examined model fit using AIC scores [64, 65] for the different frequency parameterizations, we found that the F61 parameterization was unequivocally the best-performing model, on average, for all datasets (Table 1). This result dramatically juxtaposed the substantial inaccuracy and imprecision in  $\omega$  that F61 frequently yielded. In particular, F61 had the most estimator bias for NP datasets as well as the least precision for both NP and polio datasets (Figure 3). Thus, we found that AIC could not identify the model which produced the most accurate estimates for the parameter of interest.

This finding has broad implications for practices in model selection. In particular, it appears that model fit can be confounded with model accuracy, such that the model with better model fit may produce less accurate parameter estimates. Indeed, if the data are generated by a process distinct from the inference model, AIC cannot necessarily identify which model produces the most precise and least biased parameter estimates. Good model fit, therefore, may not have any bearing on whether using that model is mechanistically justified, and selecting models based solely on fit may not guard effectively against spurious inferences but instead prove misleading. Instead, we suggest that the mechanism producing the data should be carefully considered, and an appropriate inference method which best approximates this process should then be selected.

#### **Conclusions**

By elucidating the relationship between dN/dS and scaled selection coefficients, we have shown that dN/dS-based and MutSel models convey consistent information regarding the strength of natural selection. Importantly, our proof that  $dN/dS \leq 1$  (assuming symmetric mutation and no synonymous selection) indicates that the use of MutSel models is only justified under conditions of strictly purifying selection. This restriction is in part indicated by the basic MutSel model assumption of constant selection pressures over time, or in other words a static fitness landscape [9, 22, 11, 29]. Thus, if the aim is to identify positive selection, of the two frameworks examined here, only dN/dS-based models are appropriate.

However, we have also found that dN/dS values can readily be greater than 1 when selection acts on synonymous mutations, even though the protein sequence is evolving solely under purifying selection. This seemingly paradoxical finding actually reflects an assumption violation; the assertion that

dN/dS > 1 necessarily corresponds to positive, diversifying selection requires that synonymous changes are neutral, which clearly does not hold if there are fitness differences among synonymous codons. This result contributes to a growing body of literature which has found that purifying selection can yield dN/dS > 1 if model assumptions are not met. For instance, dN/dS can be greater than 1, even under purifying selection, if sequences contain segregating polymorphisms [30, 31, 32]. Thus, it is becoming increasingly clear that the dN/dS = 1 neutral threshold typically used to distinguish purifying and positive selection is highly sensitive to violations in model assumptions. We emphasize that it is crucial to ensure that data adhere to model assumptions before conclusions from dN/dS are drawn.

We suggest several strategies to limit such false positive results under synonymous selection. For one, certain formulations of dN/dS-based methods (particularly those implemented in the HyPhy package [26]) consider dN and dSrate variation separately [6, 15, 66] rather than using a single parameter to represent dN/dS. These kinds of methods may be able to distinguish situations in which dN/dS > 1because dN is unusually large (positive selection) or dS unusually small (purifying selection). In addition, dN/dS-based models which correct dS for synonymous selection may prove fruitful in the future for teasing out the source of elevated dN/dS estimates [67]. Finally, our benchmarking approach, in which we simulate sequences according to MutSel models and infer dN/dS both analytically and using ML, may be used to benchmark these kinds of models and may help to identify circumstances under which synonymous selection confounds dN/dS interpretations.

Finally, we emphasize the utility of establishing relationships among distinct modeling frameworks to probe model behavior and evaluate model performance. Such an approach is uniquely able to reveal unrecognized behaviors and/or limitations of different modeling frameworks and can precisely reveal the circumstances in which different models are best suited. We hope that further studies in this spirit will ensure robust model development in future studies.

#### Methods

Simulation of scaled selection coefficients. To examine the relationship between dN/dS and scaled selection coefficients, we simulated 200 distributions of amino-acid scaled fitness values,  $F_a^{\rm aa}=2Nf_a^{\rm faa}$ , from a normal distribution  $\mathcal{N}(0,\sigma^2)$ , where a unique  $\sigma^2$  for each fitness distribution was drawn from a uniform distribution  $\mathcal{U}(0,4)$ . We converted these amino-acid fitnesses to codon fitnesses as follows. For 100 of the fitness distributions, we directly assigned all codons within a given amino acid family the fitness  $F_i^{\rm codon}=F_a^{\rm aa}$ , so that all synonymous codons had the same fitness. For the other 100 fitness distributions, we assigned synonymous codons different fitnesses by randomly drawing a preferred codon for each amino acid. This preferred codon was assigned the fitness of  $F_i^{\rm codon}=F_a^{\rm aa}+\lambda$ , and all non-preferred codons were given the fitness  $F_j^{\rm codon}=F_a^{\rm aa}-\lambda$ . We drew a unique  $\lambda$  for each fitness distribution from  $\mathcal{U}(0,2)$ . We then computed stationary codon frequencies as

$$P_i = \frac{e^{F_i^{\text{codon}}}}{\sum_k e^{F_k^{\text{codon}}}},$$
 [10]

where the sum in the denominator runs over all 61 sense codons [34]. Equation [10] gives the analytically precise stationary frequencies for a MutSel model, under the assumption of symmetric mutation rates [34]. We used equations [6] - [9]

to compute dN/dS for each resulting set of stationary codon frequencies. For these calculations, we assumed the HKY85 [37] nucleotide mutation model, and accordingly we set the transition mutation rate as  $\mu\kappa$  and the transversion rate as  $\mu$ . We used the value  $\mu=10^{-6}$  for all dN/dS calculations, and we drew a unique value for  $\kappa$  from  $\mathcal{U}(1,6)$  for each set of codon frequencies.

**Alignment simulations.** We simulated protein-coding sequences as a continuous-time Markov process using standard methods [7] according to the Halpern-Bruno MutSel model [9]. In brief, this model's instantaneous rate matrix  $Q = q_{ij}$  is populated by elements

$$q_{ij} = \begin{cases} m_{ij} \frac{S_{ij}}{1 - 1/S_{ij}} & \text{single nucleotide change} \\ 0 & \text{multiple nucleotide changes} \end{cases}, \quad [11]$$

for a mutation from codon i to j, where  $m_{ij}$  is the mutation rate, and the scaled selection coefficient  $S_{ij}$  is defined in equation [5]. All alignments presented here were simulated along a 4-taxon phylogeny (Figure 4), beginning with a root sequence generated in proportion to stationary codon frequencies [7]. Unless otherwise stated, all simulated alignments contained 500,000 codon positions. A single evolutionary model was applied to all positions in the simulated sequences. While this lack of site-wise heterogeneity is unrealistic for real sequence evolution, it allowed us to verify our derived relationship between scaled selection coefficients and dN/dS with a sufficiently sized data set.

Computation of stationary frequencies for experimental data sets. We used experimentally-determined site-specific amino-acid fitness parameters  $F_a^{\rm aa}$  for influenza nucleoprotein (NP), from ref. [59], in combination with experimental nucleotide mutation rates for either NP [59], yeast [60], or polio virus [61], to derive realistic distributions of stationary codon frequencies. We combined each of the 498 site-wise amino-acid preference sets reported by ref. [59] with each of the three mutation-rate matrices to construct a total of  $498 \times 3 = 1494$  unique experimental evolutionary Markov models, using the approach in refs. [59, 68]. The instantaneous rate matrix Q for each experimental model is populated by elements

$$q_{ij} = \begin{cases} F_j^{\text{codon}} / F_i^{\text{codon}} m_{ij} & \text{if } F_j^{\text{codon}} \ge F_i^{\text{codon}} \\ m_{ij} & \text{if } F_j^{\text{codon}} < F_i^{\text{codon}} \\ 0 & \text{multiple nucleotide changes} \end{cases}$$

for a substitution from codon i to codon j, where  $F_i^{\rm codon}$  is the fitness of codon i [59, 68]. We calculated  $F_i^{\rm codon}$  values by simply assigning a given amino acid's experimental fitness  $F_a^{\rm aa}$  to each of its consistituent codons; thus, all synonymous changes were neutral. We determined the stationary codon frequencies for each resulting experimental model from the matrix's eigenvector corresponding to the eigenvalue 0. Finally, we simulated alignments for each set of stationary frequencies and corresponding mutation rates according to the Halpern-Bruno model (equation [11]).

Maximum likelihood inference of dN/dS. For the 200 alignments simulated with symmetric mutation rates, we inferred dN/dS using the M0 model [2], as implemented in the HyPhy batch language [26]. The M0 model uses the GY94

instantaneous rate matrix, which is populated by elements

$$q_{ij} = \begin{cases} P_j & \text{synonymous transversion} \\ \kappa P_j & \text{synonymous transition} \\ \omega P_j & \text{nonsynonymous transversion} \\ \omega \kappa P_j & \text{nonsynonymous transition} \\ 0 & \text{multiple nucleotide changes} \end{cases} , \quad \begin{bmatrix} \textbf{13} \end{bmatrix}$$

for a substitution from codon i to codon j, where  $\kappa$  is the transition-transversion bias,  $P_j$  is the equilibrium frequency of the target codon j, and  $\omega$  represents dN/dS [5, 1]. The  $P_i$  parameters are intended to represent those codon frequencies which would exist in absence of selection pressure generated by mutation alone [5, 6, 69, 7]. Thus, when inferring  $\omega$  on datasets which used symmetric mutation rates, we assigned the value 1/61 to all parameters  $P_i$ , as all codons are equally probable in the absence of mutational bias.

Alternatively, when inferring  $\omega$  for alignments simulated with experimental fitness and mutation rates, we used several different model parameterizations, including GY-style [5] (target codon frequency) and MG-style [6] (target nucleotide frequency) parameterizations. We considered the GY-style parameterizations F61 [5], F3x4 [5], CF3x4 [62], and F1x4 [6]. We implemented two varieties of MG-style models; the first, MG1, employs four parameters for nucleotide frequencies (one per nucleotide) [6], and the second, MG3, employs twelve nucleotide frequency parameters, with four nucleotide frequency parameters for each of the three codon positions [15]. All models included the parameters  $\kappa$  and  $\omega$ .

Availability of code and data. All code used in this project is freely available from

https://github.com/clauswilke/Omega\_MutSel, and simulated alignments are available from dryad???. Alignments were simulated using the pyvolve library, which can be obtained from https://github.com/sjspielman/pyvolve.

#### Appendix 1

We prove that  $dN/dS \leq 1$  when calculated from scaled selection coefficients. We assume that mutation rates are symmetric  $(m_{ij}=m_{ji})$  and that synonymous codons have the same fitness (synonymous changes are neutral). As described in the main text, these assumptions yield dS=1, and hence we have to show that  $dN=K_{\rm N}/L_{\rm N}\leq 1$ . To this end, we note that the sums in  $K_{\rm N}$  and  $L_{\rm N}$  can be reordered such that the substitution probability from codon i to j is always added to the substitution probability from codon j to i. We can then show that the sum of each of these pairs in the expression for  $K_{\rm N}$  is smaller than the corresponding term in  $L_{\rm N}$ , and hence  $dN/dS \leq 1$ .

Without loss of generality, we consider a pair of nonsynonymous codons i and j whose respective stationary frequencies  $P_i$  and  $P_j$  satisfy  $P_i \leq P_j$  and  $P_i, P_j \geq 0$ . As follows from equations [2] and [5], the sum of the probability weights of evolving from codon i to j and from codon j to i is

$$N_e m_{ij} u_{ij} + N_e m_{ji} u_{ji} = \frac{2P_i P_j [\log(P_i) - \log(P_j)]}{P_i - P_j}$$
. [14]

This quantity represents  $K_N$  in the dN calculation. To prove  $dN \leq 1$ , we must show that this quantity is less than or equal to  $P_i + P_j$ , which represents  $L_N$  in the dN calculation. To this end, we introduce the function

$$F(x,y) = x + y - \frac{2xy[\log(x) - \log(y)]}{x - y},$$
 [15]

and we will now show that  $F(x,y) \ge 0$  for  $x \le y$  and  $y \ge 0$ . It is straightforward to show, using l'Hôpital's rule, that this

condition holds for x = y. For x < y, we show that the first derivative of equation [15] is negative throughout  $x \in (0, y)$ , which proves that the function monotonically decreases, and thus F(x, y) > 0, in this interval. We calculate the first derivative as

$$\frac{\partial F(x,y)}{\partial x} = \frac{\left[ (x-3y)(x-y) - 2y^2(\log x - \log y) \right]}{(x-y)^2} \,. \quad [\mathbf{16}]$$

We now replace the expression  $\log x - \log y$  by its Taylor expansion, yielding

$$\frac{\partial F(x,y)}{\partial x} = \frac{\left[ (x-3y)(x-y) - 2y^2 \left( \sum_{n=1}^{\infty} \frac{1}{n} (1-x/y)^n \right) \right]}{(x-y)^2}.$$

We note that the first two terms of the Taylor series equal (x-3y)(x-y), and thus expression [17] simplifies to

$$\frac{\partial F(x,y)}{\partial x} = \frac{-2y^2 \sum_{n=3}^{\infty} \frac{1}{n} \left(1 - \frac{x}{y}\right)^n}{(x-y)^2},$$
 [18]

which is clearly negative. This concludes the proof.

#### Appendix 2

GY-style matrices may be expressed in the framework of the general-time reversible (GTR) model, in which the instantaneous matrix Q can be decomposed into a  $61 \times 61$  symmetric substitution rate matrix and a 61-dimensional vector containing the equilibrium codon frequencies. The latter correspond to the stationary distribution of the Markov chain. By contrast, MG-style rate matrices are written in terms of nucleotide frequencies rather than codon frequencies. Therefore, whether these models fit into the GTR framework is

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unclear *a priori*. We now describe how the MG-style matrix can be rewritten in terms of a symmetric matrix and a vector of equilibrium codon frequencies, thus demonstrating that these matrices also fit into the GTR framework.

MG-style matrix elements, for a the substitution from codon i to j, are generally given by

$$q_{ij} = \begin{cases} \theta_{s_i t_j} \pi_{t_j} & \text{synonymous change} \\ \omega \theta_{s_i t_j} \pi_{t_j} & \text{nonsynonymous change} \\ 0 & \text{multiple nucleotide changes} \end{cases}, \quad [19]$$

where  $\omega$  is the ratio of nonsynonymous to synonymous substitution rates and the product  $\theta_{s_it_j}\pi_{t_j}$  corresponds to a nucleotide-level mutation rate  $\mu_{s_it_j}$ , where  $s_i$  is the source nucleotide in codon i, and  $t_j$  is the target nucleotide in codon j. Note that the matrix  $\theta_{s_it_j}$  is symmetric in  $s_i$  and  $t_j$ .

For a given codon i, the matrix of Eq. [19] yields the stationary frequency  $P_i = \pi_{i_1} \pi_{i_2} \pi_{i_3} C$ , where  $C = 1 - \Pi_{\text{stop}}$  and  $\Pi_{\text{stop}} = \pi_{\text{T}} \pi_{\text{A}} \pi_{\text{G}} + \pi_{\text{T}} \pi_{\text{G}} \pi_{\text{A}} + \pi_{\text{T}} \pi_{\text{A}} \pi_{\text{A}}$  [6]. Therefore, we can rewrite the term  $\theta_{s_i t_j} \pi_{t_j}$  as  $\theta_{s_i t_j} P_j C / (\pi_m \pi_n)$ , where m and n are the nucleotides which do not change in a given instantaneous codon substitution. This allows us to rewrite the rate instantaneous matrix as

$$q_{ij} = \begin{cases} \frac{C\theta_{s_it_j}}{\pi_m\pi_n}P_j & \text{synonymous change from } i \text{ to } j \\ \omega \frac{C\theta_{s_it_j}}{\pi_m\pi_n}P_j & \text{nonsynonymous change from } i \text{ to } j \\ 0 & \text{multiple nucleotide changes} \end{cases}$$

[20]

for a substitution from codon i to codon j, and this matrix clearly conforms to the GTR framework.

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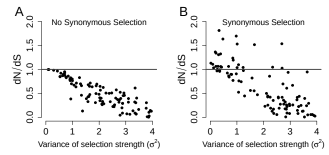
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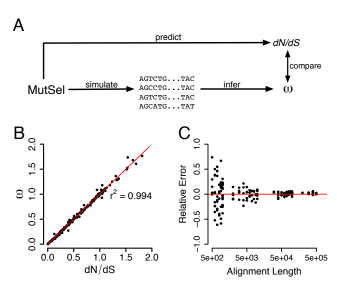
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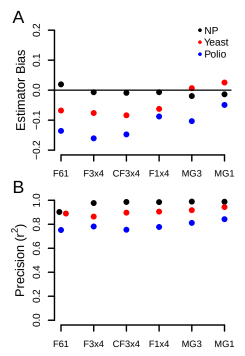
# Figures and Tables



**Fig. 1.** dN/dS decreases in proportion to amino-acid level selection strength. dN/dS is plotted against the variance  $(\sigma^2)$  of the simulated distribution of amino-acid scaled selection coefficients. Higher variances indicate larger fitness differences among amino acids, whereas the limiting value of  $\sigma^2=0$  indicates that all amino acids have the same fitness. (A) Synonymous codons have equal fitness values  $(r^2=0.83)$ . (B) Synonymous codons have different fitness values  $(r^2=0.45)$ . Note that panel B, but not A, shows dN/dS values greater than 1, in spite of the steady-state evolutionary process.



**Fig. 2.** Combined modeling approach to assess performance of dN/dS inference frameworks. (A) Protein-coding alignments are simulated in the MutSel modeling framework. dN/dS can then be calculated ("predict") from scaled selection coefficients as well as through an ML inference framework ("infer"). Comparing resulting quantities reveals the accuracy of the chosen inference framework. (B) Regression between predicted dN/dS values and inferred  $\omega$  MLEs. Each point corresponds to a single simulated alignment, and the red line is the x=y line. (C) Convergence of  $\omega$  MLEs to the true dN/dS value. The y-axis indicates the relative error of the  $\omega$  MLE, and the x-axis indicates the number of positions in the simulated alignment. As the number of positions and hence the size of the data set increases, the  $\omega$  converges to the dN/dS values calculated using equations [6]-[9]. The red line is the y=0 line, indicating no error.



**Fig. 3.** Estimator bias and precision of  $\omega$  estimates for various model frequency parameterizations. (A) Estimator bias and (B) Precision  $(r^2)$  values between dN/dS and  $\omega$  MLEs across model frequency parameterizations, for each set of nucleotide mutation rates. To calculate bias, we fit a linear model with  $\omega$  as the response and dN/dS as the predictor, with a fixed slope of 1, and the resulting intercept value represents the bias. Negative biases indicate  $\omega$  MLEs that are, on average, lower than dN/dS. Note that all standard errors for bias are smaller than the symbol size.

Table 1. Mean  $\Delta {\rm AIC}$  for datasets simulated with NP, yeast, or polio virus mutation rates.

Frequencies	NP	Yeast	Polio	
F61	0	0	0	
CF3x4	-9627.5	-7951.8	-7975.9	
MG1	-13325.5	-10042.0	-5147.6	
F1×4	-13524.5	-13658.5	-15468.3	
MG3	-14401.3	-12851.6	-8624.9	
F3x4	-14807.2	-17385.3	-19384.6	

Note: The order of frequency models shown in the table corresponds to the model ranking for NP, and the ranking differs somewhat for yeast and polio datasets. AIC is computed as  $AIC = 2(k - \ln L)$ , where k is the number of free parameters of the model, and  $\ln L$  is the log-likelihood [64, 65]. As codon and/or nucleotide frequency parameters were computed directly from the data, all models have 3 free parameters ( $\omega$ ,  $\kappa$ , and a global branch-length scaling parameter).

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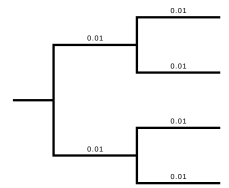


Fig. 4. Phylogeny used for all simulated alignments.

# **Supplementary Information**

Table S1. Estimator bias of  $\omega$  MLEs and the true dN/dS values, for all nucleotide mutation rates and model frequency parameterizations examined. All biases are statistically significant (different from 0), with all  $P < 2 \times 10^{-16}$  except for the estimator bias associated with yeast mutation rates for MG3, where  $P = 5.4 \times 10^{-5}$ .

Mutation rate	MG1	F1×4	MG3	CF3x4	F3x4	F61
NP	-0.014	-0.02	-0.007	-0.009	-0.007	0.019
Yeast	0.025	0.007	-0.063	-0.084	-0.076	-0.068
Polio	-0.049	-0.103	-0.088	-0.148	-0.161	-0.136

Table S2. Precision, measured as the squared correlation coefficient  $r^2$ , of  $\omega$  MLEs relative to the true dN/dS values, for all nucleotide mutation rates and model frequency parameterizations examined. All values shown are statistically significant, with all  $P < 2 \times 10^{-16}$ .

Mutation rate	MG1	F1×4	MG3	CF3×4	F3×4	F61
NP	0.988	0.989	0.985	0.986	0.977	0.902
Yeast	0.943	0.917	0.905	0.897	0.864	0.889
Polio	0.842	0.811	0.777	0.754	0.781	0.752

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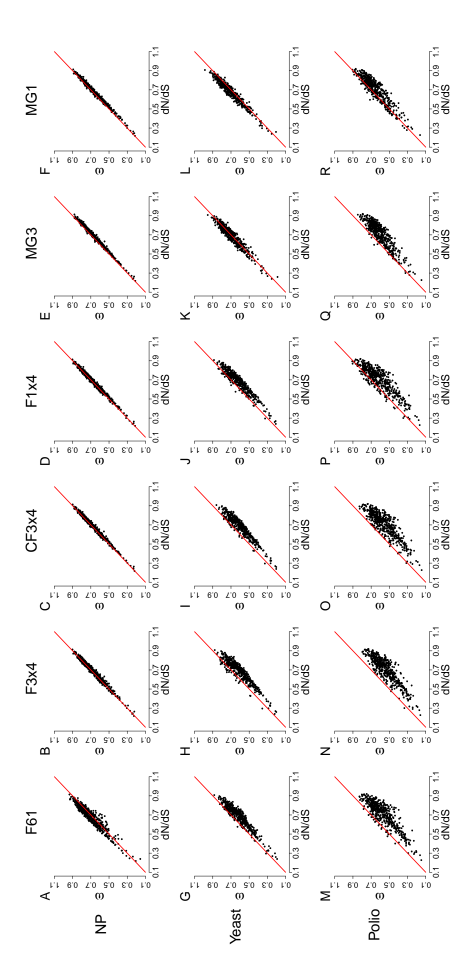


Figure S1. Regressions of  $\omega$  MLEs on the true dN/dS values, as calculated from scaled selection coefficients, for datasets simulated using experimental fitnesses and mutation rates. Each point represents an alignment, and each red line is the x = y line.