One-rate models outperform two-rate models in site-specific $dN/dS$ estimation

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Abstract

Methods that infer site-specific \( dN/dS \), the ratio of nonsynonymous to synonymous substitution rates, from coding data have been developed primarily to identify positively selected sites (\( dN/dS > 1 \)). As a consequence, it is largely unknown how well different inference methods can infer \( dN/dS \) point estimates at individual sites. In particular, \( dN/dS \) may be estimated using either a one-rate approach, where \( dN/dS \) is parameterized as a single parameter, or a two-rate approach, in which \( dN \) and \( dS \) are estimated separately. While some have suggested that the two-rate paradigm may be preferred for positive-selection inference, the relative merits of these two paradigms for site-specific \( dN/dS \) estimation remain largely untested. Here, we systematically assess how accurately several popular inference frameworks infer site-specific \( dN/dS \) values using alignments simulated within a mutation-selection framework rather than within a \( dN/dS \)-based framework. As mutation-selection models describe long-term evolutionary constraints, our simulation approach further allows us to study under what conditions inferred \( dN/dS \) captures the underlying equilibrium evolutionary process. We find that one-rate inference models universally outperform two-rate models. Surprisingly, we recover this result even for data simulated with codon bias (i.e., \( dS \) varies among sites). Therefore, even when extensive \( dS \) variation exists, modeling this variation substantially reduces accuracy. We additionally find that high levels of divergence among sequences, rather than the number of sequences in the alignment, are more critical for obtaining precise point estimates. We conclude that inference methods which model \( dN/dS \) with a single parameter are the preferred choice for estimating reliable site-specific \( dN/dS \) ratios.
Introduction

A variety of computational approaches have been developed to infer selection pressure from protein-coding sequences in a phylogenetically-aware context. Among the most commonly-used methods are those which compute the evolutionary rate ratio $dN/dS$, which represents the ratio of non-synonymous to synonymous substitution rates. Beginning in the mid-1990s, this value has been calculated using maximum-likelihood (ML) approaches (Goldman and Yang 1994; Muse and Gaut 1994), and since then, a wide variety of inference frameworks have been developed to infer $dN/dS$ at individual sites in protein-coding sequences (Nielsen and Yang 1998; Yang et al. 2000; Yang and Nielsen 2002; Yang and Swanson 2002; Kosakovsky Pond and Frost 2005; Kosakovsky Pond and Muse 2005; Murrell et al. 2012b; Lemey et al. 2012; Murrell et al. 2013).

Most commonly, the goal of $dN/dS$ inference is to identify sites subject to positive and/or diversifying selection, as indicated when $dN/dS > 1$. As a consequence, the performances of $dN/dS$ inference methods have largely been evaluated based on how well they detect if a given site evolves with a $dN/dS$ significantly above or below 1. Indeed, many positive-selection inference methods do not make a concerted attempt to calculate precise $dN/dS$ point estimates, but rather focus only on obtaining “good enough” estimates so that the value of $dN/dS$ relative to 1 can be formally tested (Murrell et al. 2012b,a; Scheffler et al. 2014).

By contrast, how accurately such methods estimate $dN/dS$ at individual sites has not been rigorously studied, and therefore it remains unclear which methods, or indeed model parameterizations, provide the most reliable $dN/dS$ point estimates. This dearth of research has hindered advancements of mechanistic studies which seek to understand the relationship between site-specific coding-sequence evolutionary rate and structural properties, such as solvent accessibility, packing density, or flexibility (Shahmoradi et al. 2014; Meyer and Wilke 2015a,b). If site-specific evolutionary rate inference is unreliable, then naturally it will be difficult to ascertain underlying mechanisms driving evolutionary rate.

We therefore seek to assess how well various $dN/dS$ inference frameworks estimate site-wise evolutionary rates from coding sequences. We adopt a robust simulation strategy through which we simulate alignments using the mutation-selection (MutSel) modeling framework. Unlike $dN/dS$ models, MutSel models use population genetics principles to model the site-specific evolutionary process as a dynamic interplay between mutational and selective forces (Halpern and Bruno 1998; Yang and Nielsen 2008). Therefore, many regard MutSel models as more mechanistically representative of real coding sequence evolution than $dN/dS$-based models, which are primarily phenomenological in nature (Thorne et al. 2007; Holder et al. 2008; Rodrigue et al. 2010; Thorne et al. 2012; Tamuri et al. 2012; Liberles et al. 2013). Indeed, substitution rate itself is not an evolutionary mechanism, but rather an emergent property of various interacting evolutionary processes.

Recently, we introduced a mathematical framework which allows us to accurately calculate a $dN/dS$ ratio directly from the parameters of a MutSel model (Spielman and Wilke 2015b) [we note that dos Reis (2015) introduced a similar framework shortly after]. This framework gives rise to a robust benchmarking strategy through which we can simulate sequences using a MutSel model, and we can subsequently infer $dN/dS$ using established approaches. Previously, we successfully used such an approach to identify biases in $dN/dS$ inference methods for whole-gene evolutionary rates (Spielman and Wilke 2015b). Here, we leverage the power of the established relationship between $dN/dS$ and MutSel models to evaluate the performance of site-specific $dN/dS$ inference approaches.

Two primary questions motivate the present study: i) How accurate are various inference methods for $dN/dS$ point estimation?, and ii) Under what conditions does $dN/dS$ capture the long-term evolutionary dynamics of site-specific coding-sequence evolution? For the first question, we focus
our efforts on distinguishing performance between two \(dN/dS\) inference paradigms: one-rate and two-rate models. One-rate models parameterize \(dN/dS\) with a single parameter for \(dN\), effectively fixing \(dS = 1\) at all sites, whereas two-rate models use separate parameters for \(dN\) and \(dS\) at each site. Some studies have suggested that the two-rate paradigm leads to more robust positive-selection inference (Kosakovsky Pond and Muse 2005; Murrell et al. 2013), whereas others have suggested that the extra \(dS\) parameter may actually confound positive selection inference (Yang et al. 2005; Wolf et al. 2009). Therefore, it remains unclear how the one-rate vs. two-rate parameterization choice influences positive-selection inferences, and consequently it is an open question how this parameterization affects \(dN/dS\) point estimation.

The second question arises naturally from our use of MutSel models, which describe the equilibrium site-specific codon fitness values. In other words, any \(dN/dS\) calculated from MutSel model parameters describes, by definition, the steady-state \(dN/dS\). As \(dN/dS\) is an inherently time-sensitive measurement (Rocha et al. 2006; Kryazhimskiy and Plotkin 2008; Mugal et al. 2014; Meyer et al. 2015), it is not necessarily true that \(dN/dS\) measured from a given dataset will reflect the equilibrium value. Therefore, our approach additionally enables us to identify the conditions under which site-specific \(dN/dS\) ratios are expected to reflect the long-term, rather than transient, evolutionary dynamics.

Results

Approach

We simulated fully heterogeneous alignments under the HB98 MutSel model (Halpern and Bruno 1998) using the simulation software Pyvolve (Spielman and Wilke 2015a). To derive site-specific MutSel model parameterizations, we simulated 100 distinct sets of amino-acid frequencies from a Boltzmann distribution (Ramsey et al. 2011), reflecting the shape of empirical amino-acid distributions observed in conserved protein sequences (see Methods for details). We ensured that these simulated distributions resulted in a range of selective pressure, from extremely stringent to nearly neutral. From each amino-acid frequency distribution, we derived two distinct stationary codon frequency distributions: one where all synonymous codons had the same fitness (i.e. no codon bias), and one where synonymous codons differed in fitness values (i.e. codon bias).

Using these derived MutSel model parameterizations, we simulated a set of fully heterogeneous alignments, with 100 sites, each for set of codon fitnesses. All simulations were conducted along balanced phylogenies with the number of sequences \(N\) set as either 128, 256, 512, 1024, or 2048 and with branch lengths \(B\) set as either 0.0025, 0.01, 0.04, 0.16, or 0.64. For each of the 25 possible combination of parameters \(N\) and \(B\), we simulated 50 replicate alignments. Importantly, the site-specific evolutionary models were the same within each simulation set, making inferences across conditions directly comparable.

We inferred site-specific \(dN/dS\) for each simulated alignment using three approaches: fixed-effects likelihood (FEL) (Kosakovsky Pond and Frost 2005), single-likelihood ancestor counting (SLAC) (Kosakovsky Pond and Frost 2005), and FUBAR (Murrell et al. 2013). Each of these methods employs a somewhat different approach when computing site-specific \(dN/dS\) values. FEL fits a unique \(dN/dS\) model to each alignment site (Kosakovsky Pond and Frost 2005), SLAC directly counts nonsynonymous and synonymous changes along the phylogeny where ancestral states are inferred with maximum likelihood (Kosakovsky Pond and Frost 2005), and FUBAR employs a Bayesian approach to determine \(dN/dS\) ratios according to a pre-specified grid of rates (Murrell et al. 2013).

For each inference method, we inferred \(dN/dS\) at each site in both a two-rate context (separate
dN and dS parameters per site) and in a one-rate context (a single dN/dS parameter per site). Although SLAC, as a counting-based method, always enumerates both dN and dS on a per-site basis, one can derive an effectively one-rate SLAC by normalizing each site-wise dN estimate by the mean of all site-wise dS estimates. We refer to one-rate inferences with these methods as FEL1, FUBAR1, and SLAC1, and similarly to two-rate inferences as FEL2, FUBAR2, and SLAC2, respectively. All inferences were conducted using the HyPhy batch language (Kosakovsky Pond et al. 2005), specifying the MG94xHKY85 model with F1x4 state frequencies. Note that we did not consider the popular random-effects likelihood methods introduced by Yang et al. (2000) (e.g. M3, M5, M8) because these methods are used predominantly in a one-rate context. Available two-rate extensions to this framework are computationally burdensome and cannot model the amount of rate heterogeneity required to calculate per-site rates (Kosakovsky Pond and Muse 2005). Finally, we computed true dN/dS values from the MutSel parameters, using the approach described in Spielman and Wilke (2015b).

Modeling synonymous rate variation reduces inference accuracy

After inferring site-wise dN/dS for all simulated alignments, we correlated the resulting estimates with true dN/dS values. In Figure 1, we show resulting Pearson correlation coefficients, averaged across all 50 replicates, between inferred and true dN/dS for each inference method. Importantly, our simulation strategy necessitates a somewhat different interpretation of results than would more traditional simulation approaches. In particular, the true dN/dS ratios calculated from the MutSel parameterizations used during simulation correspond to the dN/dS expected at steady state, which in turn indicates the signature of natural selection at evolutionary equilibrium. We can only expect to recover this true dN/dS value if the simulated data reflect the full steady-state distribution of codons. When either the simulated divergence or number of sequences analyzed is low, then, it not necessarily possible to capture this distribution. Therefore, to determine the relative performance of dN/dS inference methods, we considered the most accurate inference method to be the one with the highest dN/dS correlations within a given choice of N and B.

In the absence of codon bias, dS was equal to 1 at all sites. As such, we expected that one-rate inference methods would outperform two-rate inference methods. We indeed found that one-rate inference models showed the best performance when there was no synonymous selection (Figure 1A), in particular at low-to-intermediate divergence levels (B of 0.01 or 0.04). As the sequences became more diverged, and hence more informative, two-rate models increasingly performed as well as one-rate models did. Even so, two-rate models never outperformed one-rate models.

In the presence of codon bias, both dN and dS varied at each site. As a consequence, there are two approaches for calculating the true site-wise dN/dS ratio: One can either calculate the ratio of each site’s dN and dS values, or one can take each site’s dN value and divide by the average dS over the entire sequence. The former corresponds to a two-rate model (there are two independent rates at each site), while the latter corresponds to a one-rate model (only dN varies per site, and dS is taken as a gene-wide normalization factor). Here, we refer to these two dN/dS ratios as True2 and True1, respectively.

We correlated, for data simulated with codon bias, inferred dN/dS with both True2 and True1 dN/dS values, as shown in Figures 1B and 1C, respectively. A priori, we would expect that two-rate inference models would perform best when benchmarked against True2, and similarly one-rate inference models would perform best when benchmarked against True1. Surprisingly, however, one-rate models outperformed two-rate models across N and B conditions, regardless of whether True2 or True1 was considered. We did, however, find that two-rate models yielded higher correlations with True2 than with True1, and vice versa.
Importantly, although two-rate models appear to have outperformed one-rates models when \( B = 0.0025 \) (Figure 1), nearly all such inferences were poor; correlations from two-rate model virtually never exceeded an average of 0.4. In other words, at low divergence levels, inferred \( dN/dS \) could explain at most only \( \sim 16\% \) of the rate variation expected at equilibrium, likely indicating that, at \( B = 0.0025 \), the data was mostly uninformative. Similarly, all estimates, from both one- and two-rate models, were strongly biased at \( B = 0.0025 \) (Figure S1). As divergence increased, and hence the data became more informative, estimator bias dropped substantially for both one- and two-rate models. However, at \( B = 0.04 \), one-rate models had virtually no estimator bias, but two-rate models still strongly overestimated \( dN/dS \), indicating that two-rate models were more biased than were one-rate models.

Together, these results demonstrate, for both data with and without codon bias, that one-rate models inferred more accurate \( dN/dS \) ratios, on average, relative to two-rate models. For data simulated with codon bias, this result was robust to whether inferences were benchmarked against True1 or True2. Modeling synonymous selection with its own parameter reduced inference accuracy, especially when the data contained pervasive codon bias. Indeed, the accuracy boost achieved with one-rate inference models was far more pronounced for data with codon bias than for data without codon bias. In addition, correlations between inferred and true \( dN/dS \) were, on average, higher for data simulated without codon bias (Figure 1A) compared to data simulated with codon bias (Figures 1B and 1C). Our results therefore revealed that \( dN/dS \) inference methods were generally more reliable in the absence of codon bias.

**One-rate inference methods have minimal performance differences**

We next quantified performance differences among methods more rigorously using linear models. For each simulation set, we built mixed-effects linear models with Pearson correlation as the response, inference method as a fixed effect, and replicate as well as interaction between \( N \) and \( B \) as random effects. We performed multiple comparisons tests, with corrected P-values, to ascertain the relative performance across methods. In this analysis, we additionally tested the performance of derived one-rate inferences made with FEL2 and FUBAR2, which we called FEL2\(_1\) and FUBAR2\(_1\), respectively. These inferences represent rates calculated by normalizing site-specific \( dN \) values by the average inferred site-specific \( dS \) value, similar to how SLAC1 values were computed.

Linear model analysis confirmed observations from Figure 1 that each one-rate inference framework outperformed its respective two-rate counterpart (Figure 2). Further, FEL2\(_1\) and FUBAR2\(_1\) did not display significant performances differences from FEL1 and FUBAR1, respectively, indicating that fixing \( dS \) to 1 is essentially equivalent to normalizing all \( dN \) by an inferred average \( dS \). Across panels in Figure 2, both SLAC1 and FEL1 generally outperformed FUBAR1, with SLAC1 tending to be the most accurate method. Importantly, even when performance differences were statistically significant, the effect magnitudes were exceedingly small; mean correlations never differed by more than 0.025. Thus, whether \( dN/dS \) was modeled by one or two parameters mattered more than the specific inference method used (e.g. FEL1, SLAC1, FUBAR1) did for obtaining accurate estimates, although SLAC1 and FEL1 may be somewhat preferable to FUBAR1.

**Estimation error is higher for lower \( dN/dS \) values**

We next examined whether certain \( dN/dS \) values were more difficult to estimate. Our simulation setup ensured an evenly-spaced range of true \( dN/dS \) values, from 0.03–0.92 for simulations without codon bias and, for True2, 0.05–0.99 for simulations with codon bias. We calculated, for each simulated \( dN/dS \) value, the average relative error, across replicates, for each \( N \) and \( B \) parametr-
terization, from SLAC1 inference (Figure 3). As seen in each panel of Figure 3, error declined as \(dN/dS\) increases, indicating that it was more difficult to precisely estimate \(dN/dS\) at slowly evolving sites. Importantly, we observed this trend across \(N\) and \(B\) conditions, although the overall error decreased as datasets became more informative. Figures S2, S3, and S4 shows results for all simulation conditions and display broadly the same trends as seen in Figure 3.

We suggest that lower \(dN/dS\) values were more difficult to estimate because natural selection tolerates fewer codons at slowly-evolving sites, and there are often relatively large fitness differences among the codons that are tolerated (Spielman and Wilke 2015b). Therefore, it was less likely that slowly evolving sites reflected the full MutSel steady-state distribution of codons, compared to quickly evolving sites, which ultimately incurred higher estimation error.

Divergence is more important than is the number of sequences for identifying long-term evolutionary constraint

We observed that correlations between true and inferred \(dN/dS\) values increased both as the number of sequences \(N\) and the branch lengths \(B\) (divergence) grew (Figure 1), suggesting that large and/or highly informative datasets are necessary for the inferred \(dN/dS\) to capture the actions of natural selection at evolutionary equilibrium. However, it was not immediately clear from Figure 1 whether \(N\), \(B\), or some combination of these conditions drove this trend. Therefore, we next assessed the relative importance of \(N\) and \(B\).

We calculated the tree length (expected number of substitutions per site across the entire tree) for each \(N\) and \(B\) parameterization. If \(N\) and \(B\) served roughly equal roles in terms of providing information, then any combination of \(N\) and \(B\) corresponding to the same tree length should have produced similar \(dN/dS\) correlations. We did not, however, observe this trend; instead, all else being equal, \(B\) had a significantly greater influence than did \(N\) on resulting correlations. For example, as shown in Figure 4, we compared \(dN/dS\) correlations from SLAC1 for three combinations of \(N\) and \(B\) conditions which all had virtually the same tree lengths (162–164). Simulations with lower \(N\) and higher \(B\) yielded far more accurate \(dN/dS\) estimates, even though all simulations in Figure 4 experienced the same average number of substitutions. This increase was highly significant; for data simulated without codon bias, correlations increased an average \(\sim 20\%\) from \(B = 0.04\) to \(B = 0.64\) (\(P < 10^{-15}\)). As shown, neither codon bias nor the manner of true \(dN/dS\) calculation influenced this overarching trend (\(P > 0.17\)), although correlations between true and inferred \(dN/dS\) were generally lower when codon bias was present.

Discussion

In this study, we have examined the accuracy of different site-specific \(dN/dS\) inference approaches in the context of \(dN/dS\) point estimation. In particular, we have assessed performance differences between two \(dN/dS\) model parameterization paradigms: one-rate, where \(dN/dS\) is modeled with a single parameter, and two-rate, where \(dN\) and \(dS\) are modeled with separate parameters. We have found that one-rate inference models virtually always produce more accurate \(dN/dS\) inferences than do two-rate models. Strikingly, the presence of codon bias does not influence this result. In fact, the increased accuracy of one-rate compared to two-rate models is even more pronounced when codon bias is present (Figures 1 and 2). Therefore, our findings suggest that, even in the presence of synonymous selection, site-specific evolutionary rates should be measured using methods which estimate only \(dN\) and implicitly fix \(dS = 1\) or consider a global \(dS\) for the entire sequence. We did not, however, examine how one- and two-rate inference models compare when \(dS\) variation is driven by mutational rather than selective processes.
For this study, we simulated fully heterogeneous sequences with each site evolving according to a unique MutSel model. While MutSel models have shortcomings (e.g., they assume constant site-specific fitness values across the phylogeny), they take a far more mechanistic approach to coding-sequence evolution than dN/dS-based models do, and they have therefore been regarded as more evolutionarily realistic. A key benefit of simulating with the MutSel framework is that we are able to directly model synonymous rate variation by specifying different fitnesses for synonymous codons, instead of relying on a phenomenological rate parameter $dS$. We note that this simulation setup, however, cannot test performance accuracy on positively-selected sites ($dN/dS > 1$), as MutSel models can only correspond to sites under either purifying selection or neutral evolution ($dN/dS \leq 1$) (Spielman and Wilke 2015b). As such, we emphasize that our results here apply specifically to the question of site-specific $dN/dS$ point estimation, and not to the question of positive-selection inference. Future work may be needed to fully understand how one-rate vs. two-rate models compare for positive-selection inference.

We demonstrate that, in the context of $dN/dS$ point estimation, two-rate methods do not properly accomplish their intended goal of accounting for the effects of selection pressure on synonymous codons. Logically, one would presume that, when $dS$ differs among sites, estimating $dS$ separately across sites would produce more accurate $dN/dS$ estimates than would fixing $dS$ to a constant value. Indeed, an assumed presence of synonymous substitution rate variation is the very justification for using a two-rate $dN/dS$ model (Kosakovsky Pond and Muse 2005). However, including this additional parameter hindered accuracy under virtually all simulation conditions, and we therefore conclude that including a $dS$ parameter is not an effective way to model the presence of synonymous selection, at least on a per-site basis.

We suggest that error in $dS$ estimates may explain the relatively poor performance of two-rate models, particularly on datasets which had codon bias. Indeed, the reason that the $dN/dS$ ratio includes the $dS$ denominator is to have a suitable normalization to $dN$ that provides a baseline, neutral substitution rate (i.e., mutation rate). Statistically, the most robust way to obtain this baseline rate, if we assume that mutation rates are mostly constant across sites, is to compute an average $dS$ across all sites (as in SLAC1, FEL2.1, and FUBAR2.1). Fixing $dS$ to 1, as FUBAR1 and FEL1 do, yields $dN/dS$ values that are essentially equivalent to those returned by this procedure (Figure 2). Site-specific rate inferences necessarily have high levels of noise, due to dataset size limitations, and thus estimating a separate $dS$ at each site likely contributes substantial noise and ultimately reduces estimate reliability. One-rate methods avoid this statistical problem, and they do not appear to suffer dramatically when codon bias was present.

We additionally have found that high levels of sequence divergence are critically important for obtaining a reliable steady-state $dN/dS$ value, moreso than the number of sequences analyzed (Figure 4). This finding has important implications for data set collection: It may be preferable to include fewer, more divergent sequences rather than as many sequences as one can obtain. Measuring $dN/dS$ from thousands of sequences with low divergence may actually be less effective than analyzing fewer, more diverged sequences, even if the mean number of per-site substitutions would be the same. Increasing the number of taxa in a given analysis may only be beneficial if the new sequences are substantially diverged from the existing sequences. We emphasize that the number of taxa should still be sufficiently large ($\geq 100$) to achieve reliable estimates, due to the inherent high level of noise in site-specific inferences.

These findings additionally build on the well-documented time-dependency of the $dN/dS$ metric, a phenomenon studied largely in the context of polymorphic data (Rocha et al. 2006; Kryazhimskiy and Plotkin 2008; Wolf et al. 2009; Mugal et al. 2014; Meyer et al. 2015). Our results extend these findings, indicating that this time-dependency is more general and pertains also to circumstances where the data contain only fixed differences. This finding makes intuitive sense: As divergence
increases, sites will be more likely to visit the full range of selectively tolerated states, and therefore the long-term evolutionary constraints will become apparent. Importantly, even at exceptionally high divergence levels, inferred \( \frac{dN}{dS} \) estimates could never fully recapitulate the \( \frac{dN}{dS} \) that describes the steady-state distribution (Figure 1). For instance, the inferred \( \frac{dN}{dS} \) values for simulated alignment with the most divergence (\( N = 2048 \) and \( B = 0.64 \)) had a correlation coefficient of 0.93 with the true \( \frac{dN}{dS} \) values, thereby explaining only 86% of the variation expected at evolutionary equilibrium. Such an empirical dataset would be difficult, if even possible, to obtain, and therefore we may not be able to recover the equilibrium \( \frac{dN}{dS} \) value from empirical data. Instead, it is most likely that all \( \frac{dN}{dS} \) measurements will be biased by time to some degree, even if all differences are fixed and not polymorphic.

Finally, our study has important implications for research that seeks to relate site-specific \( \frac{dN}{dS} \) ratios to protein structural properties, such as relative-solvent accessibility or weighted contact number (Spielman and Wilke 2013; Meyer and Wilke 2013; Meyer et al. 2013; Shahmoradi et al. 2014; Meyer and Wilke 2015a,b). These metrics reflect the overarching biophysical constraints that influence protein evolutionary trajectories. Studies which have examined the correlations between site-specific \( \frac{dN}{dS} \) and such structural quantities have recovered relatively low, although significant, correlations, generally ranging from 0.1–0.6 (Shahmoradi et al. 2014; Meyer and Wilke 2015a,b). Importantly, these studies analyzed viral sequence data, which contained relatively low levels of divergence. By contrast, other studies which considered more substantially diverged enzyme proteins (albeit using protein-sequence–derived evolutionary rates instead of \( \frac{dN}{dS} \)) recovered higher correlations, ranging mostly from 0.3–0.8 (Shih and Hwang 2012; Huang et al. 2014; Yeh et al. 2014b,a), between structural measures and evolutionary rate.

Our results suggest that this discrepancy is, in fact, not unexpected, and moreover that low structure–rate correlations recovered from highly similar viral sequence data are not worrisome. Just like structural quantities do, MutSel models describe the long-term evolutionary constraints acting at specific coding-sequence positions. Indeed, for data with minimal divergence, we did not recover particularly large correlations between inferred and true \( \frac{dN}{dS} \) (Figure 1). For example, the average correlation for simulations without codon bias for \( N = 512 \) and \( B = 0.0025 \), typical values for a virus sequence alignment, was \( r = 0.414 \) (inferred with SLAC1). This correlation falls well within the range of observed structure-rate correlations in empirical viral datasets. Likewise, correlations between true and inferred \( \frac{dN}{dS} \) were higher for more diverged data, just as observed in aforementioned studies of the structure-rate relationship in enzymes. Therefore, we suggest that future work examining the relationship between protein evolutionary rate and structure should focus on obtaining highly diverged datasets, which are more likely to provide meaningful information about long-term evolutionary constraints.

**Methods**

**Alignment simulation**

We simulated heterogeneous alignments, such that each site evolved according to a distinct distribution of codon state frequencies, according to the HB98 MutSel model (Halpern and Bruno 1998) using Pyvolve (Spielman and Wilke 2015a). We began by deriving site-specific MutSel model parameterizations. We simulated 100 site-specific amino-acid frequency distributions from a Boltzmann distribution:

\[
F(a) = \frac{\exp(-\lambda_a)}{\sum_b \exp(-\lambda_b)},
\]

(1)
where $F(a)$ is the state frequency of amino-acid $a$, $a$ and $b$ index amino acids from 0–19, and the parameter $\lambda$ increases with evolutionary rate (Ramsey et al. 2011). For each frequency distribution, we sampled a value for $\lambda$ from a uniform distribution $U(0,3)$, and we selected a random fitness ranking for all amino acids.

Once amino acid frequencies were computed, we assigned frequencies to codons in two distinct ways: without selection for codon bias, and with selection for codon bias. To generate frequency distributions without codon bias, we assigned all synonymous codons the same frequency (summing to the corresponding amino-acid frequency). Alternatively, to generate frequency distributions with codon bias, we randomly selected a preferred codon for each amino acid. We assigned a state frequency of $\gamma F(a)$, where $\gamma$ was drawn from a uniform distribution $U(0.6,0.9)$, to the preferred codon, and we assigned the remaining frequency $F(a) - \gamma F(a)$ evenly to all remaining synonymous codons. In this way, the overall amino-acid state frequency was unchanged, but the synonymous codons occurred with differing frequencies.

Using these site-specific MutSel model parameterizations, we simulated two sets of heterogenous alignments, one without and one with codon bias. All sites evolved according to the HKY85 (Hasegawa et al. 1985) mutation model, with $\kappa = 4$. We simulated heterogenous alignments across an array of balanced phylogenies, containing either 128, 256, 512, 1024, or 2048 sequences. For each number of taxa, we simulated sequences with varying degrees of divergence, with all branch lengths equal to either 0.0025, 0.01, 0.04, 0.16, or 0.64. We simulated 50 alignment replicates for each combination of these conditions.

**dN/dS inference**

For each simulated codon frequency distribution, we computed $dN/dS$ according to the method outlined in Spielman and Wilke (2015b). We calculated $dN/dS$ using this method in both a two-rate manner, in which $dN$ and $dS$ were calculated individually for each site and divided to obtain $dN/dS$, and in a one-rate manner, in which each site-specific $dN$ is normalized by the mean $dS$ across all sites.

For each simulated alignment, we inferred site-specific $dN/dS$ values with the HyPhy software (Kosakovsky Pond et al. 2005) using several approaches: fixed-effects likelihood (FEL) (Kosakovsky Pond and Frost 2005), FUBAR (Murrell et al. 2013), and single ancestral counting (SLAC) (Kosakovsky Pond and Frost 2005). For all methods used, we specified the MG94xHKY85 (Muse and Gaut 1994; Kosakovsky Pond and Frost 2005) rate matrix with F1x4 codon frequencies, which has been shown to reduce bias in $dN/dS$ estimation (Spielman and Wilke 2015b). We provide customized HyPhy batchfiles which enforce the F1x4 codon frequency specification in the github repository https://github.com/sjspielman/sitewise_dnds_mutsel.

For both FEL and FUBAR, we inferred $dN/dS$ with both a one-rate model, in which $dN/dS$ is represented by a single parameter and a two-rate model, in which $dN$ and $dS$ are modeled by separate parameters (Kosakovsky Pond and Frost 2005). For the one-rate FUBAR inferences, we specified 100 grid points to account for the reduced grid dimensionality caused by ignoring $dS$ variation [as in Spielman et al. (2014)], and for the two-rate FUBAR inferences, we specified the default 20x20 grid (Murrell et al. 2013). All other settings were left as their default values. Similarly, for SLAC inference, we calculated $dN/dS$ in two ways. As SLAC enumerates $dN$ and $dS$ on a site-specific basis, there exist two ways to calculate site-wise $dN/dS$: $dS$ can be considered site-specific, or $dS$ values can be globally averaged, and this mean can be used to normalize all site-specific $dN$ estimates. The former calculations effectively correspond to a two-rate method (SLAC2), and the latter calculations correspond to a one-rate method (SLAC1). All inferences were conducted using the true tree along which each alignment was simulated.
As in Kosakovsky Pond and Frost (2005), we excluded all unreliable \( dN/dS \) inferences when correlating inferred and true \( dN/dS \) values. Specifically, estimates made by FEL were excluded if the estimated \( dN/dS \) equaled 1 and the \( P \)-value indicating whether the estimate differed significantly from 1 was equal to 1. Such measurements have been shown to indicate uninformative sites (Meyer et al. 2015). In addition, estimates made by SLAC2 were excluded if the number of synonymous mutations counted was 0, and hence the resulting \( dN/dS \) was undefined. Finally, all inferences with \( dN/dS \geq 100 \) were considered uninformative, as these high estimates likely reflect estimation error.

### Data analysis and availability

Statistics were conducted in the R statistical programming language. Linear modeling was conducted using the R package lme4 (Bates et al. 2012). We inferred effect magnitudes and significance, which we corrected for multiple testing, using `glht()` function in the R package multcomp (Hothorn et al. 2008). In particular, each mixed-effects model described in the Results subsection Modeling synonymous rate reduces inference accuracy was built in the lme4 package with the general code

\[
\text{lmer}(r \sim \text{method} + (1|\text{replicate}) + (1|N:B))
\]

where \( r \) is the Pearson correlation between inferred and true \( dN/dS \). Relative error between inferred and true \( dN/dS \) values was calculated, for each condition, as 

\[
\frac{|dN/dS_{\text{inf}} - dN/dS_{\text{true}}|}{dN/dS_{\text{true}}},
\]

where \( dN/dS_{\text{inf}} \) indicates the average inferred \( dN/dS \) and \( dN/dS_{\text{true}} \) indicates the true \( dN/dS \).

All code and simulated data are freely available from


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References

Bates D, Maechler M, Bolker B. 2012. lme4: Linear mixed-effects models using S4 classes. R package version 0.999999-0.


Figures

Figure 1: Pearson correlation coefficients between true and inferred $dN/dS$ across inference approaches and $N$-$B$ conditions. A) Correlations for alignments simulated without codon bias. B) Correlations with True2 for alignments simulated with codon bias. C) Correlations with True1 for alignments simulated with codon bias. The label above each sub-plot indicates the branch lengths $B$ of the balanced phylogeny along which sequences were simulated, and the x-axes indicate the number of sequences $N$. Each point represents the correlation coefficient averaged across 50 replicates.
Figure 2: Pairwise comparisons of correlation strength across methods, as determined through multiple comparisons tests. A) Results for data simulated without codon bias. B) Results for data simulated with codon bias, as correlated with True2. C) Results for data simulated with codon bias, as correlated with True1. Points indicate the estimated average difference between correlations for the respective methods, and lines indicate 95% confidence intervals. Black lines indicate that the performance difference between methods differed significantly from 0 (all $P < 0.01$). Gray lines indicate that the difference was not statically significant ($P > 0.01$).

Figure 3: Average relative error of inferred $dN/dS$ values by SLAC1 for a subset of $N$ and $B$ conditions. Each point represents the relative error averaged across 50 replicates. Labels above each column indicate the number of sequences $N$, and labels to the right of each row indicate the branch lengths $B$. Results are shown here for data simulated without codon bias. The horizontal line indicates an average relative error of 0, and the diagonal line is the regression line whose slope is indicated in each panel. All slopes shown are significant at $P < 10^{-15}$. 
Figure 4: The amount of divergence is more important than the number of sequences is for obtaining the equilibrium $dN/dS$ value. Each boxplot represents correlation coefficients, from SLAC1 inference, across the 50 respective replicates. From left to right, tree lengths are equal to 163.76, 163.52, and 162.56. Although mean number of per-site substitutions was therefore virtually equal among the conditions shown, higher divergence among sequences led to significantly higher accuracy than did a larger number of sequences.
Supplementary Figures

Figure S1. Estimator bias of inferred $dN/dS$ relative to true $dN/dS$. A) Estimator bias for alignments simulated without codon bias. B) Estimator bias for alignments simulated with codon bias, using True2 as a reference. C) Estimator bias for alignments simulated with codon bias, using True1 as a reference. Each point represents the correlation coefficient averaged across 50 replicates. The label above each sub-plot indicates the branch lengths $B$ of the balanced phylogeny along which sequences evolved, and the x-axes indicate the number of sequences $N$. Points not shown exist off the scale.
Figure S2. Average relative error of inferred $dN/dS$ values by SLAC1 for simulations without codon bias. Each point represents the relative error average across 50 replicates. Labels above each column indicate the number of sequences $N$, and labels to the right of each row indicate the branch lengths $B$. 
Figure S3. Average relative error of inferred $dN/dS$ values with True2 by SLAC1 for simulations with codon bias. Each point represents the relative error average across 50 replicates. Labels above each column indicate the number of sequences $N$, and labels to the right of each row indicate the branch lengths $B$. The horizontal line indicates an average relative error of 0, and the diagonal line is the regression line.
Figure S4. Average relative error of inferred $dN/dS$ values with True1 by SLAC1 for simulations with codon bias. Each point represents the relative error average across 50 replicates. Labels above each column indicate the number of sequences $N$, and labels to the right of each row indicate the branch lengths $B$. The horizontal line indicates an average relative error of 0, and the diagonal line is the regression line.