Intermediate divergence levels maximize the strength of structure-sequence correlations in enzymes and viral proteins

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Figure S1: Comparison of structure—rate correlations for the full data set of enzymes and the design set.

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Figure S2: Distribution of R² for linear models of structural predictors of evolutionary rate (ER) in enzymes.

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Figure S3: Distribution of R² for linear models of structural predictors of ER in viruses.

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Figure S4: Comparison of structure-rate correlations with Mean RSA.

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Figure S5: Comparison of structure–rate correlations with Mean WCN.

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Figure S6: Comparison of structure-rate correlations with divergence.

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Filename S7: Comparison of structure–rate correlations with mean Guidance scores of proteins.

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Structural properties such as solvent accessibility and contact number predict sitespecific sequence variability in many proteins. However, the strength and significance of these structure-sequence relationships vary widely among different proteins, with absolute correlation strengths ranging from 0 to 0.8. In particular, two recent works have made contradictory observations. Yeh et al. [Mol. Biol. Evol. 31:135–139, 2014] found that both relative solvent accessibility (RSA) and weighted contact number (WCN) are good predictors of site-wise evolutionary rate in enzymes, with WCN clearly outperforming RSA. Shahmoradi et al. [J. Mol. Evol. 79:130–142, 2014] considered these same predictors (as well as others) in viral proteins and found much weaker correlations and no clear advantage of WCN over RSA. Because these two studies had substantial methodological differences, however, a direct comparison of their results is not possible. Here, we re-analyze the data sets of the two studies with one uniform analysis pipeline, and we find that many apparent discrepancies between the two analyses can be attributed to the extent of sequence divergence in individual alignments. Specifically, the alignments of the enzyme data set are much more diverged than those of the virus data set, and proteins with higher divergence exhibit, on average, stronger structuresequence correlations. However, the highest structure-sequence correlations are observed at intermediate divergence levels, where both highly conserved and highly variable sites are present in the same alignment.

Keywords: protein evolution, protein design, relative solvent accessibility, site variability, packing density

Words (up to 75): 75

There have been numerous previous studies on the effect of structure on sequence evolution. Local packing density and solvent accessibility have emerged as the two best structural predictors of site-specific evolutionary rate (ER). Here we find that the strength of structure—rate correlations is strongly affected by the divergence of the multiple—sequence alignments (MSAs) used for calculating evolutionary rates. Proteins with MSAs that have intermediate levels of divergence have the strongest structure—rate correlations.

Accepted

Introduction

Proteins are subject to a number of biophysical and functional constraints that influence their evolutionary trajectories.^{1–4} These constraints contribute to observed patterns in both whole-gene evolutionary rate variation^{5–9} and evolutionary rate variation among sites within individual proteins.^{10–14} Such evolutionary rate variation in turn contributes to heterogeneity in site-specific sequence variability.

A number of studies have sought to understand the roles that biophysical constraints, particularly structural constraints, play in this observed site-specific variability within proteins. Structural properties such as solvent exposure and packing density have emerged as strong predictors of site-wise evolutionary rates. 11,13,15,16

Solvent exposure is typically measured with the metric relative solvent accessibility (RSA), which indicates the extent to which a given residue comes into contact with solvent (i.e., water). Residues that are exposed on the surface of the protein have high RSA, with complete exposure indicated with an RSA of one. Residues that are buried and/or in the protein core have low RSA, with completely buried residues having an RSA of zero. RSA has a significant, positive relationship with evolutionary rate, such that more buried residues tend to evolve more slowly than exposed residues do. 10,16,18-23

Alternatively, packing density indicates how tightly packed a given residue is by neighboring amino acids in a protein's tertiary structure. A residue's packing density is commonly measured as weighted contact number (WCN), which is defined as the sum of the inverse square distance of all residues in the protein to the focal amino acid.^{24,25} Recent work has suggested that WCN is a strong determinant of site-specific variability

in proteins, and that residues with high WCN evolve more slowly than do residues with low WCN.^{8,11,12,15}

However, some studies have yielded apparently contradictory results regarding the extent of the predictive power that these structural properties have on site-wise evolutionary rate (ER). For example, Yeh *et al.*¹¹ investigated structure–sequence relationships in a data set of 216 monomeric enzymes, finding that WCN is a stronger determinant of site-wise ER than RSA, although RSA was still a significant predictor. Importantly, Yeh *et al.*¹¹ recovered strong correlations between structure and ER, with WCN and RSA explaining up to ~ 41% of the variance in site-specific ER. By contrast, Shahmoradi *et al.*¹³ examined the structure–sequence relationship on a set of 9 viral proteins. While Shahmoradi *et al.*¹³ similarly found that both RSA and WCN are significant predictors of rate in proteins, the correlations Shahmoradi *et al.*¹³ observed were much smaller in magnitude. ¹³ Specifically, they found that at best, structural predictors could explain only ~15% of the variance in ER. Given these disparate findings, it remains unclear which of the two studies is the more representative one.

Although both Yeh *et al.*¹¹ and Shahmoradi *et al.*¹³ examined the relationship between sequence and structural properties, they used different methods and data sets. First, Yeh *et al.*¹¹ measured ER using the method Rate4Site,^{26,27} whereas Shahmoradi *et al.*¹³ focused on sequence entropy, which is not a rate. Second, Yeh *et al.*¹¹ used a much more comprehensive data set of monomeric enzymes, and Shahmoradi *et al.*¹³ analyzed a comparatively smaller set of viral proteins, which are subject to an additional layer of selective forces imposed by the host immune system. Finally, Shahmoradi *et al.*¹³ considered additional structural predictors, namely protein design and flexibility,

while Yeh *et al.*¹¹ focused on RSA and WCN alone. This use of different methods makes it difficult to directly compare results from the two studies.

Here, we attempt to reconcile these two studies, by re-analyzing both the enzyme data set from Yeh et al. 11 and the virus data set from Shahmoradi et al. 13 in one consistent analysis pipeline. We focus on three structural predictors from the two studies: WCN, RSA, and variability in designed proteins. We confirm that, indeed, correlations between rate and structural predictors are much smaller for the viral proteins compared to the enzymes. However, differences in structural characteristics do not appear to drive the low predictive power in the viral protein data set. Instead, we find that the enzyme and viral protein data sets primarily differ in the extent of sequence variability in the multiple-sequence alignments (MSAs) used to infer evolutionary rates. Using evolutionary models, we quantify sequence divergence for all individual MSAs, and we find that the enzyme data set displays very high levels of divergence while the viral protein data set has experienced minimal evolutionary divergence. Across both data sets, we observe that the strongest structure-sequence correlations are observed at intermediate divergence levels. We conclude that the strength of the structurestructure relationship in proteins is, in part, determined by the extent of sequence variability in the data sets analyzed.

Results

We analyzed two distinct data sets. One was a set of 208 diverse enzyme monomers selected from the prior analysis by Yeh *et al.*¹¹ The other data set was a smaller set of nine viral proteins from Shahmoradi *et al.*¹³ Note that while the viral data set from

Shahmoradi *et al.*¹³ includes some viral enzymes, in the following we will use the term "enzymes" to refer specifically to the proteins from the Yeh *et al.*¹¹ data set.

Homologous sequences for each protein were taken from Yeh et al. 11 and Shahmoradi et al. 13 For each protein we made a multiple-sequence alignment using MAFFT^{28,29} on amino-acid sequences. From these alignments we calculated sitespecific evolutionary rates using Rate4Site. 26 We measured solvent accessibility for a given residue by its relative solvent accessibility (RSA) [Fig. 1(A)]. We measured packing density in the protein structures using side-chain WCN [Fig. 1(B)]. Previous studies have used C_{α} WCN when correlating WCN with ER. 11,13,15 However, a recent study³⁰ has shown that calculating WCN using the center of mass of the side chain results in stronger WCN-ER correlations. Therefore, here we used side-chain WCN throughout. We also measured the variability in designed sequences. For each protein in the viral data set and for each enzyme less than 200 residues in length we computationally designed 500 sequences using the respective structure as a template. From these sequences we inferred a "design rate" (DR) at each site, calculated as the expected steady-state evolutionary rate for an alignment with the given amino-acid frequencies.

Structural Predictors of Evolutionary Rate

To quantify the strength of structure—rate relationships in proteins, we correlated, separately for each protein, structural properties at individual sites with site-specific ER. Unless otherwise noted, we used Spearman correlations throughout. The first structural property that we examined was relative solvent accessibility (RSA). Prior work has

shown that RSA has a positive relationship with evolutionary rate. 8,10,11,13,15 This positive relationship between solvent accessibility and ER was verified in our analysis on the two data sets. Within both data sets, residues that have high RSA evolved faster on average. However, the strength of the relationship between RSA and ER varied between the enzyme and viral protein data sets. The enzymes, on average, had larger RSA–ER correlations with a mean correlation coefficient of 0.55 compared to 0.18 for viral proteins (t test: $P = 3.324 \times 10^{-5}$) [Fig. 2(A), Table 1].

Next we investigated the relationship between ER and packing density. For both data sets, residues with more contacts evolved slower [Fig. 1(B), Table 1]. This trend was also stronger for enzymes than for viral proteins, with a mean correlation coefficient of -0.63 for enzymes and -0.21 for viral proteins (t test: $P = 2.454 \times 10^{-5}$).

Protein Design as a Structural Predictor

Using protein design to search sequence space, Kuhlman and Baker³¹ found that sequences are close to optimal for a given structure (i.e., residues found at a given site are limited for a given structure). This constraint is especially true for buried residues. Given this result, Shahmoradi *et al.*¹³ attempted to use site-wise variability in designed proteins as an additional structural predictor of ER.¹³ Likewise, here, we used protein design as a third predictor of ER. However, unlike in Shahmoradi *et al.*,¹³ we did not use design entropy at sites but instead calculated a "design rate" (DR) as our predictor. We calculated this rate by calculating a predicted nonsynonymous substitution rate (dN) from amino-acid frequencies at each site, as derived in Spielman and Wilke.³² We found that this predicted rate makes similar predictions as does design entropy (not shown).

We used design rate here because it is the more principled quantity to compare to ER. For computational feasibility, for the enzyme data set we only designed proteins that were less than or equal to 200 residues in length. This encompassed 32 enzymes. We designed proteins for all the structures in the viral protein data set. Before performing our analysis, we compared the distributions of the strength of structure–rate correlations from the full enzyme data set with that of the distributions obtained from the 32 proteins. The differences between mean of the distributions were not significant (t test: t = 0.419 for RSA, t = 0.947 for WCN, Fig. S1).

In viral proteins, DR had a mean correlation coefficient of approximately -0.02, and in enzymes the mean coefficient of correlation was approximately 0.24 (Fig. 3, Table 1). However, for viral proteins this lower mean correlation was slightly misleading because some proteins had positive correlations while others had negative correlations, for a mean near zero (Fig. 3). In both data sets, design rate was a weaker predictor of evolutionary rates compared to WCN and RSA.

Even though DR did not correlate that strongly with ER, it is possible that it could explain variance in ER not explained by either RSA or WCN. To investigate this possibility, we used DR at sites as a predictor in linear models, either individually or in combination with the two other structural predictors, and calculated the percent variance explained for each model. In general, for both enzymes and viral proteins, design rate was not a good predictor of ER at sites. However, DR, just like RSA and WCN, was better at predicting ER in enzymes than in viral proteins. For a model with design rate as a single predictor, the average R^2 was 0.01 for viral proteins and \sim 0.07 for enzymes (Fig. S2, S3). Including DR as an additional predictor along with RSA and WCN added

some additional predictive power for ER in both data sets. For example, the average R^2 of a model with RSA and WCN as predictors for enzymes was approximately 0.37 (Fig. S2). When we added DR as an additional predictor, the average R^2 increased to 0.40 (Fig. S2). This increase in predictive power was observed in the viral data set as well. In summary, although DR was poor predictor of evolutionary rate at sites, it provided a small improvement in model performance, in particular for the enzyme data set.

Effect of Divergence of Structure–Rate Relationships

We found WCN, RSA, and DR all to be poor predictors of ER in viral proteins. There could be at least two different explanations for this finding. First, there could be unique structural features found within the viral protein data set that are not in the enzymes as indicated in Tokuriki *et al.*³³ Second, the viral proteins from Shahmoradi *et al.*¹³ may have experienced unique selection pressures (such as immune escape) or different divergence times than the enzymes taken from Yeh *et al.*¹¹

We found it unlikely that biophysical differences drove observed differences in the structure–rate correlations between the two data sets. First, any differences between the distributions for mean the WCN of the proteins within the data sets were not significant (P = 0.437 for WCN, Fig. 4). Differences in the mean RSA of the proteins were significant but the means were extremely similar (t = 0.027 for RSA, Fig. 4). Second, the strength of structure–rate correlations was only weakly dependent on the mean WCN or mean RSA of a protein (Fig. S4, S5). Proteins with larger mean RSA had only slightly larger RSA–ER correlations on average and the mean WCN was not related to the magnitude of structure–rate correlations (Fig. S4, S5).

We next investigated the possibility that differences in the multiple–sequence alignments for the two data sets were driving the differences in predictive power of RSA, WCN, and DR. On average the enzymes have more sequences in their representative alignments. We examined whether this difference was causing the difference in structure–rate correlation strength. We did observe a relationship between the number of sequences and the structure–rate strength. However the strength of this relationship was modest for enzymes (ρ = -0.185, P = 7.403 x 10⁻³ for WCN–ER and ρ = 0.060, P = 0.390 for RSA–ER) and was non-significant for viral proteins (ρ = -0.433, P = 0.250 for WCN–ER and ρ = 0.633, P = 0.076 for RSA–ER).

The two data sets showed significantly different levels of evolutionary divergence (Fig. 5). We calculated the divergence for each data set using two quantities: mean root-to-tip distance and mean patristic distance. Root-to-tip distance represents the extent of evolutionary divergence from the data set's common ancestor to a given sequence. The mean root-to-tip distance for each dataset was calculated as the average branch length, which indicates the number of substitutions, from the root in the tree to each terminal edge (tip) in the tree. Patristic, or pairwise, distance is the sum of branch lengths between two tips in a tree, and indicates how distantly related two sequences are to one another. As with mean root-to-tip-distance, a higher mean patristic distance indicated more evolutionary divergence. The enzyme alignments were much more diverged than the viral protein alignments (t test: t < 2.20 x 10⁻¹⁶ for mean root-to-tip distance and t < 2.20 x 10⁻¹⁶ for mean patristic distance).

Figure S6 shows structure—rate correlation strengths as a function of divergence (here measured as mean patristic distance). For both RSA–ER and WCN–ER

correlations, proteins with MSAs that had higher levels of divergence tended to have higher structure–rate correlations in magnitude. However, the trend between RSA–ER and WCN–ER correlations and mean patristic distance was not very strong (ρ = 0.161, P = 0.017 for RSA–ER and ρ = -0.117, P = 0.086 for WCN–ER).

Because divergence correlated only weakly with the structure–rate correlations, we hypothesized that overall divergence in an alignment mattered less than did variability in divergence among sites in an alignment. To obtain strong correlations with structural quantities, we need both highly conserved and highly variable sites. To assess the variability in the alignment at each site, we next calculated Shannon entropies at each site. By plotting the variance in entropy among sites against the mean [Fig. 6(A)], we found that indeed some alignments had overall high divergence but low variability among sites while other alignments were less diverged on average but had higher variability among sites. Figure 6B–F shows specific examples of entropy distributions among sites for individual proteins. For example, consider the protein identified by PDB ID 1G24 [Fig. 6(B)]. This protein had high mean entropy while maintaining a relatively low variance of entropy. Thus, sites in this protein were uniformly highly variable. Note that the distributions of entropy varied greatly between proteins even when they were from the same data set [Fig. 6(B)].

We next plotted structure–rate correlations against the variance in entropy and found strong correlations (Fig. 7, Spearman's correlation test: ρ = -0.321, P = 1.526 x 10⁻⁶ for WCN–ER, ρ = 0.236, P = 4.746 x 10⁻⁴ for RSA–ER). Proteins that had more variance in entropy across sites had larger structure–rate correlations in magnitude. Overall, enzymes were more diverged which in turn resulted, on average, in larger

variances in entropy across proteins. The viral proteins were less diverged and as such had lower variances in site variability. However, even for the highly diverged enzymes, correlations with structural quantities were low unless the alignments showed high variation in site variability. Thus, structure—rate correlations are maximized at intermediate levels of divergence, where alignments are sufficiently diverged for a high dynamic range (both highly conserved and highly variable sites are present in the same alignment) but not overly saturated with divergence (so that all sites are highly diverged).

We also investigated the effect of alignment quality on the observed patterns. Highly diverged sequences are more difficult to align, and errors in multiple sequence alignments may propagate to yield spurious rate inferences at some sites. Such inferences may be partially responsible for the low structure–rate correlations for some proteins. To assess average alignment reliability, we calculated a reliability score using Guidance 34,35 for each multiple sequence alignment. For each alignment, we calculated a column score (CS) at each site. CS scores range from 0, indicating an unreliably-aligned site, to 1, indicating a highly reliable alignment. We averaged the Guidance CS for each multiple sequence alignment to obtain a mean Guidance score representing the overall quality of an alignment. All of the viral proteins had scores greater than 0.98, indicating that these alignments had low uncertainty. The enzyme proteins had scores that span a very wide spectrum of quality, from 0 to 1. However, in enzymes, we found that the strength of structure–rate correlations was not correlated with alignment quality (Fig. S7, Spearman's correlation test: ρ = -0.022, ρ = 0.746 for WCN–ER, ρ = -0.132, ρ

= 0.057 for RSA–ER). This finding suggests that alignment quality is not a significant factor in the observed strength of structure–rate correlations.

As a final test of the effect of divergence on structure—rate correlations, we obtained a series of more diverged viral alignments. Briefly, we used PSI-BLAST to obtain a set of homologous proteins for each of the viral proteins from Shahmoradi et al., using the UniProt90 database. This procedure was comparable to the procedure that had been used to assemble the enzyme alignments. Subsequently, we performed the same analysis using these alignments as we did on the other two data sets. Using this new methodology, we only managed to collect sufficient sequences to calculate meaningful evolutionary rates for three of the viral proteins (PDB IDs: 1RD8, 3GOL, and 3LYF). However, even though the data set was small, we could compare it to the other two data sets for consistency. We found that the new viral data set was more diverged than the original viral data set but still less diverged than the enzyme data set (Fig. S6). Despite this increased divergence in the new viral data set, the strength of WCN–ER and RSA-ER correlations were similar to the original viral data set. Additionally, the relationship between measures of divergence and the strength of structure-rate correlations was similar for both viral data sets (Fig. 7, S6). Even with the new approach it was difficult to obtain viral alignments with high divergence, which may be responsible for the lower structure-rate correlations still observed.

Discussion

The field of molecular evolution has a long history of attempting to identify the factors that affect the rate at which proteins evolve. At the level of whole-protein rates, some of

the factors identified include expression level, interactions with other protein partners, ^{5,36–38} and selection for the costs of misfolding. ³⁹ Recently, the emphasis has shifted towards explaining rate variation among sites within proteins, which seems to be driven primarily by biophysical, structural constraints. ^{10–15,22,40}

Among the structural constraints, packing density and relative solvent accessibility have emerged as the two best structural predictors of evolutionary rate. 10,11,13,15,20 Sites that are on the surface of the protein tend to evolve faster than sites in the protein interior. Similarly, sites that are densely packed and have more contacts tend to evolve slower and exhibit less sequence variability than sites with fewer contacts. However, how strongly these two structural quantities (solvent accessibility and local packing density) correlate with evolutionary rate at sites remains somewhat unclear.

Here we have examined the relationship between site variability and the strength of structure–rate relationships by performing a direct comparison of the enzyme data set from Yeh *et al.*¹¹ and the viral proteins from Shahmoradi *et al.*¹³ We have found that both WCN and RSA are significant predictors of ER in enzymes, with 37% of the variation in ER explained (on average) by WCN and 28% explained on average by RSA. In viral proteins, both quantities perform weaker, explaining on average 8% and 7% of variation in ER respectively. Therefore, when analyzed using the same methods the data sets of Yeh *et al.*¹¹ and Shahmoradi *et al.*¹³ both show that WCN performs better than RSA.

In addition to RSA and WCN, we have also considered a third predictor, protein design rate (DR). Protein design had previously been used in Shahmoradi *et al.*¹³. We

have found that protein design rate is a much poorer predictor of rates at sites than RSA and WCN are. This result could represent a limitation in current methods of sequence space sampling techniques, limitations in the scoring function used in this study, or it could be that protein design rate does not capture biophysical forces that are predictive of evolutionary rates. For example, Ollikainen and Kortemme⁴¹ published a study that examined the ability of protein design to capture naturally occurring covariation of amino acids at sites. Although flexible-backbone design was able to recapitulate some covariation from natural sequences, not all covariation could be explained by design, indicating that other forces besides structure could be involved in natural patterns of sequence covariation. Additionally, Jackson et al. 42 found that protein design did not recapture some important structure—sequence patterns observed in yeast proteins. Notably, in that study, designed proteins did not exhibit the same relationship between solvent accessibility and site variability observed in natural proteins and hydrophobic residues were often underrepresented in the protein core. These studies underscore the possibility that either current protein design methods are imperfect at mimicking natural structural constraints or that structural constraints do not capture all of the biophysical effects on sequence evolution.

In contrast to the rate predictors in the enzyme data set, for the viral data set, the structural predictors (RSA, WCN, or DR) all performed poorly. We have found that neither differences in structural features (WCN, RSA, or DR) nor differences in evolutionary rates are likely a driving factor in the difference in correlation strength. Therefore, we have investigated the possibility that there are fundamental differences in the two data sets themselves.

We have found that the lack of divergence within the viral proteins of the data set taken from Shahmoradi et al. 13 is primarily responsible for the observed low structure rate correlations. For a protein to have a high structure-rate correlation, there needs to be a high level of variability in divergence among the sites in the multiple-sequence alignment. In other words, a protein must have a combination of sites that are highly conserved and sites that are highly variable. If all sites in a protein are conserved or all sites are saturated with many substitutions, so that there is no variability within the multiple-sequence alignment, then structure-rate correlations will be low. This combination of highly conserved and highly variable sites will only occur when there is an intermediate level of divergence. This is also why absolute divergence has a much weaker relationship with the strength of structure-rate correlations as compared to variance of entropy. Although it is critical for a data set to have sufficient divergence, it is only a necessary and not a sufficient requirement for strong structure-rate correlations. The enzyme data set of Yeh et al. 11 has a variety of proteins with differing levels of divergence and, on average, has MSAs that are more diverged. The intermediate level of divergence in these enzymes results in larger structure-rate correlations.

In addition, variation in selection at sites within a protein can affect the strength of observed structure—rate correlations. Across a protein, structure may differentially affect site variability and hence the strength of structure—rate correlation strength varies.

Selection against misfolding can constrain residues within the protein core while selection for key protein-protein interactions^{43,44} and/or against nonspecific protein-protein interactions⁴⁵ may impact the variability seen on the protein surface. For

example, important binding sites on the surface of the protein might be constrained decreasing the overall variability in variance of site variability. This would result in lower observed structure–rate correlations.

Although proteins as a whole exhibit common selective pressures, depending on the type of protein there might be additional factors that affect rate. Both viral proteins and enzymes exhibit some of the same selective pressures such as selection for stability and pressure to fold and adopt the correct native conformation. Enzymes are used to catalyze chemical reactions and as such have additional constraints such as structural constraints for a proper active site for catalytic function. On the other hand, viruses use their proteins to infect and replicate within their hosts. These proteins are utilized to perform a variety of necessary functions for viral replication such as host cellular entry^{46,47} and nuclear importation. As host immune systems attack these viruses, they evolve to escape from these host mechanisms resulting in signatures of positive selection within these proteins. Because of the differences in selective pressures facing these two protein types there might be different structural constraints on sequence variability and evolutionary rate.

We would like to emphasize that even though the distributions of average WCN and average RSA among proteins are similar for both data sets, there could be other structural differences among the proteins in the two data sets that might affect structure–rate correlations. Our purpose here was not to provide a rigorous, detailed analysis of structural differences among the two data sets. We only examined two obvious structural features (i.e., average packing of residues and average residue solvent accessibility) and showed that they are likely not the cause for the major

discrepancy in correlation strengths among the two data sets. More sophisticated structural analyses may identify unique structural features among viral proteins³³, and future research will have to determine whether these features have a measurable impact on structure–rate relationships. Furthermore, our results only apply to the two data sets discussed. Any additional general conclusions about the impact of divergence on observed structure–rate correlations in other systems would need further study.

Materials and methods

Structures, sequences, and measures of sequence properties

The results presented in this work were based on two data sets. The first was a data set of 208 monomeric enzymes, taken from Echave *et al.*¹⁴ who re-analyzed the structures originally studied by Yeh *et al.*¹¹ The Echave *et al.*¹⁴ data set was slightly smaller than the original data set because Echave *et al.*¹⁴ removed proteins that had missing data at insertion sites. The data set from Echave *et al.*¹⁴ was originally comprised of 209 proteins but we removed one additional protein, 1CQQ, during our analysis (see below for details). Thus, our final enzyme data set had 208 proteins. In brief, these proteins were all enzyme monomers randomly picked from the Catalytic Site Atlas 2.2.11.⁴⁹ Proteins in this data set varied from 95 to 1287 residues in length. Each structure was accompanied by a multiple–sequence alignment of 300 homologous sequences. The second data set was taken from Shahmoradi *et al.*¹³ and consisted of nine viral proteins. The viral proteins ranged from 122 to 557 residues in length and each structure was accompanied by a multiple–sequence alignment of up to 2362 homologous sequences. Although both data sets vary in the number of sequence alignments, we did not enforce

a medium number sequences in the multiple–sequence alignments needed to be included in the study since all alignments had at least 95 sequences.

Sequence alignments for both data sets were constructed by aligning the aminoacid sequences using the alignment software MAFFT, ^{28,29} specifying the auto flag to
select the optimal algorithm for the given data set. The alignments were then used to
calculate site-specific measures of evolutionary rate for each individual protein in both
data sets. We calculated a measure of site-specific evolutionary rate for each protein
using the software Rate4Site. ²⁶ First, maximum likelihood phylogenetic trees were
inferred with RAxML, using the LG substitution matrix and the CAT model of rate
heterogeneity. ^{50,51} For each structure, we used the respective sequence alignment and
phylogenetic tree to infer site-specific substitution rates with Rate4Site, using the
empirical Bayesian method and the JTT model of sequence evolution. ²⁶

Using the alignments, we also calculated the Shannon entropy (H_i) , at each alignment column i:

$$H_i = -\sum_j P_{ij} \ln P_{ij} ,$$

where P_{ij} was the relative frequency of amino acid j at position i in the alignment. Sequence entropy is a measure of variability at each site.

Finally, we calculated the divergence of each multiple—sequence alignment, using two measures: mean root-to-tip distance and mean patristic distance. Mean root-to-tip distance counts the average number of substitutions that have occurred along the tree. The mean patristic distance of an alignment was the average patristic distance of a tree where patristic distance was defined as the sum of the branch lengths between two

nodes (i.e., sequences) within the tree.⁵² Both root-to-tip distance and patristic distance were calculated using DendroPy.⁵³

For the viral proteins we collected a second data set. Using the sequences from the nine viral proteins from Shahmoradi *et al.* ¹³ as queries, we used PSI-BLAST⁵⁴ against the Uniprot90 to obtained homologous sequences for each protein. We used MAFFT and RaxML to create alignments and build trees for each protein. Trees could not be created for three of the proteins because their alignments did not have a sufficient number of sequences. We also chose to discard proteins from the analysis that did not have at least 25 sequences. This was done to guard against inaccurate rates. We calculated evolutionary rates for the remaining three proteins (PDB IDs: 1RD8, 3GOL, and 3LYF) using Rate4Site.

We quantified MSA reliability using a re-implementation of the Guidance platform³⁴ introduced by Spielman *et al.*³⁵ Guidance quantifies how robust MSA columns are to the guide tree topology used in during a progressive alignment algorithm. For each MSA column, Guidance produces a column score ranging from 0, indicating that the column is highly unreliable, to 1, indicating that the column is highly reliable. Note that the implementation in Spielman *et al.*³⁵ differs from that in Penn *et al.*³⁴ through its use of FastTree⁵⁵ to construct perturbed guidetrees. Here, Guidance was run with 100 bootstrap replicates using the MAFFT^{28,29} alignment software, specifying the "auto" flag. We derived an overall Guidance score for each MSA by averaging its resulting Guidance column scores.

Protein Design

Using Rosetta,⁵⁶ we computationally designed 500 structures for select proteins in each data set. For the viral proteins we designed 500 structures for each of the proteins taken from Shahmoradi *et al.*¹³ For the enzymes we designed structures for each protein that was at most 200 residues in length. For each protein, we first designed 500 flexible ensembles using Backrub.⁵⁷ Backrub generates a set of flexible backbone "ensembles" onto which side-chains can then be designed.^{57,58} The Backub method takes a temperature parameter, *T*, that determines the extent of backbone flexibility during design. Higher temperatures allow for more backbone flexibility. Previous work has shown that moderate temperature parameters result in designed structures more similar to natural proteins.^{41,42} Therefore, we used 0.6 as our temperature parameter. We then used the fixed-backbone method⁵⁹ to design side-chains on these ensembles.

All designs were generated with Rosetta 3.5, 2014 week five release. To generate the series of ensembles using flexible-backbone design we used the following Rosetta commands:

```
./backrub -database rosetta_database \
-s input.pdb -resfile NATAA.res -ex1 -ex2 \
-extrachi_cutoff 0 -backrub:mc_kt 0.6 \
-backrub:ntrials 10000 -nstruct 1 -backrub:initial_pack
```

For the fixed-backbone design we used the following Rosetta commands:

```
./fixbb -database rosetta_database \
    -s input.pdb -resfile ALLAA.res -ex1 -ex2 \
    -extrachi_cutoff 0 -nstruct 1 -overwrite \
    -minimize_sidechains -linmem_ig 10
```

After design, we removed proteins that did not map back properly to the alignments. This resulted in the removal of one structure, 1CQQ, completely from the study. This resulted in a total of 32 enzymes in addition to the viral proteins.

Using the sequence alignments of designed proteins we predicted a site-wise rate, using the expression for *dN* proposed by Spielman and Wilke³² (as implemented in the software Pyvolve⁶⁰). For this calculation, we assumed that the mutation rate at all sites was equal. We called this quantity the "design rate" (DR) at sites.

Calculation of structural properties

In our analysis, we used side-chain Weighted Contact Number (WCN) as proposed by Marcos and Echave.³⁰ This quantity is defined as

$$WCN_i = \sum_{i \neq j}^{N} \frac{1}{r_{ij}^2},$$

where r_{ij} is the distance between the geometric center of the side-chain atoms of residue i and the geometric center of the side-chain atoms of residue j, and N is the length of the protein. For glycine residues the distance to the C_{α} atom was used in lieu of the geometric center of the side-chain.

To calculate Relative Solvent Accessibility (RSA), we first calculated the Accessible Surface Area (ASA) for each site in each protein, via DSSP.⁶¹ We then normalized the ASA values by the theoretical maximum ASA values found in Table 1 of *Tien et al.*¹⁷ All WCN and RSA calculations were done on the individual, monomeric protein chain of interest.

All data and analysis scripts required to reproduce the work are publicly available to view and download at https://github.com/wilkelab/rate_variability_variation.

Supplementary material

Figure S1: Comparison of structure–rate correlations for the full data set of enzymes and the designed set. (A) Comparison of Spearman correlation coefficients for WCN–ER. (B) Comparison of Spearman correlation coefficients for RSA–ER. For both WCN–ER and RSA–ER the mean of the distributions for the designed set of enzymes is the same as that of the full data set of enzymes (t test: P = 0.947 for WCN–ER, P = 0.419 for RSA–ER).

Filename: dataset comparison plot.tiff

Figure S2: Distribution of R² for linear models of structural predictors of **evolutionary rate (ER) in enzymes.** WCN, RSA, DR and all combinations were used as predictors in a linear model with ER at sites as the response. Very little variation in ER can be explained when using design rate (DR) as a single predictor. For enzymes, only 32 proteins were included.

Filename: design_r2_enzyme_violin.tiff

Figure S3: Distribution of R² **for linear models of structural predictors of ER in viruses.** WCN, RSA, DR and all combinations were used as predictors in a linear model with evolutionary rate at sites as the response. Very little variation in evolutionary rate can be predicted by RSA, WCN or DR in viral proteins.

Filename: design_r2_virus_violin.tiff

Figure S4: Comparison of structure—rate correlations with Mean RSA. (A)

Spearman correlations of WCN–ER vs. mean RSA. Proteins with residues that are more exposed on average have slightly larger WCN–ER correlations in magnitude (Spearman's correlation test: ρ = 0.181, P = 7.653 x 10⁻³). (B) Correlations of RSA–ER vs. mean RSA. Proteins with residues that are more exposed on average also have slightly larger RSA–ER correlations in magnitude (Spearman correlation test: ρ = -0.228, P = 7.241 x 10⁻³).

Filename: mean rsa cor plot.tiff

Figure S5: Comparison of structure-rate correlations with Mean WCN. (A)

Spearman correlations of WCN–ER vs. mean WCN (Spearman correlation test: ρ = -0.082, P = 0.2283). (B) Correlations of RSA–ER vs. mean WCN (Spearman correlation test: ρ = 0.077, P = 0.2585). The average WCN of a protein is not related to the strength of structure–rate correlations.

Filename: mean_wcn_cor_plot.tiff

Figure S6: Comparison of structure–rate correlations with divergence. (A)

Spearman correlations of WCN and ER vs. mean pairwise distance (Spearman's correlation test: ρ = -0.117, P = 0.086 for WCN–ER). (B) Correlations of RSA and ER vs. mean pairwise distance. Enzymes are black, the viral proteins with the original alignments are in red, and the viral proteins with the newly collected sequences are in

turquoise. Proteins that are more diverged (as represented by mean pairwise distance) have stronger RSA–ER correlations (Spearman's correlation test: ρ = 0.161, P = 0.017). Filename: divergence_cor_plot.tiff

Figure S7: Comparison of structure–rate correlations with mean Guidance scores of proteins. (A) Comparison of Spearman correlation coefficients for WCN–ER. (B) Comparison of Spearman correlation coefficients for RSA–ER. Enzymes are black and viral proteins in red. Enzymes have more variation in alignment quality among proteins and have a small, significant relationship between alignment quality and structure–rate correlations (Spearman's Correlation test: ρ = -0.023, P = 0.746 for WCN–ER and ρ = -0.132, P = 0.057 for RSA–ER). For viral proteins there is no significant relationship between alignment quality and structure–rate correlations (ρ = -0.633, P = 0.076 for WCN–ER and ρ = 0.317, P = 0.410 for RSA–ER).

Filename: alignment_quality_plot.tiff

Acknowledgements

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Tables

Table 1: Averages of Spearman correlation coefficients between structural properties and evolutionary rate (ER). The structural properties analyzed are RSA, WCN, and predicted rate of designed proteins (DR). The analysis was performed on two data sets, one comprised of 208 enzyme monomers and comprised of nine viral proteins. Structure–ER correlations are higher in absolute magnitude in enzymes.

Data set	$\langle ho_{ m ER-WCN} angle$	$\langle ho_{\mathrm{ER-RSA}} \rangle$	$\langle ho_{ m ER-DR} angle^{ m a}$	$\langle ho_{ m ER-WCN} angle^{ m a}$	$\langle \rho_{\mathrm{ER-RSA}} \rangle^{a}$
Enzyme	-0.626	0.549	0.240	-0.625	0.561
Virus	-0.207	0.184	-0.022	-0.207	0.184

^aCorrelation coefficients calculated using the 32 enzymes and nine viral proteins for which there were designed sequences.



Figure legends

Figure 1: Description of Structural Properties. (A) Visualization of Solvent Accessibility. (B) Visualization of Local Packing Density. Each colored red particle represents a residue in the protein. In A, the lower red particle represents a surface residue. The red and white molecules indicate solvent molecules (e.g., water) that are contacting the red amino acid. This residue has a larger solvent accessibility because there is a larger proportion of the residue surface exposed to solvent. The upper red particle represents a core residue. This residue is not in contact with any solvent molecules and thus has low solvent accessibility. Relative solvent accessibility is obtained by normalizing the solvent accessibility of a given residue by the maximum amount of solvent accessibility for that amino acid. In B, the arrows pointing towards each residue indicate contacts between the red focal residue and its neighboring residues. The upper red residue represents a residue that has many neighbors (represented by the arrows) and thus has a high weighted contact number. The lower red residue is a surface amino acid with few neighbors and thus has a lower weighted contact number.

Figure 2: Distribution of correlation coefficients between structural properties and evolutionary rate (ER). (A) Spearman correlation coefficients between RSA and ER for the two data sets (t test: $P = 3.324 \times 10^{-5}$). (B) Spearman correlation coefficients between WCN and ER for the two data sets. For all structural properties, on average, viral proteins show weaker correlations than do enzymes (t test: $P = 2.454 \times 10^{-5}$).

Figure 3: Correlation Coefficients of Design Rate and evolutionary rate (ER).

Distributions of Spearman correlation coefficients between design rate (DR) and evolutionary rate (ER) for the two data sets. Enzyme proteins have higher correlations on average (t test: $P = 7.50 \times 10^{-4}$).

Figure 4: Distribution of average structural properties for each protein in the two data sets. (A) Distribution of average RSA. The distribution of average RSA different are very similar for both data sets (t test: P = 0.027). (B) Distribution of average WCN. The distribution of average WCN is the same for both data sets (t test: P = 0.437).

Figure 5: Divergence of sequences within the data sets. (A) Distributions of mean patristic distances for sequences in each protein alignment. Enzymes have larger mean patristic distances (t test: $P < 2.2 \times 10^{-16}$). (B) Distributions of mean root-to-tip distances for sequences in each protein alignment. Enzymes have larger mean root-to-tip distances (t test: $P < 2.2 \times 10^{-16}$). For both measures of divergence, the proteins within the enzyme dataset are more diverged. Divergence is relatively low between the viral proteins.

Figure 6: Comparison of the mean of entropy and the variance of entropy for individual proteins. (A) Variance in entropy at sites compared against overall mean entropy for each protein. Five different enzymes are highlighted, spanning the range of different combinations of high and low mean entropy and entropy variance. The enzymes are colored in black and the virus proteins are colored red. (B)–(F) Distributions of site-wise entropy values for the five proteins highlighted in A. There are

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a variety of distributions in site entropy for different proteins. Note: The protein denoted by the PDB ID 3GOL is a viral protein.

Figure 7: Comparison of structure–rate correlations with variance of entropy at sites. (A) Comparison of Spearman Correlation Coefficients of WCN–ER and variance of entropy for proteins. (Spearman's correlation test: ρ = -0.321, P = 1.526 x 10⁻⁶ using only the original protein data sets) (B) Correlations of RSA–ER and variance of entropy for proteins (ρ = 0.236, P = 4.756 x 10⁻⁴ using only the original protein data sets). Enzymes are black, the viral proteins with the original alignments are in red, and the viral proteins with the newly collected sequences are in turquoise. Enzymes have more variance in entropy across proteins and have larger structure–rate correlations in magnitude for both RSA and WCN. Virus proteins represented by the newly curated, more diverged alignments (see Methods) have similar structure–rate correlations to the original viral protein data set.

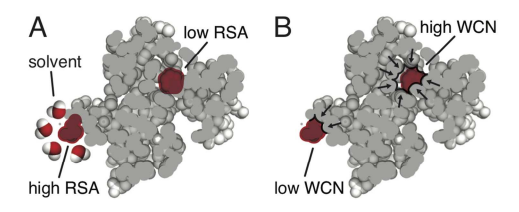


Figure 1: Description of Structural Properties. (A) Visualization of Solvent Accessibility. (B) Visualization of Local Packing Density. Each colored red particle represents a residue in the protein. In A, the lower red particle represents a surface residue. The red and white molecules indicate solvent molecules (e.g., water) that are contacting the red amino acid. This residue has a larger solvent accessibility because there is a larger proportion of the residue surface exposed to solvent. The upper red particle represents a core residue. This residue is not in contact with any solvent molecules and thus has low solvent accessibility. Relative solvent accessibility is obtained by normalizing the solvent accessibility of a given residue by the maximum amount of solvent accessibility for that amino acid. In B, the arrows pointing towards each residue indicate contacts between the red focal residue and its neighboring residues. The upper red residue represents a residue that has many neighbors (represented by the arrows) and thus has a high weighted contact number. The lower red residue is a surface amino acid with few neighbors and thus has a lower weighted contact number.

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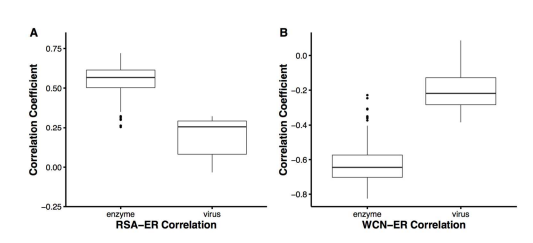


Figure 2: Distribution of correlation coefficients between structural properties and evolutionary rate (ER). (A) Spearman correlation coefficients between RSA and ER for the two data sets (t test: $P = 3.324 \times 10^{-5}$). (B) Spearman correlation coefficients between WCN and ER for the two data sets. For all structural properties, on average, viral proteins show weaker correlations than do enzymes (t test: t = 2.454 t 10⁻⁵). t 304x127mm (150 t 150 DPI)

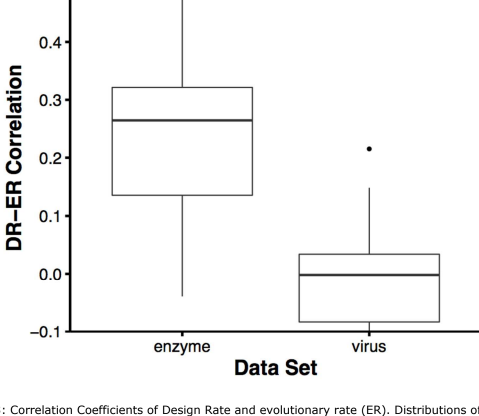


Figure 3: Correlation Coefficients of Design Rate and evolutionary rate (ER). Distributions of Spearman correlation coefficients between design rate (DR) and evolutionary rate (ER) for the two data sets. Enzyme proteins have higher correlations on average (t test: $P = 7.50 \times 10^{-4}$).

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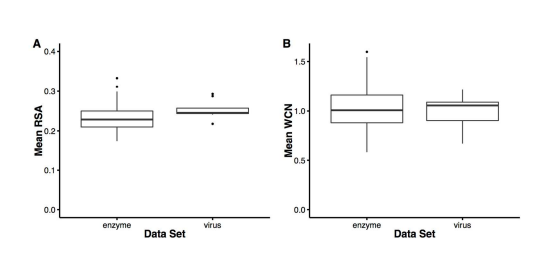


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304x127mm (150 x 150 DPI)

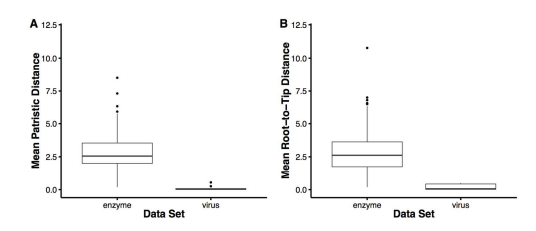


Figure 5: Divergence of sequences within the data sets. (A) Distributions of mean patristic distances for sequences in each protein alignment. Enzymes have larger mean patristic distances (t test: $P < 2.2 \times 10^{-16}$). (B) Distributions of mean root-to-tip distances for sequences in each protein alignment. Enzymes have larger mean root-to-tip distances (t test: $P < 2.2 \times 10^{-16}$). For both measures of divergence, the proteins within the enzyme dataset are more diverged. Divergence is relatively low between the viral proteins. $304 \times 127 \text{mm}$ (t 150 t 150 DPI)

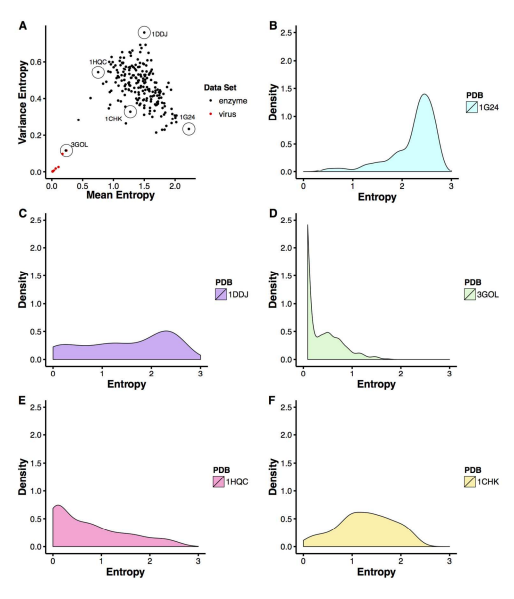


Figure 6: Comparison of the mean of entropy and the variance of entropy for individual proteins. (A) Variance in entropy at sites compared against overall mean entropy for each protein. Five different enzymes are highlighted, spanning the range of different combinations of high and low mean entropy and entropy variance. The enzymes are colored in black and the virus proteins are colored red. (B)–(F) Distributions of site-wise entropy values for the five proteins highlighted in A. There are a variety of distributions in site entropy for different proteins. Note: The protein denoted by the PDB ID 3GOL is a viral protein. 304x342mm (150 x 150 DPI)

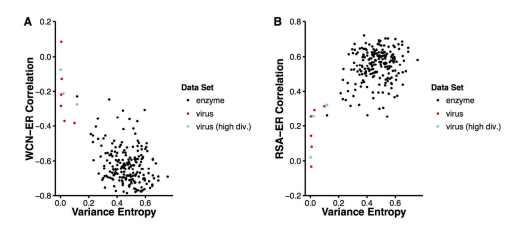


Figure 7: Comparison of structure–rate correlations with variance of entropy at sites. (A) Comparison of Spearman Correlation Coefficients of WCN–ER and variance of entropy for proteins. (Spearman's correlation test: $\rho = -0.321$, $P = 1.526 \times 10^{-6}$ using only the original protein data sets) (B) Correlations of RSA–ER and variance of entropy for proteins ($\rho = 0.236$, $P = 4.756 \times 10^{-4}$ using only the original protein data sets). Enzymes are black, the viral proteins with the original alignments are in red, and the viral proteins with the newly collected sequences are in turquoise. Enzymes have more variance in entropy across proteins and have larger structure–rate correlations in magnitude for both RSA and WCN. Virus proteins represented by the newly curated, more diverged alignments (see Methods) have similar structure–rate correlations to the original viral protein data set. 304x127mm (150×150 DPI)



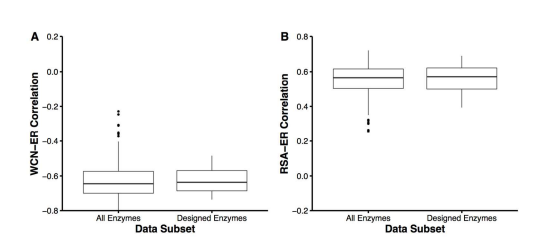


Figure S1: Comparison of structure–rate correlations for the full data set of enzymes and the designed set. (A) Comparison of Spearman correlation coefficients for WCN–ER. (B) Comparison of Spearman correlation coefficients for RSA–ER. For both WCN–ER and RSA–ER the mean of the distributions for the designed set of enzymes is the same as that of the full data set of enzymes (*t* test: *P* = 0.947 for WCN–ER, *P* = 0.419 for RSA–ER).

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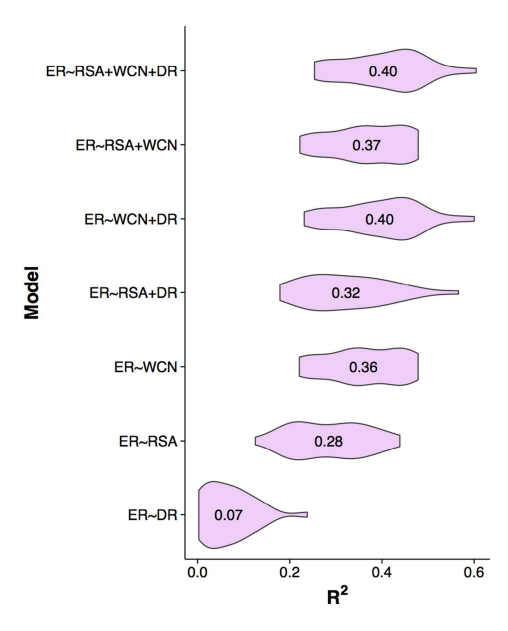


Figure S2: Distribution of R2 for linear models of structural predictors of evolutionary rate (ER) in enzymes. WCN, RSA, DR and all combinations were used as predictors in a linear model with ER at sites as the response. Very little variation in ER can be explained when using design rate (DR) as a single predictor. For enzymes, only 32 proteins were included.

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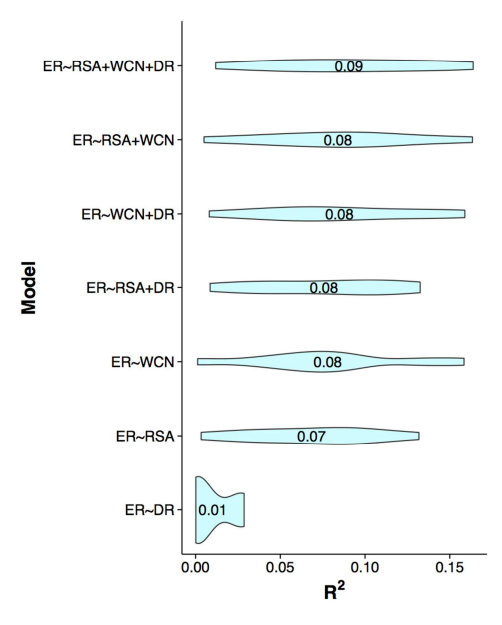


Figure S3: Distribution of R2 for linear models of structural predictors of ER in viruses. WCN, RSA, DR and all combinations were used as predictors in a linear model with evolutionary rate at sites as the response.

Very little variation in evolutionary rate can be predicted by RSA, WCN or DR in viral proteins.

165x203mm (150 x 150 DPI)

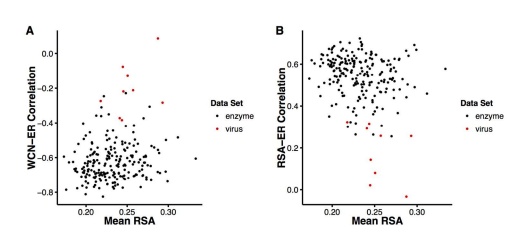


Figure S4: Comparison of structure–rate correlations with Mean RSA. (A) Spearman correlations of WCN–ER vs. mean RSA. Proteins with residues that are more exposed on average have slightly larger WCN–ER correlations in magnitude (Spearman's correlation test: $\rho = 0.181$, $P = 7.653 \times 10^{-3}$). (B) Correlations of RSA–ER vs. mean RSA. Proteins with residues that are more exposed on average also have slightly larger RSA–ER correlations in magnitude (Spearman correlation test: $\rho = -0.228$, $P = 7.241 \times 10^{-3}$). $304 \times 127 \text{mm}$ (150 x 150 DPI)

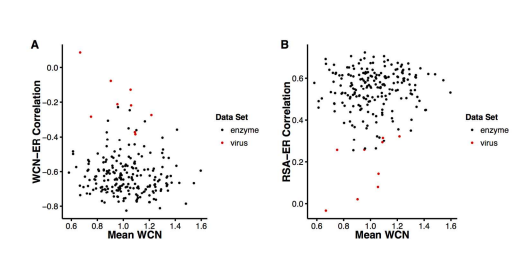


Figure S5: Comparison of structure–rate correlations with Mean WCN. (A) Spearman correlations of WCN–ER vs. mean WCN (Spearman correlation test: $\rho = -0.082$, P = 0.2283). (B) Correlations of RSA–ER vs. mean WCN (Spearman correlation test: $\rho = 0.077$, P = 0.2585). The average WCN of a protein is not related to the strength of structure–rate correlations. $304 \times 127 \text{mm}$ (150 x 150 DPI)

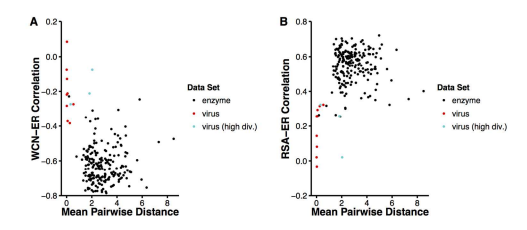


Figure S6: Comparison of structure–rate correlations with divergence. (A) Spearman correlations of WCN and ER vs. mean pairwise distance (Spearman's correlation test: $\rho = -0.117$, P = 0.086 for WCN–ER). (B) Correlations of RSA and ER vs. mean pairwise distance. Enzymes are black, the viral proteins with the original alignments are in red, and the viral proteins with the newly collected sequences are in turquoise. Proteins that are more diverged (as represented by mean pairwise distance) have stronger RSA–ER correlations (Spearman's correlation test: $\rho = 0.161$, P = 0.017).

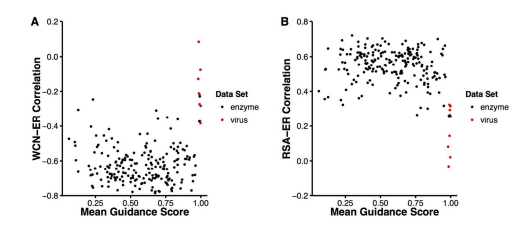


Figure S7: Comparison of structure–rate correlations with mean Guidance scores of proteins. (A) Comparison of Spearman correlation coefficients for WCN–ER. (B) Comparison of Spearman correlation coefficients for RSA–ER. Enzymes are black and viral proteins in red. Enzymes have more variation in alignment quality among proteins and have a non-significant relationship between alignment quality and structure–rate correlations (Spearman's Correlation test: ρ = -0.023, P = 0.746 for WCN–ER and ρ = -0.132, P = 0.057 for RSA–ER). For viral proteins there is no significant relationship between alignment quality and structure–rate correlations (ρ = -0.633, P = 0.076 for WCN–ER and ρ = 0.317, P = 0.410 for RSA–ER).

304x127mm (150 x 150 DPI)