Predicting and interpreting large-scale mutagenesis data using analyses of protein stability and conservation

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Graphical abstract

Highlights
- We analyze ca. 150,000 variant effects obtained by multiplexed assays
- We use protein stability and sequence conservation to predict variant effects
- About half of the loss-of-function variants lose function with loss of stability

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In brief
Missense variants in which a single amino acid is changed in a protein underlie a large number of genetic diseases. By combining experimental measurements of ca. 150,000 variant effects with analyses of protein stability and sequence conservation, Høie et al. train a machine learning model to predict variant effects.
Predicting and interpreting large-scale mutagenesis data using analyses of protein stability and conservation

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SUMMARY

Understanding and predicting the functional consequences of single amino acid changes is central in many areas of protein science. Here, we collect and analyze experimental measurements of effects of > 150,000 variants in 29 proteins. We use biophysical calculations to predict changes in stability for each variant and assess them in light of sequence conservation. We find that the sequence analyses give more accurate prediction of variant effects than predictions of stability and that about half of the variants that show loss of function do so due to stability effects. We construct a machine learning model to predict variant effects from protein structure and sequence alignments and show how the two sources of information support one another and enable mechanistic interpretations. Together, our results show how one can leverage large-scale experimental assessments of variant effects to gain deeper and general insights into the mechanisms that cause loss of function.

INTRODUCTION

The ability to predict and understand the effects of amino acid changes to protein structure, stability, and function plays central roles in a number of areas of protein science. For example, improving protein function or stability is key in many biomedical applications, and the ability to understand and predict loss of function is of central importance in disease biology. Large-scale human genome sequencing efforts are revealing millions of missense variants that change the amino acid sequences of proteins, but we do not yet know the functional consequences for most of these variants.

Studies of the effects of single amino acid changes also present opportunities to test our understanding of the protein structure-function relationships. While a large fraction of single amino acid substitutions in a given protein are relatively well tolerated, there is a subset that has significant detrimental consequences (Schaafsma and Vihinen, 2017; Gray et al., 2017). Pinpointing which variants are in the detrimental group, and the biochemical and biophysical mechanisms underlying loss of fitness, is important, for example, for assessing pathogenicity of so-called variants of uncertain significance (Richards et al., 2015) and understanding the mechanistic origins of disease.

Recent technological advances have enabled high-throughput assays that can quantify changes in activity, stability, or other protein properties of interest for thousands of variants in a single experiment. Such multiplexed assays of variant effects (MAVEs; also called deep mutational scanning experiments) also provide large sets of data with systematic fitness profiles of variants (Fowler and Fields, 2014), often providing both mechanistic insight and systematic assessment of computational models for predicting variant effects. MAVEs are often performed using customized assays, which, however, also means that functional scores (sometimes called fitness scores) extracted from different MAVEs are often not directly comparable with one another without further normalization. The data generated by MAVEs have been shown to help predict the status of pathogenic and benign variants, while also serving as useful benchmarks for computational variant classification methods (Livesey and Marsh, 2020; Frazer et al., 2021). More generally, the data can also provide detailed insight into general aspects of protein structure and function (Gray et al., 2017; Ahler et al., 2019; Stein et al., 2019; Dunham and Beltrao, 2021; Chiasson et al., 2020; Starr et al., 2020; Cagiada et al., 2021; Amorosi et al., 2021).

A given amino acid change may affect multiple properties or functions of a protein, and by combining different assays it may be possible to disentangle which substitutions affect which of those properties and functions. As most proteins need to be folded to function, assays probing protein stability and cellular abundance have received special attention. Thus, a specific type of MAVE termed VAMP-seq (variant abundance by massively parallel sequencing) has been developed to probe cellular protein abundance (Matreyek et al., 2018) and has been shown to correlate with measurements of protein stability (Matreyek et al., 2018; Suiter et al., 2020). While the detailed relationship between protein stability and abundance is complicated...
and not fully understood (Hingorani and Gierasch, 2014; Stein et al., 2019), we and others have shown that unstable proteins are often targeted for proteasomal degradation leading to lowered cellular abundance, and a large number of disease-causing variants are proteasomal targets and are found at low cellular levels (Meacham et al., 2001; Yaguchi et al., 2004; Olzmann et al., 2004; Ron and Horowitz, 2005; Yang et al., 2011, 2013; Arlow et al., 2013; Nielsen et al., 2017; Chen et al., 2017; Nielsen et al., 2017, 2020; Scheller et al., 2019; Abildgaard et al., 2019). By combining the results from a VAMP-seq experiment with a MAVE probing protein activity it is possible to distinguish between variants that cause loss of function due to lowered abundance and those that change the intrinsic activity of a protein (Jepsen et al., 2020; Chiasson et al., 2020; Cagiada et al., 2021; Amorosi et al., 2021). These experiments and analyses suggest that a relatively large fraction of variants that cause loss of function are due to loss of stability and resulting degradation in the cell.

Although it is possible to perform multiplexed assays on multiple genes in a single experiment (Després et al., 2020; Jun et al., 2020; Hanna et al., 2021; Cuella-Martin et al., 2021), we are not yet able to probe all possible variants in all proteins by experiments. Thus, computational methods are important to predict and understand variant effects, and in some cases they may be even more accurate than MAVEs for this purpose (Jepsen et al., 2020; Frazier et al., 2021). Most variant effect predictors are based on features extracted from evolutionary conservation of homologous proteins, biophysical calculations based on structure, and general knowledge of amino acid properties (Yue et al., 2005; Kumar et al., 2009; Adzhubei et al., 2010; Casadio et al., 2011; De Baets et al., 2012; Kircher et al., 2014; Choi and Chan, 2015; Ioannidis et al., 2016; Ancien et al., 2018; Wagth et al., 2018; Gerasimavicius et al., 2020; Livesey and Marsh, 2020). As some of these methods have been trained and tested on classification of clinical variants, it has been argued that comparison against data from MAVEs provides a useful and unbiased alternative to benchmark such methods (Livesey and Marsh, 2020). In such tests, it has been shown that various sequence-based approaches, including deep-learning methods, can achieve very high accuracy (Rieselman et al., 2018; Livesey and Marsh, 2020). Indeed, we and others have successfully applied sequence analysis and biophysical stability calculations for identification and analysis of pathogenic variants, although these methods were not trained on clinical variants (Pey et al., 2007; Yin et al., 2017; Nielsen et al., 2017; Gray et al., 2018; Scheller et al., 2019; Cline et al., 2019; Abildgaard et al., 2019; Jepsen et al., 2020; Frazier et al., 2021).

Experiments and computational analyses such as those discussed above are now beginning to provide a consistent picture of the effects of variants on protein stability and function. Variants that cause substantial loss of stability are generally found at low protein levels in the cell, and thus often lead to a loss-of-function phenotype and disease. Hence, when a variant is predicted to be highly destabilizing, it is likely to be non-functional. The reverse, however, does not hold true. Variants that do not perturb protein stability may still cause loss of function via other mechanisms, such as perturbing active-site residues in an enzyme or key interaction sites for binding. Such effects can often be captured by evolutionary sequence analyses, which are potentially sensitive to all conserved molecular mechanisms that lead to loss of function.

We recently quantified how variants may affect protein activity and abundance and the relationship between the two (Jepsen et al., 2020; Cagiada et al., 2021). By analyzing two MAVEs, one probing activity and another probing abundance, for two proteins (PTEN and NUDT15) we found that about one-third of all possible variants cause loss of function, and about half of the loss-of-function variants also have low cellular abundance. Predictions of protein stability and analyses of sequence conservation demonstrated that many of these effects could be recapitulated by computational methods. Specifically, we found that analyses of sequence conservation captured general aspects leading to loss of function, whereas calculations of protein stability were more strongly correlated with experimental measures of protein abundance. We also found that variants that cause loss of function, but not lowered abundance, were enriched in the active sites of PTEN and NUDT15, whereas variants that cause loss of function via decreased protein abundance were more often found in the core of the protein structure (Cagiada et al., 2021).

Here, we aim to provide further insight into the relationship between variant effects on protein stability and function, and how computational predictions of changes in thermodynamic stability and analysis of sequence conservation may be used to predict variant effects. We have collected 39 datasets previously generated by MAVEs on 29 proteins, and analyze these using analyses of sequence conservation and predictions of protein stability. We train a machine learning model that uses these features as input to predict variant effects, and show that both sequence conservation and predictions of protein stability contribute to prediction accuracy. We use our model to provide a global view of the relationship between stability and function, and help pinpoint which variants lose function due to loss of stability. Together, our results show how loss of stability is an important contributor to loss of function, and point to future improvements for predictions of variant effects.

RESULTS

Analyzing variant effects from MAVEs by calculations of stability and conservation

We first aimed to quantify how well analyses of protein stability and sequence conservation are able to capture experimental measurements of variant effects. We thus collected 39 datasets generated by MAVEs on 29 proteins from the literature (Table S1). As we aimed to use the data in a globally trained machine learning model, we used rank normalization to bring the original variant scores reported by the individual studies onto a common scale $s_{\text{exp}}$ for each of the 19 possible variants at each of the positions resolved in the experimental structures. Here, $s_{\text{exp}} = 1$ corresponds to wild-type-like activity in the experiment and $s_{\text{exp}} = 0$ corresponds to variants with low activity in the assay used in the MAVE.

For each of the 29 proteins we used Rosetta (Park et al., 2016; Leman et al., 2020; Frenz et al., 2020) to predict changes in thermodynamic stability ($\Delta G$) for each of the 19 possible variants at each of the positions resolved in the experimental structures. Here, $\Delta G = 0$ corresponds to the same stability as the wild-type protein and variants with $\Delta G > 0$ correspond to those that
We have previously shown that changes in protein stability ($\Delta \Delta G$) and multiple-sequence-alignment-based conservation scores ($\Delta \Delta E$) correlate with changes in cellular function or stability in selected proteins (Nielsen et al., 2017; Scheller et al., 2019; Abildgaard et al., 2019; Cagiada et al., 2021). This also holds for the 154,808 variants in the 29 proteins studied here (Figure 1A), so that variants with large $\Delta \Delta E$ or $\Delta \Delta G$ scores tend to show low fitness (low $s_{exp}$). In particular, variants for which both $\Delta \Delta G$ and $\Delta \Delta E$ values are high almost always show loss of function, while those with low scores for both usually show wild-type-like fitness. As expected and discussed above, variants that do not substantially perturb stability (low $\Delta \Delta G$) can both have low and high values of $s_{exp}$, because substitutions may affect function through other mechanisms than loss of stability and abundance. From the stability-conservation landscape, however, it appears that such effects are captured by the $\Delta \Delta E$ scores so that, even for variants with low $\Delta \Delta G$, conservative substitutions (low $\Delta \Delta E$) tend to be associated with high fitness, and high $\Delta \Delta E$ with low fitness.

To quantify the power of $\Delta \Delta E$ and $\Delta \Delta G$ for classifying variants, we extracted the top and bottom third of variant scores ($s_{exp}$) as the subsets that are most clearly associated with wild-type-like and loss-of-function phenotypes, respectively. Next, we divided our MAVE stability-conservation landscape into a total of 16 sectors. The normalized $\Delta \Delta E$ scores have evenly distributed thresholds of 0.25, 0.50, and 0.75, while the $\Delta \Delta G$ thresholds are at 2.0, 3.0, and 4.5 kcal/mol (Figure 1). Inspection of the sectors confirms that extreme values of $\Delta \Delta E$ and $\Delta \Delta G$ can classify variants well into the high or low fitness categories, while moderate values have lower classification power (Figure 1B). For example, when $\Delta \Delta E<0.25$ and $\Delta \Delta G<2.0$ kcal/mol, 81% of the variants are in the high fitness category, and, at the opposite end, when $\Delta \Delta E>0.75$ and $\Delta \Delta G>4.5$ kcal/mol, 93% of the variants are in the low fitness category.

More generally, the two-dimensional fitness landscapes illustrate the partial interdependency of $\Delta \Delta E$ and $\Delta \Delta G$. Most notably, it is clear that evolution selects strongly against destabilizing variants so there are almost no cases with high values of $\Delta \Delta G$ and low values of $\Delta \Delta E$ (only 4% of the variants have $\Delta \Delta E<0.25$ and $\Delta \Delta G>3.0$ kcal/mol). Also, as discussed above, low values of $\Delta \Delta G$ may be associated both with high and low values of $s_{exp}$, whereas variants with larger values of $\Delta \Delta G$ tend to have low $s_{exp}$. Thus, focusing on the stable variants ($\Delta \Delta G<2.0$ kcal/mol) and only the top/bottom third of the fitness scores, we find that 38% of these variants have $\Delta \Delta E>0.5$ (Figure 1A) and are thus more likely to be non-functional. Focusing on the variants that the GEMME analysis suggest are incompatible with what has been observed through evolution ($\Delta \Delta E>0.5$), and thus are more likely to be non-functional, we find that 47% are predicted to be unstable ($\Delta \Delta G>2.0$ kcal/mol; Figure 1B). Thus, in line with our previous analysis (Cagiada et al., 2021), these results suggest that approximately half of the non-conservative substitutions are selected against due to stability effects.

We have previously found that low-$\Delta \Delta G$ (structurally stable) but high-$\Delta \Delta E$ (evolutionarily constrained) variants are often found in clusters near active sites and other functionally important regions in the proteins PTEN and NUDT15 (Cagiada et al., 2021). We here examined two additional proteins (CBS and HMGCR) to understand the distribution and location of
Having established that there is an overall relationship between calculated values of $\Delta \Delta G$ and $\Delta \Delta E$ and experimental variant effects ($s_{\text{exp}}$) and that both independently contribute valuable information (Figure 1), we decided to train a machine learning model to predict variant effects from $\Delta \Delta G$ and $\Delta \Delta E$. Before doing so, we analyzed how well the data generated by the individual MAVEs correlate with the calculated values of $\Delta \Delta G$ and $\Delta \Delta E$ (Figure 2A).

We find a broad range of Spearman correlation coefficients ($r_s$) between the MAVE scores ($s_{\text{exp}}$) and our calculated $\Delta \Delta G$ and $\Delta \Delta E$ scores, in line with previous observations of considerable variation in correlation with variant effect predictors (Livesey and Marsh, 2020). Correlations range from low ($r_s \sim 0.1$ on BRCA1 and Calmodulin-1 [CALM1] for both $\Delta \Delta E$ and $\Delta \Delta G$) to relatively high ($r_s \sim 0.6 - 0.8$) when predicting multiple independent MAVEs on $\beta$-lactamase (bla) (Figure 2A). Overall and as expected, experimental variant effects that correlate well with $\Delta \Delta G$ also correlate well with $\Delta \Delta E$. In line with the results shown above (Figure 1) and previous analyses (Nielsen et al., 2017; Abildgaard et al., 2019; Jepsen et al., 2020; Livesey and Marsh, 2020; Gerasimavicius et al., 2020; Cagiada et al., 2021), we find that variant effects tend to be more strongly correlated with $\Delta \Delta E$ than $\Delta \Delta G$. Two notable outliers to this observation are the abundance-based MAVEs (VAMP-seq) for PTEN (Matreyek et al., 2016) and NUDT15 (Suter et al., 2020), which are more strongly correlated with our analysis of protein stability than conservation, labeled as PTEN (a) and NUDT15 (a) in Figure 2A (Cagiada et al., 2021). Interestingly, the VAMP-seq data for the protein TPMT (Matreyek et al., 2018) is slightly more strongly correlated with $\Delta \Delta G$ than $\Delta \Delta E$. PTEN and NUDT15 variants have also been assayed for their respective biochemical functions (Mighell et al., 2018; Suter et al., 2020), and the resulting $s_{\text{exp}}$ scores correlate better with $\Delta \Delta E$ and less well with $\Delta \Delta G$ than the corresponding protein abundance scores. The MAVEs of respective biochemical function are labeled as PTEN (b) and NUDT15 (b) in Figure 2A (Cagiada et al., 2021). While the reasons for the low correlation between $\Delta \Delta E$ and $\Delta \Delta G$ with several of the experiments remain unclear; possible explanations include inaccuracies in stability calculations, poor sequence alignments, or experimental assays that probe properties not directly related to stability or an evolutionary-conserved function of the protein. Indeed, similar variation in correlation between the outcome of MAVEs and $\Delta \Delta E$-like and $\Delta \Delta G$ scores have previously been observed (Hopf et al., 2017; Livesey and Marsh, 2020; Dunham and Beltrao, 2021).
Next we constructed a simple baseline model that captures effects of substituting each of the 20 protein-coding amino acids for another by averaging over the normalized scores from all variants in our dataset (\(s_{\text{exp}}\); Figure S2). Similar to previous observations (Gray et al., 2017), this substitution matrix captures well-known biochemical patterns. In particular, we observe that substitutions of hydrophobic residues—often found in the protein core—with charged or polar residues on average leads to a large loss of fitness, while changes from a polar to a hydrophobic residue on average does not cause a substantial loss of fitness. In general, we note that the \(s_{\text{exp}}\) matrix is distinctly asymmetric. In addition to the effects of polar and apolar residues, we see that substitutions to proline are substantially more detrimental to function than substitutions from proline, and substitutions from cysteine affect fitness more than substitutions to cysteine. Such asymmetry is not present in, for example, the BLOSUM62 substitution matrix (Hess et al., 2016) (Figure S3), which captures average evolutionary effects without taking directionality into account. Thus, the \(s_{\text{exp}}\) and BLOSUM62 matrices are only modestly correlated (Figure S4), and we prefer the \(s_{\text{exp}}\) matrix as our null model. Nevertheless, although \(s_{\text{exp}}\) captures the typical single amino acid effects observed in a MAVE, it lacks information about structural and sequence context, and thus is overall a poor predictor of experimental variant effects (mean \(r = 0.17\); Figure 2B, green).

With the experimental and computational variant data we proceeded to train a set of machine learning models to predict \(s_{\text{exp}}\) from our set of available \(\Delta G\), \(\Delta E\), and \(s_{\text{exp}}\) scores. We chose to use random forest models because of their robustness to outliers and noise, minimal need to adjust model hyperparameters, as well as the possibility to easily extract information about the extent to which the different features are used in the model decision process (Breiman, 2001; Bernard et al., 2009). We term the values predicted by the model \(s_{\text{calc}}\).

We used a leave-one-protein-out procedure for training, selecting one protein for validation, and training on the normalized MAVE, \(\Delta E\), and/or \(\Delta G\) data from all other proteins in our overall set. Thus, when more than one set of experiments had been performed on a single protein, we excluded the extra datasets during training. We assess the model by calculating the correlation between \(s_{\text{calc}}\) predicted from the resulting random forest with the \(s_{\text{exp}}\) values from all MAVEs on the protein that was left out in training, looping over all proteins one at a time.

First, we trained three random forest models using as inputs only the variant \(\Delta G\), only the variant \(\Delta E\), or both. In line with the variation in the correlation to the input data, we observe a range of correlation coefficients from these models, with the combined \(\Delta G\) and \(\Delta E\) model correlating with the normalized MAVE scores with a median \(r = 0.47\) (Figure 2B, orange). The \(\Delta G\)-only model (\(r = 0.25\), Figure 2B, cyan) only performs slightly better than \(s_{\text{exp}}\) (green), and the \(\Delta E\)-only model (\(r = 0.42\), Figure 2B, purple) is again closer to capturing the experimental outcomes. We note here that several of the correlation coefficients observed in these analyses (Figure 2B) are smaller than those obtained when correlating directly with the \(s_{\text{exp}}\) values (Figure 2A). While the direct correlations essentially represent 39 individual, “protein-specific” models, we aimed to build a single model generalizing across different proteins and residue contexts, capturing the relationship between, e.g., \(\Delta G\) and \(s_{\text{exp}}\) on a single scale.

As variant effects depend both on the specific substitutions but also the context within the protein, we trained a more complex “position-context” model that also takes into account the scores for other substitutions at the given protein position. Specifically, in addition to the \(\Delta E\) and \(\Delta G\) value for the variant to be predicted, we input the entire set of 20 \(\Delta G\) and 20 \(\Delta E\) values at a position, representing the stability and conservation scores for all single amino-acid variants at the given position, as well as the mean score for these. We further add three features from the baseline model, \(s_{\text{WT-Mut}}\) (the average score when mutating from the wild type to the specific variant [mutant] residue), \(s_{\text{WT-Ary}}\) (the average score for substitutions from the specific wild-type amino acid to any of the other 19), and \(s_{\text{Any-Mut}}\) (the average score when mutating each of the 19 other amino acids to the specific variant amino acid) (see more detailed discussion in STAR Methods). The resulting position-context random forest model yields an improved performance (median \(r = 0.52\), Figure 2B, brown). The model also successfully recapitulates the trends in the stability-conservation landscape (Figures S2C and S7). Looking more closely at how the model performs in different sectors of this landscape, we find that predictions generally have a similar mean absolute error (MAE) in most sectors (Figures 2D and S6) with MAE = 0.22, with somewhat more accurate predictions for highly destabilizing (\(\Delta \Delta G > 3.0\)) and non-conservative (\(\Delta \Delta E > 0.50\)) substitutions (MAE = 0.16–0.21).

### Role of stability and conservation in the predictions

With the final model in hand and having shown that it recapitulates the experimentally observed stability-conservation landscape relatively well (Figures 1A, 2C and S5), we proceeded to analyze the properties of the model. Looking at the predictions of the >150,000 variant effects from the leave-one-protein-out model, we find that the position-context model predicts fitness outcomes with a Spearman correlation coefficient of 0.50 (Figure 3A). The most dense region has a sigmoidal shape, with predictions of the experimentally derived \(s_{\text{exp}}\) range 0.5–1.0 seeming largely indistinguishable by the model. Indeed, the model is particularly good at separating the approximately one-third of the variants that cause substantial loss of function (36% have \(s_{\text{exp}} < 0.3\)) from those with activities closer to wild type. We note here that the noisy nature of data generated by many MAVEs, as well the rank-normalization that we used for each protein, likely mean that many variants with fitness scores comparable with wild type end up being spread out with \(s_{\text{exp}}\) in the range = 0.3–1.0, in line with our previous observations of about one-third of variants causing loss of function (Caggiada et al., 2021). Thus, we suggest that the model is better suited at capturing the bimodal distribution seen in many MAVEs (Gray et al., 2017). While calculations of \(\Delta G\) and \(\Delta E\) may capture more subtle effects on function in individual proteins (Nielsen et al., 2021), such effects appear difficult to extract in our global analysis. In the absence of the model’s ability to distinguish between such effects, the predictions plateau at a relatively narrow distribution of values that minimizes the mean square error between \(s_{\text{exp}}\) and \(s_{\text{calc}}\) for these variants.
To gain further insight into the relative importance of the stability and conservation parameters for predicting the variant scores, we calculated the partial dependencies in the position-context random forest model. These partial dependencies quantify how the model’s predictions of $s_{\text{calc}}$ depend on the chosen features, here the $\Delta \Delta G$ and $\Delta \Delta E$ for the specific variant, marginalizing over the remaining features in the model (Molnar, 2019). For variants that at most cause a modest change in stability ($\Delta \Delta G < 2.0 \text{ kcal/mol}$), the predictions are most strongly influenced by $\Delta \Delta E$. This observation is in line with the notion that, for stable variants, $\Delta \Delta E$ is a relatively good predictor of $s_{\text{exp}}$ (Figure 1A). In contrast, for more destabilizing variants ($\Delta \Delta G > 3.0 \text{ kcal/mol}$) the partial dependence varies with both $\Delta \Delta G$ and $\Delta \Delta E$, in line with the finding that both of these are useful quantities to help predict variant effects in this part of the stability-conservation landscape (Figures 1 and 2).

We then proceeded to examine the impact of all features in the random forest model. We stress that this so-called feature importance only quantifies how much each feature is used overall in the prediction of $s_{\text{calc}}$, but does not directly describe how and when these features are used (Strobl et al., 2007). We calculated the correlation between each pair of features and used these to cluster and build a dendrogram of the features (Figures 3C and S7). The features fall into three overall categories corresponding to $\Delta \Delta G$, $\Delta \Delta E$, and $s_{\text{exp}}$.

We here remind the reader that the model uses stability and conservation effects of all possible substitutions at a specific position even when predicting the effect of a specific variant. Thus, for example, when predicting the effect of a specific isoleucine to alanine substitution, the model will use as features the effect of changing that isoleucine residue to all other 19 protein-coding amino acids. As expected, however, the feature-importance calculations show that the model has its largest contribution from the specific substitution, with other substitutions playing a smaller role (Figure 3C). In addition to using information about the specific substitution, the average $\Delta \Delta E$ score also has a large feature importance, which we take to mean that the model uses this to determine whether a specific position is overall restrictive in the types of substitutions that have been seen during evolution. In addition to these effects, substitutions to positively charged amino acids (e.g., $\Delta \Delta E$ to arginine and histidine and $\Delta \Delta G$ to lysine) have greater than average contributions, and we speculate that these values provide additional structural context on whether, e.g., a position is buried or not (as substitutions to large positively charged residues are generally disfavored at buried positions). We note that effects of substitutions to histidine and asparagine have previously been shown to correlate most strongly with other substitutions at the same position (Gray et al., 2017).

Adding up the individual contributions of the features from the three classes ($s_{\text{exp}}$, $\Delta \Delta E$, and $\Delta \Delta G$), we find that the $\Delta \Delta E$ features have roughly twice the importance as $\Delta \Delta G$ features, with the total importance being substantially greater than from $s_{\text{exp}}$ (Figure 3C). Looking at the values for the specific variant predicted, the values are 11.8%, 5.7%, and 4.8% for $\Delta \Delta E$, $\Delta \Delta G$, and $s_{\text{exp}}$, respectively, although we note caveats in interpreting these numbers very precisely (Strobl et al., 2007). Thus, in line with analyzing simpler models using only a subset of the features (Figure 2B), this analysis shows that all three classes of features contribute to the performance of the final model, with $\Delta \Delta E$ carrying the greatest weight.

A global map of variant effects

The results above further support previous observations that analyses of sequence conservation are overall more informative than stability calculations to predict variant affects as probed by most MAVE experiments (Jepsen et al., 2020; Livesey and Marsh, 2020; Gerasimavicu et al., 2020; Cagliada et al., 2021). Nevertheless, they also show that stability calculations can help improve overall prediction accuracy and provide mechanistic insight into the relative roles of stability and conservation in explaining variant effects. Specifically, as argued previously
(Stein et al., 2019; Jepsen et al., 2020; Cagiada et al., 2021), we can use $\Delta \Delta E$ calculations as a proxy for capturing a broad range of effects on biological function, and $\Delta \Delta G$ as the subset that involves specifically stability and abundance.

We thus used the data that we collected to create global maps of variant effects, and analyzed these in terms of fitness, stability, and conservation. Inspired by previous work (Dunham and Beltrao, 2021), we visualized the 6,500 amino acid positions in our dataset with more than 15 values of $s_{\text{exp}}$, and corresponding values of $\Delta \Delta G$ and $\Delta \Delta E$ (Figure 4). Specifically, we used Uniform Manifold Approximation and Projection (UMAP) dimensionality reduction (McInnes et al., 2018) to represent all positions in a two-dimensional map where positions with similar profiles of $s_{\text{exp}}$, $\Delta \Delta G$, and $\Delta \Delta E$ are located close to one another (see STAR Methods). Finally, we color coded this map using the predicted and calculated position-averaged fitness scores, the position-averaged values of $\Delta \Delta G$ and $\Delta \Delta E$, values for $s_{\text{exp}}$, and the relative solvent accessibility (RSA) of the position (Figure 4).

As expected from the fact that the model was trained to predict the experimental data, the map colored by $s_{\text{calc}}$ (Figure 4A) and $s_{\text{exp}}$ (Figure 4B) closely resemble one another, although with a few differences highlighting the imperfection of the model. In particular, the maps reveal two regions (top left and bottom left) with amino acid positions where it appears that variants cause loss of function due to loss of stability. Comparison with the map indicating solvent accessibility (Figure 4F) shows clearly that many of these positions are buried inside the protein structure. The same comparisons also show that a group of positions near the bottom are enriched in positions where variants lose function in ways that are not due to stability but that can nonetheless be discovered using analyses of the evolutionary record. Finally, looking at the map colored by the average $s_{\text{exp}}$ scores (Figure 4E) shows similarities to both the stability map (Figure 4C) and solvent accessibility map (Figure 4F), in line with the fact that these are buried positions and that $s_{\text{exp}}$ captures basic physico-chemical aspects of amino acid chemistry and protein structure (Figure S2).

**DISCUSSION**

We have analyzed the relationship between experimental measurements of variant effects on protein function and computational analysis of protein stability and conservation. We collected over 150,000 measurements from multiplexed assays of variant effect in 29 proteins and compared them with predictions of changes in protein stability ($\Delta \Delta G$) and evolutionary conservation ($\Delta \Delta E$). Our goal was 2-fold. First, we aimed to examine how well these computed scores could predict variant effects, and, second, we wanted to shed further light on how often changes in protein stability may perturb protein function.

In general, and in line with previous observations, we find that our analysis of conservation of sequences through evolution ($\Delta \Delta E$) is more strongly correlated with the experimental measurements than predictions of changes in thermodynamic stability ($\Delta \Delta G$) (Figure 2A). We note here that, although each of the 39 experiments provides a systematic and comprehensive analysis of variant effects, they were conducted using very different assays for selection/screening and in different organisms. Together with uncertainties in the experimental and predicted values, this likely explains the substantial variation that we
observe in agreement between experimental scores and computed values. Thus, we refrain in general from analyzing individual proteins and variants and focus on the global agreement. As also previously observed (Cagiada et al., 2021), the VAMP-seq experiments that specifically probe protein abundance tend to be more strongly correlated with stability effects than measurements probing protein activity. Finally, we note that, while discrepancies between experimental measurements of variant effects and computational predictions can point to shortcomings in the prediction models, these may also reveal aspects of the protein’s function that the assay was not sensitive to.

Our global stability-conservation landscape (Figure 1A) reveals the interdependency of stability and conservation. As evolution tends to disfavor unstable proteins (Bloom et al., 2006; Echave and Wilke, 2017), we find that variants with high \( \Delta \Delta G \) tend to have large values of \( \Delta \Delta E \) and generally low fitness (low \( s_{\text{exp}} \)). The reverse relationship, however, is not true because being stable is a necessary, but not sufficient, criterion for being functional. Examining the stable variants (low \( \Delta \Delta G \)) we find that our analysis of the evolutionary record (\( \Delta \Delta E \)) can be used to predict with reasonable accuracy whether a variant retains function or not (Figure 1B). In our recent analysis of data probing activity and abundance in NUDT15 and PTEN, we showed that about half of the variants that lose function do so because they become unstable and are found at low abundance. In line with this, we find that roughly half of the variants that have high \( \Delta \Delta E \) scores also have high values of \( \Delta \Delta G \), and hypothesize that many of the remaining variants (low \( \Delta \Delta G \), high \( \Delta \Delta E \)) lose function because of substitutions in functionally important residues (Cheng et al., 2005; Echave, 2019; Cagiada et al., 2021).

Based on the partial correlation between both \( \Delta \Delta E \) and \( \Delta \Delta G \) with \( s_{\text{exp}} \), we built a prediction model with these parameters as input. The final position-context model reaches an accuracy of \( \Delta \Delta G \) and \( \Delta \Delta E \), where errors in the predicted values would have a greater impact. While our results overall conform to the expected relationship between \( \Delta \Delta E \) and \( \Delta \Delta G \), more subtle effects can modulate this relationship. For example, it has been observed that active-site residues may actually be destabilizing (Shoichet et al., 1995), and more detailed stability calculations may thus aid in detecting such effects.

The types of analyses presented here may also be used to improve or interpret experiments. For example, when constructing an experimental assay, it might be useful to compare experimental variant effects with calculations of \( \Delta \Delta E \) to examine whether the assay is sensitive to evolutionarily conserved functions. Also, it has recently been demonstrated that biophysical ambiguities prevent accurate predictions of how substitutions combine to affect phenotype (Li and Lehner, 2020), and we suggest that biophysical models and predictions of variant effects may help alleviate some of these ambiguities. Finally, a random forest model combining sequence and structural features has been used to predict pathogenicity of human missense variants (Ponzoni et al., 2020).

In summary, our large-scale analysis contributes to the mounting evidence for the important role for loss of stability in loss of function. In addition to the improvements observed when combining conservation analysis with structure-based stability predictions, our analyses also help pinpoint those variants that lose function due to stability. Such information helps provide mechanistic insight into specific variants and proteins but may also be a starting point for developing therapies. In particular, variants that lose function due to loss of abundance, whose intrinsic function is not otherwise perturbed, would be ideal targets for approaches that aim to restabilize or otherwise restore protein levels (Balch et al., 2008; Kampmeyer et al., 2017; Stein et al., 2019; Henning et al., 2021).

Limitations of the study

There are some limitations in our study. First, there may be biases and limitations arising from the specific choice of experimental data that we analyzed. We focused only on soluble
proteins, and our selection criteria further included availability of the experimental measurements and an experimentally determined structure of the protein. As proteins with known structures tend to have deeper multiple sequence alignments (Orlando et al., 2016), the latter requirement might lead to subtle biases in our analyses and represent more accurately calculated conservation scores. Second, the experimental data come from a range of assay types and organisms across the domains of life, and future analyses will be needed to examine how well our methods extrapolate to other proteins and assays. Third, our analysis of whether low-ΔΔG (structurally stable) but high-ΔΔE (evolutionarily constrained) variants are enriched in known active sites is currently based on a small number of proteins, and it remains unclear to what extent these observations hold more generally. Fourth, the method we use to predict protein stability is imperfect and has been benchmarked on a biased dataset (Frenz et al., 2020), leaving it unclear exactly how well it performs for the wider range of mutations analyzed here.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.cellrep.2021.110207.

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AUTHOR CONTRIBUTIONS


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RESEARCH AVAILABILITY

Lead contact
Further information and requests for code and data should be directed to and will be fulfilled by the Lead Contact, Kresten Lindorff-Larsen (lindorff@bio.ku.dk).

Materials availability
This study did not generate new unique reagents.

Data and code availability
- This paper analyses existing, publicly available data. A list of the datasets is available in Table S1. Data used in our analyses are available via https://doi.org/10.5281/zenodo.5647207 and https://github.com/KULL-Centre/papers/tree/main/2021/ML-variants-Hoie-et-al.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

METHOD DETAILS

\( \Delta G \) calculations using Rosetta
We searched the Protein Data Bank (Rose et al., 2010) using the BLASTp webserver (McEntyre J., 2002) with default settings using as query the sequences for which the MAVE was performed. If only a part of a protein is covered by mutagenesis data, we only searched for structures of that part. We chose structures by balancing highest available coverage and resolution, selecting structures solved using X-ray crystallography when available. All calculations were carried out using the Rosetta version with GitHub SHA 28f338acfb3bfcf87048bf38a04772486975dc83fa from July 2, 2020. We first relaxed the structures using the relax application and the following flags:
- fa_max_dis 9
- relax:constrain_relax_to_start_coords
- ignore_unrecognized_res
- missing_density_to_jump
- nstruct 1
- relax:coord_constrain_sidechains
- relax:cartesian
- beta
- score:weights beta_nov16_cart
- ex1
- ex2
- relax:min_type lbfgs_armijo_nonmonotone
- flip_hnq
- no_optth false

Subsequently, we carried out saturation mutagenesis to calculate for each single amino acid substitution using the Cartesian ΔΔG protocol and the beta_nov16_cart energy function with three iterations as previously described (Park et al., 2016; Frenz et al., 2020). Flags for the ΔΔG calculations were:

- fa_max_dis 9.0
- ddg::dump_pdb false
- ddg::iterations 3
- score:weights beta_nov16_cart
- missing_density_to_jump
- ddg::mut_only
- ddg:bbnbrs 1
- beta_cart
- ex1
- ex2
- ddg::legacy true
- optimize_proline true

Scores from the three iterations were averaged. Values of ΔΔG in Rosetta Energy Units were divided by 2.9 to bring them onto a scale corresponding to kcal/mol (Park et al., 2016).

Evolutionary conservation analysis using GEMME
We calculated evolutionary conservation scores (ΔΔE) using GEMME, a global epistatic model for predicting mutational effects (Laine et al., 2019), based on a multiple sequence alignment of homologs of each protein of interest. We used the sequence used in the MAVE experiment as input to HHblits (version 2.0.15 and settings -e 1e-10 -i 1 -p 40 -b 1 -B 20000) to search UniRef30_2020_03_hhsuite.tar.gz (Mirdita et al., 2017; UniProt Consortium, 2019; Steinegger et al., 2019). We de-gapped the alignment with respect to the MAVE sequence, removed sequences with 50% gaps and used the output alignment as input to GEMME, with default settings. Finally, we rank-normalized the output ΔΔE scores and scaled them to a [0,1] scale.

Collecting and normalizing data from MAVEs
We downloaded 39 data sets that had been generated by MAVEs from publicly available repositories including MAVEdb (Esposito et al., 2019) and a recent compilation of data (Livesey and Marsh, 2020) (see Table S1). We used the variant fitness scores as presented in the original publication including possible normalization. Three data sets (PS3, MAPK1 and Src) showed a reverse relationship between variant scores and ΔΔG and ΔΔE and were therefore reversed to match our convention of high scores for wild-type-like activity and low scores for low activity. We then rank-normalised and scaled these scores to a [0,1] range. We term these normalized scores $s_{\text{exp}}$.

We next merged each data set from the MAVEs with the corresponding ΔΔE and ΔΔG scores aligning the sequences based on using Biopython, pairwise2.align.globalds(target_seq.upper(), seq.upper(), MatrixInfo.blosum62, -3, -1).

Baseline substitution model
For each MAVE data set, we calculated a 20x20 amino-acid matrix containing the average of each the 400 possible Wild-type → Mutant variant scores (e.g. the average $s_{\text{exp}}$ for all valine to alanine substitutions). We then calculated a global substitution score matrix ($s_{\text{exp}}$) by averaging these matrices.

Features and random forest models
For each variant, we extracted in total 47 computational features. Depending on the model, each $s_{\text{exp}}$ was matched with its variant ΔΔG and ΔΔE score, as well as the 20 ΔΔG and 20 ΔΔE values corresponding to all available scores at the respective position, plus the mean ΔΔG and ΔΔE scores for the position. We note that this results in some redundancy in the use of the data, but simplifies the
We also included three global features from the baseline substitution model: (i) The mean MAVE score $\overline{s}_{\text{WT} \rightarrow \text{Mut}}$, (ii) $\overline{s}_{\text{WT} \rightarrow \text{Any}}$ was calculated as the mean for any substitution from the given wild-type amino acid, and (iii) $\overline{s}_{\text{Any} \rightarrow \text{Mut}}$ as the mean for mutating to the specified amino-acid from any wild-type. Thus, these three features correspond to (i) an entry in $\overline{s}_{\text{exp}}$ as well as the (ii) row-average, and (iii) column-average of $\overline{s}_{\text{exp}}$ (Figure S2).

We trained the Random Forest models using the RandomForestRegressor in Scikit-Learn (Pedregosa et al., 2011), with a mean-squared-error loss function, 150 trees and a minimum of 15 samples per leaf.

We first trained two models, using the variant $\Delta \Delta G$ or $\Delta \Delta E$ score only (1 feature). Next we trained a combined model using both features. Training of the position-context model was performed with all 47 available features. We iteratively trained the model and evaluated validation set performance in a leave-one-protein-out cross-validation, with the removal of all validation data sets for the selected protein from the training data set for each training round.

**Model analysis**

To extract feature importance from a globally-trained model, we trained a new random forest model using all 47 features and all available data sets. The feature co-correlation dendrogram was constructed by first calculating the Spearman correlation between each pair of features, which in turn was converted into a distance as $1 - r_S$, and used as input to build the dendrogram using SciPy (Virtanen et al., 2020). We measured partial dependence using PDPbox (Jiangchun, 2018).

**Relative solvent accessibility**

We used DSSP (Kabsch and Sander, 1983) to calculate the relative solvent accessibility (RSA) using the structures that we also used for the Rosetta calculations. The RSA was used in analysis but not in training.

**UMAP projection of MAVE positions**

To map variant scores to positions, we removed positions with fewer than 15 experimental $s_{\text{exp}}$ scores from the data set, and calculated position means for $s_{\text{exp}}$, $\Delta \Delta E$, $\Delta \Delta G$, $\overline{s}_{\text{WT} \rightarrow \text{Mut}}$ and RSA. Any missing $s_{\text{exp}}$ scores were then imputed using sklearn.impute.SimpleImputer from the amino-acid mean across all extracted positions, and all scores were then normalized to a 0–1 range. For computational efficiency we reduced the data set to 20 features using PCA before projecting the data into two dimensions (UMAP1 and UMAP2), using UMAP-learn with default settings (McInnes et al., 2018).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Quantitative analyses were performed with Python and SciPy (Virtanen et al., 2020). Stability calculations were performed using Rosetta (Leman et al., 2020) and sequence analyses using GEMME (Laine et al., 2019). All analyses and details are described in STAR Methods and linked code.