Predicting Functional Effects of Synonymous Variants: A Systematic Review and Perspectives

Zishuo Zeng1,2* and Yana Bromberg2,3*

1 Institute for Quantitative Biomedicine, Rutgers University, Piscataway, NJ, United States, 2 Department of Biochemistry and Microbiology, Rutgers University, New Brunswick, NJ, United States, 3 Department of Genetics, Rutgers University, Human Genetics Institute, Piscataway, NJ, United States

Recent advances in high-throughput experimentation have put the exploration of genome sequences at the forefront of precision medicine. In an effort to interpret the sequencing data, numerous computational methods have been developed for evaluating the effects of genome variants. Interestingly, despite the fact that every person has as many synonymous (sSNV) as non-synonymous single nucleotide variants, our ability to predict their effects is limited. The paucity of experimentally tested sSNV effects appears to be the limiting factor in development of such methods. Here, we summarize the details and evaluate the performance of nine existing computational methods capable of predicting sSNV effects. We used a set of observed and artificially generated variants to approximate large scale performance expectations of these tools. We note that the distribution of these variants across amino acid and codon types suggests purifying evolutionary selection retaining generated variants out of the observed set; i.e., we expect the generated set to be enriched for deleterious variants. Closer inspection of the relationship between the observed variant frequencies and the associated prediction scores identifies predictor-specific scoring thresholds of reliable effect predictions. Notably, across all predictors, the variants scoring above these thresholds were significantly more often generated than observed, which confirms our assumption that the generated set is enriched for deleterious variants. Finally, we find that while the methods differ in their ability to identify severe sSNV effects, no predictor appears capable of definitively recognizing subtle effects of such variants on a large scale.

Keywords: synonymous variants, effect predictors, variant frequency, variant functional effect, machine learning

INTRODUCTION

The vast majority of human genomic variation is accounted for by Single Nucleotide Variants (SNVs) (Bromberg et al., 2013). The roughly 10,000 variants in the coding region of every human genome that have no effect on the resulting product protein sequence are termed synonymous SNVs (sSNVs) (Shen et al., 2013). sSNVs are a product of the degeneracy of genetic code, where amino acids may be encoded by more than one codon. The effects of sSNVs on molecular functionality of the corresponding genes/proteins are often assumed to be minimal. However, earlier studies have argued that sSNVs are as likely to be pathogenic as non-synonymous variants (Chen et al., 2010). sSNVs have been implicated in many diseases, including pulmonary...
sarcoidosis, attention deficit/hyperactivity disorder, and cancer (Sauna and Kimchi-Sarfaty, 2011; Supek et al., 2014). Synonymous variants can disrupt transcription (Stergachis et al., 2013), splicing (Pagani et al., 2005), co-translational folding (Pechmann and Frydman, 2013), mRNA stability (Presnyak et al., 2015) (Figure 1), and cause a plethora of other functionally-relevant changes. In addition, sSNVs can affect transcription and splicing regulatory factors within protein coding regions (Plotkin and Kudla, 2011), thus modulating gene expression (Shabalina et al., 2013; Boël et al., 2016). There is also evidence of evolutionary constraint on both synonymous and non-synonymous variants, which plays a role in shaping codon bias (organism or tissues-specific codon set preference) (Stergachis et al., 2013). An informative experimental approach to evaluating functional effects of sSNVs is saturation genome editing followed by protein function assays (Findlay et al., 2014; Findlay et al., 2018). Unfortunately, there are exceedingly few reports of these experiments in the literature. While there has been a concerted effort in the field to evaluate the effects of non-synonymous single nucleotide variants (nsSNVs) (Mahlich et al., 2017) for the purposes of precision medicine, as well as improving basic understanding of concepts in molecular biology, interpretation of sSNVs is severely lacking. However, considering the significant number of observed synonymous variants, their possible effects, and the dire lack of their systematic experimental interpretations, there is a compelling need for a reliable sSNV effect computational predictor.

In this paper, we review the existing sSNV-effect predictors and apply them to a dataset containing observed and artificially generated sSNVs. Since there are few experimentally-determined SNVs with deleterious effects, and those that exist have been used as training or testing sets of the predictors, the cornerstone of this study is validating our data set assumption that deleterious sSNVs are enriched in the artificially generated set of variants. To support this assumption, in addition to previously published work, e.g., Stergachis et al., 2013, we show that the distributions of observed sSNVs by amino acids and codons are highly

FIGURE 1 | Possible mechanisms of sSNVs impact on biological function. Yellow triangles represent sSNV sites and the dashed lines indicate aberrant processes. sSNVs may affect (A) transcription factor binding, (B) splicing of pre-mRNA, (C) mRNA secondary structure and stability, (D) wobble-based tRNA binding, and (E) cotranslational folding (and thus the protein structure). Figure was created with BioRender.com.
non-random. We also demonstrate that existing predictor high-scoring variants are enriched among the artificially generated sSNVs, additionally validating of our assumption. We finally note that these predictors appear unable to definitely identify subtle effect sSNVs.

**METHODOLOGY OF THE PREDICTORS**

**SNV Predictors Vary by Targeted Variant Type, Training Data, and Descriptive Features**

We identified from the literature four sSNV-specific effect predictors: SilVA (Silent Variant Analyzer) (Buske et al., 2013), regSNPs-splicing (Zhang et al., 2017), DDIG-SN (Detecting Disease-causing Genetic SynoNymous variants) (Livingstone et al., 2017), and IDSv (Identification of Deleterious Synonymous Variants) (Shi et al., 2019). Additionally, we considered TraP (Transcript-inferred Pathogenicity) (Gelfman et al., 2017), which addresses both synonymous and intronic variants. Specifically, 1) SilVA was trained on 33 pathogenic and 785 neutral variants from 1000 Genomes Project (1000G) (Birney and Soranzo, 2015), using conservation scores, splicing, DNA, and RNA properties, 2) DDIG-SN and IDSv used positive data from the Human Gene Mutation Database (HGMD) (Cooper et al., 1998; Stenson et al., 2003; Stenson et al., 2009; Stenson et al., 2017) and negative data from 1000G (DDIG-SN) and VartiSNP (IDSv) (Schafaetsma and Vihinen, 2015) as negative data for training, described using features of translational efficiency and protein properties in addition to those used by SilVA, 3) regSNPs-splicing also used HGMD and 1000G data, but it considers sSNVs only in the context of mRNA splicing and protein function, while 4) TraP was trained on positive data combining SilVAs data with Online Mendelian Inheritance in Man (OMIM) (Hamosh, 2004) variants and negative data from control trios de novo variants. TraP uses transcript-affecting features, specific to intronic and synonymous variants.

As opposed to the sSNV-specific tools, more generic predictors, including CADD (Kircher et al., 2014), DANN (Quang et al., 2014), FATHMM-MKL (Shihab et al., 2015), and MutationTaster2 (Schwarz et al., 2014), evaluate synonymous, non-synonymous, regulatory and other kinds of variants. CADD was developed by training a support vector machine (SVM) to differentiate observed vs. simulated variants of all variant categories (Kircher et al., 2014). DANN attempts to capture nonlinear signals in (CADD-generated) variant data using a deep neural network (Quang et al., 2014). FATHMM-MKL is a Hidden Markov Model-based method integrating ENCODE (Consortium, 2012) functional annotations of SNVs to evaluate non-coding and synonymous variants (Shihab et al., 2015). MutationTaster2 (Schwarz et al., 2014) uses a naïve Bayes model trained on disease variants vs. variants from 1000G variants to evaluate all SNVs. Notably, these general-purpose predictors are heavily conservation-driven and may lack features to describe the subtle changes induced by sSNVs.

All predictors described here are machine learning-based [using random forests (RFs), SVMs, or deep neural network] and trained to predict pathogenicity, using different data and feature sets (Table 1). Supervised machine learning, often used for predicting variant effects, requires selecting a proper training/evaluation set, a number of relevant variant-, gene-, or disease-features, and an appropriate model for identifying feature patterns representative of variant effect/disease-association (Rost et al., 2016).

**Available Variant Sets Are Limited in Size and Reliability**

Association between genomic variants and diseases can be identified by carefully designed statistical tests, e.g., via Genome Wide Association Studies (GWAS) (Visscher et al., 2012). However, unequivocally identifying variants that cause disease are significantly more difficult; this is a particularly hard problem for sSNVs, which carry no corresponding protein sequence changes. Clinical or experimental validation of the causative relationships between genomic variation and disease is either infeasible altogether (as for polygenic disorders) or exceedingly difficult on a large scale due to the necessary resource and time investments. Instead, computational annotation of genomic variant pathogenicity (or simply functional effects) is a cost- and time-efficient substitute, providing a starting point for further experimental validation and discovery.

Most predictors described here (regSNPs-splicing, DDIG-SN, FATHMM-MKL, and MutationTaster2) collect variants identified as causative (positive) from HGMD. The latest version of HGMD (March 2017) comprises over 203,000 variants in over 8,000 genes, manually curated from scientific literature (Stenson et al., 2017). Despite its apparent utility, studies have questioned the reliability of HGMD data. George et al. (2007), for example, point out flaws like inconsistent mutation nomenclature and incomplete incorporation of all applicable data. Yue and Moult (2006) note that some mutations in HGMD are named causes of monogenic disease but are not fully penetrant, while Bell et al. (2011) question disease annotation of recessive variants. In a study of 1,000 exomes, Dorschner et al. (2013) note that only 16 of 585 of HGMD disease-causing variants were actually pathogenic, while in a subsequent study with 6,503 individuals, none of the identified 615 HGMD disease-causing variants were pathogenic ( Amendola et al., 2015). Other studies (Xue et al., 2012; Cassa et al., 2013) have shown that many disease-causing variants in HGMD are present in the relatively healthy 1000G individuals (Birney and Soranzo, 2015).

Other sources of positive training/testing data, including OMIM (used by TraP) and ClinVar (used by TraP, regSNPs-splicing, IDSv, CADD, MutationTaster2, and FATHMM-MKL) (Landrum et al., 2014), appear no more reliable. Notably, there is considerable inconsistency between the HGMD and OMIM (George et al., 2007). ClinVar’s entries from different sources often conflict between themselves (Landrum and Kattman, 2018), as the reliability of ClinVar’s data curation and workflow of medical interpretation has not been proven (Bauer et al., 2018). Substantial discordance between ClinVar and laboratory test results has also been reported (Gradishar et al., 2017).
Mutation databases vary drastically (George et al., 2007), not in the least because of experimental interpretation differences; e.g., roughly 17% of the variant effects reported by different laboratories carry contradictory clinical significance (Rehm et al., 2015). Labels of pathogenicity are not fixed, switching from disease to benign and back as evidence accumulates (Shah et al., 2018). As these binary labels also do not provide a quantitative measure of risk (Shah et al., 2018) or penetrance, the term “disease-causing” should be used with caution. One key problem in the field, and a reason for many of the above data limitations, is the absence of a gold standard for identifying disease-causing variants (Dorschner et al., 2013). Moreover, even the “silver-standard” available annotations are far and few between. In fact, while there are many known pathogenic mSNVs, there are currently much fewer known pathogenic sSNVs available: dbDSM (Wen et al., 2016) (including those from ClinVar, PubMed, NHGRI GWAS catalog (Welter et al., 2013), etc.) contains 1,289 pathogenic sSNVs, and HGMD contains roughly 900 pathogenic sSNVs (Livingstone et al., 2017). Arguably, this number is too small to build a generalizable model for evaluating tens of millions of the possible synonymous variants in human genome. Note that an additional problem is the absence of a true negative set of variants, i.e., those that have been verified to have no effect on protein function or no relationship to some disease (Bromberg et al., 2013).

Use of Allele Frequency to Approximate Variant Effect

SilVA was trained on 33 experimentally defined deleterious and 785 assumed neutral (observed in 1000G) variants. While the former set was very stringently selected, this small number of samples could hardly produce a generalizable model. Other predictors use less well curated data from available databases, but as such run into a problem of reliability. To supplement the lack of experimentally annotated variation, variant population frequency had been suggested as a sign of effect/pathogenicity; i.e., it is generally assumed that disease/effect variants are of low frequency (Gibson, 2012), although the precise threshold for “low” is unclear. Predictors (CADD, DANN, FATHMM-MKL, SilVA, regSNP-splicing) often filter out effect variants of higher frequency and/or neutral variants of lower frequency. CADD and DANN training data, for example, contains simulated human variants, appearing after human–chimpanzee divergence, labelled as the effect group (depleted by natural selection) and observed fixed or nearly fixed derived alleles as neutral (Kircher et al., 2014; Quang et al., 2014). Note although simulated variants are likely enriched in deleterious variants, and CADD scores have been shown useful in prioritizing variants in clinical settings (Amendola et al., 2015; Nakagomi et al., 2018; Van Der Velde et al., 2015), it is difficult to directly link the CADD predictions to pathogenicity (Kircher et al., 2014).

**TABLE 1 | Summary of sSNV-specific predictors.**

| Ref/Tool name | Training data | Model | Features | Performance |
|---------------|---------------|-------|----------|------------|
| (Buske et al., 2013) | 33 deleterious from literature, 785 neutral from one 1000 Genomes Project individual | Random forest with 1,001 trees and default number of features | 26 in total • conservation • RNA properties • DNA properties • Splicing | Dataset: 8 DM from literature and 752 NM from literature and 1000G. Result: DM’s scores ranked higher than NM’s |
| (Gelfman et al., 2017) | 75 DM from literature and OMIM and 402 de novo NM from control trios | Random forest with 1,000 trees, each with | 20 in total • Conservation • DNA properties • Splicing | Dataset: 66 DM and 4,418 NM from ClinVar. Result: AUC = 0.88 |
| (Zhang et al., 2017) | –655 DM from HGMD and –655 NM from 1000G | Random forest with 51 trees and 35 features at each node | 455 in total • Conservation • RNA properties • protein properties • splicing | Dataset: –325 DM from HGMD and 230 DM from ClinVar, –325 NM from 1000G and 4,535 NM from ClinVar. Result: For HGMD vs. 1000G data, AUC = 0.91 for variants in Splice Sites and AUC = 0.92 for all others For ClinVar data, AUC = 0.85 for variants in splice sites and AUC = 0.70 for the all others |
| (Livingstone et al., 2017) | 592 DM from HGMD and 10,925 putatively benign from 1000G | Support Vector Machine with radial function kernel | 54 in total (including all of the 26 features used in SilVA) • conservation • DNA properties • RNA properties | Dataset: 279 DM from HGMD and 4,945 NM from 1000G. Result: AUC = 0.85 |
| (Shi et al., 2019) | 300 DM from dbDSM and 300 NM from VarSNP | Random forest with 500 trees and 3 features at each split | 10 in total • Conservation • DNA properties • Splicing | Dataset: 153 DM and 5,178 NM from ClinVar. Result: AUC = 0.87 |

DM, disease/deleterious mutations; NM, neutral mutations; HGMD, human gene mutation database; 1000G, 1000 genome project; OMIM, online mendelian inheritance in man; AUC, area under the ROC curve (axes in Eqn. 1).
Allele frequency, however, is not necessarily correlated with variant effect, particularly when effect being considered is “function change” not “disease.” In an earlier study, we found that common [minor allele frequency (MAF) > 5%] non-synonymous variants were more often predicted to have a functional effect than rare (MAF < 1%) ones (Mahlich et al., 2017). Here a high-frequency allele may be beneficial/advantageous and on the way to becoming common, or slightly deleterious and on its way out (Bromberg et al., 2013). Moreover, trivially, allele frequency estimated from the sequenced genomes may be subject to change as the number of samples increases. Thus, 1) low allele frequency is not equivalent to having an effect and 2) although high frequency alleles are unlikely to be disease causing, they may have some impact. Additionally, and perhaps most fundamentally, note that the currently observed SNVs are unlikely a complete set of naturally occurring variants, i.e., many SNVs may be yet unseen.

Features Used Vary From Predictor to Predictor

A variety of features have been considered by predictors as described below. Note that the number of features used in existing predictors ranges from 26 (SilVA) to 1,281 (FATHMM-MKL).

Conservation

Evolutionary conservation, derived from multiple sequence alignments (MSAs) of homologous sequences (Niroula and Vihinen, 2016), is perhaps the most extensively used feature of variant-effect predictors. Commonly used DNA conservation scoring algorithms include GERP (Cooper et al., 2005), phastCons (Siepel et al., 2005), and PhyloP (Pollard et al., 2009) scores. GERP (Genomic Evolutionary Rate Profiling) is a statistical method identifying genomic constrained elements from MSAs. GERP uses a statistical model estimating species divergence times (Hasegawa et al., 1985) and a structural expectation maximization algorithm for phylogenetic inference (Friedman et al., 2002); the later GERP++ is a faster version of the original (Davydov et al., 2010). phastCons fits MSAs to phylogenetic hidden Markov models to identify conserved elements (Siepel et al., 2005). The major difference between phastCons and GERP is that the former models the size and distribution of conserved elements within an MSA, while the latter first individually assesses the conservation at a locus and then searches for clusters of highly conserved loci (Chen et al., 2010). PhyloP combines statistical tests and GERP to detect conservation and acceleration in nucleotide substitution rates (Pollard et al., 2009).

All variant effect predictors use at least one of these conservation scoring techniques (Tables 1, 2). DDIG-SN also additionally uses protein conservation as conserved protein positions are often structurally important (Ng, 2003), suggesting possible misfolding due to decreased rate of translation at the relevant codon. Similarly, sSNVs may lead to mistranslaction (Kramer and Farabaugh, 2006; Kramer et al., 2010; Komar, 2016) resulting in amino acid substitutions—a particularly problematic occurrence at conserved protein positions.

Conservation is a very important signature of variant effect. For example, for ClinVar’s missense dataset the solely-conservation-based component of CADD, GerpS (a derivative of GERP++), as well as PhastCons and PhyloP, attained ROC AUCs (area under the receiver operating characteristic curve) of over 0.82, while CADD’s ROC AUC was only slightly higher (0.93) (Kircher et al., 2014).

![TABLE 2 Summary of generalized SNV predictors.](image-url)
2014). In FATHMM-MKL’s cross-validation on coding variants, its ROC AUCs was 0.93 while the ROC AUCs for conservation scores alone was 0.91 (Shihab et al., 2015). Similar results are observed for DDIG-SN (DDIG-SN’s ROC AUCs = 0.85, PhyloP’s ROC AUCs = 0.76) (Livingstone et al., 2017) and TraP (TraP’s ROC AUCs = 0.88, GERF++’s ROC AUCs = 0.87) (Gelfman et al., 2017) datasets. These results suggest that over billions of years of evolution, nature’s laboratory has tested and discarded most of the detrimental variants. However, it is important to note that functional tuneability, i.e., development of new or environment-specific versions of functions is an ongoing process, which requires the presence of variants in positions of all levels of conservation, in any given snapshot of a population (Miller et al., 2017; Miller et al., 2019).

DNA Properties
The DNA properties describing the biological effects of sSNVs include but are not limited to localization to transcription factor (TF) binding sites, overall GC content of genes and genomes, and CpG island locations (cytosine followed by guanine in 5’ to 3’ direction). In more detail: many studies have shown that coding exons can serve as regulatory elements for transcription (Lang et al., 2005; Khan et al., 2012); i.e., roughly 15% of the human genome codons both code for amino acids and specify TF recognition (Stergachis et al., 2013). Thus, synonymous variants in TF-relevant codons can affect TF binding and alter gene transcription rates. Exonic and the flanking intronic region GC architectures can affect DNA methylation and exon recognition (Gellman et al., 2013). Additionally, CpG sites often host DNA methylation (Bernstein et al., 2007), playing an important role in gene transcription (Gelfman et al., 2013). As mutation rates at CpG dinucleotides are an order higher than at other sites (Nachman and Crowell, 2000), sSNVs can thus alter methylation patterns by disrupting site-specific GC architectures.

All predictors covered in this manuscript, except regSNPs-splicing, incorporate one or more of these DNA properties (Tables 1, 2).

RNA Properties
Codon bias. The preference (frequency of use) of particular codons by specific organisms or tissues is termed codon bias. Codon bias correlates with and informs gene expression levels (Coghan and Wolfe, 2000; Carbone et al., 2003; Dos Reis et al., 2003; Boël et al., 2016; Komar, 2016), translation rate (Sørensen et al., 1989), as well as protein structure (Zhou et al., 2009) and cotranslational folding (Pechmann and Frydman, 2013; Buhr et al., 2016). There are many different metrics describing codon bias including codon adaptation index (Sharp and Li, 1987), synonymous codon usage order (Angellotti et al., 2007), relative synonymous codon usage (Sharp and Li, 1987), etc. Surprisingly, only SilVA and DDIG-SN have considered codon bias as a factor in their models (Table 1).

A related factor governing translation rate is the supply of tRNA during translation. Note that tRNA concentrations are different across organisms and that some organisms lack certain tRNA altogether, supplementing the necessary functionality via third position wobble (Novoa et al., 2012). It is hypothesized that codon composition in coding regions coevolved with tRNA abundances to reach the desired translation rates (Plotkin and Kudla, 2011). tRNA adaptation index (tAI) (Reis et al., 2004), used only by IDS V (Table 1), is a measure aimed to describe codon bias from the perspective of tRNA supply and demand.

A potentially important feature also missing from all predictors is codon autocorrelation. In codon autocorrelated sequences, same codons follow each other in sequence, i.e., sequence AAABB is more autocorrelated (less anticorrelated) than sequence ABABA, where A and B are two codons of the same amino acid (Cannarozzi et al., 2010). Autocorrelated yeast transcripts are translated faster than anticorrelated ones (Cannarozzi et al., 2010) and many prokaryotes modulate translation through codon correlation (Guo et al., 2012). Thus, using codon correlation may help characterizing sSNV effect.

mRNA structure, stability, and abundance. sSNVs can alter mRNA secondary structure, thus impacting translational efficiency and mRNA decay rate (Hunt et al., 2014), which, in turn, impacts protein production (Komar, 2016) and abundance (Maier et al., 2009). mRNA sequences are more stable than random collections of nucleotides (Sefiens, 1999), suggesting that mRNA stability is evolutionarily selected to accommodate sufficient levels of translation before decay. The secondary structure of mRNAs harbors conserved elements (Meyer, 2005) and is tightly intertwined with GC content and codon usage. In fact, an earlier study found that 26% of the expressed genes display differential mRNA stability across individuals (Duan et al., 2013). In these genes, higher GC3 (G or C at the third position of the codon) percentage correlated with higher mRNA stability. This finding is in line with the fact that among the different SNVs, G and C alleles generally result in higher mRNA stability than A and T alleles (Duan et al., 2013). Furthermore, stability is enhanced in mRNA sequences enriched in optimal codons corresponding to tRNAs of higher concentrations (Presnyak et al., 2015).

A number of in silico tools have been developed to predict the mRNA structure and stability, including mFold (UNAFold) (Zuker, 2003; Markham and Zuker, 2008), remuRNA (Salari et al., 2012), KineFold (Xayaphoummine et al., 2005), and RNAfold (ViennaRNA package) (Hoefacker, 2003). Note, however, that RNA molecules are very thermodynamically flexible and can take on many possible structures. Thus, the predicted RNA structure and its stability depend on the pre-set prediction strategy, which can be aimed to find the minimum free energy structure, the structure closest to other possible structures, or to maximize expected prediction accuracy, which is difficult for RNAs longer than 500 nucleotides (Lorenz et al., 2016). Consequently, the prediction of RNA structure and stability is inherently uncertain. Among all the sSNV predictors, only SilVA and DDIG-SN use predictive tools to compute the variant-induced changes of energy and structures in pre-mRNA and mature mRNA sequences (Table 1).

Note that sSNVs, as well as other variant types (Shah and Gilchrist, 2010), are particularly relevant to functionality of highly expressed genes. Thus, the Genotype-Tissue Expression
Splicing Properties
mRNA splicing is a major predictive feature in some predictors, especially regSNPs-snipping and IDSv. It is estimated that up to 15% of disease SNVs cause aberrant splicing (Krawczak et al., 1992). SNVs can impact exonic splicing enhancers (ESEs) and silencers (ESSs), short DNA sequence motifs that promote or suppress splicing of pre-mRNA by binding to SR proteins (proteins with long repeats of serine and arginine) (Wang and Burge, 2008). Moreover, sSNVs can change the affinity of pre-mRNA to spliceosomes, leading to false recognition of exon-intron boundaries and producing abnormal mRNAs and dysfunctional proteins (Bali and Bekb, 2015). Taken together, the sSNVs’ potential of disrupting splicing is the likely reason for slower evolution at within-ESE sites (Parmley, 2005).

Predictors describe the potential impact of sSNVs on splicing by relying on the identified putative ESE and ESS motifs. Identification of these motifs and the corresponding splicing regulatory proteins has been an ongoing experimental and computational effort (Wang and Burge, 2008; Shepard and Hertel, 2009); identified motifs and regulatory proteins are available via public repositories (Desmet et al., 2009; Giulietti et al., 2013; Xing et al., 2016). Tools such as SPANR (Splicing-based Analysis of Variants) (Xiong et al., 2015), have also been developed to predict the splicing effects of SNVs. Splicing is considered by all sSNV-specific predictors, although represented via different values.

Protein Properties
One often overlooked aspect in evaluating sSNV effect is the protein structure. Rare codon variants of frequent synonymous codons may slow down the translation rate due to low concentration of tRNAs, slow or stop the elongation of the peptide chain (Zhang et al., 2009), and influence co-translational folding (Kimchi-Sarfaty et al., 2007; Pechmann and Frydman, 2013). Cotranslational folding is closely related to the formation of protein secondary and tertiary structures (Holtkamp et al., 2015); alpha-helix formation can occur in the ribosomal tunnel (Komar, 2009), while tertiary structure formation may take place before the protein completely exits the ribosome (Zhang and Ignatova, 2011). Translationally fast codons are enriched for alpha helices, while beta strands and coil regions prefer translationally slow codons (Thanaraj and Argos, 1996). Optimal codons are enriched in buried and structurally important sites but are negatively correlated with solvent accessible sites (Zhou et al., 2009). Pathogenic sSNVs are generally enriched within the buried sites, intrinsic disorder regions, and alpha-helices, as well as in exons overlapping with known or predicted protein family domains (Zhang et al., 2017). These findings suggest that protein structure should be considered when modelling the effects of sSNVs. However, only regSNPs-snipping and DDIG-SN predictors incorporate protein structural information (Table 1).

EVALUATION OF THE PREDICTORS
Collecting the Evaluation Data Set
sSNV effect predictor evaluation is hampered by three major problems: 1) there is no clear definition of neutral and effect variants and 2) available neutral/effect experimental evaluations are limited, and 3) most have been used in predictor development. Here, we created our own data set of variants for evaluation purposes as follows: we collected the observed sSNVs [all non-singleton sSNVs that have been observed in either 1000G, ExAC (Lek et al., 2016), or gnomAD (Karczewski et al., 2019)] and the generated sSNVs (all possible sSNVs in human genes, excluding observed and singleton sSNVs); we thus extracted 1,362,607 observed and 24,008,961 generated sSNVs. For evaluation purposes, we randomly selected 1,362,607 generated variants from our set to create a balanced observed/generated variant Test set (details in Supplementary Material).

There are multiple equally valid reasons for why nearly 95% of all possible sSNVs are not observed; the most obvious ones are technical, i.e., insufficient data or sequencing technology bias, and evolutionary, i.e., purifying selection, genetic drift, and genetic hitch-hiking (Smith and Haigh, 1974). As per the latter, we assume that drastically deleterious variants, which would be eliminated on a population scale due to purifying selection, are significantly more frequent in the set of generated sSNVs than in observed ones. However, the former suggests that we may have simply not (yet) sequenced many of the un-observed (generated) variants, which are actually equivalent in potential effect to observed ones. Notably, since a large proportion of discovered sSNVs are singletons (Lek et al., 2016), an equivalent proportion of similarly neutral or mild-effect variants can likely be found on the other side of the “sequencing barrier,” i.e., they have yet to be sequenced. Moreover, different categories of variants vary in the likelihood of being observed. For example, according to the ExAC project, the discovery of CpG transitions (C > T, where C is followed by G) is likely close to saturation, while additional transversion and non-CpG transitions are yet to be identified (Lek et al., 2016).

We observe that 1) most of the large effect variants are likely in the generated set and either 2a) they make up much of that set, i.e., the generated set contains mostly effect variants, or 2b) there are relatively few of them, i.e., the distribution of effect and neutral variants is roughly equivalent across the generated and observed variants. Note that if (2a) is true, we expect that a precise and sensitive sSNV effect predictor should be able to differentiate the observed sSNVs from the generated ones, while (2b) would mean that the same predictor would produce similar effect distributions.

Note that our Test set data are collected in a somewhat similar, but ultimately very different, way as CADD’s (and DANN’s) training data. CADD’s observed variants are the fixed or nearly fixed alleles at sites where human genes are different from the inferred human-chimpanzee ancestor and thus may encompass
our excluded observed singletons. CADD’s simulated variants follow an estimated de novo mutation rate since human-chimpanzee divergence, and thus are a subset of all our variants, including generated, observed, and singletons. Importantly, even with down-sampling of generated variants to create a balanced set, our Test set is much larger (~2.8 million) and more broadly defined than CADD’s strictly curated training set (~100,000).

We calculated the enrichment of observed sSNVs relative to generated sSNVs separately by amino acid (Figure 2A) and codon (Figure 2B) type. We observe that the distribution of naturally occurring sSNVs is non-random across amino acids and codons. Thus, over a fifth of all tyrosine (Y) and histidine (H) codons in our genome is affected by sSNVs, as compared to roughly 8% of lysine (K) codons. Curiously, the most mutated codons are threonine ACG, serine TCG, and proline CCG (> 43% of each is affected by an sSNV) and alanine GCG (37%). Thus, the CG end-of-codon nucleotide pair seems to indicate least stable codons. On the other hand, the isoleucine ATA codon is almost never mutated (~1%), suggesting that it is preferentially maintained as error free. Moreover, the enrichments of observed sSNVs by amino acids (or codon) are not proportional to the abundance amino acids (or codon) in human transcriptome. The amino acids (e.g., Y, H, N, D) and codons (e.g., ACG, TCG, CCG, GCG, TAC, CAC) with high enrichment of observed sSNVs are those of low abundances. This decidedly non-random distribution of variants across codons and amino acids strongly suggests that our generated and observed variants are likely indeed different from the evolutionary, and thus likely effect, perspective.

**Predictors Do Not Distinguish Observed and Generated sSNVs**

To the best of our knowledge, our collection of tools (CADD, DANN, MutationTaster2, FATHMM-MKL, SilVA, TraP, DDIG-SN, regSNP-splicing, and IDSv) make up a complete set of publicly available methods for sSNV analysis. We first evaluated (Figure S2) the ability of all predictors (except regSNP-splicing, which was not functional at the time of writing) to differentiate 50,000 observed and 50,000 generated sSNVs (Supplementary Materials). We did not include IDSv for more further analysis because its performance was similar to that of other predictors and it was not available for running it locally or online for the entire set of our variants. Unfortunately, we also had to exclude MutationTaster2, which experienced server problems when running large batches of data.

We used CADD, DANN, FATHMM-MKL, SilVA, TraP, and DDIG-SN to make predictions for our complete variant Test set. We calculated the fraction of consensus binary predictions (FCBP; i.e., the number of predictions agreed upon) for all pairs of predictors and the correlation between scores (Figure 3B). As per CADD creators (https://cadd.gs.washington.edu/info), it is hard to threshold its raw scores, while the recommended neutral/deleterious cutoff for phred-scaled scores is 15. For the rest of the predictors, we used 0.5 as the binary threshold (> 0.5 is deleterious). We observed (Figure 3A) that the CADD and other sSNV-specific predictors agree with each other because their scores are mostly low (Figures 3F–H). Scores from general-purpose predictors do not have high correlation with
Evaluating Synonymous Variant-Effect Predictors

Zeng and Bromberg

October 2019 | Volume 10 | Article 914

Frontiers in Genetics | www.frontiersin.org

sSNV-specific predictors. Meanwhile, DANN and FATHMM-MKL did not agree with others or between themselves. This lack of agreement across the Test set indicates that, in the best case, predictors are orthogonal, correctly identifying a different subset of variants each or, in the worst case, they are mostly unable to recognize effect. Curiously, for each predictor, the distributions of sSNV scores of observed and generated variants were very similar (Figure 3), i.e., predictors disagreed between themselves and with our dataset labels. Note that since the data is large, statistical tests to establish their difference could easily achieve significance and may not be meaningful (Kim and Bang, 2016). Instead, we directly evaluated predictor ability (Table 3) to differentiate the two types of variants using ROC AUCs (Eqn. 1).

\[
TPR = \frac{TP}{TP + FN} \quad FPR = \frac{FP}{FP + TN}
\]

FIGURE 3 | Predictor scores correlate somewhat, but do not differentiate observed vs. generated sSNVs. Panel (A) shows the amount of agreement (i.e., FCBP) for any pair of predictors. High FCBP values indicate that two predictors agree in assigning binary (neutral/deleterious) predictions to variants. Panel (B) shows the Pearson correlations among the prediction scores. (C-I) Violin/box plots of prediction score distributions across predictors: CADD raw, CADD phred-scaled, DANN, FATHMM-MKL, SIWA, TraP, and DDIG-SN, respectively.

TABLE 3 | AUCs of the predictors on sSNVs and nsSNVs.

|          | Observed vs. generated sSNVs | Observed vs. generated nsSNVs |
|----------|-----------------------------|-------------------------------|
|          | AUC on Test set | Average of AUCs ±SD * |          |                         |                       |
| CADD raw score | 0.518               | 0.517±0.00012 | 0.564         |
| CADD phred-scaled score | 0.518               | 0.518±0.0013 | 0.564         |
| DANN      | 0.506               | 0.506±0.0023 | 0.491         |
| FATHMM-MKL | 0.540               | 0.540±0.0013 | 0.555         |
| SIWA      | 0.527               | 0.527±0.0009 |                       |
| TraP      | 0.496               | 0.496±0.0038 |                       |
| DDIG-SN   | 0.535               | 0.535±0.0012 |                       |

*Test set was sampled 20 times (each with 100,000 samples) and calculated the resulting standard errors of ROC AUCs (Table 3).
Predictor Performance Is Only Slightly Better for nsSNVs Than for sSNVs

As mentioned previously, the unexpected inability of predictors (Figure 3) to differentiate observed and generated variants may indicate either the inappropriateness of the data set for the evaluation task or limited predictor abilities. The latter may be related to the specific variant type; i.e., general-purpose predictors, such as CADD and FATHMM-MKL, are good at analyzing non-synonymous variants (Kircher et al., 2014; Shihab et al., 2015), but they may be less sensitive to effects of synonymous variants. To evaluate this possibility, we randomly selected 500,000 each observed and generated non-synonymous variants from dbNSFP (Liu et al., 2011; Liu et al., 2016) and extracted their associated predictor scores (see Supplementary Material). Briefly, an nsSNV was considered observed if it was reported by either 1000G, ExAC, or gnomAD; otherwise it was deemed a generated nsSNV. While some of the predictors were slightly better at differentiating generated from observed nsSNVs (Figure 4, Table 3) than sSNVs, their performance was still not up to the expectations. We also calculated FCBP (Figure 4A; cutoffs as above) and score correlation (Figure 4B) to find that CADD, DANN, and FATHMM-MKL have a considerably higher agreement on nsSNVs than on sSNVs (Figure 3A).

Inferring a Predictor Scoring Threshold From Prediction of Common Variant Effects

The predictor inability to differentiate observed and generated variants may also be due to the difficulty of defining effect threshold; i.e., variants of low effect are harder to precisely annotate, both computationally and experimentally, and can be equally well classified as effect or neutral. In an effort to increase resolution between the two, predictors often link high allele frequency to absence of effect. In fact, CADD, DANN, FATHMM-MKL, SiV A, and regSNP-splicing effectively label high allele frequency variants as neutral. Taken further, TraP scores were reported (Gelfman et al., 2017) to have negative correlation (~0.51) with bin-average ExAC allele frequencies (Lek et al., 2016). Note that, as mentioned above, this reasoning side-steps evolutionary flow where common (not yet fixed or removed) variants may be advantageous or damaging. To further elaborate on allele frequency relationship with effect predictions, we obtained frequency data from multiple sources (1000G, ExAC, and gnomAD, see Supplementary Material) for our observed variants. Notably, we saw no correlation, positive or negative, between allele frequency and any predictor score (Figure 5). This observation highlights predictor binary classification abilities rather than a continuous spectrum of effect.

For some of the predictors (CADD, SiV A, TraP, DDIG-SN) high scoring variants were overwhelmingly of low frequency. At the same time, many of the low frequency variants were low scoring. Assuming that the predictor scores can be used as reliable indicators of common variant neutrality (low scoring), this result reinforces the idea that low frequency variants are as likely to be pathogenic/effect as neutral/benign. Furthermore, common variant score distributions could help approximate the predictor thresholds of effect. Thus, while variants scoring above a certain threshold can be considered to have an effect, below this threshold binary predictor resolution is questionable.

Predictor thresholds were chosen as the score below which most (99%) of the common variants (allele frequency >0.01) reside (Figure 5). Thus, scores above this threshold indicate effect, while scores below the threshold could be effect or neutral.
We further applied the selected thresholds to both observed and generated sSNVs (Table 4). We define resolution (Eqn 2, where “N” stands for number) as a predictor’s ability to capture the enrichment of deleterious variants above threshold.

\[
\text{resolution} = \frac{N_{\text{sSNVs above the threshold}}}{N_{\text{observed sSNVs}}} \times \frac{N_{\text{generated sSNVs above the threshold}}}{N_{\text{generated sSNVs}}} \tag{2}
\]

The resolutions were greater than one for all the predictors, with CADD attaining the highest resolution (> 2). Note that only a small fraction of variants in both sets scored above the threshold, but since the total number of generated variants is nearly 18 times higher than the number of observed variants, the estimated number of potential identifiable-deleterious sSNVs is only an order of magnitude less than ALL observed sSNVs (~475K vs. ~1.3M). These results suggest that the generated set indeed contains many more deleterious variants than the observed set and that a new predictor train to recognize these differences may identify deleterious variants more reliably than existing methods.

|                         | % Above-the-threshold sSNVs in observed | % above-the-threshold sSNVs in generated | Resolution |
|-------------------------|----------------------------------------|-----------------------------------------|------------|
| CADD raw score          | 0.871                                  | 1.981                                   | 2.274      |
| CADD phred-scaled score | 0.868                                  | 1.979                                   | 2.280      |
| DANN                    | 1.594                                  | 2.156                                   | 1.352      |
| FATHMM-MKL              | 1.639                                  | 2.522                                   | 1.538      |
| SIVA                    | 4.902                                  | 6.015                                   | 1.227      |
| TraP                    | 1.764                                  | 2.414                                   | 1.368      |

**CONCLUSION**

Training data is perhaps the most critical component for a machine learning-based variant-effect-predictor. Most sSNV effect predictors we reviewed, retrieved training data from disease mutation databases, such as HGMD and ClinVar. Disease-causing variants can be thought of as severely functionally deleterious, although non-disease variants could also be deleterious or beneficial. Moreover, identifying an sSNV as disease causing, as opposed to associated with disease, is extremely difficult, if not impossible. In fact, studies have revealed flaws of existing disease mutation databases, which may further undermine the reliability of the contained data. Progress in saturation genome mutagenesis may improve data availability in the near future. Currently, however, there is no publicly available, sufficiently large collection of variants with experimentally validated effect annotations that can be used for building a generalizable sSNV effect-predictor.

The lack of gold standard data also prevents proper evaluation of the predictors. Here, we proposed a Test set of observed and generated sSNVs. We assumed that the generated set is enriched for deleterious sSNVs due to purifying selection and expected the predictors to differentiate these from the observed variants. However, the predictor performance on this data was below our expectations. Note that predictor scores for the variants in our set were poorly correlated and the amount of binary prediction agreement was limited. This observation suggests that predictions may be biased by shared input features, but do not sufficiently well indicate variant effect. We proposed a scoring threshold to separate reliable predictions from the highly uncertain ones for each of the predictor. With the thresholds identified, we further observed that all predictors had significantly more reliably identified sSNVs in the generated set than in observed set, in line with our earlier expectations of the quality and contents of the
Test set. However, the inability of the predictors to clearly identify effect variants below the severity threshold, suggests that more work is necessary to understand sSNV effects.

We note that our Test set is not a gold-standard testing set and is only appropriate for predictor testing only if our underlying biological/data distribution assumptions hold. Thus, we cannot make concrete recommendations of best-practice prediction tools. However, our results clearly indicate that the predictions are highly correlated across sSNV-specific methods, i.e., there is little difference between using SiVA, DDIG-SN, or TraP. On the other hand, outputs of general purpose-predictors (CADD, DANN, and FATHMM-MKL) do not correlate as well. Of these, CADD phred-scaled scores are least likely to classify common variants as having a large effect; i.e., CADD high scores may be deemed reliably non-neutral. Note, however, that this does not mean that CADD low scores indicate variant neutrality – a necessary distinction that evades much of the variant effect literature.

Looking forward to a future sSNV effect-predictor, we note that comparing observed vs. generated, rather than effect vs. no-effect, variants drastically increases the amount of data useful for training. We also note that this variant collection will need further filtering to address the problem of false positives, i.e., the yet-to-be-observed generated variants. Moreover, the transition from observed to no-effect and from generated to effect annotations will not be trivial. As mentioned earlier, while severe effect variants are likely predominantly confined to the generated set, the mild effect variation is probably distributed throughout both observed and generated collections. Despite these difficulties, the observation that existing predictors identify more higher-scoring effect variants in the generated data, suggests that the effect signal can indeed be learnable by models trained to differentiate observed vs generated variants. Thus, a model using the previously mentioned set of features, possibly in combination with an ensemble of (orthogonal, as evaluated above) existing classifiers, may provide a more reliable description of variant effects.

DATA AVAILABILITY STATEMENT

Public available datasets were analyzed in this study. Human transcripts and genomic coordinate information (GRCh37) can be found at https://gwr37.ensembl.org/biomart/martview/e15159959ac5f1b72adec301bb7e02e59. DANN scores can be found at https://cbl.ics.uci.edu/public_data/DANN/. TraP scores can be found at http://innovation.columbia.edu/technologies/cu17233_supplementary-material. FATHMM-MKL scores can be found at https://github.com/HASihhab/fathmm-MKL. ANNOVAR annotation tool can be found at http://annovar.openbioinformatics.org/en/latest/. dbNSFP annotation tool can be found at https://sites.google.com/site/jpopen/dbNSFP/. DDISN-SN server is at http://compbio.cs.toronto.edu/silva/. MutationTaster2 server is at http://www.mutationtaster.org/StartQueryEngine.html. IDSv can be found at http://bioinfo.ahu.edu.cn:8080/IDSv. Our observed/generated data with predicted scores can be downloaded at https://doi.org/10.5281/zenodo.3471642.

AUTHOR CONTRIBUTIONS

ZZ and YB contributed to the idea conception, analysis design, literature review, and manuscript writing. ZZ conducted data collection, analysis, and visualization.

FUNDING

ZZ and YB Were Supported by the NIH U01 GM115486 Grant.

ACKNOWLEDGMENTS

We would like to thank to thank Dr. Junfeng Xia (Anhui University) for providing IDSv predictions, Dr. Martin Kircher (Berlin Institute of Health) for providing CADD training data, Dr. Jana Marie Schwarz and Dr. Dominik Seelow (both from Charité Berlin) for technical support with MutationTaster2, and Dr. Yunlong Liu (Indiana University) for technical support with DDIG-SN. Particular thanks go to all Bromberg lab members (Dr. Chengsheng Zhu, Dr. Maximillian Miller, Dr. Ariel Aptekmann, Dr. Adrienne Hoarfrost, Dr. Kenneth McGinness, Yannick Mahlich, and Yanran Wang, all Rutgers) for their constructive discussion and advice. We also acknowledge people of the Rutgers Office of Advanced Research Computing (OARC), and particularly Kevin Abbey and Galen Collier, for providing technical support with and access to the compute cluster and associated research computing resources necessary for the work reported here. Finally, we would like to thank all researchers that deposit their data into public databases.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2019.00914/full#supplementary-material

REFERENCES

Amendola, L. M., Dorschner, M. O., Robertson, P. D., Salama, J. S., Hart, R., Shirts, B. H., et al. (2015). Actionable exomic incidental findings in 6503 participants: challenges of variant classification. Genome Res. 25, 305–315. gr. 183483.114. doi: 10.1101/gr.183483.114
Angelotti, M. C., Bhuiyan, S. B., Chen, G., and Wan, X.-F. (2007). CodonO: codon usage bias analysis within and across genomes. Nucleic Acids Res. 35, W132–W136. doi: 10.1093/nar/gkm392
Bali, V., and Bebok, Z. (2015). Decoding mechanisms by which silent codon changes influence protein biogenesis and function. Int. J. Biochem. Cell Biol. 64, 58–74. doi: 10.1016/j.biocel.2015.03.011
Bauer, P., Karges, E., Oprea, G., and Rolfs, A. (2018). Unmet needs in human genomic variant interpretation. Genet. Med. 20, 376–377. doi: 10.1038/gim.2017.187
Bell, C. J., Dinwiddie, D. L., Miller, N. A., Hateley, S. L., Ganusova, E. E., Mudge, J., et al. (2011). Carrier testing for severe childhood recessive diseases by next-generation sequencing. Sci. Transl. Med. 3, 65ra4–65ra4. doi: 10.1126/scitranslmed.3001756
Landrum, M. J., Lee, J. M., Riley, G. R., Jiang, W., Rubinstein, W. S., Church, D. M., et al. (2014). ClinVar: public archive of relationships among sequence variation and human phenotype. Nucleic Acids Res. 42, D980–D985. doi:10.1093/nar/gkt1113

Lang, G., Gombert, W. M., and Gould, H. J. (2005). A transcriptional regulatory element in the coding sequence of the human Bcl-2 gene. Immunology 114, 25–36. doi:10.1111/j.1365-2657.2004.02073.x

Lek, M., Karczewski, K. J., Minikel, E. V., Samocha, K. E., Banks, E., Fennell, T., et al. (2016). Analysis of protein-coding genetic variation in 60,706 humans. Nature 536, 285–291. doi:10.1038/nature19057

Liu, X., Jian, X., and Boerwinkle, E. (2011). dbNSFP: a lightweight database of human nonsynonymous and splice-site SNVs. Hum. Mutat. 37, 235–241. doi:10.1002/humu.22932

Livingstone, M., Folkman, L., Yang, Y., Zhang, P., Mort, M., Cooper, D. N., et al. (2017). Investigating DNA-, RNA-, and protein-based features as a means to discriminate pathogenic synonymous variants. Hum. Mutat. 38, 1336–1347. doi:10.1002/humu.23283

Maier, T., Güell, M., and Serrano, L. (2009). Correlation of mRNA and protein expression in complex biological samples. FEBS Lett. 583, 3966–3973. doi:10.1016/j.febslet.2009.10.036

Markham, N. R., and Zuker, M. (2008). UNAFold. Bioinformatics 3-31. doi:10.1093/nar/gkh923

Miller, M., Bromberg, Y., and Swint-Kruse, L. (2017). Computational predictors fail to identify amino acid substitution effects at rheostat positions. Sci. Rep. 7, 41329. doi:10.1038/srep41329

Nachman, M. W., and Crowell, S. L. (2000). Estimate of the mutation rate per nucleotide in humans. Genetics 156, 297–304.

Nakagomi, H., Hoshikawa, H., Inoue, M., Hirotsu, Y., Amemiya, K., Sakamoto, L., et al. (2018). Combined annotation-dependent depletion score for BRCAl/2 variants in patients with breast and/or ovarian cancer. Cancer Sci. 109, 453–461. doi:10.1111/cas.13464

Ng, P. C. (2003). SIFT: predicting amino acid changes that affect protein function. Nucleic Acids Res. 31, 3812–3814. doi:10.1093/nar/gkg509

Nisula, A., and Vihinen, M. (2016). Variation interpretation predictors: principles, types, performance, and choice. Hum. Mutat. 37, 579–597. doi:10.1002/humu.22987

Novoa, E. M., Povon-Eternod, M., Pan, T., and De Poulplana, L. R. (2012). A role for tRNA modifications in genome structure and codon usage. Cell 149, 202–213. doi:10.1016/j.cell.2012.01.050

Pagni, F., Raponi, M., and Baralle, F. E. (2005). Synonymous mutations in CTRF exon 12 affect splicing and are not neutral in evolution. Proc. Natl. Acad. Sci. 102, 6368–6372. doi:10.1073/pnas.0502288102

Parmley, L. J. (2005). Evidence for purifying selection against synonymous mutations in mammalian exonic splicing enhancers. Molecular biology and evolution 23, 301–309. doi:10.1093/molbev/msj035

Pechnmann, S., and Frydman, J. (2013). Evolutionary conservation of codon optimality reveals hidden signatures of cotranslational folding. Nat. Struct. Mol. Biol. 20, 237. doi:10.1038/nsmb.2446

Plotkin, J. B., and Kudla, G. (2011). Synonymous but not the same: the causes and consequences of codon bias. Nat. Rev. Genet. 12, 32. doi:10.1038/nrg2899

Pollard, K. S., Huizins, M. J., Rosenblom, K. R., and Siepel, A. (2009). Detection of nonneutral substitution rates on mammalian phylogenies. Genome Res. 20, 110–121. doi:10.1101/gr.097485.109

Presnyak, V., Alhusaini, N., Chen, Y.-H., Martin, S., Morris, N., Kline, N., et al. (2015). Codon optimality is a major determinant of mRNA stability. Cell 160, 1111–1124. doi:10.1016/j.cell.2015.02.029

Quang, D., Chen, Y., and XIE, X. (2014). DANN: a deep learning approach for annotating the pathogenicity of genetic variants. Bioinformatics 31, 761–763. doi:10.1093/bioinformatics/btu703

Rehm, H. L., Berg, J. S., Brooks, L. D., Bustamante, C. D., Evans, J. P., Landrum, M. J., et al. (2013). ClinGen—the clinical genome resource. New Engl. J. Med. 372, 2235–2242. doi:10.1056/NEJMsr1406261

Rost, B., Radiuojas, P., and Bromberg, Y. (2016). Protein function in precision medicine: deep understanding with machine learning. FEBS Lett. 590, 2327–2341. doi:10.1016/j.febslet.2015.12.037

Schlabalina, S. A., Spiridonov, N. A., and Kashina, A. (2013). Sounds of silence: synonymous nucleotides as a key to biological regulation and complexity. Nucleic Acids Res. 41, 2073–2094. doi:10.1093/nar/gku205

Shah, N., Hou, Y.-C. C., Yu, H.-C., Sainger, R., Caskey, C. T., Venter, J. C., et al. (2018). Identification of misclassified clivar variants via disease population prevalence. Am. J. Hum. Genet. 102, 609–619. doi:10.1016/j.ajhg.2018.02.019

Shah, P., and Gilchrist, M. A. (2010). Effect of correlated RNA abundances on translation errors and evolution of codon usage bias. PLoS Genet. 6, e1001128. doi:10.1371/journal.pgen.1001128

Sharp, P. M., and Li, W. H. (1987). The codon adaptation index—a measure of directional synonymous codon usage bias, and its potential applications. Nucleic Acids Res. 15, 1281–1295. doi:10.1093/nar/15.3.1281

Shi, F., Yao, Y., Bin, Y., Zheng, C.-H., and Xia, J. (2019). Computational identification of deleterious synonymous variants in human genomes using a feature-based approach. BMC Med. Genom. 12, 12. doi:10.1186/s12920-018-0455-6

Shihab, H. A., Rogers, M. E., Gough, J., Mort, M., Cooper, D. N., Day, I. N., et al. (2015). An integrative approach to predicting the functional effects of non-coding and coding sequence variation. Bioinformatics 31, 1536–1543. doi:10.1093/bioinformatics/btv009

Siepel, A., Bejerano, G., Pedersen, J. S., Hinrichs, A. S., Rosenbloom, K., et al. (2005). Evolutionarily conserved elements in vertebrate, insect, worm, and yeast genomes. Genome Res. 15, 1034–1050. doi:10.1101/gr.371505

Smith, J. M., and Haigh, J. (1974). The hitch-hiking effect of a favourable gene. Genet. Res. 23, 23. doi:10.1017/S0016673X00014634

Sorensen, M. A., Kurland, C., and Pedersen, S. (1989). Codon usage determines translation rate in Escherichia coli. J. Mol. Biol. 207, 365–377. doi:10.1016/0022-2836(89)90260-X
Evaluating Synonymous Variant-Effect Predictors

Zeng and Bromberg

Copyright © 2019 Zeng and Bromberg. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.

Stenson, P. D., Ball, E. V., Mort, M., Phillips, A. D., Shiel, J. A., Thomas, N. S., et al. (2003). Human gene mutation database (HGMD®): 2003 update. Hum. Mutat. 21, 577–581. doi: 10.1002/humu.10212

Stenson, P. D., Mort, M., Ball, E. V., Evans, K., Hayden, M., Heywood, S., et al. (2017). The human gene mutation database: towards a comprehensive repository of inherited mutation data for medical research, genetic diagnosis and next-generation sequencing studies. Hum. Genet. 136, 665–677. doi: 10.1007/s00439-017-1779-6

Stenson, P. D., Mort, M., Ball, E. V., Howells, K., Phillips, A. D., Thomas, N. S., et al. (2009). The human gene mutation database: 2008 update. Genome Med. 1, 13. doi: 10.1186/gm13

Stergachis, A. B., Haugen, E., Shafer, A., Fu, W., Vernot, B., Reynolds, A., et al. (2013). Exonic transcription factor binding directs codon choice and affects protein evolution. Science 342, 1367–1372. doi: 10.1126/science.1243490

Supek, F., Mišanina, B., Valcárcel, J., Gabaldón, T., and Lehner, B. (2014). Synonymous mutations frequently act as driver mutations in human cancers. Cell 156, 1324–1335. doi: 10.1016/j.cell.2014.01.051

Thanaraj, T., and Argos, P. (1996). Protein secondary structural types are differentially coded on messenger RNA. Protein Sci. 5, 1973–1983. doi: 10.1002/pro.5560051003

Van Der Velde, K. J., Kuiper, J., Thompson, B. A., Plasser, J. P., Van Valkenhoef, G., De Haan, M., et al. (2015). Evaluation of CADDS score in curated mismatch repair gene variants yields a model for clinical validation and prioritization. Hum. Mutat. 36, 712–719. doi: 10.1002/humu.22798

Visscher, P. M., Brown, M. A., McCarthy, M. I., and Yang, J. (2012). Five years of GWAS discovery. Am. J. Hum. Genet. 90(1), 7–24. doi: 10.1016/j.ajhg.2011.11.029

Wang, Z., and Burge, C. B. (2008). Splicing regulation: from a parts list of regulatory elements to an integrated splicing code. RNA 14, 802–813. doi: 10.1261/rna.876308

Welter, D., MacArthur, J., Morales, J., Burdett, T., Hall, P., Junkins, H., et al. (2013). The NHGRI GWAS catalog, a curated resource of SNP-trait associations. Nucleic Acids Res. 42, D1001–D1006. doi: 10.1093/nar/gkt1122

Wen, P., Xiao, P., and Xia, J. (2016). dbDSM: a manually curated database for deleterious synonymous mutations. Bioinformatics 32, 1914–1916. doi: 10.1093/bioinformatics/btw086

Xayaphoummine, A., Bucher, T., and Isambert, H. (2005). Kinefold web server for RNA/DNA folding path and structure prediction including pseudoknots and knots. Nucleic Acids Res. 33, W605–W610. doi: 10.1093/nar/gki447

Xing, Y., Zhao, X., Yu, T., Liang, D., Li, J., Wei, G., et al. (2016). MiasDB: a database of molecular interactions associated with alternative splicing of human pre-RNAs. PLoS One 11, e0155443. doi: 10.1371/journal.pone.0155443

Xiong, H. Y., Alipanabi, B., Lee, L. J., Bretschneider, H., Merico, D., Yuen, R. K. C., et al. (2015). The human splicing code reveals new insights into the genetic determinants of disease. Science 347, 1254806–1254806. doi: 10.1126/science.1254806

Xue, Y., Chen, Y., Ayub, Q., Huang, N., Ball, E. V., Mort, M., et al. (2012). Deleterious- and disease-allele prevalence in healthy individuals: insights from current predictions, mutation databases, and population-scale resequencing. Am. J. Hum. Genet. 91, 1022–1032. doi: 10.1016/j.ajhg.2012.10.015

Yue, P., and Moult, J. (2006). Identification and analysis of deleterious human SNPs. J. Mol. Biol. 356, 1263–1274. doi: 10.1016/j.jmb.2005.12.025

Zhang, G., Hubalewska, M., and Ignatova, Z. (2009). Transient ribosomal attenuation coordinates protein synthesis and co-translational folding. Nat. Struct. Mol. Biol. 16, 274. doi: 10.1038/nsmb.1554

Zhang, G., and Ignatova, Z. (2011). Folding at the birth of the nascent chain: coordinating translation with co-translational folding. Curr. Opin. Struct. Biol. 21, 25–31. doi: 10.1016/j.sbi.2010.10.008

Zhang, X., Li, M., Lin, H., Rao, X., Feng, W., Yang, Y., et al. (2017). regSNPs-splicing: a tool for prioritizing synonymous single-nucleotide substitution. Hum. Genet. 136, 1279–1289. doi: 10.1007/s00439-017-1783-x

Zhou, T., Weems, M., and Wilke, C. O. (2009). Translationally optimal codons associate with structurally sensitive sites in proteins. Mol. Biol. Evol. 26, 1571–1580. doi: 10.1093/molbev/msp070

Zuker, M. (2003). Mfold web server for nucleic acid folding and hybridization prediction. Nucleic Acids Res. 31, 3406–3415 doi: 10.1093/nar/gkg595

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2019 Zeng and Bromberg. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.