Decoding the effects of synonymous variants

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Abstract

Synonymous single nucleotide variants (sSNVs) are common in the human genome but are often overlooked. However, sSNVs can have significant biological impact and may lead to disease. Existing computational methods for evaluating the effect of sSNVs suffer from the lack of gold-standard training/evaluation data and exhibit over-reliance on sequence conservation signals. We developed synVep (synonymous Variant effect predictor), a machine learning-based method that overcomes both of these limitations. Our training data was a combination of variants reported by gnomAD (observed) and those unreported, but possible in the human genome (generated). We used positive-unlabeled learning to purify the generated variant set of any likely unobservable variants. We then trained two sequential extreme gradient boosting models to identify subsets of the remaining variants putatively enriched and depleted in effect. Our method attained 90% precision/recall on a previously unseen set of variants. Furthermore, although synVep does not explicitly use conservation, its scores correlated with evolutionary distances between orthologs in cross-species variation analysis. synVep was also able to differentiate pathogenic vs. benign variants, as well as splice-site disrupting variants (SDV) vs. non-SDVs. Thus, synVep provides an important improvement in annotation of sSNVs, allowing users to focus on variants that most likely harbor effects.

Availability: synVep webserver for online query: https://services.bromberglab.org/synvep; For local runs Python script (https://bitbucket.org/bromberglab/synvep_local) and prediction database (https://zenodo.org/record/4763256) are also available.
Introduction

The recent increase in accessibility of sequencing has facilitated a rise in precision medicine efforts focused on the interpretation of the effects of individual-specific genome variation (1). Genome-wide association studies (GWAS) have identified multiple variants marking specific phenotypes (2). However, the evaluation of variants in terms of their functional contributions to molecular pathogenicity mechanisms holds promise for both a better understanding of disease and drug discovery/optimization (3). SNVs (single nucleotide variants) are the most common variants in the human genome (4). Three types of SNVs are of particular interest – regulatory (i.e. changing the quantity/production of the gene product, e.g. transcription or splice site variants), non-synonymous (i.e. altering product protein sequence), and synonymous (i.e. variants in protein-coding regions that, due to the degeneracy of the genetic code, do not alter the protein sequence). Many computational tools have been developed to evaluate the functional effects of regulatory and non-synonymous variants (5,6). However, while an individual genome carries as many synonymous as non-synonymous SNVs (7), the former are often disregarded as functionally irrelevant. Still, sSNVs can cause disease (8) and affect gene function via multiple mechanisms, including binding of transcription factors (9), splicing (10), mRNA stability (11-13), co-translational folding (14-16), etc., as reviewed in our earlier work (17).

Existing methods for predicting sSNV effects are either (1) sSNV-specific tools, including SilVA (18), reg-SNP-splicing (19), DDIG-SN (20), TraP (21), and IDSV (22), or (2) general-purpose ones, including CADD (23,24), DANN (25), FATHMM-MKL (26), and MutationTaster2 (27). The number of computational sSNV effect predictors is limited in comparison to that of nsSNV effect predictors, as reviewed in (6,17). Partially, this paucity is due to the limited available experimental data evaluating variant effects, which could be used for training or testing of such methods. In fact, all existing predictors, except CADD and DANN, are trained using "pathogenic" variants from databases such as Human Gene Mutation Database (HGMD) (28) and ClinVar (29). Here we note that "pathogenicity" is not equivalent to "functional effect" (30,31) and inferring variant-disease causality is complicated by this inequality. Moreover, the experimental disease variant annotations are often unreliable (17), as it is difficult to distinguish causative variants from simply associated ones. Thus, the pathogenic label is inconsistent across databases, and
possibly over time/database releases. Finally, even these labeled effect variants are few; fewer still, if any, are experimentally labeled neutral polymorphisms. Thus, predictors trained on these variants are likely insufficient to predict the effects of tens of millions of possible sSNVs in human genome.

Here we had inferred, using positive-unlabeled learning (32-34), a subset of human sSNVs that could be used for training a predictor of sSNV molecular effect. We then developed, synVep (Synonymous Variant Effect Predictor), a machine learning-based method for scoring said effect for each possible human sSNV. Notably, synVep does not use conservation as an input feature but is more precise than other predictors (which do use conservation) in discriminating experimentally validated pathogenic sSNVs from randomly sampled observed variants. Its predictions also display the expected trend (35), where the sSNVs that change the human sequence nucleotide to one found in orthologs of evolutionarily-close human relatives, e.g. chimp, have lower effect scores than those that change to the nucleotides of further-removed organisms, e.g. fly. Furthermore, nucleotides that are not identified in any of the species evaluated here are deemed to have most effect when substituted into the human reference. However, many of the sSNVs that are not observed in the human population, tend to be scored very high (most effect), regardless of their appearance in other species.

In line with our earlier observations (17), we find that the variant frequency in the population is poorly correlated with the effect score. By our definition, common variants have little effect on gene function, but we find that rare variants are about equally likely to be neutral (65% common vs 69% rare neutral variants). To the best of our knowledge, synVep is the first variant-effect predictor that does not rely on conservation and is developed without an experimental or explicitly evolutionarily-estimated gold standard training/development set. Its success also suggests the feasibility of a similar approach for the development of a training set for other variant types, e.g. nsSNVs or indels. We expect that synVep sSNV effect predictions will greatly contribute to our understanding of pathogenicity pathways and to the prioritization of synonymous variants in disease.
Materials and Methods

Data Collection. We extracted all 93,437 human protein-coding transcripts from the Ensembl BioMart (36) GRCh37 p.13 assembly (37) and discarded the ones containing unknown nucleotides, lacking a start/stop codon, or having patched (https://grch37.ensembl.org/Homo_sapiens/Info/Annotation) chromosome IDs. We then generated all possible sSNVs for the remaining 72,400 transcripts. We further used ANNOVAR (38) (installed Aug 5, 2019) to extract sSNVs in these transcripts, and their allele-count based frequencies, from the Genome Aggregation Database exome subset (gnomAD exome) (39). An sSNV present in gnomAD was labeled a singleton if it was seen in only one individual and otherwise labeled observed. Generated sSNVs were those in the set of all possible variants in the 72,400 transcripts that were not singleton or observed. Thus, we collected 4,160,063 observed, 3,438,470 singleton, and 57,208,450 generated sSNVs; https://zenodo.org/record/4763256. Note that these correspond to 1,520,334 observed, 1,233,878 singleton, and 21,314,668 generated sSNVs with unique genomic coordinates and reference/alternative alleles, i.e. in one transcript per gene.

To evaluate and compare the performance of our predictor to other predictors, we manually curated a dataset of 42 curated-effect sSNVs with known biological effects, including the 33 pathogenic variants from the Buske et al. study (18). We required that all sSNVs in this set were strongly associated with disease and that there was experimental evidence of their molecular effects. These 42 sSNVs (Supplementary Table 1) mapped to 170 transcript-based sSNVs and were excluded from model training throughout this manuscript.

Variant Features. We collected 35 variant and sequence features (Supplementary Table 2), grouped into six categories: codon bias and autocorrelation (10 features), protein structure (3), mRNA stability (8), distance to regulatory factors (4), expression profile (3), and miscellaneous (7).

1. Transcript expression profiles. We downloaded the GTEx (40) “Transcript TPMs” dataset (dbGaP Accession phs000424.v7.p2) and standardized the transcript expression across tissue samples. We then used the average expression of each transcript over all samples from the same tissue as the representative transcript expression for that tissue.
Calculations of some of the codon bias metrics described below require a reference set of coding sequences, which are typically a set of highest expressed transcripts (41). To identify these references, we collected the maximum expression values for all transcripts across the 53 tissues. We then selected the transcripts within the highest 1% expression per tissue. We also used log₁₀ (minimum expression per tissue), log₁₀ (median expression per tissue), and log₁₀ (maximum expression per tissue) for each transcript as features.

2. Codon bias and autocorrelation. A variety of measures and/or their “Δ” form (difference in measure value after mutation vs. value before mutation) are adopted as features to characterize the codon bias of transcripts (see Supplementary Methods for more details), including the Codon Adaptation Index (CAI, Supplementary Methods Eqn. 1) (41), Fraction of Optimal Codons (fracOpt, Supplementary Methods Eqn. 2) (42), Codon Usage Bias (CUB, Supplementary Methods Eqn. 3) (43), Intrinsic Codon Deviation Index (ICDI, Supplementary Methods Eqn. 4) (44), Synonymous Codon Usage Order (SCUO, Supplementary Methods Eqn. 5) (45), and tRNA Adaptation Index (tAI, Supplementary Eqn. 6) (46). The calculation of these values was performed in R (47) and is available as an R package in https://github.com/zengzishuo/codonBiasMetrics.

These measures describe codon bias from different perspectives. CAI, fracOpt, and CUB rely on a reference set of optimal codons, found in highly expressed genes (48). CAI computes the geometric mean of relative usage of a codon compared to the most frequently used codon for the same amino acid (41). fracOpt is the fraction of optimal codons in a sequence of a certain length. CUB weights the frequency of amino acids in calculating codon bias. ICDI is independent of a reference set of genes (44). SCUO borrows the idea of entropy from Shannon information theory to describe codon usage bias of sequences (45). tAI focuses on translational efficiency by taking tRNA levels into account (46).

We also considered codon autocorrelation – a feature that has not yet been used by any sSNV predictors. In autocorrelated sequences same codons cluster together, whereas they are separated in anticorrelated sequences (e.g. XXXYYY is more autocorrelated than XYXYXY, where X and Y are two different codons) (49). Cannarozzi et al. noted the association between codon autocorrelation and translation dynamics and proposed the
tRNA pairing index (TPI) to describe a sequence’s codon autocorrelation. Autocorrelated sequences benefit from rapid translation due to the recycling of isoaccepting tRNAs (49). However, we note that the significance of recycling is likely weaker if the interval between two isoaccepting codons is larger – a feature that is not accounted for in TPI. Therefore, we proposed a new measure, Codon Autocorrelation Measure (CAM, Supplementary Eqn. 7), to describe the variant-specific codon autocorrelation impact penalized by the distance between the synonymous codons.

Finally, we also introduced the change of frequency measure (CF, Supplementary Methods Eqn. 8), to describe the amount of impact on codon’s frequency in a sequence due to the introduction of the variant.

3. Distance to regulatory and splicing sites. We used as features the distances to the nearest splice sites, transcription factor binding site (TFBS), RNA-binding protein (RBP) motif, and exonic splicing regulator (ESR). Their genomic coordinates were obtained from different sources as described below. We then computed the distance of a variant (in nucleotides) to all regulatory sites and selected the minimum value as the feature distance. We categorized these distances ($d$) into six categories as feature inputs: $d=0$, $0<d\leq3$, $3<d\leq5$, $5<d\leq10$, $10<d\leq20$, and $d>20$.

Genomic coordinates of regulatory regions were inferred as follows: (1) Splice sites were inferred from the “Genomic coding start” and “Genomic coding end” of all human protein-coding transcripts annotated in Ensembl BioMart GRCh37 p.13 assembly. (2) We downloaded the Gene Transcription Regulation Database (GTRD, version 18.06) (50) and identified the genomic coordinates of TFBS, using hg38 to hg19 conversion via CrossMap (51) for correspondence with our transcript coordinates. (3) We downloaded the ATTRACT database of RNA binding proteins and AssoCiated moTifs (52) and mapped the human RPB motifs to our set of transcript sequences. (4) We also downloaded the supplementary data of Cáceres et al (53) gold standard ESR motif set and mapped these to our transcripts.

4. Protein structure. We ran PredictProtein (54), a collection of tools for protein structure predictions, on all of the translated transcript sequences. We were particularly interested in protein secondary structure (PSS), residue solvent accessibility (SS), and disorder (PD)
predictions; in PredictProtein, PROFphd (55) predicts PSS and SS, while Meta-disorder (MD) (56) predicts PD.

5. mRNA stability, structure, and structural changes. We ran RNAfold (57) to predict (with calculation of partition function and base pairing probability matrix) the secondary structure and stability of all transcripts. We extracted the frequency of the structure with minimum free energy (MFE) in the structure ensemble, the free energy of the centroid structure, and its distance to the structure ensemble, as well as the local mRNA structure (strongly paired, strongly up/down -stream paired, weakly paired, weakly up/down -stream paired, or unpaired bases).

We also used RNAsnp (58) to predict the variant-induced local secondary structure changes for all sSNVs. The “mode” and “winsizeFold” parameters should be assigned according to the length in nucleotides (L) of the input sequence. We assigned the parameters as follows: (1) for L <= 200, mod=1 and winsizeFold=100; (2) for 200 < L <= 500, mod=1 and winsizeFold=200; (3) for L > 500, mod=2 and winsizeFold=500. We recorded the local structure dissimilarity, global structural dissimilarity and their statistical significance (p-values).

Model construction

Selecting a classifier. We standardized all continuous features and label-encoded categorical features. We compared two classifiers for differentiating observed and generated variants: deep neural network (DNN) (59) and XGBoost (60). DNN and XGBoost were implemented in Python (v3.6.4) using Keras (61) (https://keras.io/) and the xgboost package (v0.8.2) integrated with sci-kit learn (0.20.3) (62) (https://xgboost.readthedocs.io/en/latest/python/python_api.html), respectively. We split a balanced set (as described below) of observed and generated sSNVs into training, validation, and testing set in 8:1:1 ratio. Of the two methods, XGBoost was more efficient and achieved higher accuracy in testing and thus was selected as the classifier for future use.

Balancing variant data by transcript. The generated set of sSNVs is much larger than the observed set, but the number of observed sSNVs per transcript varies greatly. Moreover, some classifier input features are transcript-specific. Thus, a predictor may “memorize”
transcripts that have more observed sSNVs, and preferentially assign its variants observed status, instead of finding variant-specific differences between observed and generated. To avoid this, we assigned sampling likelihood weights for the generated set, i.e. the sampling likelihood weight of a generated variant is the number of observed sSNVs in the corresponding transcript. In all further balancing of data sets, generated sSNVs were probabilistically added to the set on the basis of their weights. Thus, the number of generated sSNVs on a transcript that were selected for a particular training set was correlated with the number observed sSNVs on this transcript.

**Positive unlabeled learning (PUL) to identify unobservable sSNVs.** PUL is a semi-supervised approach applicable to scenarios where only positive data points are labeled and the rest can be positive or negative (32-34). We employed the modified version of PUL (34) to separate the generated sSNVs into unobservable and not-seen sets. To prevent overfitting, we adopted relatively conservative hyperparameters of XGBoost (100 trees [n_estimators], 5 maximum depth [max_depth], 30% of the features per tree [colsample_bytree], 30% subsamples per tree [subsample]). We left out from PUL a fraction of observed as a test set, aiming to reach <5% incorrect predictions for this set at the end of the PUL.

In one epoch of PUL, a classifier was trained to differentiate the observed sSNVs from the same number of unlabeled ones (generated; selected via transcript-based set balancing as described above). All unlabeled sSNVs, including the ones not used in training, were evaluated with the resulting model and those classified as observed (scoring below 0.5) were added to the not-seen pool. The PUL process was repeated until convergence (Supplementary Methods). sSNVs scoring >0.5 in prediction from the last PUL model were further excluded from our data set.

**Differentiating the observable from not-seen using an intermediate model.** We trained a model to differentiate the observable sSNVs from the not-seen sSNVs (termed intermediate model from here on). We excluded 10% (9,274) of the common sSNVs (MAF > 0.01; excluded set) and all curated-effect sSNVs (170) from the construction of intermediate model for testing and final model parameter optimization. We split the observed sSNVs into subsets of 9: 0.5: 0.5 size ratio for training (3,631,441 variants),
validation and testing (201,746 variants each). We then randomly sampled the \textit{not-seen} variants to match the \textit{observed} validation and test set sizes; this left 47,923,258 \textit{not-seen} sSNVs for training. We then up-sampled the 3.6M \textit{observed} variants in the training set to create a balanced set of 47,923,258 \textit{observed} and \textit{not-seen} variants, each). We tuned the model hyperparameters by optimizing the F1 measure (Eqn. 3) of performance on the validation set and evaluated the resulting model on the test set.

\[
\text{Precision} = \frac{TP}{TP+FP} \quad \text{(Eqn. 1)}
\]

\[
\text{Recall} = \frac{TP}{TP+FN} \quad \text{(Eqn. 2)}
\]

\[
F1 \text{ measure} = \frac{2 \times \text{Precision} \times \text{Recall}}{(\text{Precision} + \text{Recall})} \quad \text{(Eqn. 3)}
\]

where TP, TN, FP, FN are respectively, true positive, \textit{i.e.} \textit{observed} sSNVs predicted to be \textit{observed}; true negative, \textit{not-seen} sSNVs predicted to be \textit{not-seen}; false positive, \textit{not-seen} sSNVs predicted to be \textit{observed}; false negative, \textit{observed} sSNVs predicted to be \textit{not-seen}.

**Final model (synVep) training.** We used the intermediate model to score the excluded common and \textit{curated-effect} sSNVs, as well as all \textit{observed} and \textit{not-seen} sSNVs. Here we assumed that common variants should be enriched in no-effect/neutral variation. Based on the scores of excluded sSNVs, we defined \textit{effect} and \textit{no-effect} synVep development sets, where sSNVs (both \textit{observed} and \textit{not-seen}) scoring above the median of the \textit{curated-effect} predictions were deemed \textit{effect}; while sSNVs (both \textit{observed} and \textit{not-seen}) scoring below the median of the excluded common sSNV predictions were labeled \textit{no-effect}. We thus collected 7,385,137 \textit{no-effect} and 32,117,625 \textit{effect} sSNVs.

We split the \textit{no-effect} and \textit{effect} sSNVs into subsets of 9: 0.5: 0.5 size ratio (in the same way as for the intermediate model) for training, validation, and test sets (62,758,222: 735,194: 735,194 variants per set). We sampled equal numbers of \textit{effect} sSNVs to match the \textit{no-effect} sSNVs in validation and test sets. We trained the final model on the training set using the hyperparameters optimized (F1 measure; Eqn. 3) on the validation set. We finally evaluated the model on the test set. Note that none of the \textit{curated-effect}, the \textit{excluded} common sSNVs, or the ClinVar (described below) dataset variants were included in our model training.
Performance comparison with other predictors. To evaluate synVep in comparison with other predictors, we used the 170 (transcript-based; 42 genomic coordinate-based) curated-effect sSNVs and the 9,274 (transcript-based; 7,957 genomic coordinate-based) excluded common sSNVs. Here we again assumed that common variants should be enriched in no-effect/neutral variation.

Other predictors in this comparison included CADD (phred-like scaled scores) (23), DANN (25), FATHMM-MKL (26), DDIG-SN (20), and EIGEN (63). EIGEN scores were collected using ANNOVAR (38) annotations; for other predictors, the scores were collected with default parameters as described in our earlier work (17). We did not include SiLVA (18) or TraP (21) in this comparison because 33 of 42 of the curated-effect sSNVs are in their training sets.

Note that synVep scores are produced per variant per transcript, while other predictors use the genomic coordinates, i.e. one reference sequence per variant. For the purposes of our comparison, we randomly re-sampled each tool's predictions of the effect set (42 variant scores) to produce 170 scores. Furthermore, as the common sSNVs (putatively no-effect) outnumbered the effect set, we randomly sampled 170 common variant scores in 100 comparison iterations. For each re-sampling, we performed a one-sided permutation test (null hypothesis: mean of common variant scores is equal to mean of effect scores; alternative hypothesis: mean of common variant scores is lower than mean of effect scores) and recorded the p-value and the difference of the common and effect variant score distribution medians (Δmedians). To assure that all predictor scores fall into the same [0,1] range, we standardized CADD and EIGEN Δmedians to their maximum non-outlier score (20 for CADD and 3 for EIGEN).

We also computed the Spearman correlation across predictor scores and the Fraction of Consensus Binary Predictions (FCBP; i.e. the number of binarized predictions agreed upon by all predictors, divided by total number of predictions) (17). An effect/no-effect scoring threshold for the FCBP computation is required; we used the default value of score=0.5 for DANN, FATHMM-MKL, and DDIG-SN. For CADD, we used score=15 as the threshold recommended by its online documentation (https://cadd.gs.washington.edu/info). As there was no recommended cutoff in the EIGEN
publication (63), we selected the cutoff (score=1.35) at the 75-percentile of EIGEN scores of 1,000 randomly sampled observed sSNVs.

**ClinVar data analysis.** We downloaded all ClinVar (64) submissions from the FTP site ([https://ftp.ncbi.nlm.nih.gov/pub/clinvar/](https://ftp.ncbi.nlm.nih.gov/pub/clinvar/)) and identified the sSNVs among these. We only considered the sSNVs with the "reviewed by expert panel" review status. From these we selected the (1) pathogenic and pathogenic/likely pathogenic variants as the pathogenic set and (2) benign and benign/likely benign as the benign set. There were 51 benign (genomic coordinate-based; 254 transcript-based) and 17 pathogenic (genomic coordinate-based; n=68 transcript-based) sSNVs (Supplementary Table 3). We also mapped to these ClinVar sSNVs to the precomputed GERP++ scores ([http://mendel.stanford.edu/SidowLab/downloads/gerp/](http://mendel.stanford.edu/SidowLab/downloads/gerp/)) (65).

**Cross-species sequence variation (CSV) analysis.** Cross-species variation (CSV) describes the nucleotide difference between the human reference sequence and the ortholog reference sequence of another species. In this study, we selected 20 species to generate CSVs: yeast (Saccharomyces cerevisiae), worm (Caenorhabdiis elegans), fruitfly (Drosophila melanogaster), zebrafish (Danio rerio), xenopus (Xenopus laevis), anole lizard (Anolis carolinensis), chicken (Gallus gallus), platypus (Ornithorhynchus anatinus), opossum (Monodelphis domestica), dog (Canis familiaris), pig (Sus scrofa), dolphin (Tursiops truncatus), mouse (Mus musculus), rabbit (Oryctolagus cuniculus), tree shrew (Tupaia belangeri), tarsier (Carlito syrichta), gibbon (Nomascus leucogenys), gorilla (Gorilla gorilla), bonobo (Pan paniscus), and chimpanzee (Pan troglodytes).

To represent the evolutionary distance of the CSV species to human, we obtained the value in million years since divergence from the TimeTree database (66). Given a human transcript T and its corresponding human gene G, we queried Ensembl BioMart for G's orthologs in the 20 species, \( \mathbf{G_{orthologs}} = [G_{yeast}, G_{worm}, G_{fruitfly}, \ldots, G_{chimpanzee}] \). We downloaded all coding DNA sequences (CDS) for these orthologs from Ensembl (release-94) (67). For each gene in \( \mathbf{G_{orthologs}} \), we identified its longest transcript per organism, \( \mathbf{T_{orthologs}} = [T_{yeast}, T_{worm}, T_{fruitfly}, \ldots, T_{chimpanzee}] \). We then used PRANK (68) to generate a multiple sequence alignment (MSA) for each T. PRANK aligns CDSs by first translating them into protein sequences so that gaps tend to be placed between codons, instead of
within codons. For each codon in each human transcript, we could identify if other organisms carried the same codon or another, even if the amino acid remained the same. If the codon was different, the corresponding human sSNV was termed a CSV.

**Evaluation of synVep predictions and splicing effects.** We downloaded and analyzed a dataset of SNV splicing effects (69) ([https://github.com/KosuriLab/MFASS](https://github.com/KosuriLab/MFASS)), referenced by genomic coordinates and Ensembl transcript IDs.

**Evaluation of synVep predictions according to constraint on coding regions.** Constrained regions (70), referenced by genomic coordinates, were downloaded from [https://s3.us-east-2.amazonaws.com/ccrs/ccr.html](https://s3.us-east-2.amazonaws.com/ccrs/ccr.html). The constraint of human coding region is measured by percentile (of residuals from a linear regression for distance-to-mutation prediction as computes in (70)), where a high percentile indicates a more constrained the region. We annotated the sparsity of sSNVs, *i.e.* the fraction of observed sSNVs among all possible sSNVs in a region of a certain constraint level, and the median synVep prediction of variants in these regions.

**Analysis of sSNVs identified in Qatari Genome.** We downloaded all VCF files containing variants identified from Qatari Genome project (QTRG) (71) from NCBI Sequence Read Archive ([https://trace.ncbi.nlm.nih.gov/Traces/sra/?study=SRP061943](https://trace.ncbi.nlm.nih.gov/Traces/sra/?study=SRP061943)). We then parsed these VCF files, extracted the variants, and mapped the sSNVs to our *observed, singleton, not-seen*, and *unobservable* sets.
Figure 1. Pipeline of predictor construction. Starting with 4,160,063 observed and 57,208,450 generated sSNVs, 63 epochs of positive unlabeled learning (PUL) was conducted to separate the generated set into not-seen and unobservable set (Supplementary Methods). An intermediate model was trained using the observed and not-seen sets (observed set was up-sampled to equal amount of not-seen variants). Intermediate model’s predictions for common and pathogenic sSNVs were used as guideline to set cutoffs assigning no-effect and effect set. The final predictor was trained using the no-effect and effect sets (no-effect set was up-sampled to equal amount of effect sets).
Figure 2. Extraction of cross-species sequence variants (CSV). For each human protein coding transcript \( T \), codon-oriented multiple sequence alignment was performed with 20 species’ longest coding sequencing of the same ortholog as \( T \). The CSV are represented as “codon>codon” format for specific transcript positions and may coincide with human sSNVs.

Results and Discussion

Generated sSNVs may be observable in the future. In the absence of a gold-standard experimentally validated data set describing sSNV functional effects, we sought an alternative for the development of our method. We had previously proposed to use sSNVs that have been observed in major sequencing projects vs. all other possible human genome sSNVs (the generated set) for method evaluation (17). We collected (Methods) 72,400 human transcripts with 4,160,063 \((n=1,520,334\) genomic coordinate-based) observed sSNVs and 3,438,470 \((n=1,233,878\) genomic coordinate-based) singletons, observed in only one individual, from the exome sequencing data of the Genome Aggregation Database (gnomAD exome) (39). We then created a generated set of 57,208,450 \((n=21,314,668\) genomic coordinate-based) all possible sSNVs in these transcripts that were not found in gnomAD data. Note that only \(~12\%\) of all sSNVs in our set were ever observed. We annotated these sSNVs with 35 transcript- and variant-specific features, including codon bias, codon autocorrelation, transcript stability,
expression level, distance to regulatory sites, predicted protein secondary structures, etc. (Methods and Supplementary Table 2). Notably, we did not include any conservation-related features.

While the observed sSNVs are not necessarily functionally neutral, they are at least compatible with life. The generated sSNVs, on the other hand, are likely split of two subtypes: the not-seen sSNVs, which may or may not become observed with more sequencing, and the unobservable ones, which cannot be observed given the contemporary variant-discovery capability. Note that the unobservable character of sSNVs may be due to a broad range of technical and biological reasons such as sequencing (73,74), molecular functional constraints (75), and analytical biases or extreme deleteriousness resulting in early embryonic incompatibility with life (76,77). We also note that in our modelling, the unobservable set may simply be poorly described by our selection of variant features, which capture the observed and not-seen sSNVs.

We used observed sSNVs as positives in positive-unlabeled learning (PUL) (32-34) to differentiate the not-seen sSNVs (similar to observed) from unobservable ones in the generated (unlabeled) set. At convergence (epoch 63, Methods; Supplementary Fig. 1), PUL partitioned all generated sSNVs into unobservable (n=6,278,254 transcript-based and 2,764,229 genomic coordinate-based; 11%) and not-seen (n=50,930,196 transcript-based and 19,730,623 genomic coordinate-based; 89%). Additionally, 8% (n=266,192) of singletons were deemed unobservable by the PUL model, as were 2% of the observed sSNVs (n=79,639). The latter result highlights the possible insufficiency of our variant descriptors for capturing the complete observable variant diversity, while the former may also indicate sequencing errors. The difference in percentages of variants misidentified by the model (11% of generated vs 2% of observed), however, suggests that deleteriousness of variants also plays a role in defining unobservable variants.

**Observed and not-seen variant sets contain both no-effect and effect sSNVs.** We trained the intermediate model (Methods) to recognize observed vs. not-seen sSNVs. The model accurately (F1=0.71; Eqn. 3) recognized the two classes in a previously unseen variant test set (Supplementary Fig. 2). The model also predicted 9% (9,282,542) of the not-seen
sSNVs to be *observed* (scoring < 0.5), implying that these may be sequenced in the future.

While observed sSNVs are likely depleted in large functional effects, they are not all functionally neutral; for example, pathogenic sSNVs are, in fact, observed. Similarly, not all *not-seen* sSNVs have a functional effect; rather, they may be observable in the future. To build a model for evaluating the effects of sSNVs, we leveraged the intermediate model's predictions on common variants *excluded* from training and the experimentally validated *effect* sSNVs (*curated-effect*; Methods; 170 transcript-based sSNVs). While these *curated-effect* sSNVs are, in fact, observed, their prediction scores were higher than those of the *excluded* common set (Supplementary Fig. 2, Mann-Whitney U test p-value<2.2e-16). This observation is likely due to the fact that the *not-seen* set is enriched while the common variant set is depleted, in large effect sSNVs. We assume this for common variants because large-effect deleterious variants would not become common and large-effect advantageous variants would tend to become wild-type.

We excluded 10% (7,957) of the common sSNVs from training of the intermediate model for selecting the cutoff of *effect/no-effect* variants as next described. For training of the final model, we selected as *no-effect* those sSNVs (both *observed* and *not-seen*) scoring below the intermediate model prediction median (0.38) of the excluded common sSNVs; variants scoring above the intermediate model median of the *curated-effect* sSNVs (score = 0.63) were labeled *effect* (Supplementary Fig. 2). We thus obtained 7,385,137 (2,580,540 *observed* and 4,804,597 *not-seen*) *no-effect* and 32,117,625 (405,170 *observed* and 31,712,455 *not-seen*) *effect* sSNVs. We trained the final model (*synVep*) to differentiate the *no-effect* and *effect* sSNVs (in balanced class training), using a 9: 0.5: 0.5 split of data for training, validation, and testing purposes (Methods). *synVep* was accurate (F1 measure=0.90; binary score cutoff=0.5) in evaluating the previously unseen test set (369,257 *no-effect* and 369,257 *effect*). For the same set, the F1 measures of the CADD, DANN, DDIG-SN, FATHMM-MKL, and EIGEN performance were 0.03, 0.48, 0.03, 0.46, and 0.46, respectively.
**Figure 3.** Predictor performance on common vs. curated-effect sSNVs. Panels A-F show the differential predictions on sets of curated-effect (n=170) and common sSNVs (randomly selected n=170) for CADD (phred-like scaled scores), DANN, DDIG-SN, FATHMM-MKL, EIGEN, and synVep, respectively. Gray line indicates scoring cutoff suggested by tool authors. Neither the common set nor the curated-effect set were included in synVep training. In panel (G), this comparison was repeated 100 times to compute the Δ medians (i.e., median_{curated-effect} − median_{common}). Permutation tests show that all of the predictors give significantly different scores between the two sets in every iteration, except for DANN where two of 100 comparisons were not significant (p-value>0.05 after Bonferroni correction).

**synVep outperforms other predictors in identifying effect sSNVs.** In order to evaluate the performance of our method, we needed a gold standard set of designated effect and no effect variants. However, since there are no experimentally validated "neutral" sSNVs, we used the common sSNVs never seen in training as neutrals (*no effect*), *i.e.* as described above we assumed that the majority of common sSNVs have little effect. On the level of molecular function, the *effect* of a variant is not directional (neither advantageous nor deleterious). On the level of individual fitness (evolutionary), however, variants could be advantageous, neutral, or deleterious. Note that evolutionarily *effect* variants must have a molecular *effect*, but not vice versa. The neutral theory of molecular evolution, as well as some experimental work, suggest that advantageous mutations are rare, and most effect mutations are deleterious (78,79). Variants with large deleterious effects on evolutionary fitness would have been purified out and will not be seen in the population. Neutral or weak *effect* mutations could, depending on the population size, become
common or even fixed due to genetic drift (78). Thus, in the absence of experimentally neutral variants, common variation may be considered a reference point for neutral or weak effect variants.

We used the set of curated-effect sSNVs as the effect group. Note that these sSNVs, as well as any other sSNVs in their corresponding transcripts, were never used in our model training. Since the test set of no effect common variants (9,274 sSNVs) greatly outnumbered the curated-effect set (170), we randomly sampled 170 sSNVs from the common set 100 times (Methods). Predictor scores were significantly different (p-value < 0.05 after Bonferroni correction for multiple hypothesis testing; Fig. 3A-E; Methods) between variant groups in all 100 rounds; the sole exception was DANN, where in 11 of 100 rounds the differences were not significant (p-value > 0.05). Notably, the difference in median scores between the effect and no-effect variants was at least two-fold greater for synVep than for other methods (Fig. 3F; CADD and EIGEN scores normalized to 0-1 scale). Furthermore, while all predictors identified most curated-effect sSNVs as having more effect than common sSNVs, most failed to differentiate the two at the default method binary prediction threshold. synVep’s predictions fulfilled this criterion, while other predictors either placed both sets of variants below (CADD, DDIG-SN) or above the default threshold (DANN, FATHMM-MKL).

We further examined the correlation of the predicted scores (Fig. 4A) and the Fraction of Consensus Binary Prediction (17) (FCBP; Fig. 4B) on all 4,160,063 observed sSNVs for all predictors (synVep, CADD, DANN, FATHMM-MKL, DDIG-SN, and EIGEN). synVep’s scores were poorly correlated with other predictor scores (Pearson correlation ranging from -0.1 [DANN] to 0.21 [FATHMM-MKL]), while binary classification was more similar (FCBP ranging from 0.37 [DANN] to 0.69 [CADD and DDIG-SN]).
Figure 4. Comparing predictions made by synVep and other tools. Panel (A) Spearman correlation and (B) Fraction of Consensus Binary Prediction (FCBP) highlight similarity and lack thereof among the predictors for all observed sSNVs (n=4,160,063).

Singletons are more likely than observed to be effect. Singletons were not included in our training because it is difficult to estimate how many of them are artifacts due to the 0.1-0.6% error rates of next-generation sequencing (80). If the singletons are not artifacts, then they are likely to be individual or ultrarare variants. These are more likely to be effect than higher frequency variants (81,82). An excess burden of ultrarare variants (although not necessarily synonymous) is also often seen in diseases, such as schizophrenia (83-85), Parkinson disease (86), and bipolar disorder (87). In line with these expectations, we found that singletons were, on average, scored higher than observed sSNVs (Supplementary Fig. 3), suggesting that singletons are more likely to be effect than the observed.

Variant effect predictors differentiate benign and pathogenic variants. Among the predictors considered in this work, only two (FATHMM-MKL and DDIG-SN) are explicitly predictors of variant pathogenicity. To investigate whether predictors of variant effect (not pathogenicity) can identify pathogenic sSNVs, we obtained from ClinVar (reviewed by
expert panel) 17 pathogenic (genomic coordinate-based, 68 transcript-based) sSNVs and 51 benign sSNVs (genomic coordinate-based, 254 transcript-based). Of these, one benign and one pathogenic variant (genomic coordinate-based, 13 transcript-based) were deemed unobservable by our model and were removed from consideration. Note that, for fairness of evaluation, these sSNVs were excluded from training synVep.

Figure 5. Evaluating variant effect predictors using ClinVar data. Benign are variants labeled “Benign” and “Benign/Likely benign” in ClinVar, with “Review by expert panel” as review status. Pathogenic are those labeled “Pathogenic” and “Pathogenic /Likely Pathogenic” in ClinVar, with “Review by expert panel” as review status or with “research” method and at least one publication experimentally validating the effect. Panel A-E show the predictions from for CADD (phred-like scaled scores), DANN, EIGEN, GERP++ score, and synVep, respectively. Grey dashed lines show the method author-recommended cutoffs, where available.

All variant-effect predictors, including synVep, assigned higher scores to pathogenic than benign variants (Fig. 5A-C and E, all statistically significant, one-sided permutation test p-value = 0). Importantly, note that all variant-effect predictors except synVep use GERP++ (65) as a feature. GERP++ identifies constrained elements in multiple sequence alignments and is thus a feature reflective of conservation. Highly conserved genomic
positions often have experienced extensive purifying selection (88). Therefore, conservation is understandably a commonly used feature for disease variant prioritization (89). The area under the precision-recall curve (PR AUC, Eqn. 1,2) measures for CADD, DANN, EIGEN, and synVep were 0.60, 0.62, 0.70, and 0.73, respectively, suggesting that synVep provides the best discriminative power on this rigorous expert-reviewed disease variants dataset. Note that GERP++ attained 0.62 PR AUC, meaning that the performance of the specialized predictors did not substantially improve or was worse than that of one of their features, i.e. GERP++. Also note that at the default/recommended cutoff, only synVep placed the majority of benign vs. pathogenic variants on opposite sides of the cutoff (precision=0.32, recall=0.60). Here, the method’s low precision indicates that many of ClinVar labeled “benign” variants likely carry functional effects, which may not manifest as disease. Moreover, in the scenario of disease variant prioritization synVep offers discriminative power independent of conservation, so it may be used in combination with a conservation score or other predictors.

One major challenge in disease variant prioritization is that for complex diseases, causality can rarely be explained by a single variant (90). The utility of variant pathogenicity score is thus questionable: does a high score suggest a high likelihood of an individual developing a disease or a high likelihood of this variant contributing to a disease? Also, would an individual with many predicted-pathogenic variants carry many diseases or be very certain to carry at least one disease? One potential way to bypass this puzzle is to establish the variant-disease relationship with the collective effect from the whole variome, instead of a single or a few variants. One way to use the effect information could be to optimize the polygenic risk scoring methods (91,92) to only focus on effect variants. However, this approach would be limited by the location of most SNPs in non-coding regions. Attempts to unite variant effects across the coding region of the genome (e.g. (93), (94)) measure every gene product’s change by aggregating all variants per gene (as a burden or as function changes) to predict disease predisposition. As a future application, synVep predictions (as well as other variant effect) may be plugged in these pipelines to explore the contribution of sSNVs to complex diseases.

*synVep highlights correlation between conservation and effect.* We annotated all sSNVs as CSVs (cross-species variation) or not (Fig. 2; Methods). CSVs are codon differences
between the human reference sequence and another species’ ortholog. For example, if the proline-coding codon in a human transcript T is CCC, while the aligned proline codon on T’s chimp ortholog is CCT, then the human sSNV CCC->CCT is considered a chimp-CSV. We thus annotated 15,618,155 unique (only exists in one species) and 35,102,565 non-unique (overlapping across species) CSVs (Supplementary Fig. 4). Since less than 10% (7,026 of 72,400) of the human transcripts can be mapped to orthologs in all 20 species, we analyzed separately the CSVs in (1) all transcripts (n=32,264,860) and (2) only the transcripts that have orthologs in all 20 species (n=3,321,574) and that are likely ancient (ancient genes) (95).

Figure 6. Variant effect prediction from the perspective of cross-species variation (CSV). Panels A-C show synVep-predicted scores for variants grouped by the number of species carrying the mutant nucleotide; separately for observed, singletons, and not-seen sets. The red dashed line is synVep’s default cutoff for effect and no-effect. The number in each box indicates the number of variants of that group (in thousands). Panels D-F show the median score (y-axis) across species at log2 million years since divergence from common ancestor with human (x-axis) and linear regression trendline (red line) between the two. The Spearman correlations between median synVep score and log2(million years) for panel D-F are 0.68, 0.64, and 0.66, respectively.
The distribution of synVep prediction scores for CSVs in the ancient genes and for those in all transcripts were similar ($\Delta$ mean=0.05, Mann-Whitney U test $p$-value<2.2e-16), suggesting that synVep’s evaluation of variants does not discriminate by gene age. For all transcripts, observed sSNVs had more CSVs (67%, n=2,823,142) than did the not-seen variants (53%, n=26,976,016; Supplementary Fig. 5). CSVs overall were predicted less likely to be effect than non-CSV for both ancient and all transcripts (Fig. 6 A-C; Supplementary Fig. 6 A-C). While this is in line with the scoring trends of the observed and not-seen variants overall, it also mirrors earlier findings of few CSV nsSNVs corresponding to a known human disease (96-99). synVep scores also trended lower for CSVs whose substituting nucleotide was found in more species, for both ancient (Fig. 6 A-C) and all transcripts (Supplementary Fig. 6 A-C). Since the number of CSV species is somewhat indicative of codon conservation, this trend suggests that, although synVep was trained without using conservation descriptors, its predictions still identify conserved codons that are often functionally relevant (100).

To further elucidate the effect of sequence conservation across species, we calculated codon mutation fraction (CMF, Supplementary Eqn. 9) to describe how common a human’s alternative codon is, compared to the reference codon, among the 20 species included for CSV analysis. For example, if in a multiple sequence alignment of the 20 species orthologs, the human CCC codon is aligned to 10 CCC, 5 CCT, and 5 other codons, then the CMF of the corresponding synonymous variant, CCC>CCT, is 5/15=0.33. We observed that predicted scores generally decrease with higher CMF (supplementary Figure 7 A-C), indicating that sSNVs with alternative codons commonly present as reference codons among other species have less effect.

We additionally investigated the relationship between the evolutionary distance of CSV species from human and the effects of the corresponding sSNVs. Since one sSNV can correspond to multiple species CSVs, we only considered CSVs that are uniquely found in one species for this evaluation. The medians of synVep scores of these species-exclusive CSVs in both ancient genes (Fig. 6 D-F) and all transcripts (Supplementary Fig. 6 D-F) correlated with the evolutionary distance of the corresponding species to human. However, for ancient genes, the median scores of observed variants unique to further related (i.e. beyond Tarsier) species were in the effect range (synVep>0.5). Arguably, this
means that human sSNVs that introduce nucleotides likely present in recent ancestors tend to be *no-effect*, while similarity to further removed relatives carries no such benefit (Fig. 6 D). These findings agree with our recent work on nsSNV CSV analysis (35). We note that species relationship had much less impact on binary *effect* classification for singleton variants and none for not-seen variants (Fig. 6E-F). The same observations could not be made for the *all transcript* set of variants, where *observed* and *singleton* CSVs were predicted to be *no-effect* for a large portion of species (Supplementary Fig. 6D-F). This observation suggests that ancient genes are functionally crucial and have been sufficiently optimized over time to only permit minor levels of variation without impact on functionality.

**synVep differentiates splice-disrupting variants.** Cheung et al. (69) measured the splice-disrupting effects of genomic variants (3,297 transcript-specific sSNVs) and defined a group of large-effect splice-disrupting variants (140 SDV sSNVs). As expected, synVep scores of SDVs were on average higher than those of non-SDVs (Fig. 7 A-C). Curiously, 140 SDVs comprised only six *observed* (4.6%) and 18 *singleton* (13.7%) variants; nine were deemed *unobservable* (6.4%) and 107 were *not-seen* (76.4%). The fact that most of SDVs are *not-seen* reinforces our assumption that *not-seen* sSNVs are enriched for large-effect deleterious sSNVs that may have been purified.

![Figure 7](image)

**Figure 7.** synVep predictions for large-effect splice-disrupting variants (SDVs) and non-SDVs. SynVep predictions are higher scoring for a set of experimentally determined SDV than non-SDV variants across *observed*, *singleton*, and *not-seen* data sets. Note that non-SDVs may
still carry other functional effects. The mispredicted neutral SDVs highlight the limitations of our training data, which is limited in differentiating between effect and no-effect observed variants; although two of the five observed SDVs are correctly annotated as effect.

Splicing disruption is a well-known and well-studied mechanism of sSNV effect (101). In fact, most of the experimental validations of our curated-effect and ClinVar pathogenic variants refer to elucidating splicing effects (Supplementary Table 1 and 3). Moreover, many cancer driver mutations are found to be splice-disrupting synonymous variants (102). Aside from splicing, experimental validation of variant effect is rare, arguably due to technical challenges (103). Perhaps, since the experimental evidence for splicing disruption is more abundant than non-splicing effects', the former is considered a major factor in clinical consideration for sSNVs. For example, according to the guidelines from American College of Medical Genetics and Genomics (104), an sSNV is clinically benign if it is not in a conserved position and is predicted to be non-impacting to a splice site (e.g. via GeneSplicer (105), NNsplice (106)). Thus, synVep’s ability to identify effect and score sSNVs regardless of their splice effects or conservation makes it an ideal tool for prioritization of all possible variants, regardless of their mechanism or evolutionary evidence of effect.

sSNV effects are limited by genomic constraints. Havrilla et al. developed the concept of “coding constrained regions” (CCR) to describe the regional scarcity of protein-changing (missense or loss-of-function) variants in the human genome (70). Here, a region with fewer of these variants observed in the human population has a higher CCR percentile score. For our set of variants, the fraction of observed (number of observed sSNVs divided by all possible sSNVs in this region) negatively correlated (Pearson $\rho=-0.61$) with CCR percentile (Fig. 8 A); i.e. higher constrain indicates fewer sSNVs. Furthermore, synVep predictions positively correlated with CCR percentiles for observed ($\rho=0.58$, Fig. 8 B), i.e. lower CCR percentile (less constrained regions) indicated lower (no-effect) synVep scores.
Figure 8. sSNV effect measured by region constraint. Coding constrained regions (CCR) describe the regional scarcity of nsSNVs; higher percentile regions represent have fewer observed nsSNVs. Observed sSNVs are relatively scarce in constrained regions (A), while their median synVep scores are higher (B). Pearson correlation are indicated in blue.

The negative correlation between the fraction of sSNVs and CCR indicates a positive correlation between synonymous mutation rate and missense or loss-of-function mutation rate. This observation is in line with earlier studies (107,108), but raises a question of the utility of Ka/Ks ratio (nonsynonymous divided by synonymous mutation rate), which is widely used to measure the strength of evolutionary selection at certain genomic sites (109). The application of the Ka/Ks ratio is based on the assumption that synonymous mutations are neutral and thus Ks can serve as a baseline for Ka. However, it has been demonstrated that a high Ka/Ks can also result from a low Ks due to strong negative selection at the synonymous sites (10,11,110-112). Efforts have been made to improve the utility of Ka/Ks by incorporating codon preference (113-115), but the question remains: how often is the selection at synonymous sites sufficiently underestimated so that Ka/Ks is no longer accurate? Lawrie et al. found that 22% of the fourfold synonymous sites (where the amino acid can be encoded by four codons) in the fruitfly genome are under strong selection (116). Lu and Wu estimated that 90% synonymous mutations in human and chimp are deleterious (117). Hellmann et al. estimated that 39% mutations at
the human-chimp-diverged non-CpG fourfold synonymous sites have been purified (118). Zhou et al. showed that 9% of all yeast genes and 5% all worm genes undergo purifying selection on synonymous sites (115). In turn, our results show that about 67% of all possible human sSNVs (excluding unobservable) are effect (synVep score>0.5), but we cannot estimate the strength of selection at these sites. Taken together, these observations suggest that Ka/Ks measures of genomic site constraints may be underpowered.

**synVep sheds light on future variant discovery and interpretation.** Whenever a human genome variant is sequenced, it will automatically be reassigned a class in our collection. Thus, a newly sequenced variant will first become a singleton and may, eventually, be a member of the observed group. An enrichment in observed variants will likely come from large-scale sequencing. Here, it is important to remember that although only 16% of global population are of European descent (119), 45% of the subjects whose exomes are recorded in the gnomAD exomes database (120) are; i.e. there is a significant underrepresentation of sSNVs from other ethnicities. When more diverse genomes are sequenced, will there be a significant addition to the observed set (i.e. significant reduction of the not-seen set)?

To answer this question, we obtained all variants from the Qatar Genome (QTRG) project (71) and mapped them to our set of sSNVs. QTRG comprises 1,376 individuals and may serve as a representative pool of genomic variants in Middle East and north Africa (MENA) area (71); thus, this set is complimentary to gnomAD. We identified 526,616 transcript-based sSNVs (n=192,246 genomic coordinate-based) from QTRG sequencing. Importantly, only 0.6% of the Qatari sSNVs mapped to our unobservable set – a fraction that is lower than the misprediction rate (5%) that we allowed during PUL. Moreover, two thirds of these variants were singletons in QTRG. This observation suggests that our unobservable variants are indeed unlikely to be ever observed in future sequenced human populations. The majority of QTRG sSNVs (81.9%) mapped to our observed set; 4.6% and 12.8% were singleton and not-seen, respectively, in our set (Fig. 9 A). Interestingly, 63.5% and 64.6% QTRG sSNVs mapping to our singleton and not-seen sets, respectively, were singletons in the Qatari cohort.
Figure 9. Distributions of the Qatar Genome sSNVs. In both panels, our gnomAD-based observed (orange), singleton (blue) and not-seen (dark orange) sets are highlighted. (A) represents the fraction of the QTRG sSNVs mapped our observed, singleton, not-seen, and unobservable (gray) sets. (B) synVep scores for our (gnomAD-based) variant sets, as well as the scores for QTRG sSNVs (white) mapping to the corresponding gnomAD-classes. Importantly, the synVep scores of QTRG variants that were previously classified as singletons or not-seen score much lower than other variants in the corresponding groups.

How many of the previously not-seen sSNVs are effect? New sSNVs are likely to come from clinical sequencing and could thus could often be deemed disease-associated. We expect, however, that these variants will carry little or no effect. In other words, currently not-seen no-effect observable sSNVs (n=14,259,180 transcript-based and n=5,975,076 genomic coordinate-based) are more likely to be discovered in the future than an effect ones – even if a sample is taken from a sick individual. Recall our assumption that the not-seen set is composed of those sSNVs that carry a large effect and have been purified, as well as those that are putatively neutral and will be seen in the future if more sequencing is performed. The synVep scores of the QTRG sSNVs mapping to our not-seen set were, on average, much lower than those of the entire not-seen set (Fig. 9 B, average synVep score 0.49 vs. 0.70, Mann-Whitney U test p-value<2.2e-16). This result confirms our assumption, as these newly identified sSNVs are actually observable (not
purified) and thus they are generally less likely to have large effect (and thus lower synVep scores). It may also be that the newly identified predicted effect variants (from QTRG, and other sequencing efforts in the future) are the ethnicity-differentiating, i.e. not necessarily affecting overall fitness, but contributing to individual differences (as in e.g. (121)).

Conclusion

We developed synVep—a machine learning-based model for evaluating the effect of human sSNVs. Our model does not use disease/deleteriousness-labeled training data or rely on conservation. Instead, we used the signals derived from observed (and corresponding generated) sSNVs from large sequencing projects. Our model successfully distinguishes sSNVs with experimentally validated effect, e.g. splice-site disrupters, as well as pathogenic sSNVs. Moreover, our model’s predictions of cross-species variants (CSVs) correlate with the evolutionary distance between human and CSV-species. While further experimental validations of effect prediction are necessary, synVep’s evaluation on sSNV effect will greatly contribute to our understanding of biological molecular pathways in general, and of pathogenicity pathways in particular.

Author contributions

ZZ and YB designed the study, evaluated the results, and wrote the manuscript; ZZ conducted the study; ZZ and AA built the webserver.

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Conflicts of interest
None declared.

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