BENCHMARKING DNA SEQUENCE MODELS FOR CAUSAL VARIANT PREDICTION IN HUMAN GENETICS

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Paper under double-blind review

Abstract

Machine learning holds immense promise in biology, particularly for the challenging task of identifying causal variants for Mendelian and complex traits. Two primary approaches have emerged for this task: supervised sequence-to-function models trained on functional genomics experimental data and self-supervised DNA language models that learn evolutionary constraints on sequences. However, the field currently lacks consistently curated datasets with accurate labels, especially for non-coding variants, that are necessary to comprehensively benchmark these models and advance the field. In this work, we present TraitGym, a curated dataset of genetic variants that are either known to be causal or are strong candidates across 113 Mendelian and 83 complex traits, along with carefully constructed control variants. We frame the causal variant prediction task as a binary classification problem and benchmark various models, including functionalgenomics-supervised models, self-supervised models, models that combine machine learning predictions with curated annotation features, and ensembles of these. Our results provide insights into the capabilities and limitations of different approaches for predicting the functional consequences of genetic variants. We find that alignment-based models CADD and GPN-MSA compare favorably for Mendelian traits and complex disease traits, while functional-genomicssupervised models Enformer and Borzoi perform better for complex non-disease traits. All curated benchmark data, together with training and benchmarking scripts, will be made publicly available upon publication.

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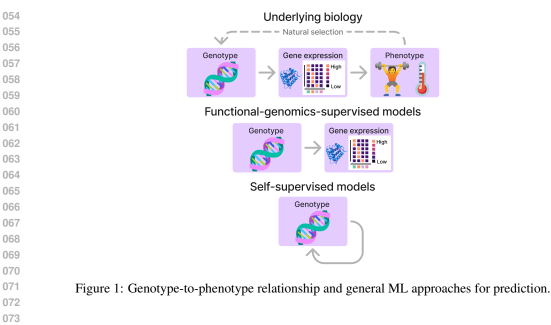
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1 INTRODUCTION

034 Machine learning is increasingly transforming the fields of genomics, human genetics, and healthcare by offering new avenues for predicting the impact of genetic variants on phenotypes and by po-035 tentially improving the accuracy of trait or disease risk predictions from individual human genomes. A major challenge in these domains is determining which among millions of intercorrelated genetic 037 variants are causal for Mendelian and complex traits, including diseases. (A Mendelian trait is typically influenced by a single gene, whereas the variation of a complex trait is shaped by multiple mutations across several genes, each contributing a small individual effect.) Tackling this challenge, 040 which has profound implications for human health, requires robust and scalable methods that can 041 decode the biological syntax of the human genome and how it drives molecular functions across 042 different cells and tissues.

043 Three major classes of approaches have been developed to model DNA sequences and predict the ef-044 fects of genetic variants. The first approach utilizes supervised machine learning models, commonly referred to as sequence-to-function models, which are trained to predict genome-wide functional 046 genomics experimental data from DNA sequences (Eraslan et al., 2019); we refer to these models 047 as functional-genomics-supervised. These models predict the functional effects of specific variants 048 by assessing how changes in the DNA sequence influence experimental outcomes. The second approach involves self-supervised genomic language models (gLMs), such as masked or autoregressive language models, which are trained only on DNA sequences from one or multiple species without 051 relying on experimental data (Benegas et al., 2024). Models that utilize sequences from multiple species take advantage of evolutionary conservation to gain functional insights. Variant effects in 052 such models are assessed by comparing the log-likelihood between the alternative and reference alleles of the variant, as well as by quantifying changes in the latent representations. Another class of



methods includes *integrative* approaches, which combine machine learning predictions with curated 074 annotation features to improve the accuracy of variant effect prediction (Schubach et al., 2024). 075

076 Despite its importance, the field currently lacks consistently processed and comprehensively curated 077 datasets of putative causal genetic variants with reliable labels. Furthermore, there is a pressing need for establishing a common ground for systematically benchmarking state-of-the-art models based on functional-genomics-supervised, self-supervised and integrative approaches, in order to 079 help advance the field.

081 In this article, we present TraitGym, a curation of two benchmark datasets from human genetics: 082 one comprising causal variants for 113 Mendelian traits, and another consisting of strong causal 083 variant candidates across 83 complex traits, along with carefully constructed control sets matching 084 relevant summary statistics (such as minor allele frequencies, variant types, distances from transcrip-085 tion start sites, and linkage disequilibrium scores) of putative causal variants. We frame the task as binary classification between putatively causal and non-causal variants, allowing to evaluate several 086 state-of-the-art functional-genomics-supervised and self-supervised models, alongside integrative 087 methods and their ensembles. We find that alignment-based integrative and self-supervised mod-088 els compare favorably for Mendelian traits and complex disease traits, while functional-genomics-089 supervised models do better on complex non-disease traits. The classification of variants is substan-090 tially harder for complex traits, but consistent improvement is observed by ensembling input and 091 predicted features from different models. Additionally, we introduce a new gLM trained specifi-092 cally on regulatory regions and demonstrate that it compares favorably with other alignment-free self-supervised language models.

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2 BACKGROUND

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One of the essential quests in biology is to understand the genotype-to-phenotype relationship (Fig-098 ure 1). The genotype is the genetic makeup of an organism, i.e., the set of DNA sequences composing each genome. The phenotype is the collection of observable traits of an individual, such as 100 height or cholesterol levels. Phenotypic variance can be decomposed into components attributed 101 to genetic and environmental factors. The influence of non-coding genetic variants on phenotype 102 is mediated via the expression of genes in different tissues and cell types. Functional-genomics-103 supervised models attempt to learn the relationship between DNA sequence and gene expression, 104 leveraging genome-wide experimental data (Eraslan et al., 2019). Natural selection closes the loop 105 by impacting which genotypes are favored over time, based on the fitness of the phenotype on a given environment. Therefore, the space of observed DNA sequences contains rich information about the 106 underlying biology; this is precisely the signal leveraged by self-supervised DNA language models 107 (Benegas et al., 2024).

The are two classes of phenotypic traits: Mendelian and complex (Figure 2). Mendelian traits, such as hemophilia, can be strongly affected by a single mutation in a single gene. On the other hand, complex traits, such as the risk to dayalan

complex traits, such as the risk to develop 111 Alzheimer's disease, are affected by several 112 mutations in multiple genes, each typically with a small individual effect. The fact that variants 113 affecting Mendelian traits have larger pheno-114 typic effect sizes than variants affecting com-115 plex traits makes the former relatively eas-116 ier to predict, as they tend to have larger ef-117 fects on gene expression (the signal picked 118 up by functional-genomics-supervised models) 119 and tend to be subject to stronger purifying se-120 lection (the signal picked up by self-supervised 121 models). 122

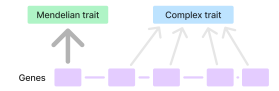


Figure 2: Mendelian vs. complex traits. A single gene typically controls a Mendelian trait, whereas a complex trait is influenced by multiple mutations across several genes, each contributing a small individual effect.

3 RELATED WORK

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126 GeneticsGym (Finucane et al., 2024) evaluates the prediction of causal variants for human com-127 plex traits, but limited to protein-coding variants. Dey et al. (2020) evaluate the prediction of non-128 coding causal variants for human complex traits, but limited to a previous generation of functional-129 genomics-supervised models. Concurrent work (Fabiha et al., 2024) also evaluates the prediction of causal variants for complex traits, but does not cover self-supervised models nor Mendelian traits. 130 Benegas et al. (2023a) evaluate the prediction of non-coding causal variants for human Mendelian 131 traits, but with a much larger, non-subsampled negative set of 2.6 million variants, which makes it 132 less practical to evaluate some of the latest, computationally expensive models. 133

134 Tang et al. (2024) benchmark the ability of functional-genomics-supervised and self-supervised 135 models to predict non-coding variant effects on gene expression, but they cover neither Mendelian nor complex traits. BEND (Marin et al., 2024) and GV-Rep (Li et al., 2024) evaluate self-supervised 136 models for the prediction of disease-associated variants from ClinVar (Landrum et al., 2020). While 137 not documented, it is likely that these variants mostly cover Mendelian rather than complex diseases. 138 Furthermore, expert-reviewed pathogenic variants in ClinVar are highly skewed towards coding and 139 splice region variants, containing only a single promoter variant and no intergenic variants (Ta-140 ble A.2). Neither of these benchmarks establishes adequate baselines for this task. BEND includes 141 a single early-generation functional-genomics-supervised model (Zhou & Troyanskaya, 2015), but 142 does not include any conservation-based model, which are usually strong for this task (Benegas 143 et al., 2023a). GV-Rep does not include any baseline. 144

Thus, TraitGym is the only benchmark of causal non-coding variant prediction for both Mendelian and complex human traits. Furthermore, it is the only available framework to evaluate both the latest functional-genomics-supervised and self-supervised models, as well as strong non-neural baselines.

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4 BENCHMARK DATASETS

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TraitGym consists of two curated datasets of non-coding genetic variants affecting Mendelian and complex traits (Table 1). We focus on non-coding variants since understanding their impact is a particularly important use case for DNA sequence models, compared to coding variants which are more commonly interpreted using protein sequence models. Further, we focus on single-nucleotide variants, the most common form of genetic variation, which is still challenging to interpret. Our data curation process is outlined in Figure 3 and additional details are provided in Appendix A.1.

Mendelian traits. Curated causal non-coding variants for 113 Mendelian diseases were collected
from Online Mendelian Inheritance in Man, OMIM (Smedley et al., 2016). For additional stringency, we filtered out a small percentage of variants with minor allele frequency (MAF) greater
than 0.1% in the Genome Aggregation Database, gnomAD (Chen et al., 2024). We used gnomAD common variants (MAF > 5%) as controls.

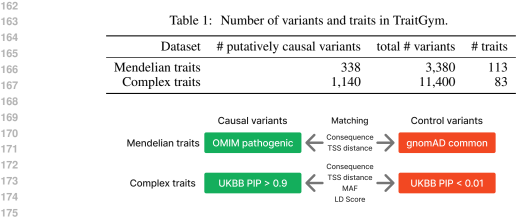


Figure 3: Matching putatively causal and control variants. Nine matched control variants are used for each putatively causal variant, within each chromosome. See the text for the details.

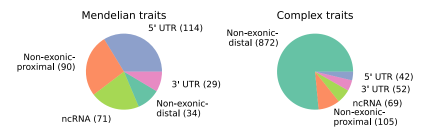


Figure 4: Distribution of consequence classes of putative causal non-coding variants.

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Complex traits. Putative causal and control non-coding variants for 83 complex traits were obtained by processing statistical fine-mapping results (Kanai et al., 2021) from association studies in the UK BioBank data (Bycroft et al., 2018). Specifically, we used variants with posterior inclusion probability (PIP) in the credible set greater than 0.9 in any trait as positives and variants with PIP < 0.01 in all traits as controls. We additionally filtered the positive set to genome-wide significant variants ($p < 5 \times 10^{-8}$).

Variant type (or consequence) annotation. We annotated the consequence (e.g., intergenic, intronic, 5' UTR, 3' UTR, etc.) of each variant using Ensembl (McLaren et al., 2016), and refined this annotation by overlapping with candidate *cis*-regulatory elements from ENCODE (Epstein et al., 2020). Distal non-exonic variants (potential enhancers) comprise a small proportion (10%) in the Mendelian traits dataset but the vast majority (76%) in the complex traits dataset (Figure 4).

201 Matching positives and negatives. For each putative causal non-coding variant, we sampled 9 non-202 coding variants from the control set, matching chromosome, consequence, and distance to transcrip-203 tion start site (TSS). For complex traits, we additionally matched MAF and linkage disequilibrium 204 (LD) score (Bulik-Sullivan et al., 2015) in the UK BioBank. We sampled only 9 controls per positive variant in order to be able to evaluate even the most computationally demanding models. However, 205 we also provide a larger version of the dataset with millions of negative controls per positive variant, 206 for which we evaluate a subset of the models. This expanded version of the dataset for Mendelian 207 traits does not require any subsampling of negatives, but for complex traits we do subsample to 208 match the MAF distribution (Finucane et al., 2024), while still keeping millions of variants. 209

Task definition. The task is to classify whether a variant is putatively causal for any trait or not.
The input data consist of the reference and alternate allele together with the DNA sequence context.
As evaluation metric, we calculate the area under the precision recall curve (AUPRC) for each chromosome (for a model trained on the remaining chromosomes), and then compute a weighted average across chromosomes based on sample size, together with a standard error estimated via bootstrapping (described in Appendix A.2.4). The baseline AUPRC is 0.1, which is the proportion of positives.

Model	Dependencies		# params	Context size	# extracted features	Source	
	Functional genomics	Alignment	Population data				
Functional-geno	mics-supervis	ed models					
Enformer	Yes	No	No	246M	196K	5,138	Avsec et al. (2021)
Sei	Yes	No	No	890M	4K	41	Chen et al. (2022)
Borzoi	Yes	No	No	186M	524K	7,617	Linder et al. (2023)
Self-supervised	models						
GPN-MSA	No	Yes	No	86M	128	770	Benegas et al. (2023a)
NT	No	No	No	2.5B	6K	2,562	Dalla-Torre et al. (2023
HyenaDNA	No	No	No	14M	160K	258	Nguyen et al. (2023)
Caduceus	No	No	No	8M	131K	514	Schiff et al. (2024)
gLM-Promoter	No	No	No	152M	512	770	This work
Integrative mod	els						
CADD	Yes	Yes	Yes	N/A	N/A	114	Schubach et al. (2024)

Table 2: Benchmarked models.

Table 3: Extracted features and zero-shot scores for each model type.

Model type	Extracted features	Zero-shot score
Functional-genomics supervised (Enformer/Borzoi)	ℓ_2 scores: change in activity in each track ℓ_2 of ℓ_2 scores: aggregation of ℓ_2 scores across several tracks (all + within each assay type)	ℓ_2 of ℓ_2 scores (all tracks)
Functional-genomics supervised (Sei)	Change in sequence class scores	Max absolute change in se- quence class scores
Self-supervised	LLR, abs(LLR) Embeddings inner product for each hidden di- mension	LLR, $abs(LLR)$ Embeddings inner product, ℓ_2 distance, cosine distance
Integrative	CADD input features, CADD score	CADD score

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5 MODELS

We benchmark functional-genomics-supervised models, self-supervised gLMs and integrative models (Table 2). We introduce a new gLM, called gLM-Promoter, trained using the genomes of 434 animal species, following the training objective of GPN (Benegas et al., 2023b) and the ByteNet convolutional architecture (Kalchbrenner et al., 2017; Yang et al., 2024). It is only trained on promoters as an attempt to focus on regulatory regions (we would have liked to train on enhancers as well but no annotation exists for non-model organisms). Additional details on models are provided in Appendix A.2.

We evaluate zero-shot model scores as well as ridge logistic regression classifiers (linear probing)
trained using extracted features (Table 3). We use a number of folds equal to the number of chromosomes. In each fold, we test on a single chromosome using a model trained on the remaining chromosomes, and the regularization hyperparameter is chosen based on cross-validation on the training chromosomes (detailed in Appendix A.2.4).

262 Functional-genomics-supervised models. Sequence-to-function models predict activity in thou-263 sands of different functional genomic tracks, covering different assays, such as gene expression or 264 chromatin accessibility, in different tissues and cell types. As variant effect prediction features, we 265 calculate the norm (across spatial positions) of the predicted log-fold-change in activity between the 266 reference and the alternate sequence, for each separate track (referred to as " ℓ_2 score" in Linder et al. 267 (2023)). As zero-shot score, we aggregate the ℓ_2 scores of different tracks by taking their ℓ_2 norm (" ℓ_2 of ℓ_2 scores"). Sei (Chen et al., 2022) adopts a different variant scoring approach; it maps each 268 sequence into discrete classes, such as promoters or brain-specific enhancers, and scores a variant 269 according to how much it impacts the relative scores of different classes.

270 Self-supervised models. For self-supervised gLMs, a popular zero-shot score is the log-likelihood 271 ratio (LLR) between the alternate and reference allele¹, which has been shown to reflect learned 272 functional constraints, such as transcription factor binding sites (Benegas et al., 2023b). Good re-273 sults have also been obtained comparing the embeddings of the alternate and reference alleles (Dalla-274 Torre et al., 2023; Mendoza-Revilla et al., 2024). We evaluate these different scoring approaches for each model (Table A.8) and choose the best performing one when benchmarking against other mod-275 els (Table A.9). We additionally obtain a high-dimensional featurization of a variant by calculating 276 the inner product (across genomic positions in a given window) between contextual embeddings of 277 the alternate and reference sequences, for each hidden dimension separately. 278

Integrative models. CADD (Schubach et al., 2024) is built on top of a broad set of curated annotations, including conservation, biochemical activity, population-level data as well as predictions from several machine learning models. Utilizing this rich set of input features, CADD is a logistic regression model trained to distinguish proxy-deleterious from proxy-neutral variants. The output of the model is called the CADD score, which we use as zero-shot score. In this paper, we also train our own models using the broad set of CADD *input* features, which we refer to as CADD features even though they are the input, not the output, of CADD.

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288 289 RESULTS

290 Mendelian traits. Among zero-shot scores, CADD and GPN-MSA perform the best, but a super-291 vised model trained using CADD input features achieves the best performance when using linear probing (Figure 5). GPN-MSA is a gLM for the human genome that leverages whole-genome 292

sequence alignments across diverse 293 multiple species. Among the models studied in this paper, CADD and 295 GPN-MSA are the only ones explic-296 itly incorporating conservation fea-297 tures, which might be particularly 298 helpful to predict causal variants for 299 Mendelian traits, expected to be un-300 der relatively strong purifying se-301 lection. Next come the functional-302 genomics-supervised models Borzoi and Enformer. Alignment-free gLMs 303 come last, with our new gLM-304 Promoter model clearly performing 305 the best among them. Only 19 ad-306 ditional positive variants are included 307 when using a more relaxed MAF cut-308 off of 1%, with very similar results 309 (Figure A.1). We explored match-310 ing negatives from the same gene 311 rather than from the same chromo-312 some, which required dropping many 313 variants that could not be properly matched, but with similar overall 314 conclusions (Figure A.2). 315

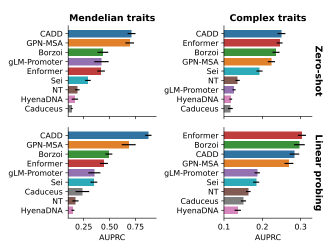
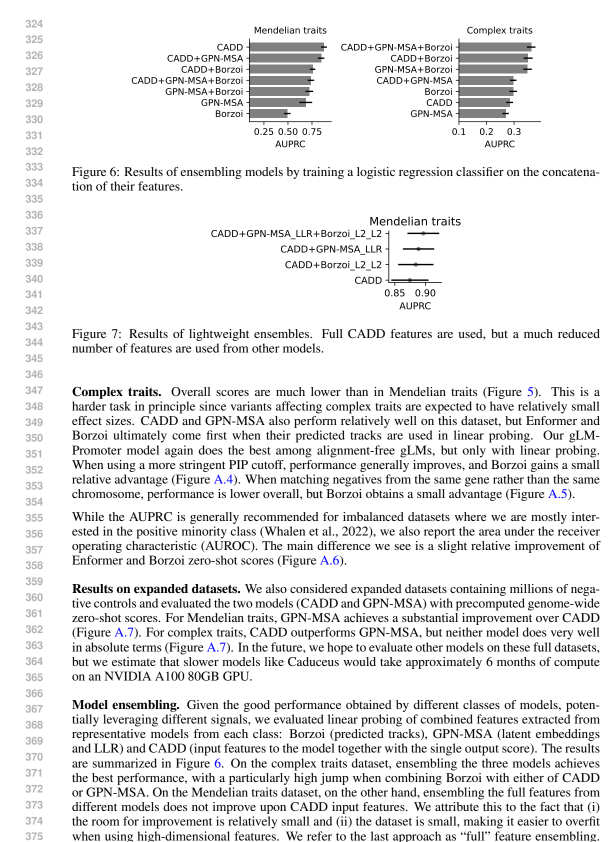


Figure 5: Results on each dataset with zero-shot and linear probing approaches. Zero-shot scores are described in Table 3. For linear probing, we use 113 CADD input features, together with the single CADD output score, while for the other models we only use output features (predicted tracks, LLR or embedding similarity).

316 CADD is the only model trained on variants and its training variants overlap with around 1% of 317 the variants in our datasets (Table A.10). However, CADD's positives and negatives are not de-318 fined based on causal variant annotations (Schubach et al., 2024), and they do not exhibit a clear 319 association with the positive or negative sets in our datasets (Table A.10). We repeated our analysis 320 upon removing this small amount of overlapping variants and found that the aforementioned results remain stable (Figure A.3). 321

¹The absolute value of the LLR is more appropriate when we want scores to be invariant to swapping ref. and alt., as in the case of association studies.



However, we do see small improvements when ensembling CADD with a reduced number of features from other models (LLR for GPN-MSA and " ℓ_2 of ℓ_2 scores" for Borzoi), which we refer to as "lightweight" feature ensembling (Figure 7).

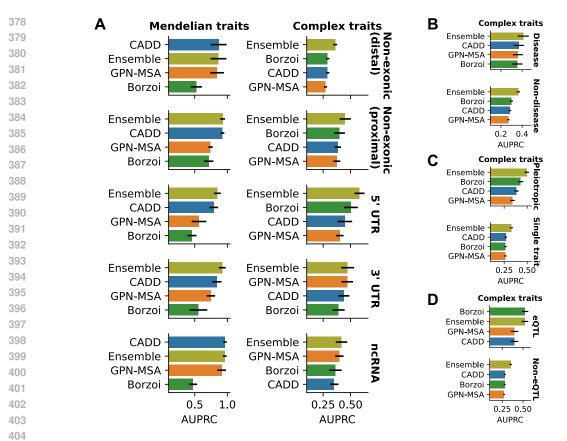


Figure 8: Stratified results. (A) Results by consequence (variant type). The best score is reported
between zero-shot and linear probing. Full feature ensemble is evaluated for complex traits, but
lightweight feature ensemble is evaluated for Mendelian traits. (B) Results for disease vs. nondisease complex traits. The best score is reported between zero-shot and linear probing. (C) Results
for pleiotropic vs. non-pleiotropic variants. The best score is reported between zero-shot and linear probing. (D) Results for complex traits variants stratified by whether or not they overlap with finemapped eQTLs. The best score is reported between zero-shot and linear probing.

Results by consequence (variant type). We also evaluated the performance stratified by variant consequence classes (Figure 8A). The most important insight here is that the advantage of ensembling for complex traits holds within each consequence class, so it is not simply that different models are experts on different consequences. Second, we note that distal (TSS distance > 1 kb) non-exonic variants for complex traits (which make up the majority) are the hardest class overall. Lastly, while Borzoi performs the worst for Mendelian traits, the gap is the smallest for proximal non-exonic variants.

We also inspected the performance of gLM-Promoter on different consequences, given that it was trained only on promoters (Figure A.8). gLM-Promoter's zero-shot scores perform better on proximal non-exonic and 5' UTR variants, which lie in the regions of the gene with the highest overlap with the model's training data (256 bp around the TSS). Except for the aforementioned classes in Mendelian traits, linear probing outperforms zero-shot scores.

Results by trait. We also report performance (Table A.11) for specific traits with sufficiently many putative causal variants and not overlapping too much with each other; specifically, traits with at least 10 causal variants and less than 10% overlap of causal variants with other traits. Ensembling wins in the majority of these traits. Among the 1,140 putative causal variants for complex traits, only 53 affect a disease trait (Table A.1). We evaluated the results stratified by disease vs. non-disease complex traits, pooled given the small sample size (Figure 8B)—for example, our dataset only contains 3 non-coding variants affecting the risk of developing Alzheimer's disease. We note that causal variants for disease traits are easier to classify overall than for non-disease traits, and that

Dataset	Category	Feature	AUPRC	Description
Mendelian traits	Alignment Functional genomics	ZooVerPhyloP EncodetotalRNA-max	0.673 0.348	Conservation in mamm Max. RNA-seq level
	Population data	(-)Freq100bp	0.509	# common variants 100bp
Complex traits	Alignment Functional	ZooPriPhyloP EncodeDNase-max	0.225 0.145	Conservation in prima Max. DNase-seq level
	genomics Population data	(-)Freq10000bp	0.131	# common variants 10kb
	RN CAC ATA CH	GE - CHIP		-
	CAC	GE - CHIP - AC - DNASE - IIP - CAGE -		-
	CAC ATA CH	GE - CHIP - AC - DNASE - DNASE - CAGE - CAGE - RNA - 0.25 0.50 0	.1 0.2	
	CAC ATA CH	GE - CHIP - CHIP - CHIP - CHIP - CAGE	.1 0.2 AUPRC	
Figure	CAC ATA CH DNAS	GE - CHIP - AC - DNASE - CAGE - CAGE - SE - RNA - 0.25 0.50 0 AUPRC	AUPRC	
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C	CAC ATA CH DNAS 9: Results o	$\begin{array}{c} \text{GE} & \text{CHIP} \\ \text{DNASE} \\ \text{DNASE} \\ \text{CAGE} \\ \text{RNA} \\ 0.25 & 0.50 & 0 \\ \text{AUPRC} \\ \text{f} ``\ell_2 \text{ of } \ell_2 \text{ scores'' aggregat} \end{array}$	AUPRC ting differe	ent assays (Borzoi).
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р 0 such overlap for Mendelian trait variants, as expected given their low allele frequencies. The low 464 overlap of complex trait and eQTL variants is well known and Mostafavi et al. (2023) discuss several 465 hypotheses for the cause. We found that eQTL-overlapping variants are much easier to predict 466 than non-eQTL-overlapping variants, across all model types (Figure 8D). We also note that Borzoi 467 achieves a wide margin compared to other models and little is gained from ensembling. We observed 468 that eQTL-overlapping variants are enriched in exonic variants (Fisher's exact $p = 8 \times 10^{-8}$) and, among non-exonic variants, they have lower TSS distances (Mann Whitney $p = 4 \times 10^{-4}$), all of 469 which could explain their increased predictability. 470

Interpreting CADD features. CADD contains informative features from three orthogonal categories: alignment, functional genomics, and population genetic data (Table 4). Conservation features are the most predictive overall. Conservation in mammals is most predictive for Mendelian traits, whereas conservation in primates is most predictive for complex traits. This might be due to the fact that enhancer-like regions, where most causal variants for complex traits lie, tend to only be alignable over shorter evolutionary distances than other functional regions (Phan et al., 2024).

Interpreting Borzoi features. We evaluated the performance of aggregated Borzoi scores across specific experimental assays (Figure 9). Of note, gene expression tracks (RNA and CAGE) perform the best on Mendelian traits, while epigenetic tracks (ATAC, CHIP and DNASE) perform the best on complex traits. It has been shown that models such as Borzoi tend to particularly struggle with finding causal variants affecting gene expression when these are distal as opposed to proximal (Karollus et al., 2023). In the case of distal causal variants for complex traits (which make up the majority, see Figure 4), epigenetic tracks might instead be more informative.

485 A key feature of functional-genomics-supervised models such as Borzoi is that their features are associated with a specific tissue or cell type, which can help interpret disease pathways as well as de-

Table 5: Top three tissue/cell types for different traits, ranked by the highest AUPRC of Borzoi
 predicted tracks from such tissue/cell type.

Trait	Tissue/cell type/cell line	AUPRC
Mendelian traits		
Beta-thalassemia	aorta	0.997
	stomach	0.988
	adrenal gland	0.986
Hemophilia B	liver	1.0
	HepG2	1.0
	hepatocyte	1.0
Hypercholes-	CD8+ T cell	0.983
terolemia-1	HepG2	0.975
	CD4+ T cell	0.972
Complex traits		
Monocyte count	neutrophil	0.559
2	CD14+ monocyte	0.559
	HL-60	0.559
Hemoglobin A1c	K562	0.449
	erythroblast	0.423
	hematopoietic progenitor	0.412
High density	liver	0.44
lipoprotein	abdominal adipose tissue	0.42
cholesterol	adrenal gland	0.417

sign therapeutics. For traits where Borzoi achieved a good performance, we inspected the tissue/cell
type of the top features, and found that they are usually well aligned with previous knowledge (Table 5). For example, the top tissues for high density lipoprotein cholesterol are liver, abdominal
adipose tissue and adrenal gland.

7 DISCUSSION

Conclusion. TraitGym allows to benchmark DNA sequence models on the challenging task of pre-dicting causal variants in human genetics. Alignment-based, conservation-aware models compare favorably on Mendelian traits and complex disease traits, while functional-genomics-supervised models achieve the best performance on complex non-disease traits. A reason for hope in the par-ticularly challenging complex traits dataset is that ensembling predictions and input features from different models yields consistent improvements. We find that alignment-free gLMs are not com-petitive on causal variant prediction. The best performing model among them-gLM-Promoter, developed in this work—is not the largest gLM, nor does it have a long context. However, one of its defining characteristics is that it was trained only on functional regions; this suggests that, as previ-ously proposed (Tang et al., 2024; Benegas et al., 2024), data curation may warrant more research than architectures. We leave this as promising future work.

Limitations and future extensions. The major limitation for benchmarking causal variant predic-tion for human traits is that the number of known causal variants is small, especially for non-coding regions. One implication is that TraitGym lacks power to distinguish subtle, incremental differences in performance, and is more suited to recognize larger improvements in this task. In the long term, we expect the number of known causal variants to increase as experimental and statistical techniques improve, together with larger and more diverse patient cohorts. In the short term, we hope to expand the dataset to include variants from other cohorts such as FinnGen (Kurki et al., 2023) and BioBank Japan (Nagai et al., 2017). One of the challenges is that, while many fine-mapping results are pub-licly available, it is still hard to get access to other quantities such as LD scores, which are important for constructing a rigorous control set.

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- 756 A APPENDIX
- 758 A.1 DATASETS

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760 A.1.1 MENDELIAN TRAITS

Non-coding pathogenic OMIM variants were obtained from Table S6 in Smedley et al. (2016). Common variants were obtained from gnomAD (Chen et al., 2024) (version 3.1.2).

764 A.1.2 COMPLEX TRAITS

UK BioBank fine-mapping results (Kanai et al., 2021) were downloaded from https://www.
finucanelab.org/data (version: Dec. 3rd, 2019). As recommended to increase fine-mapping
accuracy (Kanai et al., 2021), we averaged the posterior inclusion probability (PIP) from FINEMAP
(Benner et al., 2016) and SuSiE (Wang et al., 2020), and excluded variants where the two methods
disagreed by more than 5%.

Table A.1: Complex traits in our dataset which are considered diseases or disorders.

774	Trait
775	Atrial fibrillation
776	Autoimmune disease (Phecode + Self-reported)
777	Alzheimer disease (LTFH)
778	Asthma
779	Blood clot in the lung
780	Breast cancer
781	Coronary artery disease
782	Colorectal cancer
783	Cholelithiasis
784	Seen doctor (GP) for nerves, anxiety, tension or depression
785	Blood clot in the leg
	Fibroblastic disorders
786	Glaucoma (Phecode + Self-reported)
787	Hypothyroidism
788	Inflammatory bowel disease
789	Inguinal hernia
790	Insomnia
791	Migraine (Self-reported)
792	Prostate cancer
793	Type 2 diabetes
794	Type 2 diabetes (adjusted by BMI)
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A.1.3 VARIANT ANNOTATION

Consequences were annotated using Ensembl VEP (McLaren et al., 2016) (release 109.1), using flags --most_severe and --distance 1000 (used to distinguish upstream and downstream from intergenic variants). We only kept non-coding consequences (Table A.3). We discarded splice region variants, such as splice donor variants, as these were very few in number. Coding variants, as well as non-coding variants with a very high expected impact such as in splice donors, are excluded from our analysis.

We refined the annotation of non-exonic variants by checking overlap with each of five different ENCODE candidate *cis*-regulatory element (cCRE) categories (Epstein et al., 2020) (Table A.4).
We additionally refined the annotation if a variant overlapped not a cCRE but the 500-bp flank of a cCRE, similar to Finucane et al. (2015). When we match negative controls, we make sure to keep the exact same proportion of consequences, including the distribution of cCRE elements and their flanks. For the analysis of performance by consequence, however, we simplify the categorization of non-exonic variants into proximal (TSS dist. ≤ 1 kb) and distal (TSS dist. > 1 kb).

811 Table A.2: ClinVar "Pathogenic" variant consequences (reviewed by expert panel). ClinVar release:
812 20240909.

Consequence	Count
stop_gained	1683
missense_variant	980
splice_donor_variant	173
splice_acceptor_variant	156
splice_region_variant	35
start_lost	33
splice_donor_5th_base_variant	21
<pre>splice_donor_region_variant</pre>	13
<pre>splice_polypyrimidine_tract_variant</pre>	8
synonymous_variant	5
intron_variant	5
stop_lost	3
3_prime_UTR_variant	1
upstream_gene_variant	1

Table A.3: Selected consequences in this study.

Consequence
Non-exonic
intergenic_variant
upstream_gene_variant
downstream_gene_variant
intron_variant
Exonic
5_prime_UTR_variant
3_prime_UTR_variant
<pre>non_coding_transcript_exon_variant</pre>

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TSS distance was computed with respect to protein coding transcripts only. MAF and LD scores for the UK Biobank computed by the Pan-UK Biobank initiative (Karczewski et al., 2024) were downloaded from s3://pan-ukb-us-east-1/ld_release/UKBB.EUR.ldscore.ht.

GTEx fine-mapping results where downloaded from https://www.finucanelab.org/ data. We used a similar PIP cutoff of 0.9 in any tissue, combined between FINEMAP and SuSiE, to define putative causal eQTL variants.

A.1.4 MATCHING CONTROLS

853 Nine negative control variants were sampled for each positive causal variant. Chromosome and 854 consequence were matched exactly. We matched variants with the most similar TSS distance, as 855 well as MAF and LD score in the complex traits dataset. More precisely, we defined a vector space of (TSS distance, MAF, LD score) tuples, applied scikit-learn's robust scaler (Pedregosa et al., 2011), 856 and selected negative variants minimizing the euclidean distance to the positive variant. Table A.5 857 shows that the matched features have minimal predictive power, as intended. For special cases 858 where there were not enough negative controls to match positive variants for a given chromosome 859 and consequence, we subsampled the positive variants until we had at least nine controls per positive 860 variant. 861

For the full version of the complex traits dataset, we created 100 equal-size MAF bins and subsampled the negative set until the proportion of variants in each bin was equal to that of the positive set.

Table A.4:	ENCODE	cCRE	categories.
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Category
PLS (promoter-like signature) pELS (proximal enhancer-like signature) dELS (distal enhancer-like signature) DNase-H3K4me3 CTCF-only

Table A.5: Global AUPRC of matched features, close to baseline (0.1).

Dataset	Feature	AUPRC
Mendelian traits	(-) TSS distance	0.115
Complex traits Complex traits	(-) TSS distance MAF	$0.104 \\ 0.101$
Complex traits	(-) LD score	0.104

A.2 MODELS

A.2.1 PUBLISHED MODELS

We downloaded several models from Hugging Face Hub (Wolf et al., 2020) (Table A.6). We downloaded Enformer and Borzoi from gReLU's Model Zoo (Lal et al., 2024). Sei scores were obtained via their web server: https://hb.flatironinstitute.org/sei. We obtained CADD v1.7 scores and annotations from https://krishna.gs.washington.edu/download/ CADD/v1.7/GRCh38/whole_genome_SNVs_inclAnno.tsv.gz.

Table A.6: Hugging Face Hub models.

Model	Hugging Face Hub path	
GPN-MSA	songlab/gpn-msa-sapiens	
NT	InstaDeepAI/nucleotide-transformer-2.5b-multi-species	
HyenaDNA	LongSafari/hyenadna-medium-160k-seqlen-hf	
Caduceus	kuleshov-group/caduceus-ps_seqlen-131k_d_model-256_n_layer-16	

A.2.2 OUR GLM-PROMOTER MODEL

gLM-Promoter was trained on 512-bp sequences centered at TSSs of protein-coding genes from reference genomes of animal species. TSS coordinates were obtained from the gene annotations available at NCBI Datasets (O'Leary et al., 2024). Species available at NCBI Datasets were sub-sampled, among those with gene annotations, to keep at most one per family. This resulted in 434 reference genomes. gLM-Promoter's training objective follows GPN: base-pair-level tokenization and masked language modeling of local windows of 512-bp with downweighting of repeat positions (soft-masked in the reference genome). gLM-Promoter's architecture follows ByteNet (Kalchbren-ner et al., 2017; Yang et al., 2024), consisting of blocks alternating dilated convolutions and feed-forward layers. Hyperparameters are displayed in Table A.7. Training took approximately 2 weeks using 4 NVIDIA A100 40GB GPUs.

- A.2.3 FEATURE EXTRACTION
- **Functional-genomics-supervised models.** Let $y_i \in \mathbb{R}^L_+$ be the predicted activity for genomic track *i* in each of *L* spatial positions. The " ℓ_2 score" (Linder et al., 2023) is defined as the norm of the

919	Table A.7: gLM-Promoter training hyperparameters		
920	Window size	512	
921	Repeat weight	0.01	
922	Embedding dimension	1024	
923	Slim	True	
924	Convolutional blocks	64	
925	Convolutional kernel size (first block)	9	
926	Convolutional kernel size (remaining blocks)	5	
927	Convolutional dilation schedule	$1, 2, 4, 8, 16, 32, 64, 128, 1, \dots$	
928	Optimizer	AdamW	
929	Weight decay	0.01	
930	Batch size	2048	
931	Steps	370 K	
932	Learning rate	10^{-3}	
933	Learning rate warmup	1 K steps	

Table A 7. al M Descretes too in in a house measure of

log-fold-change between the predicted activity for the reference vs. alternate sequences:

$$\ell_2 \operatorname{score}_i := \left\| \log_2 \left(1 + \boldsymbol{y}_i^{(\operatorname{alt})} \right) - \log_2 \left(1 + \boldsymbol{y}_i^{(\operatorname{ref})} \right) \right\|$$
(1)

We define the " ℓ_2 of ℓ_2 score" as the norm of the ℓ_2 scores across tracks in a set A (e.g. all genomic tracks, or all genomic tracks from the same experimental assay):

$$\ell_2 \text{ of } \ell_2 \text{ score}(\mathbb{A}) := \|(\ell_2 \text{ score}_i, i \in \mathbb{A})\|$$
(2)

For Sei we used the official scores provided in their web server https://hb. flatironinstitute.org/sei.

Self-supervised models. We compute the log-likelihood ratio between the reference and alternate alleles:

$$\log \frac{\mathbb{P}(\text{alt})}{\mathbb{P}(\text{ref})} \tag{3}$$

For masked language models, it can be computed from the output probabilities when the variant position is masked. For autoregressive models (HyenaDNA), it can be computed from the likelihood of the entire reference and alternate sequences. We also compute similarity in the embedding space. Let $Z \in \mathbb{R}^{D \times L}$ be the sequence embedding with D hidden dimensions and L spatial positions. For HyenaDNA, an autoregressive model, we take the embedding of the rightmost position (could be interpreted as L = 1). We compare the reference and alternate embedding using the Euclidean distance:

$$\left\| \boldsymbol{Z}^{(\text{ref})} - \boldsymbol{Z}^{(\text{alt})} \right\|_{F}$$
(4)

cosine distance:

$$1 - \frac{\langle \boldsymbol{Z}^{(\text{ref})}, \boldsymbol{Z}^{(\text{alt})} \rangle_F}{\left\| \boldsymbol{Z}^{(\text{ref})} \right\|_F \left\| \boldsymbol{Z}^{(\text{alt})} \right\|_F}$$
(5)

and innner product:

$$\langle \boldsymbol{Z}^{(\mathrm{ref})}, \boldsymbol{Z}^{(\mathrm{alt})} \rangle_F$$
 (6)

To obtain a high-dimensional featurization of a variant we calculate the inner product separately for each individual hidden dimension d:

$$\langle \boldsymbol{Z}_{d:}^{(\text{ref})}, \boldsymbol{Z}_{d:}^{(\text{alt})} \rangle \tag{7}$$

For both functional-genomics-supervised and self-supervised models, we always average the pre-dictions using the forward vs. reverse strand, to ensure reverse-complement invariance.

972 A.2.4 LINEAR PROBING 973

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We train a ridge logistic regression classifier pipeline using scikit-learn (Pedregosa et al., 2011), using default arguments as much as possible (Listing 1). The pipeline starts with imputation (only relevant for CADD input features) and standardization. To choose the regularization hyperparameter, we do a grid search using group K-fold cross-validation, with the groups consisting of the training chromosomes. Since we are in a high-dimensional regression setting, We use the default number of grid points (10), but shift the range to allow for heavier regularization given our regression setting is very high-dimensional.

We repeat the entire pipeline training on all but one chromosome and predicting on the held-out chromosome. At the end we obtain predictions for all chromosomes, but each from a separate logistic regression model. Therefore, instead of calculating a global AUPRC, we calculate the AUPRC within each chromosome, and then perform a weighted average based on sample size. To obtain a standard error, we calculate the standard deviation of the distribution of weighted means performed on 1000 bootstrap samples of chromosomes. To allow easy comparison, we also use the weighted average AUPRC to evaluate zero-shot scores, even though it is not strictly necessary.

We only evaluate zero-shot scores on the full version of the datasets. We obtain standard errors from
 100 bootstrap samples within the positive and negative sets, in order to maintain the proportion of
 positives.

```
from sklearn.impute import SimpleImputer
992
      from sklearn.linear model import LogisticRegression
993
      from sklearn.model_selection import GroupKFold, GridSearchCV
994
      from sklearn.pipeline import Pipeline
995
      from sklearn.preprocessing import StandardScaler
996
      def train_logistic_regression(X, y, groups):
997
           pipeline = Pipeline([
998
                ('imputer', SimpleImputer(
999
                    missing values=np.nan, strategy='mean',
1000
                    keep_empty_features=True,
1001
               )),
1002
                ('scaler', StandardScaler()),
1003
                ('linear', LogisticRegression(
1004
                    class weight="balanced",
                    random state=42,
               ))
           ])
           Cs = np.logspace(-8, 0, 10)
1008
           param_grid = {
1009
                'linear__C': Cs,
1010
           }
1011
           clf = GridSearchCV(
1012
               pipeline,
1013
               param_grid,
1014
               scoring="average_precision",
1015
               cv=GroupKFold(),
1016
               n jobs=-1,
1017
           )
           clf.fit(X, y, groups=groups)
           return clf
1020
                   Listing 1: Logistic regression classifier (the default penalty is \ell_2).
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```

1026 A.3 ADDITIONAL RESULTS

Table A.8: AUPRC for different gLM zero-shot scores. In boldface: scores within 1% of best score (for a given model).

		LLR	abs(LLR)	L2 dist.	Cosine dist.	Inner prod.
Dataset	Model					-
Mendelian traits	GPN-MSA	0.694	0.654	0.207	0.208	0.301
	gLM-Promoter	0.422	0.379	0.345	0.263	0.169
	NT	0.120	0.098	0.188	0.186	0.185
	HyenaDNA	0.115	0.106	0.117	0.116	0.165
	Caduceus	0.108	0.088	0.135	0.135	0.131
Complex traits	GPN-MSA	0.212	0.224	0.150	0.150	0.177
1	gLM-Promoter	0.112	0.110	0.126	0.126	0.125
	NT	0.101	0.100	0.118	0.119	0.136
	HyenaDNA	0.110	0.111	0.102	0.102	0.118
	Caduceus	0.098	0.097	0.115	0.115	0.117

Table A.9: Selected zero-shot approach for each gLM.

	Mendelian traits	Complex traits
GPN-MSA	LLR	abs(LLR)
gLM-Promoter	LLR	L2 dist.
NT	L2 dist.	Inner prod.
HyenaDNA	Inner prod.	Inner prod.
Caduceus	L2 dist.	Inner prod.

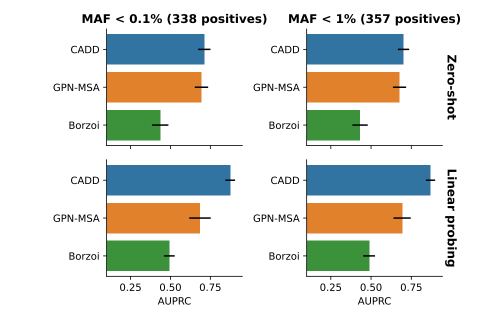
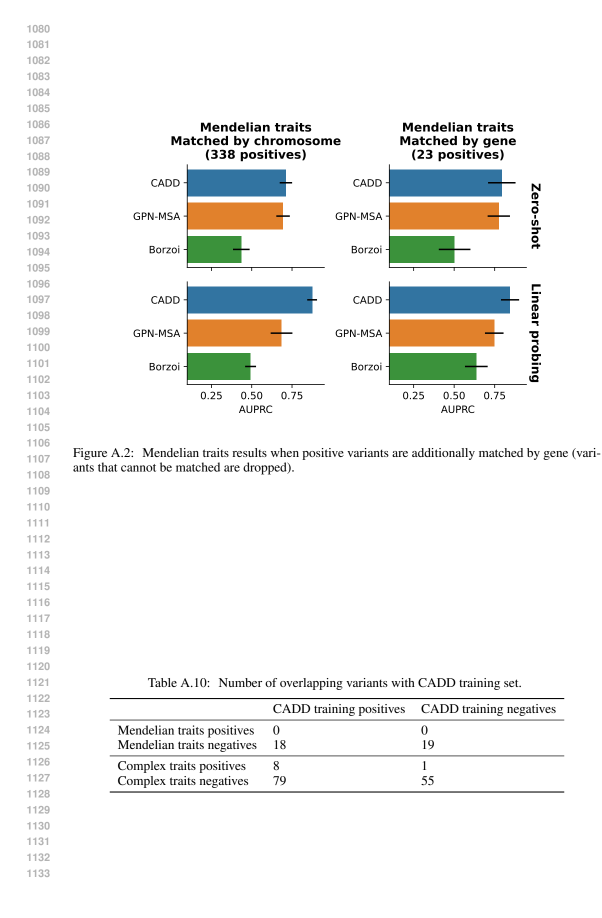
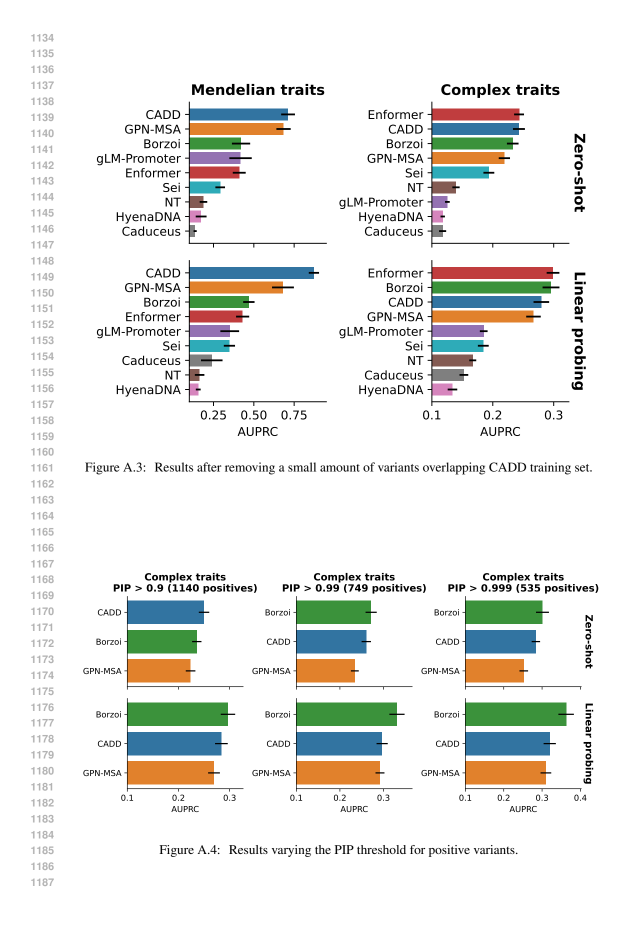


Figure A.1: Ablation of MAF cutoff for positive variants in Mendelian traits dataset.





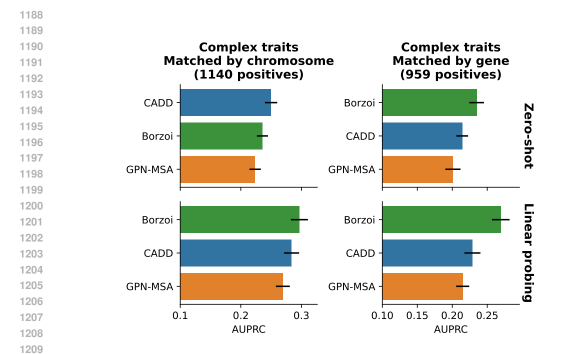
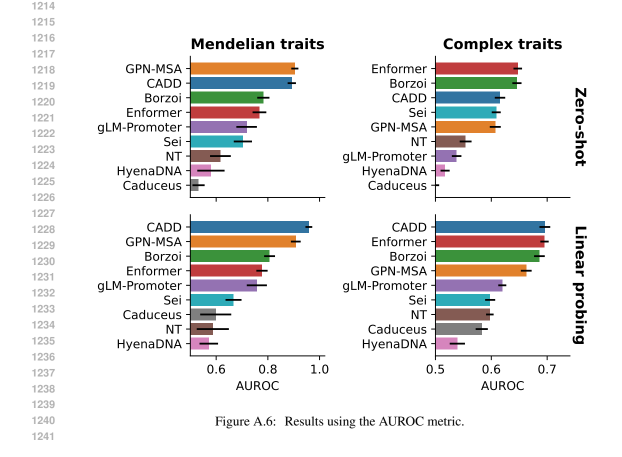


Figure A.5: Complex traits results when positive variants are additionally matched by gene (variants that cannot be matched are dropped).



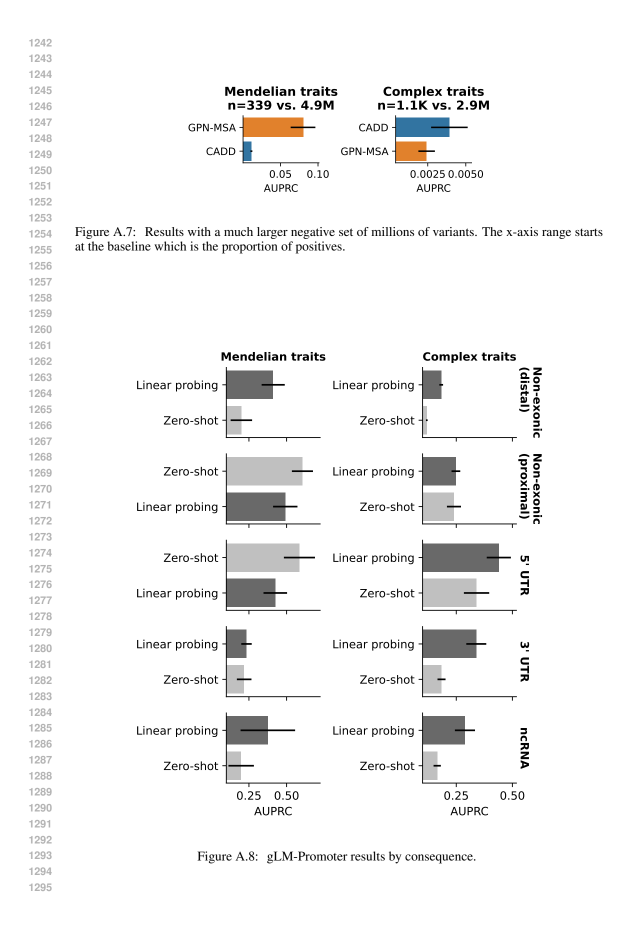


Table A.11: AUPRC for selected traits (at least 10 causal variants and less than 10% overlap of causal variants with other traits). The best score is reported between zero-shot and linear probing.
Full feature ensemble is evaluated for complex traits, but lightweight feature ensemble is evaluated for Mendelian traits. In boldface: scores within 1% of best score.

	Borzoi	GPN-MSA	CADD	Ensemble
Mendelian traits				
Hyperferritinemia	0.315	0.965	0.981	0.985
Beta-thalassemia	0.927	0.796	0.926	0.955
Pulmonary fibrosis	0.564	0.948	1.000	1.000
Hemophilia B	0.914	0.709	1.000	0.991
Cartilage-hair hypoplasia	0.594	0.987	0.923	0.918
Preaxial polydactyly II	0.546	0.959	0.969	0.967
Hypercholesterolemia-1	0.844	0.974	0.887	0.938
Dwarfism (MOPD1)	0.484	1.000	1.000	1.000
Complex traits				
Adult height	0.292	0.383	0.407	0.339
Platelet count	0.426	0.309	0.397	0.478
Estimated heel bone mineral density	0.308	0.432	0.422	0.406
Mean corpuscular volume	0.434	0.319	0.391	0.454
Monocyte count	0.561	0.404	0.375	0.535
Hemoglobin A1c	0.475	0.375	0.426	0.517
Albumin/Globulin ratio	0.455	0.431	0.516	0.559
High density lipoprotein cholesterol	0.521	0.362	0.425	0.554
Estimated glomerular filtration rate (cystain C)	0.457	0.456	0.421	0.470
Alkaline phosphatase	0.492	0.292	0.352	0.446
Gamma-glutamyl transferase	0.515	0.382	0.460	0.527
FEV1/FVC ratio	0.430	0.494	0.505	0.487
Pulse pressure	0.457	0.435	0.420	0.489
Calcium	0.468	0.433	0.425	0.408
Albumin	0.615	0.544	0.480	0.602
Body mass index	0.344	0.514	0.436	0.499
Balding Type 4	0.459	0.536	0.414	0.625
Blood clot in the leg	0.574	0.551	0.498	0.565