

SPECIAL ARTICLE

Assessing computational predictions of the phenotypic effect of cystathionine-beta-synthase variants

Laura Kasak^{1,2} | Constantina Bakolitsa¹ | Zhiqiang Hu¹ | Changhua Yu¹ | Jasper Rine³ | Dago F. Dimster-Denk³ | Gaurav Pandey¹ | Greet De Baets^{4,5} | Yana Bromberg⁶ | Chen Cao^{7,8} | Emidio Capriotti⁹ | Rita Casadio¹⁰ | Joost Van Durme^{4,11} | Manuel Giollo¹² | Rachel Karchin¹³ | Panagiotis Katsonis¹⁴ | Emanuela Leonardi¹⁵ | Olivier Lichtarge¹⁴ | Pier Luigi Martelli¹⁰ | David Masica¹³ | Sean D. Mooney¹⁶ | Ayodeji Olatubosun¹⁷ | Predrag Radivojac^{18,19} | Frederic Rousseau^{4,5} | Lipika R. Pal⁷ | Castrense Savojardo¹⁰ | Joost Schymkowitz^{4,5} | Janita Thusberg¹⁶ | Silvio C.E. Tosatto¹² | Mauno Vihinen¹⁷ | Jouni Väliäho¹⁷ | Susanna Repo¹ | John Moulton^{5,20} | Steven E. Brenner¹ | Iddo Friedberg^{21,22}

¹Department of Plant and Microbial Biology, University of California, Berkeley, California

²Institute of Biomedicine and Translational Medicine, University of Tartu, Tartu, Estonia

³California Institute for Quantitative Biosciences, University of California, Berkeley, California

⁴Switch Laboratory, VIB Center for Brain and Disease Research, Leuven, Belgium

⁵Department of Cellular and Molecular Medicine, KU Leuven, Leuven, Belgium

⁶Department of Biochemistry and Microbiology, Rutgers University, New Brunswick, New Jersey

⁷Institute for Bioscience and Biotechnology Research, University of Maryland, Rockville, Maryland

⁸Computational Biology, Bioinformatics and Genomics, Biological Sciences Graduate Program, University of Maryland, College Park, Maryland

⁹Department of Bioengineering, Stanford University, Stanford, California

¹⁰Biocomputing Group, Department of Pharmacy and Biotechnology, University of Bologna, Bologna, Italy

¹¹VIB Switch Laboratory, Vrije Universiteit Brussel, Brussels, Belgium

¹²Department of Biomedical Sciences, University of Padua, Padua, Italy

¹³Department of Biomedical Engineering and Institute for Computational Medicine, Johns Hopkins University, Baltimore, Maryland

¹⁴Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, Texas

¹⁵Department for Woman and Child Health, University of Padua, Italy

¹⁶Computational Biology Group, Buck Institute for Research on Aging, Novato, California

¹⁷Institute of Medical Technology, University of Tampere, Tampere, Finland

¹⁸School of Informatics and Computing, Indiana University, Bloomington, Indiana

¹⁹Khoury College of Computer Sciences, Northeastern University, Boston, Massachusetts

²⁰Department of Cell Biology and Molecular Genetics, University of Maryland, College Park, Maryland

²¹Department of Microbiology, Miami University, Oxford, Ohio

²²Department of Veterinary Microbiology and Preventive Medicine, Iowa State University, Ames, Iowa,

Correspondence

Iddo Friedberg, Department of Veterinary Microbiology and Preventive Medicine, Iowa State University, Ames, IA.
Email: idoerg@iastate.edu

Abstract

Accurate prediction of the impact of genomic variation on phenotype is a major goal of computational biology and an important contributor to personalized medicine.

Present Address

Dago F. Dimster-Denk, Pionyr Immunotherapeutics, San Francisco, CA; Gaurav Pandey, Department of Genetics and Genomic Sciences and Icahn Institute for Data Science and Genomic Technology, Icahn School of Medicine at Mount Sinai, New York, NY; Emidio Capriotti, BioFolD Group, Department of Pharmacy and Biotechnology (FaBiT), University of Bologna, Bologna, Italy; Emanuela Leonardi, Department for Woman and Child Health, University of Padua, Italy; David Masica, AbbVie Inc., Redwood City, CA; Sean D. Mooney, Department of Biomedical Informatics and Medical Education, University of Washington, Seattle, WA; Predrag Radivojac, Khoury College of Computer Sciences, Northeastern University, Boston, Massachusetts; Janita Thusberg, Invitae, San Francisco, CA, USA; Susanna Repo, ELIXIR, Wellcome Genome Campus, Hinxton, Cambridge, UK; Mauno Vihinen, Department of Experimental Medical Science, Lund University, Lund, Sweden; Susanna Repo, ELIXIR, Wellcome Genome Campus, Hinxton, Cambridge, UK

Funding information

National Institutes of Health, Grant/Award Numbers: R01 GM120374, R01 GM079656, R01 LM009722, R01 GM071749, R13 HG006650, R01 GM114434, R01 GM066099, U01 GM115486, U41 HG007346; Fondazione Istituto di Ricerca Pediatrica – Città della Speranza, Grant/Award Number: 18-04; Estonian Research Council, Grant/Award Number: IUT34-12; Marie Curie International Outgoing Fellowship, Grant/Award Number: P10F-GA-2009-237751; The Ministry of Education, Universities and Research (MIUR), Grant/Award Number: FFABR grant; Italian Ministry of Health, Grant/Award Number: GR-2011-02347754; The Interuniversity Attraction Pole (IAP) Network from the Science Policy of the Federal Government of Belgium (Belspo), Grant/Award Number: P7/16; National Science Foundation, Grant/Award Number: ABI 1458359; The University of Leuven (KUL); Flanders Institute for Biotechnology (VIB); The Flanders Institute for Science and Technology (IWT) (PhD grant to GDB); The Funds for Scientific Research Flanders (FWO), Grant/Award Number: G.0509.13

Computational predictions can lead to a better understanding of the mechanisms underlying genetic diseases, including cancer, but their adoption requires thorough and unbiased assessment. Cystathionine-beta-synthase (CBS) is an enzyme that catalyzes the first step of the transsulfuration pathway, from homocysteine to cystathionine, and in which variations are associated with human hyperhomocysteinemia and homocystinuria. We have created a computational challenge under the CAGI framework to evaluate how well different methods can predict the phenotypic effect(s) of CBS single amino acid substitutions using a blinded experimental data set. CAGI participants were asked to predict yeast growth based on the identity of the mutations. The performance of the methods was evaluated using several metrics. The CBS challenge highlighted the difficulty of predicting the phenotype of an *ex vivo* system in a model organism when classification models were trained on human disease data. We also discuss the variations in difficulty of prediction for known benign and deleterious variants, as well as identify methodological and experimental constraints with lessons to be learned for future challenges.

KEYWORDS

CAGI challenge, critical assessment, cystathionine-beta-synthase, machine learning, phenotype prediction, single amino acid substitution

1 | INTRODUCTION

One of the central challenges in biology is to determine the impact of genomic variation on the phenotype(s) of an organism. As the amount of genomic data increases and accumulates at an exponential rate, comprehensive, and accurate prediction algorithms are needed when biological experiments are expensive or difficult to execute (Fernald,

Capriotti, Daneshjou, Karczewski, & Altman, 2011). Missense mutations are the most studied class of protein-altering variants; however, even today the algorithms often disagree on the pathogenicity prediction of the variants (Ioannidis et al., 2016). To determine the optimal use of each algorithm in different tasks, a thorough and unbiased methodological assessment is required. The ultimate aim is to attain a better understanding of the complex

genotype–phenotype relationship, and, most importantly, provide the basis for clinical application to improve human health (Rost, Radivojac, & Bromberg, 2016). Since 2010, the Critical Assessment of Genome Interpretation (CAGI) experiment has been seeking to address these needs by evaluating bioinformatics tools developed for phenotype prediction from genomic variation data (Hoskins et al., 2017).

Cystathionine-beta-synthase (CBS, MIM# 613381) is an extensively studied vitamin-dependent enzyme involved in cysteine biosynthesis via the transsulfuration pathway. The molecular architecture of human CBS comprises an N-terminal heme-binding domain (residues 1–70), followed by the catalytic domain (residues 71–381), and a C-terminal regulatory domain (residues 412–551; Majtan et al., 2018). The heme domain, which is found only in mammalian forms of CBS, lacks any significant structural elements and exhibits significant sequence diversity. Changes in the heme's coordination environment can be transmitted to the active site ~20 Å away, leading to inhibition of CBS activity (Weeks, Singh, Madzlan, Banerjee, & Spiro, 2009). The central domain belongs to the family of pyridoxal-5'-phosphate (PLP)-dependent enzymes, with the PLP cofactor bound via a Schiff base to K119 in the CBS active site. The C-terminal domain, also known as the Bateman module, contains two consecutive CBS-motifs (residues 412–471 and 477–551) that form distinct binding sites for S-adenosyl-methionine (AdoMet) and enable CBS homotetramerization. Two high-affinity and four low-affinity AdoMet binding sites have been identified per CBS tetramer, with distinct roles proposed in the regulation and activation (Pey, Majtan, Sanchez-Ruiz, & Kraus, 2013). Catalytic and regulatory domains are joined by a flexible linker (residues 382–411) that is sensitive to proteolysis. Targeted proteolysis of CBS results in a truncated, dimeric, and more active form of the enzyme, adding another layer of CBS regulation (Skovby, Kraus, & Rosenberg, 1984; Zou & Banerjee, 2003).

Homocystinuria due to CBS deficiency (MIM# 236200) is an autosomal recessive inborn error of sulfur amino acid metabolism, characterized by increased levels of homocysteine in the urine (Mudd, Levy, & Kraus, 2001), myopia, osteoporosis, or other skeletal abnormalities. The estimated worldwide prevalence of homocystinuria is approximately 1 in 100,000 (Moorthie, Cameron, Sagoo, Bonham, & Burton, 2014). More than 160 different disease-associated variants have been identified in the CBS gene (<http://cbs.lf1.cuni.cz/index.php>). The majority of these are substitutions that do not involve catalytic residues, suggesting that their effect resides in structural or conformational perturbations leading to a misfolded protein (Majtan et al., 2018). About one-half of homocystinuric patients respond to high doses of pyridoxine, the soluble form of PLP (Mudd et al., 2001) and several mutations are clearly pyridoxine remediable (B6-responsive homocystinuria): p.A114V, p.R266K, p.R369H, p.K384E, p.L539S, and the most common substitution p.I278T, which accounts for ~20% of all homocystinuric alleles (Dimster-Denk, Tripp, Marini, Marqusee, & Rine, 2013; Moat et al., 2004; Skovby, Gaustadnes, & Mudd, 2010).

Since CBS is well studied and its ex-vivo mutation effects are easily quantified, it is a tractable system for investigating phenotype–genotype relationships, making it an attractive target for the CAGI

challenges. Here we present an assessment of computational predictions on the effects of single amino acid substitutions in the function of CBS. In the CAGI1 (2010), CBS challenge, participants were asked to predict yeast growth rates when compared with wild-type yeast based on amino acid substitution information. This data set comprised 51 synthetic single amino acid substitutions within the human CBS coding region. In the CAGI2 (2010), CBS challenge, a larger set of variants (78 amino acid substitutions) that had been observed in patients with homocystinuria was provided for the predictors. In both challenges, participants were asked to submit predictions on the effect of the variants in the function of CBS at high (400 ng/ml) and low (2 ng/ml) cofactor (pyridoxine) concentration. Both CBS predictions were blinded experiments. At the time these challenges took place, the experimental effects of the mutations had not yet been published. Altogether 38 predictions from CAGI1 and CAGI2 were assessed. Methods employed varied from the purely structural to ones combining both structural and sequence conservation information or annotation, and from meta-predictors (models that use the predictions of other methods as features) to methods driven mainly by sequence, while one submission was a set of random predictions.

In general, deleterious mutations were better predicted than variants with minor or no effects on phenotype, with the hardest to predict effects often involving variants with weak sequence and structural signals. The use of distinct assessment criteria revealed differences in performance between methods, with methods integrating sequence, structure, and functional features performing best overall.

2 | METHODS

2.1 | Dataset

The CAGI1 CBS challenge included 51 synthetic single amino acid substitutions within the human CBS (Table 1), while the CAGI2 CBS challenge included 78 single amino acid variants that had been observed in patients with homocystinuria. The experimental data used in CAGI1 and CAGI2 were published after the challenges were closed by Dimster-Denk et al. (2013) and Mayfield et al. (2012), respectively. Only one variant, p.N228K, overlapped between CAGI1 and CAGI2. In addition, four positions (p.H65, p.L154, p.V354, and p.V371) overlap between the two challenges but the amino acid substitutions are different. The variant nomenclature refers to the human CBS cDNA (Refseq NM_000071.2).

The functionality of the variants was tested in an in vivo yeast complementation assay, where the human CBS allele is expressed in yeast and functionally complements the yeast ortholog, CYS4, which was removed from the chromosome. In this assay, growth is dependent upon the level of mutant human CBS function, and growth rates are expressed as a percentage relative to wild-type grown with the same amount of exogenous pyridoxine supplementation. An experimental standard deviation is also available based on 3–4 repeated assays. The assay was performed in the presence of high (400 ng/ml) and low (2 ng/ml) pyridoxine concentrations. For a detailed description of this assay, see Mayfield et al. (2012) and Dimster-Denk et al. (2013). Participants were

TABLE 1 CAGI1 prediction data set: 51 single amino acid substitutions within the human CBS coding region. CAGI2 prediction data set: 78 single amino acid variants that had been observed in patients with homocystinuria. Current (2019) ClinVar pathogenic (P) or likely pathogenic (LP) status of the CAGI2 variants in relation to homocystinuria are shown

CAGI1		CAGI2		
Nucleotide variant	Protein variant	Nucleotide variant	Protein variant	ClinVar (2019)
c.194A>T	p.H65L	c.194A>G	p.H65R	
c.250A>G	p.I84V	c.205G>C	p.A69P	
c.353T>G	p.V118G	c.209C>T	p.P70L	U
c.370C>G	p.L124V	c.233C>G	p.P78R	LP
c.370C>G, c.371T>C	p.L124A	c.253G>C	p.G85R	
c.379A>G	p.I127V	c.260C>A	p.T87N	
c.424A>G	p.I142V	c.262C>T	p.P88S	
c.424A>G, c.425T>C	p.I142A	c.302T>C	p.L101P	P/LP
c.425 T>A	p.I142N	c.304A>C	p.K102Q	
c.460C>G, c.461T>C	p.L154A	c.306G>C	p.K102N	LP
c.461T>C	p.L154P	c.325T>C	p.C109R	P/LP
c.529A>G	p.K177E	c.341C>T	p.A114V	P/LP
c.541C>G, c.542T>C	p.L181A	c.346G>A	p.G116R	LP
c.562A>G	p.I188V	c.361C>T	p.R121C	C
c.566T>C	p.V189A	c.362G>A	p.R121H	LP
c.629T>A	p.L210Q	c.362G>T	p.R121L	
c.640A>G	p.I214V	c.376A>G	p.M126V	
c.659T>G	p.L220R	c.384G>T	p.E128D	
c.684C>A	p.N228K	c.393G>C	p.E131D	U
c.718A>G	p.I240V	c.415G>A	p.G139R	P
c.718A>G, c.719T>C	p.I240A	c.429C>G	p.I143M	
c.721C>G	p.L241V	c.430G>A	p.E144K	P/LP
c.742C>G, c.743T>C	p.L248A	c.434C>T	p.P145L	P
c.755T>C	p.V252A	c.442G>C	p.G148R	LP
c.772G>C	p.G258R	c.451G>A	p.G151R	
c.800A>T	p.K267M	c.457G>C	p.G153R	U
c.799A>G	p.K267E	c.461T>A	p.L154Q	
c.811A>G	p.K271E	c.463G>A	p.A155T	
c.829A>C, c.830T>C	p.I277P	c.473C>T	p.A158V	
c.839T>C	p.V280A	c.494G>A	p.C165Y	P
c.856A>G, c.857T>C	p.I286A	c.502G>A	p.V168M	C
c.877C>G	p.L293V	c.539T>C	p.V180A	U
c.931A>G	p.I311V	c.572C>T	p.T191M	P
c.1012C>G, c.1013T>C	p.L338A	c.593A>T	p.D198V	
c.1023A>T	p.Q341H	c.671G>A	p.R224H	
c.1034T>C	p.L345P	c.676G>A	p.A226T	LP
c.1061T>G	p.V354G	c.683A>G	p.N228S	U
c.1067T>C	p.V356A	c.684C>A	p.N228K	
c.1073T>C	p.V358A	c.700G>A	p.D234N	P
c.1112T>C	p.V371A	c.715G>A	p.E239K	
c.1115T>C	p.V372A	c.770C>T	p.T257M	P/LP
c.1120C>G, c.1121T>C	p.L374A	c.775G>A	p.G259S	U

(Continues)

TABLE 1 (Continued)

CAG1		CAG2		
Nucleotide variant	Protein variant	Nucleotide variant	Protein variant	ClinVar (2019)
c.1147A>T	p.T383S	c.785C>T	p.T262M	P/LP
c.1153T>C, c.1154T>A, c.1155C>A	p.F385Q	c.796A>G	p.R266G	
c.1153T>C	p.F385L	c.797G>A	p.R266K	P/LP
c.1223G>T	p.W408L	c.824G>A	p.C275Y	
c.1268T>C	p.L423P	c.833T>C	p.I278T	P
c.1298A>T	p.H433L	c.862G>A	p.A288T	U
c.1370G>A	p.G457E	c.862G>C	p.A288P	
c.1468A>C	p.I490L	c.904G>A	p.E302K	LP
c.1646A>G	p.D549G	c.919G>A	p.G307S	P
		c.959T>C	p.V320A	LP
		c.992C>A	p.A331E	LP
		c.992C>T	p.A331V	
		c.1007G>A	p.R336H	LP
		c.1039G>A	p.G347S	LP
		c.1046G>A	p.S349N	
		c.1055G>A	p.S352N	
		c.1058C>T	p.T353M	P/LP
		c.1060G>A	p.V354M	
		c.1063G>C	p.A355P	
		c.1081G>A	p.A361T	
		c.1105C>T	p.R369C	U
		c.1106G>A	p.R369H	U
		c.1106G>C	p.R369P	
		c.1109G>A	p.C370Y	LP
		c.1111G>A	p.V371M	LP
		c.1126G>A	p.D376N	
		c.1150A>G	p.K384E	P
		c.1173G>A	p.M391I	
		c.1265C>T	p.P422L	U
		c.1301C>A	p.T434N	U
		c.1304T>C	p.I435T	U
		c.1316G>A	p.R439Q	C
		c.1330G>A	p.D444N	P/LP
		c.1367T>C	p.L456P	
		c.1397C>T	p.S466L	U
		c.1572C>A	p.Q526K	

Note: Positions are based on Refseq NM_000071.2. C denotes conflicting interpretations of pathogenicity, U uncertain significance.

asked to submit predictions on the effect of the variants on the function of CBS both in high and low-pyridoxine concentrations. The submitted prediction was requested as the percent of growth when compared with wild-type yeast, with a standard deviation. The predictions were then assessed against the percent of growth values actually measured for each substitution in the yeast assay.

2.2 | Prediction assessment

The correlation between the predicted effects of the mutations and the actual effects serves as an initially simple but powerful measure to assess the accuracy of the prediction methods. Because the mutation data are not derived from a normal distribution, nonparametric tests were used to assess the methods. Both Spearman's rank

correlation and Kendall's Tau correlation (KCC) were used to assess each algorithm's predictions against the observed growth rates. The root-mean-square deviation (RMSD) was also calculated to estimate the difference between experimental and predicted values. To assess the accuracy of the algorithms in a clinical sense, evaluation was also conducted against a binary version of the experimental growth rates. A threshold of 0 was chosen for the binary version and the performance was evaluated in terms of area under the ROC curve (AUC), sensitivity, specificity, accuracy, and positive/negative predictive value (Lever, Krzywinski, & Altman, 2016). For experimental data, the total number of negatives (no growth substitutions) were defined as $N = TN + FP$ and the total number of experimental positives (growth detected) as $P = TP + FN$. All the performance indices are shown in Table S1. An overall ranking of the methods was defined as a mean ranking of three different measures (KCC, AUC, and RMSD) shown in Table S2. These three measures were chosen since they assess complementary aspects of prediction performance.

2.3 | Bootstrapping

To assess the robustness of our comparison of different prediction models, and to identify any possible performance bias generated by outliers, we performed bootstrap simulation on measurements. 10,000 random samples were generated using sampling with replacement of the experimental data. The resulting evaluation metrics (KUC, AUC, and RMSD) were plotted as bar plots with error bars representing one standard deviation below and above the mean value for each metric generated through iterations of bootstrap sampling of data-points.

To estimate the statistical significance of pairwise prediction comparisons, we applied bootstrap simulation (as described above) to the metrics of interest (KCC, AUC, and RMSD). We then performed an all-vs-all comparison recording the p -values, and corrected for multiple hypothesis errors using the Bonferroni method.

2.4 | Characterization of easy and hard to predict variants

We examined the mutation effects that were easiest and hardest to predict to determine whether they shared any common features. We first identified these mutations by summing the binary predictions (0 for no growth, > 0 for growth) at low-pyridoxine conditions across all methods for each variant. The variants with the lowest and highest summed scores were individually examined in terms of sequence, solvent-accessibility, and location within the CBS structure.

To determine the likelihood of a variant being benign or deleterious through sequence analysis, scores for amino acid substitutions were taken from the BLOSUM90 substitution matrix (Henikoff & Henikoff, 1992). Scores of -1, 0, or 1 were classified as moderate substitutions, that is, substitutions with a likelihood of arising by chance in terms of evolution and therefore of unknown effect on CBS function. Scores > 1 were classified as conservative substitutions with a projected benign effect on CBS, while scores < -1 were classified as nonconservative,

indicating a potentially deleterious effect on CBS function. Solvent accessible surface area (SASA) was calculated for the human CBS monomer (PDB id 4COO) using GetArea (Fraczkiewicz & Braun, 1998) and when different, dimer SASA results were noted. Secondary structure assignments and analysis were according to PDB id 4COO and visual inspection of the structure.

2.5 | Method uniqueness in prediction results

For CAGI2, evaluation of the specific contribution of each prediction to the variance with experimental results was addressed using a multiple linear regression model. First, a multiple linear regression model was built with the best methods from each group. The top method from each group was chosen based on the highest adjusted R^2 values of every single method, to exclude predictions using modified versions of the same methods. The final methods included in the model were SID#16, 23, 26, 27, 29, 34, 36, and 41. Subsequently, methods were removed one at a time, and the linear regression equation was recalculated. The contribution of each method to the model was estimated from the delta adjusted R^2 values. SID#25 was excluded from the model as it lacked predictions for 10 substitutions.

3 | RESULTS

3.1 | CAGI1 challenge

In the CAGI1 CBS challenge, participants were asked to submit predictions to assess the impact of 51 single amino acid substitutions upon the function of the human CBS enzyme in both high (400 ng/ml) and low (2 ng/ml) pyridoxine concentrations. The function of the variants had been experimentally tested in an in vivo yeast complementation assay (Dimster-Denk et al., 2013). Twenty predictions from 13 groups were submitted to this challenge (Table 2), which were assessed blindly. A summary of each method is described in Supporting Information. Of the 13 participating groups, nine submitted one prediction, two contributed two, one submitted three and one provided four different submissions. Some methods used sequence-only or structure-only information, some employed meta-predictors, and others combined sequence, structural and annotation data. SID#17 (submission identification number) submitted only raw data without predictions and was excluded from the assessment. Most participants (18/19) provided predictions for both high and low-pyridoxine concentration; however, seven predictions did not distinguish between the different cofactor levels. The majority of the predictions did not include standard deviations (13/19), and most of the methods that included estimates of reliability for each prediction appeared to be arbitrary (constant values like $\sigma = 5, 10, \text{ and } 15; n = 5/7$).

3.1.1 | Assessment across different performance metrics

No single evaluation measure can capture a method's performance; thus, various measures were used to assess the phenotype prediction

TABLE 2 Overview of the phenotype prediction programs used to generate predictions for the CAG1 and CAG12 CBS challenges

Submission ID	Group ID	Program name	Program features	Reference
CAG1				
SID#1	Lichtarge lab	Evolutionary action working version	Sequence and structure	
SID#2	Bromberg lab	SNAP	Sequence, structure, and annotation	Bromberg and Rost (2007)
SID#3	Wei lab	SAPRED	Structure	Ye et al. (2007)
SID#4	Switch lab	FoldX	Structure	Schymkowitz et al. (2005)
SID#5	Vihinen lab	PON-P	Meta-predictor	Olatubosun, Valiaho, Harkonen, Thusberg, and Vihinen (2012)
SID#6	Vihinen lab	PolyPhen2	Sequence, structure	Adzhubei et al. (2010)
SID#7	Vihinen lab	SNPanalyzer	Sequence	Yoo, Lee, Kim, Rha, and Kim (2008)
SID#8	Vihinen lab	Panther	Sequence	Thomas et al. (2006)
SID#9	Casadio lab	IMutant3	Structure and thermal stability	Capriotti, Fariselli, Rossi, and Casadio (2008)
SID#10	Casadio lab	IMutant4	Structure and thermal stability	
SID#11	Casadio lab	IMutant baseline	Structure and thermal stability	
SID#12	Forman lab	SDM	Sequence and structure	
SID#13	BioFold Unit	IMutant3	Sequence and structure	Capriotti et al. (2008)
SID#14	Karchin lab		Sequence and structure	
SID#16	Mooney lab		Meta-predictor	
SID#18	Forman lab	SDM	Sequence and structure	
SID#19	Tavtigian lab	AlignGVGD	Sequence	Tavtigian, Byrnes, Goldgar, and Thomas (2008)
SID#20	Tavtigian lab	AlignGVGD	Meta-predictor	
CAG12				
SID#16	Bromberg lab	SNAP	Sequence and structure	Bromberg and Rost (2007)
SID#19	Tosatto lab		Structure	
SID#20	Tosatto lab		Structure	
SID#21	Tosatto lab		Structure	
SID#22	Tosatto lab		Structure	
SID#23	Tosatto lab		Structure	
SID#24	Tosatto lab	D100 roll	Random	
SID#25	Switch lab		Structure	
SID#26	Lichtarge Lab	Evolutionary Action	Sequence and structure	Katsonis and Lichtarge (2014)
SID#27	Vihinen lab	PON-P	Meta-predictor	Olatubosun et al. (2012)
SID#28	Vihinen lab	PON-P	Meta-predictor	Olatubosun et al. (2012)
SID#29	Shatsky lab		Meta-predictor	
SID#30	Shatsky lab		Meta-predictor	
SID#32	Shatsky lab		Meta-predictor	
SID#33	Mooney lab		Meta-predictor	
SID#34	Sunyayev lab		Sequence and structure	
SID#35	Moult lab	SNPs3D SVM	Sequence and structure	Yue and Moult (2006)
SID#36	Moult lab	SNPs3D SVM	Sequence, structure, and annotation	
SID#41	Mooney lab		Meta-predictor	

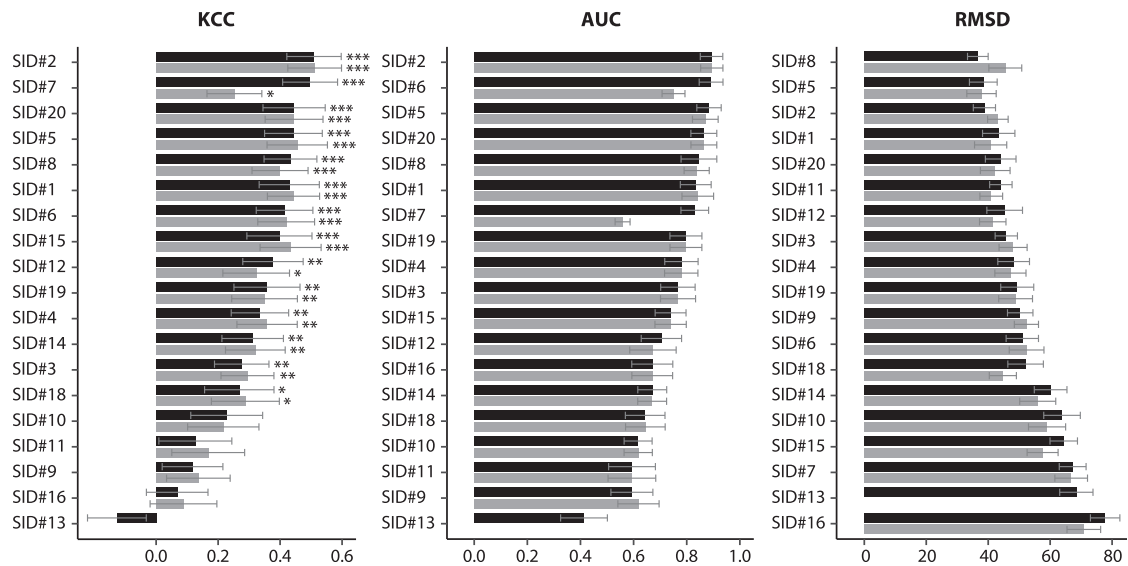


FIGURE 1 Kendall's Tau correlation coefficient (KCC), area under the ROC curve (AUC), and root-mean-square deviation (RMSD) for the phenotype prediction programs at high (black) and low (gray) cofactor concentration in the CAG1 CBS challenge. Statistical significance of correlation scores is indicated with asterisks (* $p \leq .05$, ** $p \leq .01$, *** $p \leq .001$). Error bars represent one standard deviation below and above the mean value for each metric generated through iterations of bootstrap sampling of data-points

programs, including Kendall's Tau correlation coefficient (KCC), precision/recall, accuracy, and root-mean-square deviation (RMSD), inter alia (Table S1). For KCC, most of the prediction methods display statistically significant correlation with experimental data at both pyridoxine concentrations (Figure 1). Methods SID#2, SID#5, and SID#20 showed strong correlation at both high and low cofactor concentrations. SID#7 was the second best at high pyridoxine concentration ($\tau = 0.50$, $p = 4.87 \times 10^{-6}$); however, it showed little correlation at low cofactor concentration ($\tau = 0.25$, $p = .034$). For RMSD, SID#5, which is a meta-predictor, was the best and second best at low and high cofactor concentration, respectively. The highest accuracy of 82.4% was achieved by three methods: SID#1, 6, and 20 (Table S1). The first two combined sequence and structure information, while SID#20 is a meta-predictor, integrating several prediction methods. SID#2, 3, 9, and 11 achieved 100% sensitivity, whereas other methods had the highest specificity (94–100%, SID#7 and SID#14) at both cofactor concentrations. The most sensitive models were mostly structure-based. The majority of the methods had good results for PPV, where the values varied from 65% to 100%; however, for NPV, the values displayed a much wider range (0–90%), meaning that the methods are better at predicting benign than deleterious variants.

3.1.2 | Overall ranking

To carry out an overall performance assessment, ranks of the prediction methods based on KCC, AUC, and RMSD were averaged to obtain the overall ranks of the methods (Table S2). This revealed SID#2 as the best performing method, having a mean rank of 2.2 across all measures, with SID#5 close behind (mean rank of 2.3). The first method is based on sequence information integrated with

functional and structural annotations, while the other is a meta-predictor. One of the methods that did not perform as well, SID#16, was biased toward the prediction of low growth variants, whereas SID#13 was more conservative, with moderate to high growth predicted for most of the substitutions.

To estimate the robustness of the rankings, bootstrapping was performed (see Methods). Error bars for each metric were obtained by random resampling of the 51 variants 10,000 times. The rankings of the KCC, AUC, and RMSD for both high and low concentrations have only minor fluctuations, indicating that the prediction models are relatively robust (Figure 1). More than half of the predictions did better than the baseline method, SIFT (SID#15), which ranked 11th overall. Statistical significance estimation was performed for the metrics of interest (KCC, AUC, and RMSD) also using bootstrap simulation. None of the methods were significantly better than SIFT for KCC and AUC after Bonferroni correction (Figure S1 and S2). However for RMSD, SID#5, 11, and 20 outperformed SIFT at both cofactor concentrations even after Bonferroni correction (Figure S3).

3.1.3 | Easy and hard to predict variants

We examined variants that were the easiest or hardest to predict based on the consensus output of all methods to determine whether they shared any common features (Figure 2b, Figure S7). At low cofactor concentration, there were overall 12% (2/17) deleterious variants and 18% (6/34) benign variants whose effects were predicted incorrectly by more than half of the methods, our definition of consensus.

The deleterious mutations that were easiest to predict at a low cofactor concentration were p.L154P, p.N228K, p.G258R, and p.G457E. The majority of these are nonconservative substitutions and are located

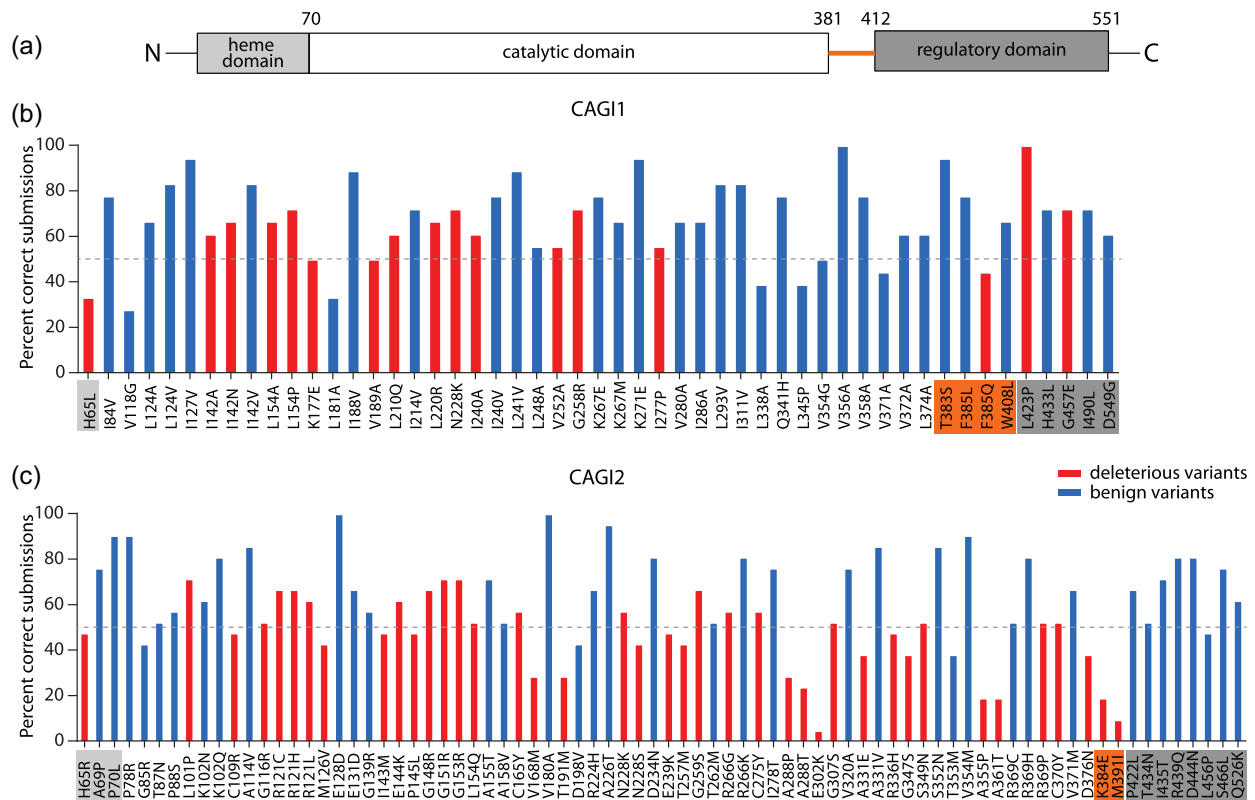


FIGURE 2 Consensus predictions for CBS. (a) CBS domain diagram, (b) CAGI1, (c) CAGI2. The percentage of correct predictions for deleterious (red) and benign (blue) variants is shown for each experimentally determined variant at low-pyridoxine concentration. Residues are shaded in the color of the corresponding domain, with the linker region highlighted in orange

within helices in the CBS structure (Table S3). The easiest to predict benign variants (low pyridoxine) were p.K271E, p.V356A, and p.T383S, with moderate conservation scores and were again mostly located within helices. Among the better predicted benign variants, the majority were partially or fully exposed to the solvent.

The hardest to predict deleterious variants at both high and low-pyridoxine concentrations were p.H65L and p.F385Q. p.H65 is located in the H1–H2 loop and axially coordinates the iron atom on one side of the heme plane, with C52 on the other, and mutation of either of these residues results in low-catalytic activity (Ojha, Wu, LoBrutto, & Banerjee, 2002). Interestingly, the functional impact of these variants was not easy to assess from sequence and structure comparisons. Although p.H65 and the sequence flanking it are locally conserved, the heme-binding domain itself (comprising approximately the first 70 N-terminal residues), with the exception of a short 5-residue helix, has no secondary structure elements and does not resemble other heme proteins in either primary sequence or tertiary structure (Kumar et al., 2018). p.F385Q is located in the H17–H18 loop that forms part of the linker connecting the N-terminal catalytic domain with the C-terminal regulatory domain, and lies within an aromatic cluster of residues p.Y381, p.F332, p.F334, p.F385, p.W390, and p.F396 enclosed by salt bridges p.R336–D388, p.K394–E302, and p.K384–E302 connecting helices H12–H14, H17, and H18. Erroneous coordination between aromatic residues can

disrupt the extended pi-pi networks formed by aromatic clusters, thereby affecting protein stability and folding (Madhusudan Makwana & Mahalakshmi, 2015). In addition, both these variants involve nonconservative substitutions, and thus could be expected to have a deleterious effect on CBS function.

The hardest to predict benign variants (low pyridoxine) were surprising in that they involved nonconserved substitutions, so they could be expected to disrupt CBS function. In addition, within the structure, some were implicated in functionally relevant regions of CBS, such as the dimer interface (p.L345P) and the active site (p.V118G, adjacent to the PLP-ligating p.K119). All inaccurate consensus predictions of benign variants at low pyridoxine were for variants that confer sensitivity to reduced pyridoxine levels relative to the major allele (Dimster-Denk et al., 2013). The methods that correctly predicted all of these variants (SID#2, SID#3, and SID#9) displayed a broad spread both in features used (sequence, structure, and thermodynamics, respectively) and in overall performance (Table 2). In addition, SID#2 and SID#9 did not distinguish between high and low pyridoxine.

3.2 | CAGI2 challenge

In the CAGI2 CBS challenge, 84 single amino acid variants that had been observed in patients with homocystinuria were collected and

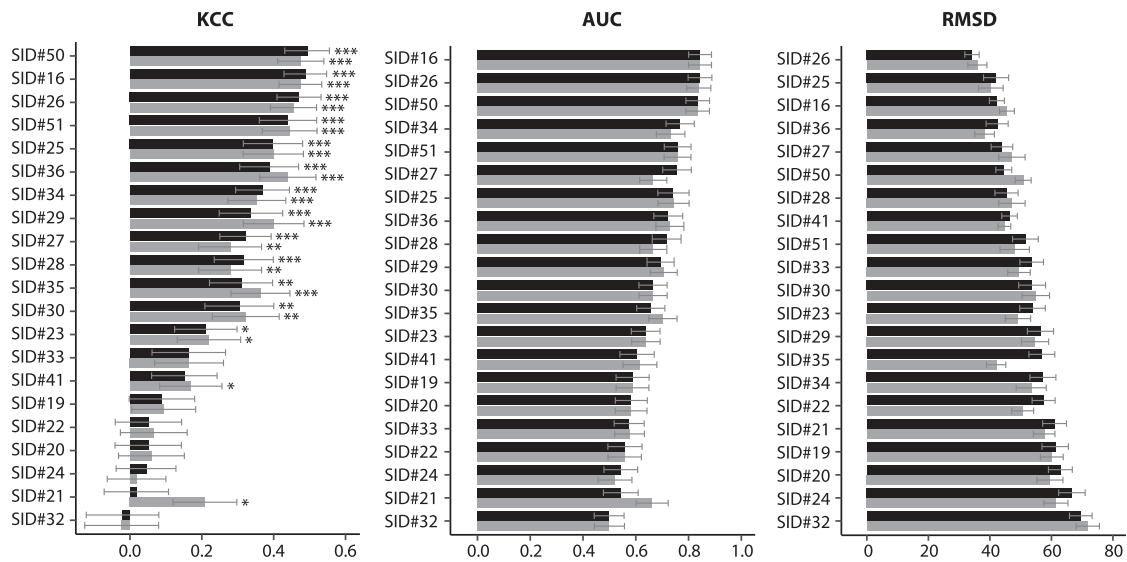


FIGURE 3 Kendall's Tau correlation coefficient (KCC), area under the ROC curve (AUC) and root-mean-square deviation (RMSD) for the phenotype predictions at high (black) and low (gray) cofactor concentration in the CAGI2 CBS challenge. Statistical significance of correlation scores is indicated with asterisks ($*p \leq .05$, $**p \leq .01$, $***p \leq .001$). Error bars represent one standard deviation below and above the mean value for each metric generated through iterations of bootstrap sampling of data-points

functionally tested in an *in vivo* yeast complementation assay (Mayfield et al., 2012). Seventy-eight had experimental values for both pyridoxine concentrations; 6 “hem1 rescue” variants were left out from the assessment due to absent/conflicting data (Table 1). Participants were again asked to submit predictions of the effect of the variants on the function of CBS both in high and low cofactor concentration. This challenge attracted 20 submissions from nine groups (Table 2) that were assessed without knowledge of the identity of the predictors. An overview of the methods is provided in Supporting Information. Four groups submitted one submission each, three groups submitted two each, and one group each contributed four and six predictions, respectively. Four groups participated in both CAGI1 and CAGI2 CBS challenges. As in CAGI1, features used to generate the predictions ranged from sequence- or structure-only information to meta-predictors and methods combining sequence, structural, and functional annotation data. SID#31 was excluded from the assessment due to its constant growth rate prediction of 100 for all substitutions. Almost all groups (17/19) provided distinct values for high/low cofactor concentrations. For this challenge, most submitters also provided standard deviations (13/19). Only one of the methods had arbitrary standard deviation values for all predictions (SID#26, $\sigma = 10$). In addition to prediction programs, reference results were obtained by submitting the mutations to the SNAP (SID#50) and SIFT (SID#51) public servers (Bromberg, Yachdav, & Rost, 2008; P. Kumar, Henikoff, & Ng, 2009).

3.2.1 | Assessment across different performance metrics

The same assessment measures as in CAGI1 were used in CAGI2 (Methods). Looking at KCC, over half of the predictions were highly

significant (Figure 3); however, even the best deviated substantially from experimental values. At both high and low-pyridoxine concentrations, methods SID#16 and SID#26 showed the strongest correlation with the experimental data and had also high AUC values (Figure 3). The latter was also the top predictor in terms of RMSD. In terms of accuracy, a structure-based method (SID#25) had the highest value (72%) at high and low-pyridoxine concentration (Table S2). At both cofactor concentrations, methods SID#16, 26, 34, and 41 had the highest sensitivity (100%). Most of these methods employed integrated sequence and structure information. SID#23 was the top method (high pyridoxine) with a 75% specificity, SID#27 and SID#28 scored 83% (low pyridoxine). SID#23 used structural data, whereas the other two are meta-predictors. In contrast to the CAGI1 CBS challenge, NPV showed higher median values than PPV (65 vs. 57%), implying that the probability of loss of function was slightly better predicted than the probability of having no or minor effects on the phenotype.

3.2.2 | Overall ranking

As in the CAGI1 CBS challenge, we computed the overall ranks representing the performance of the submissions. Based on this criterion, the top methods in the CAGI2 CBS challenge were SID#26 and SID#16 with overall ranks of 1.8 and 2.3, respectively. Both methods utilized combined evolutionary information and structural features and ranked higher than the best baseline method (SID#50). For CAGI2, random resampling of the 78 variants 10,000 times generated error bars for each metric. As in CAGI1, the overall performance ranking remained unchanged. Methods that performed well generally resulted in smaller error bars, whereas methods that had only a few correct predictions exhibited a larger variance in performance as assessed by resampling (Figure 3).

Almost all of the methods performed better than the random predictor (SID#24), with the exception of SID#32 that ranked lower according to all assessment measures at both cofactor concentrations (Table S2). SID#32 is a random forest classifier that provided only binary outputs for both concentrations (no growth or growth, with values of 0 or 100, respectively). According to the bootstrap simulation, SID#16 and 26 performed significantly better than random at high pyridoxine concentration for all metrics (Figure S4–6). Two of the baseline methods, SNAP (SID#50) and SIFT (SID#51), had an overall ranking of third and sixth, respectively. None of the methods outperformed the baseline methods for KCC and AUC after Bonferroni correction (Figure S4, 5). For RMSD on the other hand, SID#26 did significantly better than SIFT at high cofactor concentration and SID#16 outperformed SNAP at low cofactor concentration.

3.2.3 | Easy and hard to predict variants

The hardest and easiest to predict variants (Figure 2c, Figure S8) showed similar trends to those observed in the CAG1 CBS challenge. As observed before, the inaccurately predicted benign variants (low pyridoxine) involved nonconservative substitutions and were located in loop regions of the CBS structure (Table S3). The most accurately predicted deleterious variants (low pyridoxine) involved a majority of nonconservative mutations of residues located within stable secondary structure elements (helices), while the hardest to predict deleterious predictions involved residues with moderate conservation scores mostly located within loops.

Within this last group, a number of mutants have been indirectly implicated in CBS function through involvement in homooligomerization, redox sensing, and regulation. p.A355 lies within helix H15, which is in turn sandwiched between H4 and strand beta3 at the CBS homodimerization interface. By introducing a kink in H15, the p.A355P mutant could potentially disrupt the folding in this region of the protein thereby impacting CBS function. Similarly, p.V168 is positioned at the homodimer interface while p.M391 is located within helix H18, a region of putative involvement in CBS homotetramer formation (Ereno-Orbea, Majtan, Oyenarte, Kraus, & Martinez-Cruz, 2013). p.A288 packs against p.W323 on strand beta6, and next to it, p.F324 packs against p.A360 in helix H15. p.A361 lies within interacting distance of p.C370, a residue that has been implicated in homocystinuria (Kraus et al., 1999) and proposed to modulate CBS function through interaction with an endogenous regulator such as nitric oxide (Eto & Kimura, 2002). The p.A361T mutant could therefore potentially interfere with a functionally relevant modification (e.g., S-nitrosylation) of p.C370. Similarly, modification of p.A288 could disrupt the pairing or orientation between beta5 and beta6, thereby potentially impacting the 272-CxxC-275 oxygen sensing motif of CBS, a redox active disulfide bond that allosterically controls CBS activity (Niu et al., 2018).

The most inaccurately predicted deleterious mutant, p.E302K, lies within interacting distance of one of the two active site loops (situated between helices H6 and H7). Recent studies have

highlighted the importance of conformational flexibility of the loops defining the entrance to the catalytic site (Majtan et al., 2018).

3.2.4 | Correlation between methods and unique contribution of different methods

To have a better understanding of the strengths and weaknesses of the different methods, we investigated the correlation between their predictions (Figure 4). Correlation heatmaps for high and low-pyridoxine concentration had negligible differences. The strongest predictor for method correlation appeared to be the relation to a single group (Table 2). For example, SID#27–28, SID#33&41, and all (SID#19–22) except one method (SID#23) were highly correlated among each other. However, SID#29–32 and 35–36 had higher correlations with other methods than among their own group. SID#29–32 were the only predictors that used simply two states (growth rates of 0 or 100). Interestingly, two of the best ranking methods (SID#16 and SID#26) were strongly correlated. In addition, SID# 27, 28, and 34 showed a strong correlation with the top prediction methods, although were based on different features.

Baseline methods SNAP (SID#50) and SIFT (SID#51) showed strong correlation with the best performing predictions, which is partly expected as SID#16 was based on a version of the SNAP algorithm.

To assess the specific contribution of each method to the variance with experimental results in CAG12, we applied a multiple linear regression model as described in Methods above. For high pyridoxine concentration, this revealed SID#16 and SID#36 as the most significant contributors (Δ adjusted R^2 values of 0.053 and 0.041, respectively). At the same time, for low-pyridoxine concentration, SID#36 and SID#27 contributed the most (0.054 and 0.053, respectively). SID#36 is based on protein structure, sequence homology, and included functional information, whereas SID#16 combined evolutionary information with structural features, and SID#27 is a meta-predictor (Figure 5).

4 | DISCUSSION

4.1 | Prediction features in relation to performance

In terms of prediction features, different methods performed well in distinct assessment measures. We observed that methods integrating sequence and structural information performed the best overall, ranking first or second (SID#2 in CAG1 and SID#16 and SID#26 in CAG12). Methods that used only structural information (SID#9 and 11 in CAG1 and SID#19–20 in CAG12) did not perform as well as those combining additional features. However, in terms of individual evaluation metrics, a structure-based method showed the highest accuracy of 72% (SID#25) in CAG12. In CAG1, SID#1 and 20 were the best at both cofactor concentrations, reaching accuracy of 82%. The first one combined structure and sequence data while SID#20 is a meta-predictor. In terms of sensitivity, most of the top-performing methods were structure-based in CAG1 (SID#3, SID#9, and SID#11), while in CAG12, the most sensitive algorithms combined structure with sequence data (SID#16, SID#26, and SID#34). At the same time,

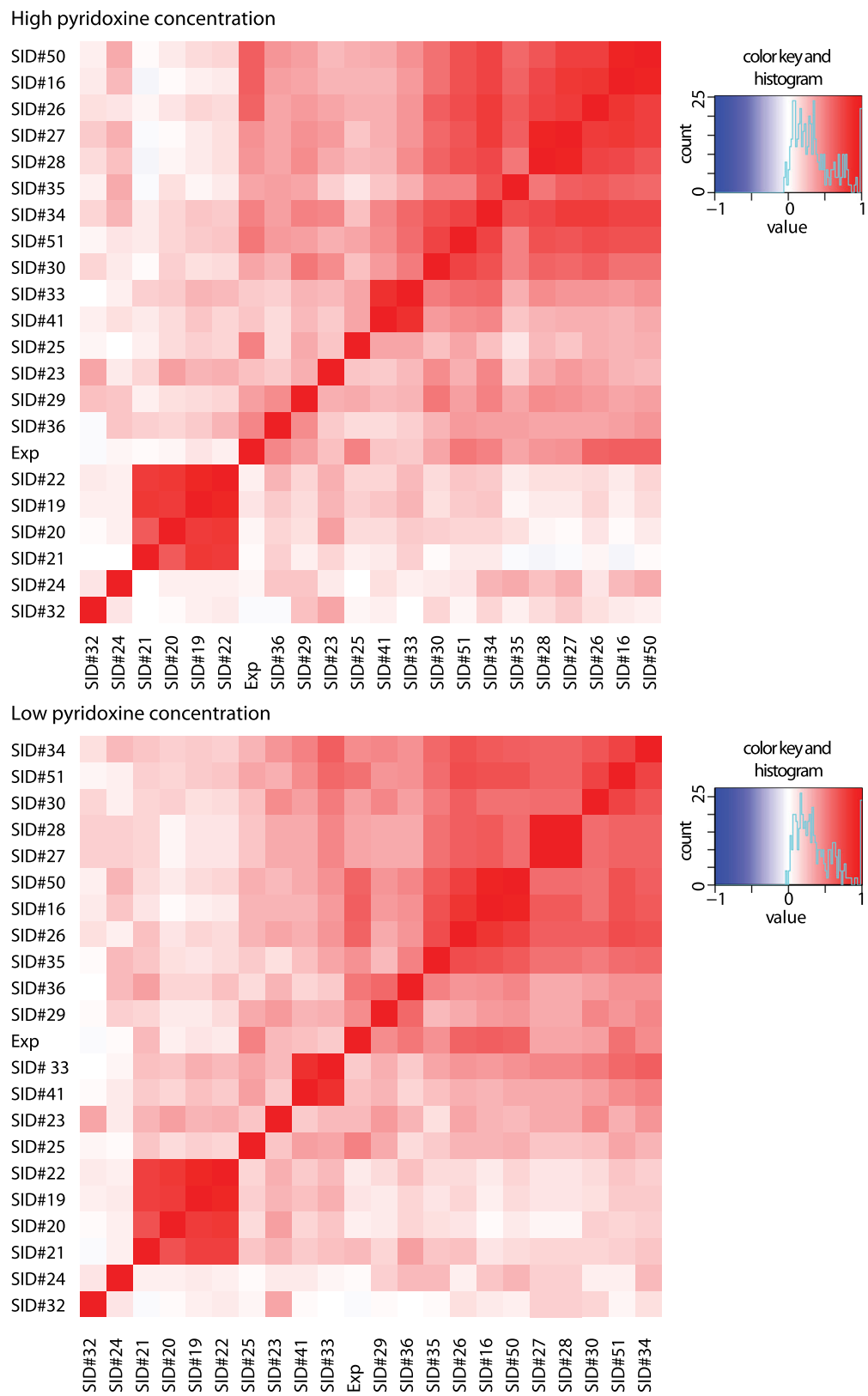


FIGURE 4 Spearman's rank correlation among methods and with experimental data (Exp) for high and low cofactor concentration. Each cell shows the correlation between two methods, with a color scale ranging from red (perfect correlation) to white (no correlation) and blue (perfect anti-correlation)

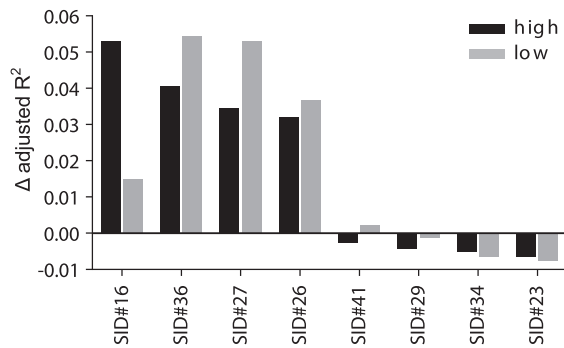


FIGURE 5 Δ adjusted R^2 values of the methods from the linear regression model for high and low cofactor concentration, quantifying the contribution of each method to the proportion of total variance explained

for specificity, almost all of the top methods were meta-predictors in CAGI2 (SID#27–28). These observations suggest that combining different features and methods would yield the best results, as has been indicated previously (Grimm et al., 2015; Tang & Thomas, 2016). Some methods are tailored to predict whether a variant affects the function of the protein in hand and others are optimized to determine whether a variant is pathogenic or benign in the clinical sense (Grimm et al., 2015; Katsonis et al., 2014; Pejaver, Mooney, & Radivojac, 2017).

The importance of integrating information from different sources is reflected in the most inaccurately predicted mutants that tended towards nonconserved substitutions, structural uncertainty, or both. The power of combining structural, sequence, and functional information was visible in CAGI2, where the overall performance of a structure and sequence combined method (SID#35) was improved significantly (by five ranks) with the inclusion of functional annotation data (SID#36). The latter was also the method that uniquely contributed the most predictive power of all methods at low-pyridoxine concentration. Another structure-based method (SID#25) that incorporated functional information (trained on the CAGI1 data set) also performed strongly. Methods trained on HGMD (Human Gene Mutation Database) mutations (SID#35–36) would be expected to perform well (Dong et al., 2015; Ioannidis et al., 2016; Pejaver et al., 2017). Interestingly, however, the best methods in CAGI2 CBS challenge showed variable training data, from no training to training on PMD (the Protein Mutant Database), HGMD, and CAGI1 CBS variant data (Supporting Information).

4.2 | Limitations of the challenges

In terms of methodological limitations, most methods were developed to predict pathogenicity in humans or enzyme activity, not yeast growth or the effect of cofactor concentration on growth rate, something that could at least in part explain the difficulties they encountered in identifying the remediable class of variants (Supporting Information). So, while a meaningful distinction could be made in these challenges between growth and no growth under low-pyridoxine conditions, this was not the case for distinguishing

between rescue (high pyridoxine) and no rescue (low pyridoxine) variants (Figure S9-12). Similarly, only qualitative comparisons could be made between CAGI1 and CAGI2, since the data sets differed in size and type, and only four groups participated in both challenges. Among these, one group used different versions of their method (SID#1 in CAGI1 and SID#26 in CAGI2), while two others did not make use of the CBS training data.

Another limitation of the assessment involved requesting standard deviation as estimates of reliability from predictors, as opposed to the more commonly employed confidence levels that most prediction methods provide. Consequently, some predictors did not provide these values, chose them arbitrarily, or provided large values with the result that they could not be reasonably used in these assessments.

These challenges also revealed a number of experimental limitations. Yeast CBS lacks the heme domain and is not regulated by AdoMet (Jhee, McPhie, & Miles, 2000), thereby engaging different pathways in the enzyme's regulation and physiological roles. In addition, overexpression can result in nonphysiological effects, including protein aggregation. These differences could help explain some of the inconsistencies observed in the experimental study in which yeast growth phenotypes did not match the clinical data (Mayfield et al., 2012). Although, only three positions from the heme domain were part of these two CBS challenges, with merely one position being problematic for the predictors (Figure 2). In a similar study, several variants identical to the ones used in these experiments resulted in contrasting yeast growth phenotypes (Wei, Wang, Wang, Kruger, & Dunbrack, 2010).

In addition, the clinical assessment of the majority of variants explored in this study has since changed. Of the 78 alleles described in CAGI2 as having been observed in patients with homocystinuria, only 30 are currently classified as pathogenic or likely pathogenic in ClinVar (<https://www.ncbi.nlm.nih.gov/clinvar/>, accessed March 25, 2019), with an additional 16 annotated as being of uncertain significance or with conflicting interpretations of pathogenicity (Table 1). Eight substitutions (p.P78R, p.K102N, p.D234N, p.R266K, p.V320A, p.T353M, p.V371M, and p.D444N) out of 22 that showed experimental growth rates of $\geq 85\%$ in CAGI2 are currently annotated as pathogenic or likely pathogenic. Most of the participants made accurate predictions for these “benign” variants (Figure 2). It is important to mention here that ClinVar had not yet been launched during the first two CAGI challenges. Also, not all the (likely) pathogenic CBS variants currently present in ClinVar have been collected from clinical testing, some are based on literature with no assertion criteria provided. Ideally, different functional assays should be applied, to increase the confidence in the observed phenotypic effect of the studied variant, because the function of a gene can differ in distinct organisms. Finally, mutations in *cis* with the ability to either suppress other pathogenic missense mutations or increase the severity of the clinical phenotype continue to be reported (de Franchis, Kraus, Kozich, Sebastio, & Kraus, 1999; Shan, Dunbrack, Christopher, & Kruger, 2001), raising the possibility that the incidence of double mutant alleles may be underestimated in homocystinuric patients.

5 | CONCLUSION

CBS is a multifunctional enzyme with complex biology and intricate regulation that remains the object of much study. Our assessment of the CAGI1 and CAGI2 CBS challenges highlighted the strengths and weaknesses of different prediction features and approaches, as well as the need to address issues of methodological and experimental limitations. Both computational and experimental methods need to be tailored to the particular biological question under investigation to improve the predictive potential of the variant effect. It is hoped that future iterations of CAGI will see improvements on all these fronts.

ACKNOWLEDGMENTS

The authors are grateful to the CAGI organizers, data providers, participants, and would like to thank Pauline C. Ng for her contribution in the initial analysis of CAGI1. The CAGI experiment coordination is supported by NIH U41 HG007346 and the CAGI conference by NIH R13 HG006650. We are grateful to Robin Peters-Petrulewicz for a careful reading of the manuscript.


CONFLICT OF INTERESTS







The authors declare no conflicts of interest.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available to registered users from the CAGI web site <https://genomeinterpretation.org/content/cagi-2011-results>.

ORCID

Laura Kasak  <http://orcid.org/0000-0003-4182-2396>
 Constantina Bakolitsa  <http://orcid.org/0000-0002-6980-9831>
 Zhiqiang Hu  <http://orcid.org/0000-0001-8854-3410>
 Jasper Rine  <http://orcid.org/0000-0003-2297-9814>
 Gaurav Pandey  <http://orcid.org/0000-0003-1939-679X>
 Greet De Baets  <http://orcid.org/0000-0001-8844-800X>
 Yana Bromberg  <http://orcid.org/0000-0002-8351-0844>
 Emidio Capriotti  <http://orcid.org/0000-0002-2323-0963>
 Rita Casadio  <http://orcid.org/0000-0002-7462-7039>
 Manuel Giollo  <http://orcid.org/0000-0001-9434-2383>
 Rachel Karchin  <https://orcid.org/0000-0002-5069-1239>
 Panagiotis Katsonis  <http://orcid.org/0000-0002-7172-1644>
 Emanuela Leonardi  <http://orcid.org/0000-0001-8486-8461>
 Olivier Lichtarge  <http://orcid.org/0000-0003-4057-7122>
 Pier Luigi Martelli  <http://orcid.org/0000-0002-0274-5669>
 Sean D. Mooney  <http://orcid.org/0000-0003-2654-0833>
 Predrag Radivojac  <http://orcid.org/0000-0002-6769-0793>
 Frederic Rousseau  <http://orcid.org/0000-0002-9189-7399>
 Lipika R. Pal  <http://orcid.org/0000-0002-3390-110X>

Castrense Savojardo  <http://orcid.org/0000-0002-7359-0633>
 Joost Schymkowitz  <http://orcid.org/0000-0003-2020-0168>
 Janita Thusberg  <http://orcid.org/0000-0001-8028-4426>
 Silvio C.E. Tosatto  <http://orcid.org/0000-0003-4525-7793>
 Mauno Vihinen  <http://orcid.org/0000-0002-9614-7976>
 Susanna Repo  <http://orcid.org/0000-0003-3488-4767>
 John Moulton  <http://orcid.org/0000-0002-3012-2282>
 Steven E. Brenner  <http://orcid.org/0000-0001-7559-6185>
 Iddo Friedberg  <http://orcid.org/0000-0002-1789-8000>

REFERENCES

- Adzhubei, I. A., Schmidt, S., Peshkin, L., Ramensky, V. E., Gerasimova, A., Bork, P.,... Sunyaev, S. R. (2010). A method and server for predicting damaging missense mutations. *Nature Methods*, 7(4), 248–249. <https://doi.org/10.1038/nmeth0410-248>
- Bromberg, Y., & Rost, B. (2007). SNAP: Predict effect of non-synonymous polymorphisms on function. *Nucleic Acids Research*, 35(11), 3823–3835. <https://doi.org/10.1093/nar/gkm238>
- Bromberg, Y., Yachdav, G., & Rost, B. (2008). SNAP predicts effect of mutations on protein function. *Bioinformatics*, 24(20), 2397–2398. <https://doi.org/10.1093/bioinformatics/btn435>
- CAGI (2010/2011). CBS challenge dataset; Dataset available to registered users from the CAGI web site: <https://genomeinterpretation.org/content/cagi-2011-results>.
- Capriotti, E., Fariselli, P., Rossi, I., & Casadio, R. (2008). A three-state prediction of single point mutations on protein stability changes. *BMC Bioinformatics*, 9(Suppl 2), S6. <https://doi.org/10.1186/1471-2105-9-S2-S6>
- Dimster-Denk, D., Tripp, K. W., Marini, N. J., Marqusee, S., & Rine, J. (2013). Mono and dual cofactor dependence of human cystathionine beta-synthase enzyme variants in vivo and in vitro. *G3 (Bethesda)*, 3(10), 1619–1628. <https://doi.org/10.1534/g3.113.006916>
- Dong, C., Wei, P., Jian, X., Gibbs, R., Boerwinkle, E., Wang, K., & Liu, X. (2015). Comparison and integration of deleteriousness prediction methods for nonsynonymous SNVs in whole exome sequencing studies. *Human Molecular Genetics*, 24(8), 2125–2137. <https://doi.org/10.1093/hmg/ddu733>
- Ereno-Orbea, J., Majtan, T., Oyentearte, I., Kraus, J. P., & Martinez-Cruz, L. A. (2013). Structural basis of regulation and oligomerization of human cystathionine beta-synthase, the central enzyme of transsulfuration. *Proceedings of the National Academy of Sciences of the United States of America*, 110(40), E3790–E3799. <https://doi.org/10.1073/pnas.1313683110>
- Eto, K., & Kimura, H. (2002). A novel enhancing mechanism for hydrogen sulfide-producing activity of cystathionine beta-synthase. *Journal of Biological Chemistry*, 277(45), 42680–42685. <https://doi.org/10.1074/jbc.M205835200>
- Fernald, G. H., Capriotti, E., Daneshjoui, R., Karczewski, K. J., & Altman, R. B. (2011). Bioinformatics challenges for personalized medicine. *Bioinformatics*, 27(13), 1741–1748. <https://doi.org/10.1093/bioinformatics/btr295>
- Fraczkiewicz, R., & Braun, W. (1998). Exact and efficient analytical calculation of the accessible surface areas and their gradients for macromolecules. *Journal of Computational Chemistry*, 19(3), 319–333. [https://doi.org/10.1002/\(sici\)1096-987x\(199802\)19:3<319::aid-jcc6>3.0.co;2-w](https://doi.org/10.1002/(sici)1096-987x(199802)19:3<319::aid-jcc6>3.0.co;2-w)
- de Franchis, R., Kraus, E., Kozich, V., Sebastio, G., & Kraus, J. P. (1999). Four novel mutations in the cystathionine beta-synthase gene: Effect of a second linked mutation on the severity of the homocystinuric phenotype. *Human Mutation*, 13(6), 453–457. [https://doi.org/10.1002/\(SICI\)1098-1004\(1999\)13:6<453::AID-HUMU4>3.0.CO;2-K](https://doi.org/10.1002/(SICI)1098-1004(1999)13:6<453::AID-HUMU4>3.0.CO;2-K)
- Grimm, D. G., Azencott, C. A., Aicheler, F., Gieraths, U., MacArthur, D. G., Samocha, K. E.,... Borgwardt, K. M. (2015). The evaluation of tools

- used to predict the impact of missense variants is hindered by two types of circularity. *Human Mutation*, 36(5), 513–523. <https://doi.org/10.1002/humu.22768>
- Henikoff, S., & Henikoff, J. G. (1992). Amino acid substitution matrices from protein blocks. *Proceedings of the National Academy of Sciences of the United States of America*, 89(22), 10915–10919. <https://doi.org/10.1073/pnas.89.22.10915>
- Hoskins, R. A., Repo, S., Barsky, D., Andreoletti, G., Moul, J., & Brenner, S. E. (2017). Reports from CAGI: The critical assessment of genome interpretation. *Human Mutation*, 38(9), 1039–1041. <https://doi.org/10.1002/humu.23290>
- Ioannidis, N. M., Rothstein, J. H., Pejaver, V., Middha, S., McDonnell, S. K., Baheti, S., ... Sieh, W. (2016). REVEL: An Ensemble method for predicting the pathogenicity of rare missense variants. *American Journal of Human Genetics*, 99(4), 877–885. <https://doi.org/10.1016/j.ajhg.2016.08.016>
- Jhee, K. H., McPhie, P., & Miles, E. W. (2000). Yeast cystathionine beta-synthase is a pyridoxal phosphate enzyme but, unlike the human enzyme, is not a heme protein. *Journal of Biological Chemistry*, 275(16), 11541–11544. <https://doi.org/10.1074/jbc.C000056200>
- Katsonis, P., Koire, A., Wilson, S. J., Hsu, T. K., Lua, R. C., Wilkins, A. D., & Lichtarge, O. (2014). Single nucleotide variations: Biological impact and theoretical interpretation. *Protein Science*, 23(12), 1650–1666. <https://doi.org/10.1002/pro.2552>
- Katsonis, P., & Lichtarge, O. (2014). A formal perturbation equation between genotype and phenotype determines the evolutionary action of protein-coding variations on fitness. *Genome Research*, 24(12), 2050–2058. <https://doi.org/10.1101/gr.176214.114>
- Kraus, J. P., Janosik, M., Kozich, V., Mandell, R., Shih, V., Sperandio, M. P., ... Gaustadnes, M. (1999). Cystathionine beta-synthase mutations in homocystinuria. *Human Mutation*, 13(5), 362–375. [https://doi.org/10.1002/\(SICI\)1098-1004\(1999\)13:5<362::AID-HUMU4>3.0.CO;2-K](https://doi.org/10.1002/(SICI)1098-1004(1999)13:5<362::AID-HUMU4>3.0.CO;2-K)
- Kumar, A., Wissbrock, A., Goradia, N., Bellstedt, P., Ramachandran, R., Imhof, D., & Ohlenschlager, O. (2018). Heme interaction of the intrinsically disordered N-terminal peptide segment of human cystathionine-beta-synthase. *Scientific Reports*, 8(1), 2474. <https://doi.org/10.1038/s41598-018-20841-z>
- Kumar, P., Henikoff, S., & Ng, P. C. (2009). Predicting the effects of coding non-synonymous variants on protein function using the SIFT algorithm. *Nature Protocols*, 4(7), 1073–1081. <https://doi.org/10.1038/nprot.2009.86>
- Lever, J., Krzywinski, M., & Altman, N. (2016). Classification evaluation. *Nature Methods*, 13, 603–604. <https://doi.org/10.1038/nmeth.3945>
- Madhusudan Makwana, K., & Mahalakshmi, R. (2015). Implications of aromatic-aromatic interactions: From protein structures to peptide models. *Protein Science*, 24(12), 1920–1933. <https://doi.org/10.1002/pro.2814>
- Majtan, T., Pey, A. L., Gimenez-Mascarell, P., Martinez-Cruz, L. A., Szabo, C., Kozich, V., & Kraus, J. P. (2018). Potential pharmacological chaperones for cystathionine beta-synthase-deficient homocystinuria. *Handbook of Experimental Pharmacology*, 245, 345–383. https://doi.org/10.1007/164_2017_72
- Mayfield, J. A., Davies, M. W., Dimster-Denk, D., Pleskac, N., McCarthy, S., Boydston, E. A., ... Rine, J. (2012). Surrogate genetics and metabolic profiling for characterization of human disease alleles. *Genetics*, 190(4), 1309–1323. <https://doi.org/10.1534/genetics.111.137471>
- Moat, S. J., Bao, L., Fowler, B., Bonham, J. R., Walter, J. H., & Kraus, J. P. (2004). The molecular basis of cystathionine beta-synthase (CBS) deficiency in UK and US patients with homocystinuria. *Human Mutation*, 23(2), 206–206. <https://doi.org/10.1002/humu.9214>
- Moorthie, S., Cameron, L., Sagoo, G. S., Bonham, J. R., & Burton, H. (2014). Systematic review and meta-analysis to estimate the birth prevalence of five inherited metabolic diseases. *Journal of Inherited Metabolic Disease*, 37(6), 889–898. <https://doi.org/10.1007/s10545-014-9729-0>
- Mudd, S. H., Levy, H. L., & Kraus, J. P. (2001). Disorders of transsulfuration. In C. R. Scriver, A. Beaudet, W. Sly, & D. Valle (Eds.), *The Metabolic Basis of Inherited Disease* (pp. 2007–2056). New York: McGraw-Hill.
- Niu, W., Wang, J., Qian, J., Wang, M., Wu, P., Chen, F., & Yan, S. (2018). Allosteric control of human cystathionine beta-synthase activity by a redox active disulfide bond. *Journal of Biological Chemistry*, 293(7), 2523–2533. <https://doi.org/10.1074/jbc.RA117.000103>
- Ojha, S., Wu, J., LoBrutto, R., & Banerjee, R. (2002). Effects of heme ligand mutations including a pathogenic variant, H65R, on the properties of human cystathionine beta-synthase. *Biochemistry*, 41(14), 4649–4654.
- Olatubosun, A., Valiaho, J., Harkonen, J., Thusberg, J., & Vihinen, M. (2012). PON-P: Integrated predictor for pathogenicity of missense variants. *Human Mutation*, 33(8), 1166–1174. <https://doi.org/10.1002/humu.22102>
- Pejaver, V., Mooney, S. D., & Radivojac, P. (2017). Missense variant pathogenicity predictors generalize well across a range of function-specific prediction challenges. *Human Mutation*, 38(9), 1092–1108. <https://doi.org/10.1002/humu.23258>
- Pey, A. L., Majtan, T., Sanchez-Ruiz, J. M., & Kraus, J. P. (2013). Human cystathionine beta-synthase (CBS) contains two classes of binding sites for S-adenosylmethionine (SAM): Complex regulation of CBS activity and stability by SAM. *Biochemical Journal*, 449(1), 109–121. <https://doi.org/10.1042/BJ20120731>
- Rost, B., Radivojac, P., & Bromberg, Y. (2016). Protein function in precision medicine: Deep understanding with machine learning. *FEBS Letters*, 590(15), 2327–2341. <https://doi.org/10.1002/1873-3468.12307>
- Schymkowitz, J., Borg, J., Stricher, F., Nys, R., Rousseau, F., & Serrano, L. (2005). The FoldX web server: An online force field. *Nucleic Acids Research*, 33(Web Server issue), W382–W388. <https://doi.org/10.1093/nar/gki387>
- Shan, X., Dunbrack, R. L., Jr., Christopher, S. A., & Kruger, W. D. (2001). Mutations in the regulatory domain of cystathionine beta synthase can functionally suppress patient-derived mutations in cis. *Human Molecular Genetics*, 10(6), 635–643. <https://doi.org/10.1093/hmg/10.6.635>
- Skovby, F., Gaustadnes, M., & Mudd, S. H. (2010). A revisit to the natural history of homocystinuria due to cystathionine beta-synthase deficiency. *Molecular Genetics and Metabolism*, 99(1), 1–3. <https://doi.org/10.1016/j.ymgme.2009.09.009>
- Skovby, F., Kraus, J. P., & Rosenberg, L. E. (1984). Biosynthesis and proteolytic activation of cystathionine beta-synthase in rat liver. *Journal of Biological Chemistry*, 259(1), 588–593.
- Tang, H., & Thomas, P. D. (2016). Tools for predicting the functional impact of nonsynonymous genetic variation. *Genetics*, 203(2), 635–647. <https://doi.org/10.1534/genetics.116.190033>
- Tavtigian, S. V., Byrnes, G. B., Goldgar, D. E., & Thomas, A. (2008). Classification of rare missense substitutions, using risk surfaces, with genetic- and molecular-epidemiology applications. *Human Mutation*, 29(11), 1342–1354. <https://doi.org/10.1002/humu.20896>
- Thomas, P. D., Kejarawal, A., Guo, N., Mi, H., Campbell, M. J., Muruganujan, A., & Lazareva-Ulitsky, B. (2006). Applications for protein sequence-function evolution data: MRNA/protein expression analysis and coding SNP scoring tools. *Nucleic Acids Research*, 34(Web Server issue), W645–W650. <https://doi.org/10.1093/nar/gkl229>
- Weeks, C. L., Singh, S., Madzalan, P., Banerjee, R., & Spiro, T. G. (2009). Heme regulation of human cystathionine beta-synthase activity: Insights from fluorescence and Raman spectroscopy. *Journal of the American Chemical Society*, 131(35), 12809–12816. <https://doi.org/10.1021/ja904468w>
- Wei, Q., Wang, L., Wang, Q., Kruger, W. D., & Dunbrack, R. L., Jr. (2010). Testing computational prediction of missense mutation phenotypes: Functional characterization of 204 mutations of human cystathionine beta synthase. *Proteins*, 78(9), 2058–2074. <https://doi.org/10.1002/prot.22722>
- Ye, Z. Q., Zhao, S. Q., Gao, G., Liu, X. Q., Langlois, R. E., Lu, H., & Wei, L. (2007). Finding new structural and sequence attributes to predict possible disease

- association of single amino acid polymorphism (SAP). *Bioinformatics*, 23(12), 1444–1450. <https://doi.org/10.1093/bioinformatics/btm119>
- Yoo, J., Lee, Y., Kim, Y., Rha, S. Y., & Kim, Y. (2008). SNPAnalyzer 2.0: A web-based integrated workbench for linkage disequilibrium analysis and association analysis. *BMC Bioinformatics*, 9, 290. <https://doi.org/10.1186/1471-2105-9-290>
- Yue, P., & Moul, J. (2006). Identification and analysis of deleterious human SNPs. *Journal of Molecular Biology*, 356(5), 1263–1274. <https://doi.org/10.1016/j.jmb.2005.12.025>
- Zou, C. G., & Banerjee, R. (2003). Tumor necrosis factor- α -induced targeted proteolysis of cystathionine beta-synthase modulates redox homeostasis. *Journal of Biological Chemistry*, 278(19), 16802–16808. <https://doi.org/10.1074/jbc.M212376200>

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

How to cite this article: Kasak L, Bakolitsa C, Hu Z, et al. Assessing computational predictions of the phenotypic effect of cystathionine-beta-synthase variants. *Human Mutation*. 2019;40:1530–1545. <https://doi.org/10.1002/humu.23868>