

# 1 Predicting the impact of rare variants on RNA splicing in CAGI6

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45

46

47 **Abstract**

48 **Background:** Variants which disrupt splicing are a frequent cause of rare disease that have been  
49 under-ascertained clinically. Accurate and efficient methods to predict a variant's impact on splicing  
50 are needed to interpret the growing number of variants of unknown significance (VUS) identified by  
51 exome and genome sequencing. Here we present the results of the CAGI6 Splicing VUS challenge,  
52 which invited predictions of the splicing impact of 56 variants ascertained clinically and functionally  
53 validated to determine splicing impact.

54 **Results:** The performance of 12 prediction methods, along with SpliceAI and CADD, was compared  
55 on the 56 functionally validated variants. The maximum overall accuracy achieved was 82% from two  
56 different approaches, one weighting SpliceAI scores by minor allele frequency, and one applying the  
57 recently published Splicing Prediction Pipeline (SPiP). SPiP performed optimally in terms of  
58 sensitivity, while an ensemble method combining multiple prediction tools and information from  
59 databases exceeded all others for specificity.

60 **Conclusions:** Several challenge methods equalled or exceeded the performance of SpliceAI, with  
61 ultimate choice of prediction method likely to depend on experimental or clinical aims. One quarter  
62 of the variants were incorrectly predicted by at least 50% of the methods, highlighting the need for  
63 further improvements to splicing prediction methods for successful clinical application.

64

65 **Introduction**

66 The diagnosis of rare disorders has been revolutionised in recent years thanks to the availability and  
67 widespread adoption of next generation sequencing technologies capable of detecting disease-  
68 causing variants. With the ever-decreasing prices of whole-exome sequencing (WES) and whole-  
69 genome sequencing (WGS) comes an increased use of these approaches, leading to the detection of  
70 more genetic variants than ever before. This brings with it a major challenge in understanding what  
71 these variants do, since our ability to detect them has far outstripped our ability to meaningfully  
72 interpret their effects, particularly outside of protein coding regions. As a result, even with WGS,  
73 around half of patients with rare disorders do not get a diagnosis (Turro et al. 2020; Stranneheim et  
74 al. 2021).

75 While estimates vary widely (Lord and Baralle 2021), it is thought somewhere between 15-60% of  
76 disease causing variants affect splicing (Krawczak et al. 1992; López-Bigas et al. 2005). Generally  
77 speaking, in diagnostic and research variant prioritisation pipelines, variants which fall within the  
78 2bp canonical splice acceptor or donor sites will be classed as splice-affecting, while variants outside

79 of those small regions are often not assessed for splicing impact. It is common for intronic and  
80 synonymous variants to be filtered out, while missense variants are generally assessed for their  
81 impact on protein structure and function without consideration for the role they may play in  
82 splicing. All of these variant types, however, can and do impact splicing and cause disease. This  
83 approach has led to an under-ascertainment of splice-affecting variants clinically (Lord et al. 2019).  
84 What is needed, particularly with the increasing use of WGS over WES enabling the detection of far  
85 more intronic variants than before, is a way to effectively triage which variants are splice-affecting  
86 and which are not.

87 Currently, under ACMG/AMP guidelines (Richards et al. 2015), *in silico* splicing prediction  
88 approaches may be used as supporting evidence for genetic diagnosis if multiple independent tools  
89 predict an impact on splicing. Experimental validation of splicing effects using RT-PCR, mini-genes or  
90 RNAseq is often required to confidently establish a variant's impact on splicing, but such approaches  
91 are time-consuming and expensive to perform at scale. Recent years have seen an explosion of  
92 innovative new approaches to splicing prediction, with many new tools being generated, often  
93 utilising machine learning. If a high degree of accuracy and reliability can be obtained from *in silico*  
94 approaches, we may be able to move away from requiring experimental confirmations, or at the  
95 least, have an efficient method to triage variants most in need of validation. This would require  
96 highly accurate algorithms and extensive testing in the clinical setting to give sufficient confidence in  
97 these optimal approaches.

98 The Splicing Variants of Unknown Significance (VUS) challenge in the 6<sup>th</sup> Critical Assessment of  
99 Genome Interpretation (CAGI6) sought to assess splicing prediction accuracy on a set of clinically  
100 ascertained, functionally validated variants. This enabled performance comparison of many cutting-  
101 edge splicing prediction approaches and gave insights into the types of variants not currently well  
102 captured by these methods.

## 103 **Methods**

### 104 Variant selection and validation

105 As previously described in Wai et al. 2020 (Wai et al. 2020), a total of 64 variants were ascertained  
106 through Wessex Regional Genetics Laboratory in Salisbury (52 variants) or the Splicing and Disease  
107 research study (12 variants) at the University of Southampton, ethically approved by the Health  
108 Research Authority (IRAS Project ID 49685, REC 11/SC/0269) and by the University of Southampton  
109 (ERGO ID 23056). Informed consent was provided for all patients for splicing studies to be  
110 conducted. All variants had been, or were undergoing RT-PCR analysis to determine their impact on

111 splicing using RNA from whole blood collected in PAXgene tubes, again as previously described (Wai  
112 et al. 2020).

113 Eight variants were excluded from the final analysis (unable to establish splicing impact before  
114 analysis period (n=3), incorrect gene/variant annotations given in the dataset distributed (n=3),  
115 variant found to impact gene expression rather than splicing (n=2)), giving a total of 56 variants in  
116 the final assessment set (**Supplementary Table 1**), which span a wide range of rare disease and  
117 cancer predisposition associations, none of which had had their impact on splicing published  
118 previously.

119 The Splicing VUS challenge

120 Variants were distributed as a tab delimited text file including the following information: HGNC  
121 identifier, chromosome, position, reference allele, alternative allele, gene and strand. Entrants also  
122 had access to 256 previously published variants (Wai et al. 2020) obtained and validated by the same  
123 approach to aid in method development/testing.

124 Challenge participants submitted their entries in the form of tab delimited text files including the  
125 variant information, a binary prediction of whether a variant affected splicing or not (1=yes, 0=no),  
126 along with a score for the probability of the variant affecting splicing and the level of confidence in  
127 the prediction given. All assessments were based on the binary splice-affecting prediction alone.

128 Challenge assessment

129 The performance of each prediction model was assessed by calculating and comparing a series of  
130 metrics: overall accuracy, area under the receiver operating characteristic curve (AUC), sensitivity,  
131 specificity, positive predictive value (PPV) and negative predictive value (NPV). AUC and confidence  
132 intervals (2000 stratified bootstrap replicates) were calculated using the pROC package (Robin et al.  
133 2011) in R v3.5.1 (R Core Team 2018), and plots made with ggplot2 (Wickham 2009). Performance of  
134 each method was compared across binned splicing locations – Near Acceptor (acceptor +/- 10bp),  
135 Near Donor (donor +/- 10bp), Exonic Distant (exonic, 11bp or more from either splice site), Intronic  
136 Distant (intronic, 11bp or more from either splice site. For grouped analyses, exonic distant and  
137 intronic distant variants were grouped together due to low numbers). These scores were based on  
138 the concordance of the binary classification of the variants provided by each team/model (1=splice-  
139 affecting and 0=not splice-affecting) with the experimental validation of the splicing impact.

140 SpliceAI (Jaganathan et al. 2019) and CADD v1.6 (Kircher et al. 2014) (which incorporates SpliceAI  
141 predictions) were included in the assessment alongside the challenge models as a comparison to  
142 emerging industry standards. CADD-phred scores were obtained by uploading a VCF to the CADD

143 webserver (<https://cadd.gs.washington.edu/score>). SpliceAI scores were obtained from Ensembl's  
144 Variant Effect Predictor (VEP) web interface (McLaren et al. 2016) (44 variants scored) or using the  
145 SpliceAI webserver from the Broad Institute (<https://spliceailookup.broadinstitute.org/>, 11 variants  
146 that were not scored by VEP; options: hg38, masked scores, max distance 50bp). A cut-off of 0.2 was  
147 used for SpliceAI scores, and 18 for CADD.

148

## 149 **Results**

### 150 Variant characteristics of challenge set

151 Of the 56 variants in the final analysis, the majority (n=49, 87.5%) were SNVs, with 7 indels (12.5%).  
152 The variants fell within 42 different genes, broadly representative of clinical genetics referrals in the  
153 UK, with the majority of genes having a single variant in the set, and only 7 genes with >1 variant  
154 (*BRCA1* n=6, *FBN1* n=4, *MYBPC3* n=3, *BRCA2* n=2, *SCN5A* n=2, *APC* n=2, *USP7* n=2). 37 variants (66%)  
155 were found to affect splicing, while 19 (34%) had no observable impact.

156 Variants were divided into 5 groups by their positions relative to intron-exon boundaries. There were  
157 16 variants within 10bp of a splice acceptor site (NearAcc), and 23 within 10bp of a splice donor site  
158 (NearDon). 10 exonic variants >10bp from either splice site were classed as Exonic>10. Intronic  
159 variants >10bp from their nearest splice site were termed Intronic Distant (six upstream of the  
160 acceptor, one downstream of the donor). The locations of all variants relative to the intron-exon  
161 boundary and whether the variants were determined to be splice disrupting or not are given in **Fig1**.

### 162 Challenge participants

163 Eight teams submitted predictions for the challenge, with two teams submitting predictions from  
164 multiple models, giving 12 models altogether. **Table 1** gives a summary of the approach taken by  
165 each model, which was provided by the challenge entrants upon submission of their predictions, but  
166 blinded to the assessors until after the assessment period.

### 167 Model performance across 56 variants

168 **Table 2** summarises the performance metrics of the 12 models, along with CADD and SpliceAI. Full  
169 variant information, scores and binary predictions for the 12 models, SpliceAI and CADD and  
170 experimental outcome of splicing status are given in **Supplementary Table 1**. The ROC plots for each  
171 model are shown in **Fig2**, and **Supplementary Fig1** shows the performance of each method on each  
172 variant across the splicing region.

173 **Table 1** – Summary of the prediction approaches of the 12 models from 8 entrants. Additional information on Teams 4 and 5 given in the **Supplementary**  
 174 **Methods**.

Team	Authors	Prediction approach
1	YW, ZH	<p>Models were built based on reported pathogenic splicing variants from the literature and benign variants from ClinVar(Landrum et al. 2018). The models were trained and tuned using Gradient Boosting Machine (GBM) with R package “caret” and “gbm”, considering 80 annotation features, including conservation, distance to exon-junctions, population allele frequencies, epigenetic states and prediction scores from SpliceAI(Jaganathan et al. 2019), CADD(Kircher et al. 2014), SCAP(Jagadeesh et al. 2019) and dbSCNV(Jian et al. 2014).</p> <p>Model 1 - Full model which uses all 80 features</p> <p>Model 2 - Five existing prediction scores as features</p> <p>Model 3 - As Model 2, plus distance to splice site and the splice site type as two additional features.</p>
2	ZZ	Positive predictions from CADD-Splice(Rentzsch et al. 2021) (>15), SpliceAI(Jaganathan et al. 2019) (>0.5), MMsplice(Cheng et al. 2019) (>2), and Ensembl Variant Effect Predictor(McLaren et al. 2016) variant consequence (splice region) ranked as “1”, negative predictions as “0”. Mean of the four ranks calculated, and mean $\geq 0.5$ classed as positive overall.
3	DD	Super Quick Information-content Random-forest Learning of Splice variants (SQUIRLS)(Danis et al. 2021) applied to data using default thresholds
4	PK, AW, OL	SpliceAI(Jaganathan et al. 2019) adjusted with minor allele frequency(Karczewski et al. 2020), with scores $>0.25$ classified as splice affecting
5	YC, RDB	<p>Combined information from ClinVar(Landrum et al. 2018), gnomAD(Karczewski et al. 2020), established splicing tools (SpliceAI(Jaganathan et al. 2019) (&gt;0.5), MaxEntScan(Yeo and Burge 2004) (&gt;4)), branchpoint/enhancer locations, distance to exon, splice site database.</p> <p>Model 1 – Base model for prediction</p> <p>Model 2 – Same as Model 1 but using different in-silico prediction score thresholds (SpliceAI(Jaganathan et al. 2019) (&gt;0.5), MaxEntScan(Yeo and Burge 2004) (&gt;6), MMsplice(Cheng et al. 2019) (&gt;2))</p> <p>Model 3 - Required well-scoring compatible site (e.g. for donor loss, a well-scored donor within 300bp of the existing acceptor), adding branchpoint/enhancer locations as extra features</p>
6	SMM, BM, CL	SpliceAI(Jaganathan et al. 2019) applied, with scores $\geq 0.21$ classified as splice affecting
7	TvOH	Alamut splicing software (Sophia Genetics) utilised – consensus of 3 programs with at least 10% difference between reference and alternative score predicted to be splice affecting and ACMG splicing guidelines (BRCA1/BRCA2 – ENIGMA).
8	RL, AM, CH, SK	Splicing Prediction Pipeline (SPiP)(Leman et al. 2022) applied (>0.18 for exonic variants, >0.035 for intronic variants)

175 **Table 2** – Summary statistics on predictive performance of the 12 competition entrants plus SpliceAI and CADD on the 56 challenge variants. Maximum  
176 value for each metric indicated in bold.

	<b>T1_1</b>	<b>T1_2</b>	<b>T1_3</b>	<b>T2</b>	<b>T3</b>	<b>T4</b>	<b>T5_1</b>	<b>T5_2</b>	<b>T5_3</b>	<b>T6</b>	<b>T7</b>	<b>T8</b>	<b>SpliceAI</b>	<b>CADD</b>
<b>AUC (binary)</b>	0.813	0.826	0.786	0.720	0.708	<b>0.839</b>	0.718	0.717	0.731	0.813	0.731	0.775	0.826	0.537
<b>AUC (score)</b>	0.883	0.903	0.883	0.780	0.788	0.912	<b>0.770</b>	0.770	0.770	0.910	0.801	0.874	<b>0.919</b>	0.543
<b>95% CI (bootstrap n=2000)</b>	0.771- 0.969	0.805- 0.976	0.771- 0.970	0.658- 0.891	0.652- 0.909	0.827- 0.977	0.637- 0.891	0.648- 0.883	0.642- 0.883	0.819- 0.974	0.693- 0.907	0.754- 0.964	0.841- 0.964	0.386- 0.706
<b>Accuracy</b>	0.804	0.804	0.768	0.714	0.732	<b>0.821</b>	0.661	0.643	0.679	0.804	0.679	<b>0.821</b>	0.804	0.625
<b>Sens</b>	0.784	0.757	0.730	0.703	0.784	0.784	0.541	0.486	0.568	0.784	0.568	<b>0.919</b>	0.757	0.811
<b>Spec</b>	0.842	0.895	0.842	0.737	0.632	0.895	0.895	<b>0.947</b>	0.895	0.842	0.895	0.632	0.895	0.263
<b>PPV</b>	0.906	0.933	0.900	0.839	0.806	0.935	0.909	<b>0.947</b>	0.913	0.906	0.913	0.829	0.933	0.682
<b>NPV</b>	0.667	0.654	0.615	0.560	0.600	0.680	0.500	0.486	0.515	0.667	0.515	<b>0.800</b>	0.654	0.417

177 AUC = Area Under the Curve; CI = Confidence Interval; Sens = Sensitivity; Spec = Specificity; PPV = Positive Predictive Value; NPV = Negative Predictive Value

178 No single approach performed optimally on all assessment metrics (**Table 2**). Overall accuracy was  
179 joint highest in Teams 4 and 8 at 0.82, with Team 4 also achieving the highest binary outcome AUC  
180 at 0.839 (**Fig2**). Team 8 ranked highest on the related metrics for sensitivity (0.919) and NPV (0.800),  
181 indicating its permissive prediction approach. Conversely, Team 5's Model 2 performed the best in  
182 terms of specificity (0.947) and PPV (0.947), with the lowest proportion of false positive findings. All  
183 three models by Team 1, plus Team 4 and Team 6 achieved over 70% in both sensitivity and  
184 specificity, indicating more balanced performance.

185 Included as comparators were SpliceAI with a cut-off of 0.2 and CADD with a cut-off of 18. SpliceAI  
186 was competitive with the challenge entrants, ranking near-top but not top on all metrics, and indeed  
187 top in the AUC when measured using prediction score rather than binary prediction outcome. CADD,  
188 however, performed poorly on the challenge set with specificity in particular being very low (0.263).

189 Performance comparison by variant type

190 In order to get an overall impression of the performance of the methods on different types of  
191 variants, variants were grouped by location relative to their nearest splice site (**Fig3**), as described in  
192 Methods. All methods performed better on exonic distant variants than intronic distant variants,  
193 with the exception of SpliceAI, which correctly predicted all seven intronic distant variants. Across  
194 methods, there was a high degree of consistency in the proportion of variants correctly predicted in  
195 the near acceptor region, and a high degree of variance in performance in the intronic distant set.  
196 The types of error differed across regions, with the near acceptor region and exonic distant region  
197 having very few false positive predictions across all methods, while almost all methods gave false  
198 positive predictions in the near donor and intronic distant regions (**Supplementary Fig2**).

199 We also compared the performance of the approaches on SNVs vs indels, and found all methods  
200 except CADD had higher accuracy on SNVs than indels (**Supplementary Fig3**).

201 Some variants are consistently mispredicted

202 21 of the variants (37.5%) were correctly predicted by all 12 submitted prediction methods. None of  
203 the variants were incorrectly predicted by all methods, but 14 variants (25%) were predicted  
204 correctly by <=50% of the methods, with two variants only being correctly predicted by a single  
205 method. These were a splice-affecting single nucleotide deletion 4bp from a splice acceptor site in  
206 *KANSL1* (correctly predicted by Team 3) and an SNV in the last base of an exon in *TRPM6* which  
207 despite altering the conserved last G nucleotide did not affect splicing in functional testing (correctly  
208 predicted by Team 4).

209

210 **Discussion**

211 The CAGI6 Splicing VUS challenge assessed the performance of 14 prediction approaches on a set of  
212 56 clinically relevant variants whose impact on splicing had been functionally tested using RT-PCR. A  
213 variety of approaches were adopted, and several methods equalled or exceeded the performance of  
214 the emergent field leader, SpliceAI.

215 While Teams 4 and 8 had joint highest overall accuracy, there was no single optimal method for the  
216 Splicing VUS challenge, since several different models performed optimally on different metrics.

217 Choice of approach may therefore be dependent on the specific nature of the predictions required.

218 Seeking a molecular diagnosis for a particular family may favour sensitivity over specificity, since  
219 overlooking a causal variant would prevent this aim, so Team 8's approach with almost 92%  
220 sensitivity may be preferred. Seeking confident splice disrupting candidates for functional validation  
221 or mechanistic research may call for greater specificity than sensitivity to avoid wasting resources on  
222 false positive variants that do not have an impact, in which case Team 5's model 2 with almost 95%  
223 specificity may be the strategy of choice.

224 SpliceAI and CADDv1.6 were chosen as comparators for the entrants to the Splicing VUS challenge  
225 and were run by the assessors on the 56 challenge variants. SpliceAI has been emerging as a field  
226 leader in recent years, with accuracies >90% attained in several studies (Wai et al. 2020; Ha et al.  
227 2021; Strauch et al. 2022), although variable performance reported by some (Riepe 2020) which is  
228 more consistent with our observed 80.4% overall accuracy in this study.

229 CADD did not perform well on the challenge variants, achieving an overall accuracy of 62.5%.  
230 However, this was predominantly driven by a very low specificity, which is to be expected from  
231 CADD, since it is not only the impact on splicing being assessed by the tool, but overall  
232 deleteriousness. For example, missense variants which were not found to affect splicing in the  
233 challenge set may still have been pathogenic through impact on protein structure and/or function.  
234 For such variants, CADD would accurately classify them as deleterious in general, but in our  
235 assessment solely of splicing impact, this would appear as a false positive, lowering CADD's  
236 specificity. Notably, the version of CADD included in the assessment (v1.6) includes SpliceAI and  
237 additional splicing prediction tools in its underlying model (Rentzsch et al. 2021). Scoring the  
238 challenge variants with CADD v1.5 which did not include these splicing metrics resulted in an overall  
239 accuracy around 44.6% (data not shown). From these values it is clear that the explicit inclusion of  
240 splicing prediction methods within CADD's underlying model has improved its ability to predict  
241 variants that impact splicing. CADD's broad approach makes it a versatile tool for prediction of

242 deleteriousness for many different variant types. At present, however, if predicting a variant's  
243 splicing impact is the sole aim, the use of more specialised splicing tools is more appropriate.

244 Of note, SpliceAI featured heavily across the predictive strategies, being the sole predictive method  
245 for Team 6 and contributing heavily to the predictions of Team 4, which were weighted by MAF, as  
246 well as being run as a comparator by the assessors. Differences in the performance of these  
247 approaches highlight that even with the same nominal method, there can be variance in predictions  
248 depending on how the scores are obtained, and the thresholds that are used to determine positive  
249 predictions. There were just three approaches that did not include SpliceAI as part of their  
250 predictions, two utilising instead recent machine learning based prediction tools SQUIRLS (Danis et  
251 al. 2021) and SPiP (Leman et al. 2022), and one based on the splicing prediction tools available  
252 within the Alamut software, which has been widely used in clinical practice. Of the three, SPiP was  
253 the only method to achieve greater accuracy than SpliceAI.

254 A major strength of the challenge in terms of providing a real-world assessment of the performance  
255 of these tools is the ascertainment of the challenge variants from genuine clinical practice, where  
256 potential splice altering variants in genes relevant to the patient's presentation were referred for  
257 validation. This is precisely the type of variant splicing prediction models should be tested on when  
258 assessing their suitability for clinical application in rare disorders. It highlights that even in  
259 exceptionally well-studied genes, such as the BRCA genes, challenges in variant interpretation  
260 remain, since 3 of 8 variants across *BRCA1* and *BRCA2* were incorrectly predicted by over half of  
261 challenge methods, and only two of these were accurately predicted by all methods. However, the  
262 relatively small sample size makes it difficult to draw any major inferences and is a significant  
263 limitation of the study. Apparent variance in performance may be stochastic at such a sample size,  
264 and may not be fully reflective of overall performance in a wider context. It also made drawing firm  
265 conclusions about performance in subsets of the data, e.g. split by location, variant type, or disease  
266 group challenging. However, ascertaining a large body of clinical variants, validating the splicing  
267 impact and keeping that private, as is needed for a blinded challenge such as the CAGI6 Splicing VUS  
268 challenge, raises ethical concerns. Accurate and timely variant interpretation is reliant on sharing of  
269 data, and withholding a large body of functionally validated variants from resources such as ClinVar  
270 (Landrum et al. 2018) which are heavily used in clinical assessment of variants does not represent  
271 good practice.

272 This small but highly clinically relevant challenge assessed the performance of 12 prediction methods  
273 plus SpliceAI and CADD on 56 clinically ascertained variants and found SpliceAI weighted by allele  
274 frequency and SPiP to be the most accurate overall, while other methods had particular strengths in

275 their sensitivity or specificity. A quarter of variants were incorrectly predicted by half or more of the  
276 methods, showing there is still improvement to be made. Furthermore, this challenge was limited to  
277 a binary outcome – whether or not splicing was disrupted, but did not address the nature of that  
278 disruption, which may present an even greater challenge. A larger assessment set that would enable  
279 further investigation of the types of variants that are consistently incorrectly predicted may help  
280 direct efforts for refinement of models moving forwards.

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## 296 **Author contributions**

297 DB and JL conceived of the challenge. AGLD, DJB and JL selected variants to include in the set, which  
298 had been functionally validated by HAW and DJB. JL assessed challenge entrants and conducted data  
299 analysis. CJO conducted additional analyses and presented the findings at the CAGI6 conference. All  
300 further authors submitted prediction methods in response to the challenge. JL drafted the  
301 manuscript, with revision suggestions and final approval from all other authors.

## 302 **Data availability**

303 All data generated or analysed during this study are included in this published article [and its  
304 supplementary information files].

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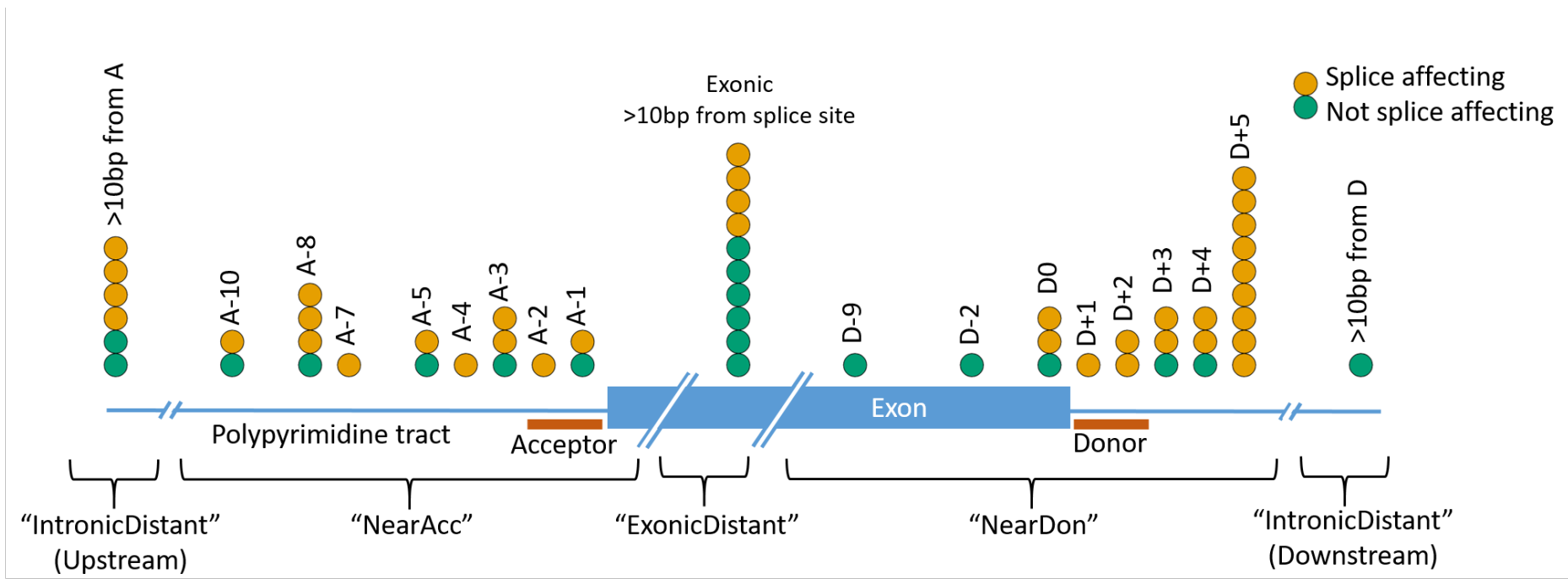
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373 **Figures**

374 **Fig1.** Schematic diagram showing locations of the 56 challenge variants in relation to their nearest splice site, with colour indicating whether (yellow) or not  
375 (green) each variant was determined experimentally to impact splicing.

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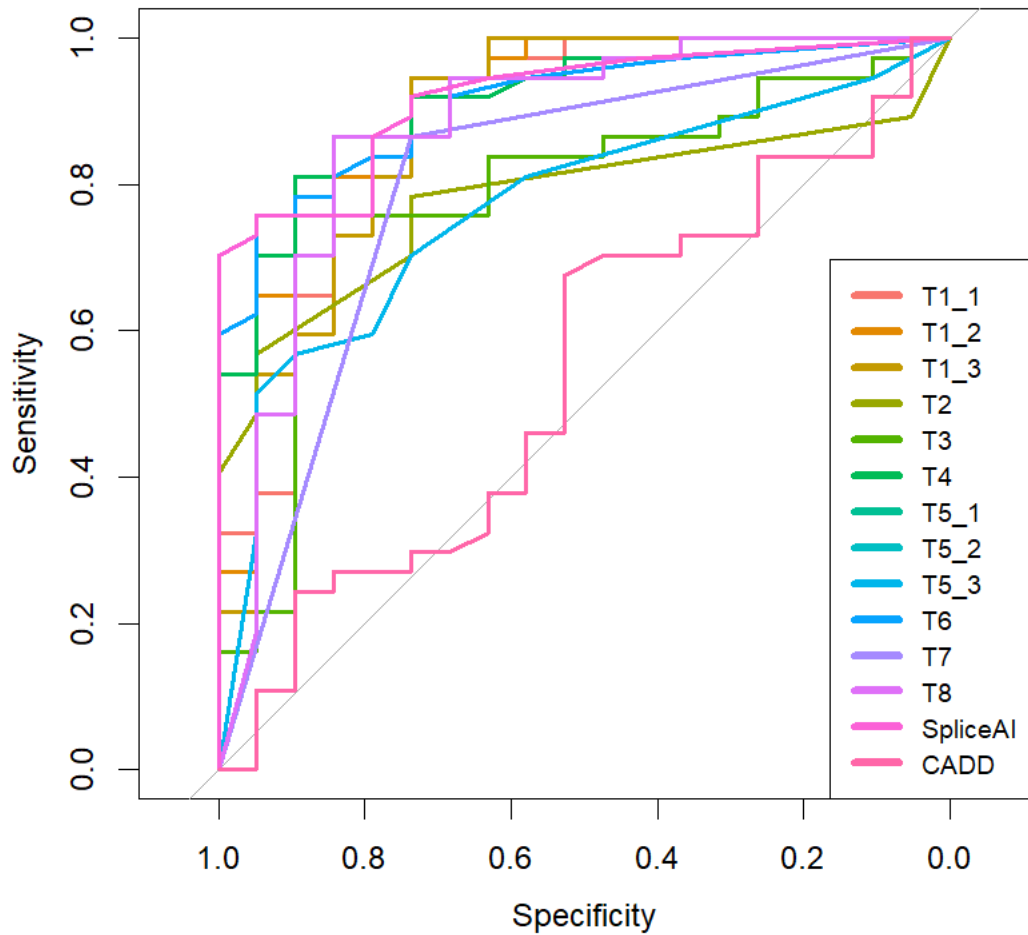
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**Fig2.** Receiver operating characteristic (ROC) curves of model performance based on prediction scores. For Area Under Curve (AUC), see **Table 2**.

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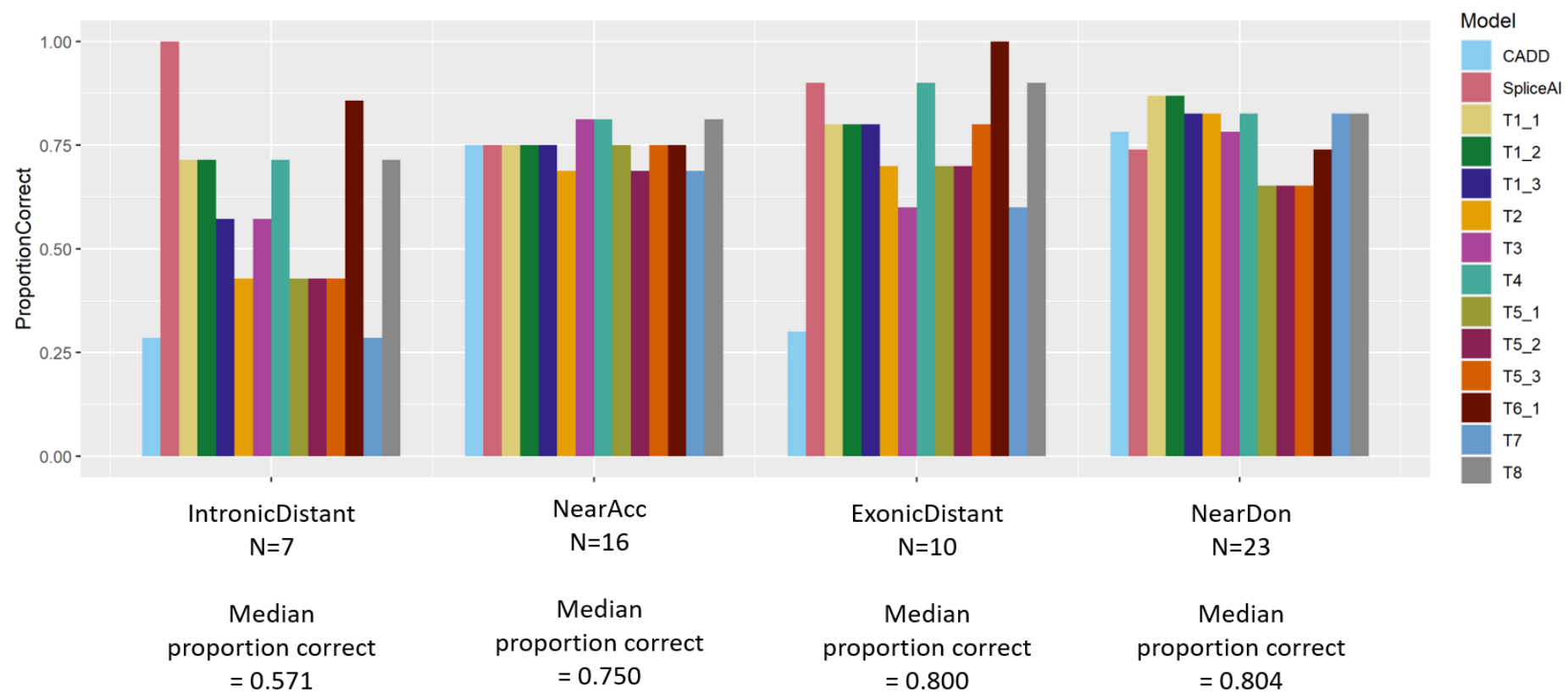


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383 **Fig3.** Proportion of variants correctly predicted by each method in the different regions (near acceptor, near donor, exonic and intronic distant).

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387 **Fig4.** Variants across the splicing region coloured by the number of prediction methods (out of the 12 challenge entrants) that correctly predicted the  
388 splicing outcome.

