

# **Predicting Pathogenicity using Aberrant Splicing Predictions of Single Nucleotide Variants in Autism Spectrum Disorder**

**Bioinformatics Laboratory, BIMM 185**

**Kevin Chau**

University of California, San Diego  
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## **Abstract**

Distributions of splicing likelihood scores for single nucleotide variants (SNVs) were analyzed with the hopes that these scores could be used to predict their pathogenicity, defined as a pathogenicity score multiplied by a risk score for that gene. SNVs implicated in autism as well as those found in control phenotypes were gathered and scored for their likelihood to cause cassette exon skipping and their pathogenicity. Gene coexpression networks were constructed for affected genes per eight developmental periods using publicly available RNA-seq data. Genes were scored for risk, per period, based on relative numbers of coexpression partners. Figures show that there is no difference in the distribution of splicing scores between SNVs implicated in autism versus control, nor was there any correlation between the splice score and pathogenicity (modified by the gene risk score).

# 1 Introduction

Prediction of pathogenicity of genetic variants is paramount to current understanding of diseases with heavy genetic influence and, by extension, development of treatments. In addition, it has been shown that aberrant splicing can have major contributions to expression of disease phenotypes, likely due to a propensity to disrupt biological pathways through alteration of gene products. Therefore, one might deduce that variants predicted to cause alternative splicing in genes central to biological processes are likely to result in the expression of the pathogenic phenotype. With numerous tools currently available to the public for academic use, a predictive model may be constructed with the hopes that the likelihood that a genetic variant causes aberrant splicing can be used to predict the contribution of that variant to development of a given disease. This knowledge could provide new insight into therapeutic targets at the gene or even isoform level.

## 2 Data and Processing

A list of variants classified by expressed phenotype was downloaded from DenovoDB, a database of nucleotide variants. Only variants tagged with the autism phenotype and control phenotype were downloaded. Overall, the taken from DenovoDB included the phenotype, the nucleotide variant, chromosome, affected gene, and starting position of the variant. This table was uploaded to a locally hosted MySQL database for simplified querying. Since only SNVs were analyzed, the dataset was filtered for only mutations that substitute one nucleotide for another. The SPIDEX splicing scoring tool was downloaded in order to score the nucleotide variants for their likelihood to cause cassette exon skipping.

### 2.1 Splicing Scores with SPIDEX

The SPIDEX tool was used to score each SNV for the likelihood that it causes cassette exon skipping. The scoring metric used was delta PSI (dpsi), the change in percent inclusion rate; negative values correspond to decreased inclusion (exon skipping) and positive values correspond to increased inclusion (exon retention). The criteria for scoring was that the SNV must have occurred within 300 nucleotides from a splice junction; that is, within 300 nu-

cleotides from an exon-to-intron or intron-to-exon transition position. Since not all variants fell within that distance, only scorable variants were retained for further analysis.

The SPIDEX tool is downloaded as a tab-indexed gzip-compressed file. The contents of the file was queried using the *tabix* utility provided by the htslib C library (formerly packaged with samtools). Example query and output follows:

```
tabix spidex_public_noncommercial_v1.0/
    spidex_public_noncommercial_v1.0.tab.gz chr1:0 | head -10 |
    cut -f1,2,3,4,5,7
chr1    861181    G          A          0.5983    SAMD11
chr1    861181    G          C          0.3903    SAMD11
chr1    861181    G          T          0.6564    SAMD11
chr1    861182    T          A          0.4921    SAMD11
chr1    861182    T          C          0.4723    SAMD11
chr1    861182    T          G          0.7517    SAMD11
chr1    861183    G          A          -0.0870    SAMD11
chr1    861183    G          C          -0.0874    SAMD11
chr1    861183    G          T          0.1979    SAMD11
chr1    861184    G          A          -0.1115    SAMD11
```

with fields as Chromosome, Position, Reference Allele, Mutant Allele, dPSI, and Gene.

## 2.2 Pathogenicity Scores with UMD Predictor

Pathogenicity of the given variants were scored with the web tool UMD Predictor. This online tool takes in a list of formatted variants and positions and returns a score for pathogenicity of the variant for each transcript affected. These scores ranged from 0 to 100, with 0 being non-pathogenic and 100 being confirmed pathogenicity. These scores were converted to percentages in order to more closely conform with the splicing scores.

## 2.3 Gene Networks and Risk Analysis

Gene coexpression networks were constructed in order to take into account gene risk factors, with the reasoning that genes with many coexpression partners are more likely to play an important role in their respective biological pathways; disruption in their splicing would thus lead to pathogenicity. These

coexpression networks were created using RNA-seq expression data from the BrainSpan database. A single network was developed for each of eight developmental periods, since expression values are likely to vary depending on the given stage of life. These developmental periods are as follows: 8 weeks postconception (PCW) to 12PCW, 13PCW to 18PCW, 19PCW to 23PCW, 24PCW to 37PCW, 0 months after birth (M) to 11M, 1 year (Y) to 11Y, 12Y to 19Y, and 21Y<sup>+</sup>. Pairwise comparisons were performed using the Pearson correlation metric and all gene-gene coexpression pairs with Pearson correlation coefficients of  $\geq 0.7$  were retained. The number of coexpression partners for each gene was calculated and divided by the maximum of the set, yielding the risk score for that gene. These scores were appended to all variants that affect that gene; the relevant variant information along with the calculated scores were all uploaded to a local MySQL database.

### 3 Data Analysis

The final dataset to analyze was compiled into a MySQL database with the following layout:

```
mysql> DESCRIBE scored_denovo_db;
```

Field	Type	Null	Key	Default
PrimaryPhenotype	<b>varchar</b> (125)	<b>NO</b>	MUL	<b>NULL</b>
Gene	<b>varchar</b> (125)	<b>NO</b>	MUL	<b>NULL</b>
Transcript	<b>varchar</b> (125)	<b>NO</b>	MUL	<b>NULL</b>
Chromosome	<b>varchar</b> (125)	<b>NO</b>		<b>NULL</b>
<b>Position</b>	<b>bigint</b> (15)	<b>NO</b>		<b>NULL</b>
Variant	<b>varchar</b> (500)	<b>NO</b>		<b>NULL</b>
SpliceScore	double	<b>NO</b>		<b>NULL</b>
PathogenScore	double	<b>NO</b>		<b>NULL</b>
P1Risk	double	<b>NO</b>		<b>NULL</b>
P2Risk	double	<b>NO</b>		<b>NULL</b>
P3Risk	double	<b>NO</b>		<b>NULL</b>
P4Risk	double	<b>NO</b>		<b>NULL</b>
P5Risk	double	<b>NO</b>		<b>NULL</b>
P6Risk	double	<b>NO</b>		<b>NULL</b>
P7Risk	double	<b>NO</b>		<b>NULL</b>
P8Risk	double	<b>NO</b>		<b>NULL</b>

From this data, distributions could be plotted. First, a kernel density estimate was performed for the frequencies of splice scores for both the autism phenotype and control phenotype.

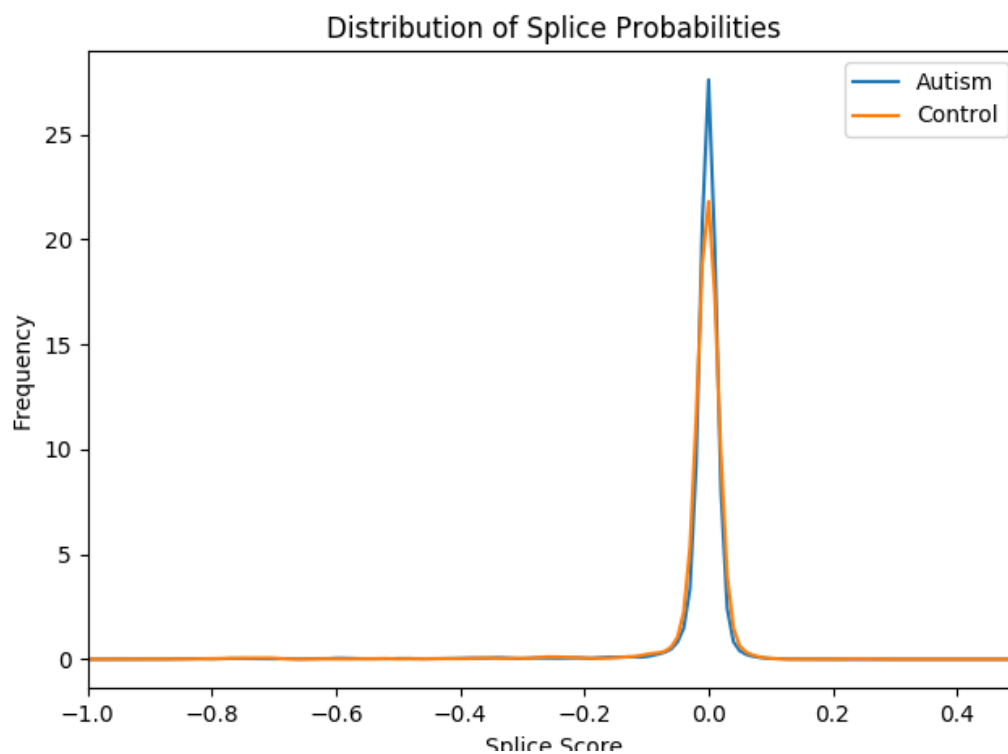


Figure 1: Kernel density estimates of autism SNV splice scores and control SNV splice scores

Scatter plots between splicing scores and pathogenicity scores may also be drawn in order to illustrate any correlation between the two over each developmental period and in each condition.

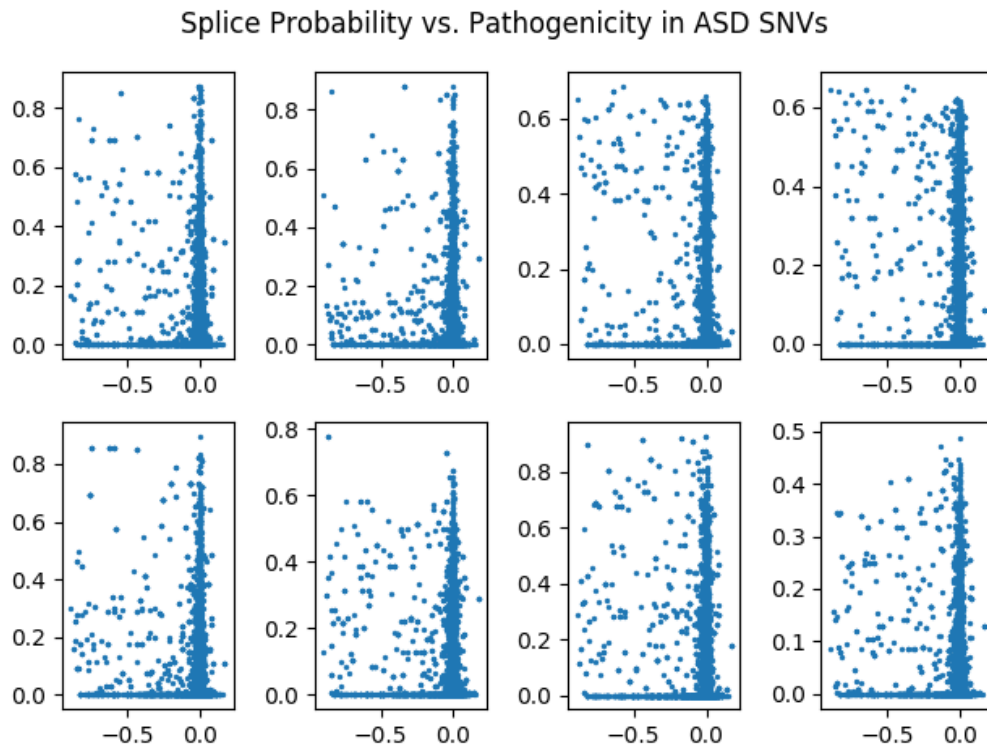


Figure 2: Splicing probabilities (x-axis) plotted against pathogenicity probabilities (y-axis) for each developmental period in SNVs implicated in autism

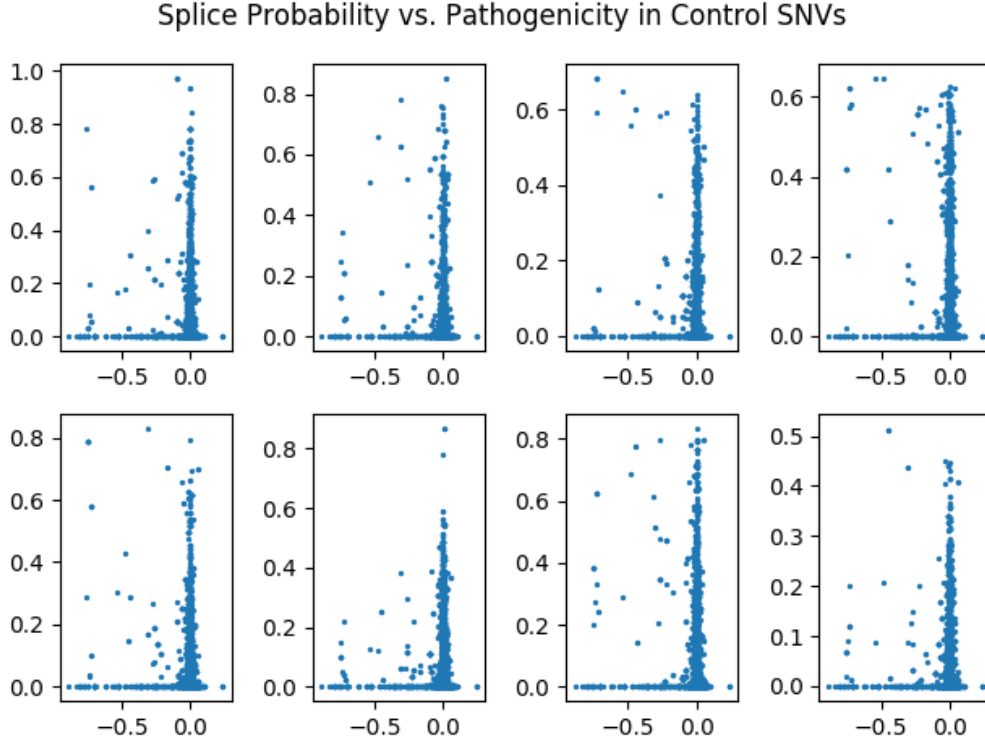


Figure 3: Splicing probabilities (x-axis) plotted against pathogenicity probabilities (y-axis) for each developmental period in control SNVs

## 4 Discussion

As clearly shown by the kernel density graph, there appears to be no difference in the distribution of autism-implicated SNVs and control SNVs; it would appear that there is no association between splicing predictions and disease phenotype expression. This conclusion is further supported by the splice versus pathogen score plots that follow. In this specific context, there is no correlation between splicing prediction scores and pathogenicity scores.

Since there is no difference between the distributions of SNV splice likelihoods between the autism phenotype and control phenotypes, an inference model incorporating these values would be completely unreliable. However, the data and the methods used to process that data are highly specific to this



particular context. That is, aberrant splicing of gene transcripts may contribute more so to other genetic diseases besides autism spectrum disorder. Additionally, this analysis was performed against only single nucleotide variants; that is, only nucleotide substitutions were observed. Therefore, there is likely a lower chance for actual protein modifications relative to mutations that induce frameshifts. There was no differentiation between synonymous and nonsynonymous SNVs; the inclusion of synonymous SNVs could very well skew the resulting distributions toward similarity between the phenotypes. Furthermore, this entire process is based on the assumption that the given tools can reliably predict the required values. Further analysis is required using alternative datasets and prediction tools.

Although the results show that an inference model based on splice scores is not feasible, there are still a number of tangential observations that can be made. Interestingly, most splice scores are gathered near 0; that is, most of the given SNVs yield a change-in-exon-inclusion percentage of only slightly less than 0, indicating that most of these variants do not actually induce a change in splicing, assuming the reliability of the SPIDEX splice predictor. With regards to the gene risk assessment, period four exhibited the highest interconnectedness among its genes, which is reasonable given that this period represents the time interval directly before birth, giving evidence of high rates of neurodevelopment during this time period.

## 5 Supplementary Materials

Relevant code:

<https://github.com/kkchau/BIMM-185-FINAL-PROJECT/>