The contribution of non-canonical splicing mutations to severe dominant developmental disorders

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Abstract

Accurate and efficient pre-mRNA splicing is crucial for normal development and function, and mutations which perturb normal splicing patterns are significant contributors to human disease. We used exome sequencing data from 7,833 probands with developmental disorders (DD) and their unaffected parents to quantify the contribution of splicing mutations to DDs. Patterns of purifying selection, a deficit of variants in highly constrained genes in healthy subjects and excess de novo mutations in patients highlighted particular positions within and around the consensus splice site of greater disease relevance. Using mutational burden analyses in this large cohort of proband-parent trios, we could estimate in an unbiased manner the relative contributions of mutations at canonical dinucleotides (73%) and flanking non-canonical positions (27%), and calculated the positive predictive value of pathogenicity for different classes of mutations. We identified 18 likely diagnostic de novo mutations in dominant DD-associated genes at non-canonical positions in splice sites. We estimate 35-40% of pathogenic variants in non-canonical splice site positions are missing from public databases.
Introduction

Pre-mRNA splicing in humans is mediated by the major and minor spliceosomes, highly dynamic, metalloenzyme complexes comprised of five key small nuclear RNAs (snRNA), along with over 100 protein components and accessory molecules\(^1-3\). Accurate recruitment and function of the complex is reliant on a plethora of cis-acting regulatory elements encoded within the pre-mRNA itself, including the canonical acceptor and donor (or 3’ and 5’ splice) dinucleotides, the branchpoint, which offers a “tether” for lariat formation, the polypyrimidine tract (PolyPy), and a catalogue of exonic and intronic splice enhancers and silencers (ESE/ISE/ESS/ISS). Whilst our understanding of the underlying mechanistic processes regulating splicing has greatly increased in recent years, our ability to predict whether or not a mutation will affect splicing remains limited. However, with estimates that up to 50% of monogenic disease-causing variants may affect splicing\(^4;5\), a better understanding and more coherent approach to interpretation of variants affecting splicing is badly needed\(^6;7\). With a plethora of *in silico* splicing pathogenicity predictors available, there is little consensus on what a “gold standard” for splicing pathogenicity prediction would be\(^8-10\). Whilst many of these methods perform well within the canonical splice site dinucleotides (CSS, the two highly-conserved bases flanking the acceptor and donor sites), their utility for other splice relevant regions is less clear\(^8\). In the clinical setting, often multiple algorithms and expert judgment are used to predict pathogenicity, while for large scale gene discovery research projects, classification of variants is often binary, with CSS mutations typically classified as likely splice affecting, whilst mutations in other splicing regulatory components are typically overlooked\(^11-13\). Both clinical and research interpretation of potential splice-disrupting variants has lacked a robust quantitative foundation.

We sought to assess the relative contribution of pathogenic, splice-altering mutations between the CSS and other, near-splice positions using both population-based and disease-focussed analyses utilising large scale exome sequencing data from the Deciphering Developmental Disorders (DDD) project\(^12\) and ExAC\(^14\).

We recruited 7,833 probands with undiagnosed developmental disorders, along with their parents, from clinical genetics centres across the UK and Ireland to the DDD study. Exome sequencing was performed to find likely genetic diagnoses underlying their conditions. Likely diagnostic *de novo* mutations (DNMs) in known developmental disorder (DD) genes have been found for ~25% of probands, while a smaller contribution of recessive disorders has also been observed\(^12;13;15\), meaning over half the cohort currently lacks a molecular diagnosis.
Our analyses focus on near-splice site positions across a set of 148,244 stringently defined exons well covered (median coverage >15X at both CSS) across the DDD cohort (see Methods), including 25bp intronic and 11bp exonic sequence at the splice acceptor site, and 10bp intronic and 11bp exonic sequence at the splice donor site. For exonic positions, non-synonymous variants were removed to minimise the chances of observing effects not due to splicing regulation. To investigate selective constraint within the splicing region, we utilise exome sequencing data from 13,750 unaffected parents within the DDD study, as well as >60,000 aggregated exomes from ExAC. To investigate near-splice positions in a disease-centric way, we analysed >16,700 high confidence DNMs identified within coding and near-coding regions well covered by exome sequencing in the 7,833 probands.

Materials and methods

Cohort and sequencing

For full description of cohort and analytical methodology, see previous DDD publications. Briefly, 7,833 patients with severe, undiagnosed developmental disorders were recruited to the DDD study from 24 clinical genetics centres from across the UK and Ireland. Whole exome sequencing was conducted on the proband and both parents, with exome capture using SureSelect RNA baits (Agilent Human All-Exon V3 Plus with custom ELID C0338371 and Agilent Human All-Exon V5 Plus with custom ELID C0338371) and sequencing using 75 base paired-end reads using Illumina’s HiSeq. Mapping was conducted to GRCH37 using the Burrows-Wheeler aligner (BWA, v0.59) and variant identification was conducted using the Genome Analysis Toolkit (GATK, v3.5.0). Variant annotation was conducted with Ensembl’s Variant Effect Predictor (VEP), using Ensembl gene build 76. DNMs were identified using DeNovoGear (v0.54), and filtered using an in house pipeline - denovoFilter - developed by Jeremy F. McRae (see web resources).

Defining exons of interest

We took exons from gencode v19 which met the following criteria: annotation_type = “exon”, gene_type = “protein_coding”, gene_status = “KNOWN”, transcript_type = “protein_coding”, transcript_status = “KNOWN”, annotation != “level 3” (automated annotation), and tag = “CCDS”, “appris_principal”, “appris_candidate_longest”, “appris_candidate”, or “exp_conf” (n = 255,812 exons). We removed a small subset of exons which no longer met these criteria in the more recent, GRCH38 based gencode v22 release (leaving 253,275 exons). We removed any exons where the median coverage at the canonical acceptor or donor positions was <15X in two sets of DDD data.
which used different exon capture methods (Agilent Human All-Exon V3 Plus with custom ELID
C0338371 and Agilent Human All-Exon V5 Plus with custom ELID C0338371). 148,244 exons passed
these criteria.

We annotated individual genomic positions relative to the acceptor and donor sites, removing any
exons <14bp, and any positions which had multiple potential annotations. At the acceptor end, we
considered 25bp of intronic sequence (acc-25 to acc-1) and 11bp exonic sequence (acc to acc+10). At
the donor end, we considered 10bp of intronic sequence (don+1 to don+10) and 11bp exonic
sequence (don to don-10). This yielded ~6.9 million near-splice positions of interest.

We define the polypyrimidine tract (PolyPy) region as acc-3, and acc-5 to acc-17, based on
pyrimidine content > 70% in our exons of interest. We assess changes from a pyrimidine to a purine
(PyPu) adjusting for the strand the exon is on.

**Mutability adjusted proportion of singletons (MAPS)**

In 13,750 unaffected parents enrolled as part of the DDD study, as well as >60,000 aggregated
exomes from ExAC v0.3.1, we calculated the MAPS metric using code developed in house by
Patrick J. Short (see web resources). This was done for all splice positions, the last base of the exon
split by reference nucleotide, and the PolyPy, split by PyPu vs all other changes, as well as VEP
ascertained synonymous, missense and nonsense sites across autosomal regions. To establish
whether the MAPS metric was significantly different between PolyPy PyPu vs all other changes, a
bootstrap resampling method was run with 1000 iterations.

**Parental variants in high pLI genes**

We looked at all variant positions overlapping with our splicing sites of interest within the
unaffected parental data, and calculated the proportion of these sites which fell within genes with
high probability of loss of function intolerance (pLI) scores (> 0.9) for each position of the splice
region, grouped sites, the last base of the exon split by reference allele, and the PolyPy split by PyPu
vs all other changes, as well as VEP annotated synonymous, missense, and nonsense sites across
autosomal regions.

**De novo mutations**

DNMs were identified using DeNovoGear as described in McRae et al, 2017, and a stringent
certainty threshold (posterior probability > 0.8) was applied. We used triplet-based mutation
rates to calculate the expected number of DNMs across autosomal regions in the 7,833 probands,
depth >1 and <50, exp*(0.119+0.204*log(depth))). We used the Poisson test to examine differences in the observed and expected values, and a 5% FDR correction to control for multiple testing (R v3.1.3). We stratified this analysis into known dominant, known recessive and non-DD associated genes using the DDG2P gene list, downloaded in June 2016. Genes with both recessive and dominant modes of inheritance were restricted to the dominant list.

We compared the relative distributions of CSS position variants with other protein truncating variants in the DDD and ExAC data.

Positive predictive values were calculated ((observed - expected) / observed) for CSS positions, combined and individually, don+5 sites, PolyPy PyPu, PolyPy other, other near splice exonic and intronic variants, as well as VEP defined missense and stop gained mutations.

We divided our exons into sextiles based on the pLI metric\textsuperscript{14}, and calculated the observed and expected number of DNMs in each group for don+5, PolyPy PyPu and synonymous variants (as above) to see if the enrichment of don+5 and PolyPy PyPu changes was concentrated in genes more likely to be intolerant of loss of function (LoF) mutations.

**Potential diagnostic variants**

DNMs overlapping with our near-splice positions of interest within dominant DDG2P genes were identified in DDD probands currently lacking a likely diagnosis (n = 5907). The HPO encoded\textsuperscript{23} phenotypes of the probands were assessed by consultant clinical geneticist Helen V. Firth, along with the patient’s recruiting clinician, and compared to the known clinical presentation of individuals with LoF mutations within those genes, classifying each variant as likely diagnostic, unlikely diagnostic, or unsure, depending on the strength of similarity between the proband and the disorder, and the specificity of the phenotype.

**Validation of putative splicing variants**

Eight variants were selected for validation via a minigene vector system. These comprised six likely diagnostic variants from the PolyPy, a PolyPy variant of uncertain clinical significance, and a likely diagnostic don+5 variant. Additionally, two untransmitted variants identified in unaffected parents within the same PolyPys as test variants were selected as negative controls. Details of all variants selected for validation are shown in Table S1.

**Cloning splicing vectors**
The minigene splice assay vector was adapted from that used in Singh et al., 201624, by replacing intron 1 with the first intron from the rat insulin 2 gene (Ins-2)(Rnor_6.0 Chr1:215857148-215857695). To generate individual assay vectors, either the 5’ most 231bp (for the don+5 variant) or the 3’ most 274bp (for PolyPy variants) of this vector was replaced with the appropriate endogenous intronic sequence encompassing the DNM of interest (Figure S3a and S3b), as described below.

First, proband genotypes (Table S1) were verified by capillary sequencing of genomic PCR products. Genomic regions containing the reference and alternate sequences were then either amplified by nested PCR, generated by site directed mutagenesis, or generated using gene synthesis (IDT). These fragments were sub-cloned, by Gibson Assembly (NEB), into our minigene vector. The regions assayed in our vectors are detailed by genomic coordinates in Table S1. Full sequence for all plasmids and primers available upon request.

**In vitro splicing assay**

HeLa cells were seeded into 12-well plates at a density of 160,000 cells per well, grown for 24 hours and transfected with 1 microgram of plasmid vector using Lipofectamine 3000 (Invitrogen). All transfections were carried out in duplicate and cultured for 48 hours. HeLa cells were cultured in DMEM (10% FCS + 1% pen/strep) at 37°C in a humidified incubator. Total RNA was extracted using a Micro RNeasy Qiagen kit and mRNA converted into cDNA using superscript IV (Invitrogen). RT-PCR was carried out using primers designed to span from exon 1 to exon 2, exon 2 to exon 3 and exon 1 to exon 3 and amplified on a thermocycler for either 25 or 35 cycles. Amplicons were capillary sequenced (GATC). For amplicons showing more than one splice variant (mixed capillary traces, for CHD7-Alt and MBD5-Alt), we cloned the PCR amplicons (Zero Blunt PCR cloning kit, Invitrogen) and sequenced individual colonies by capillary sequencing to identify the splice variants present. All PCR and sequencing primers available upon request.

Chromatograms were generated in R from .ab1 files using the sangerseqR25 package (R v3.1.3), and likely consequences on the protein primary structure were generated using reference and alternative RNA sequences with the ExPASy Nucleotide Sequence Translation tool26.

**Splicing pathogenicity scores**

Since our region of interest spanned >6 million individual positions, each with three potential single nucleotide changes, we were restricted in the choice of splicing pathogenicity prediction tools we could utilise, as many function primarily through a low throughput web interface model. We...
identified three resources recently published which provide “genome wide” splicing pathogenicity
scores. Two methods, dbscSNV’s AdaBoost and RandomForest are based on ensemble learning
combining predictions from multiple other splice prediction tools as well as conservation and CADD
scores\(^1\). The targeted region at the acceptor end spans 14 bases (12 intronic, 2 exonic) and at the
donor end spans 11 bases (8 intronic, 3 exonic). Spidex utilises deep learning methods trained on
RNA sequencing data to estimate the consequence of variants on the “percent spliced in” of an
exon, relative to the reference sequence\(^2\). Spidex scores positions up to 300bp from intron/exon
boundaries, so provides greater coverage of our splicing region of interest. We also utilised the
longer standing, and widely used MaxEntScan (MES)\(^3\), for which perl scripts were available, allowing
the tool to be run locally for all alternative alleles of all positions of interest. The metric used for MES
was the percent difference between the scores for the reference and alternative alleles, with the
greatest reduction in score classed as most pathogenic. All sites were also scored with CADD\(^4\).

To allow cross-tool comparison, we ordered positions by increasing pathogenicity from each metric,
and split positions into 20 brackets, such that the triplet based mutation rate for each bracket was
equal, and the 20\(^{th}\) bracket contained the positions with the most pathogenic scores. We calculated
MAPS and the proportion of parental variants falling in high pLI genes for each bracket for all five
metrics, as above, and looked at the number of DNMs in known dominant genes which fell in each
bracket for the five metrics. Each of these analyses was conducted including and excluding CSS
dinucleotide positions.

**Splice region variants in the ClinVar database**

We extracted all ClinVar\(^5\) variants using the UCSC table browser\(^6\) on 02.05.2017 and matched these
against our splicing positions of interest, removing exonic sites with non-synonymous consequences.
This resulted in 3603 positions with clinical significance recorded as pathogenic or likely pathogenic.
We calculated the ratio of canonical to non-canonical splice positions within this data. Since each
variant is present in this data only once, we used number of submissions as a proxy for allele count,
and calculated the ratio of canonical to non-canonical variants adjusting for this. Differences
between these observed values and our expectations, based on 27% of splice affecting mutations
being in non-canonical positions, were assessed using Fisher’s exact test (R v3.1.3).
Results

Signatures of purifying selection around the splice site

Since purifying selection acts to keep deleterious alleles rare, population variation data can be used to identify and assess the relative strengths of signals of purifying selection. To assess selective constraint acting on positions around the canonical splice site we used the MAPS metric\textsuperscript{14} in 13,750 unaffected parents enrolled in the DDD study as well as >60,000 aggregated exomes from ExAC (Figure 1a). There is a high level of consistency between the data from the DDD and ExAC cohorts. The canonical splice acceptor and donor dinucleotides show a clear signal of purifying selection in both datasets. In the DDD data, we observed a difference between the strength of selection for the two positions within the canonical donor site, with the donor+1 site showing a signal of selection akin to stop-gained mutations, while the donor+2 site is lower, intermediate between the signal of selection observed at missense and nonsense sites. Concordant with this observation of apparently weaker purifying selection acting on variants at the CSS compared to nonsense variants, we identified that CSS variants represent 21.7% of likely protein truncating variants in highly constrained (pLI>0.9) genes in ExAC individuals without overt monogenic disorders, but only 13.9% of likely protein truncating DNMs in DDD probands. These proportions are significantly different (p = 4.21x10\textsuperscript{-6}), and support the idea that, on average, CSS mutations have lower pathogenicity than other predicted truncating variants.

Outside of the CSS, other positions clearly show a signal of purifying selection beyond the background level, including the donor site (last base of the exon, which is particularly pronounced when the reference allele is G (Figure 1a)), and the intronic positions proximal to the canonical donor site, peaking at the don+5 position, which exhibits a signal of purifying selection intermediate between missense and nonsense variants. Although no sites within the PolyPy show a signal of purifying selection individually, when these sites are grouped together (Methods) and stratified by changes from a pyrimidine to a purine (PyPu), versus all other changes, there is a clear difference between the two types of variants, with PyPu changes exhibiting an increased signal of purifying selection when compared to non-PyPu changes (p < 0.001, Figure 1a, Figure S1).

Deficit of splicing variants in highly constrained (pLI>0.9) genes in healthy individuals

We also examined the distribution of variants of different classes among genes that are known to be under different levels of selective constraint. Highly constrained genes should contain fewer deleterious variants than less constrained genes. We investigated the proportion of variants observed in the 13,750 unaffected parents which fell within highly constrained genes with pLI > 0.9
Figure 1 – Signals of purifying selection around splice sites

A. Selective constraint across splicing region in 13,750 unaffected parents of DDD probands and >60,000 aggregated exomes from ExAC. Mutability adjusted proportion of singletons (MAPS) shown for VEP annotated exonic sites, extended splice acceptor and splice donor regions, the last base of the exon, split by reference nucleotide, and grouped sites in the polypyrrimidine tract region (PolyPy), split by changes from a pyrimidine to a purine (PyPu) vs all other changes. B. Proportion of variants in 13,750 unaffected parents of DDD probands which fall within genes with high pLI (>0.9) across VEP annotated exonic sites, extended splice acceptor and splice donor regions, the last base of the exon, split by reference nucleotide, and grouped sites in the polypyrrimidine tract region, split by changes from a pyrimidine to a purine (PyPu) vs all other changes.
in our splicing regions of interest (Figure 1b). In the near splice positions at which the highest MAPS values were seen (CSS, donor, donor+5), we also observed a stronger depletion of variants in high pLI genes within the unaffected parents, again supporting the potential pathogenicity of variants at these positions. The proportion of parental variants in high pLI genes also recapitulates the signals of purifying selection seen in the MAPS analyses with regard to the donor position split by reference allele (Figure 1b) and the PolyPy region (Figure 1b, Figure S1), with the lowest proportions in high pLI genes observed for sites with the highest MAPS values.

**Assessing the significance of mutational burden for different classes of splicing mutations**

We identified 871 autosomal high confidence DNMs (non-synonymous consequences excluded) within canonical and near-splice regions of interest well covered by exome data in the 7,833 probands, allowing us to test for enrichment of DNMs relative to expectations based on a trinucleotide null model of mutation rate across different sets of genes (DD-associated with dominant or recessive mechanisms, and non-DD associated, see Methods). Across recessive DD and non-DD associated genes, no enrichment of DNMs beyond the null expectation was observed (Figure 2a). In dominant DD genes, a significant cumulative excess of DNMs was noted across the full splicing region (Poisson p (FDR adjusted) = 2.60x10^{-14}, fold enrichment = 3.47), which remained significant upon exclusion of the canonical dinucleotide positions (Poisson p (FDR adjusted) = 0.0041, fold enrichment = 1.86). Individually, the four canonical splice site positions each showed a significant excess of DNMs (Poisson p (FDR adjusted), fold enrichment: acc-2 = 7.68x10^{-12}, 26.61; acc-1 = 6.02x10^{-8}, 16.57; don+1 = 2.60x10^{-14}, 20.06; don+2 = 0.0045, 9.99), as did the don+5 site (0.0001, 9.29). The greater fold enrichment in don+1 over don+2 sites suggests variants at don+2 may be less damaging, while the similar level of enrichment between don+5 and don+2 implies these positions harbour comparable levels of splice disrupting mutations. No individual positions within the PolyPy region showed an individual excess of DNMs, however, when the positions were considered cumulatively and split between PyPu and non-PyPu changes (Figure 2b), a significant excess of DNMs was observed in the PyPu group for dominant DD genes (Poisson p (raw) = 0.0048, fold enrichment = 3.46). These results are highly concordant with the signatures of purifying selection identified using the MAPS metric, and the deficit of parental variants in high pLI genes, providing multiple independent lines of evidence that the canonical positions do not contribute equally to splice site recognition, and that mutations in positions outside of the CSS can disrupt normal splicing.
Figure 2 – *De novo* mutations around splice sites

Enrichment of *de novo* mutations (DNMs) across the splicing region in 7,833 DDD probands. A. Numbers of observed and expected DNMs across splicing region, in known dominant and recessive DD genes, as well as non-DD associated genes, with FDR corrected Poisson p-values. B. Aggregation of observed and expected numbers of DNMs in the polypyrimidine tract (PolyPy) region, with changes from a pyrimidine to a purine and all other changes shown separately for known dominant and recessive DD genes, as well as non-DD associated genes. C. Positive predictive values (PPVs) for *de novo* mutations in dominant DD-associated genes in positions across the splicing region, as well as VEP annotated stop gained and missense changes, calculated from observed and expected numbers of DNMs.
Estimating positive predictive values for different classes of splice mutation

We used the fold-enrichment of the numbers of observed DNMs in dominant DD genes in the DDD cohort over the number expected under the null mutation model to calculate positive predictive values (PPV) for groupings of near splice site positions, independent of any prior clinical classification. We compared these with positive predictive values for other, more commonly disease-associated variant classes within the same exons of the same genes (Figure 2c). We observe minor differences in PPV for the individual positions of the canonical acceptor and donor sites, with don+2 showing the lowest PPV at 0.90, which is approximately the same as for the don+5 positions (PPV 0.89). Variants within the PolyPy region which change a pyrimidine for a purine have a PPV of 0.71, which is below the PPV for missense mutations (0.79), but still predicts a substantive number of pathogenic mutations arising from disruption of the PolyPy.

Despite the modest number of observed DNMs used to make these PPV estimates, we see striking concordance with the population based metrics described above (MAPS and deficit of splicing variants in high pLI genes in unaffected parents of DDD patients – Figure S2), suggesting these estimates are robust.

Identifying diagnostic non-canonical splice mutations

After exclusion of probands with likely diagnostic protein-coding or canonical splice site variants, 38 DNMs in our near splice site positions of interest in dominant DD genes were identified. The clinical phenotypes of patients carrying these mutations were reviewed by a consultant clinical geneticist, blinded to the precise mutation and PPVs estimated above, and the patient’s recruiting clinician, to assess the phenotypic similarity between the proband and the disorder expected from a LoF mutation in that gene. The 38 variants were classified as likely diagnostic (Table 1), or unlikely diagnostic/unknown (Table 2), depending on the strength of phenotypic similarity. The clinical review resulted in 18 variants (47%) being classified as likely diagnostic, highly concordant with the number predicted from the overall PPV of non-canonical sites of 46%, moreover, a higher proportion of likely diagnostic variants were classified at sites with higher PPVs, calculated as described above (Figure 3). With 48 CSS DNMs observed within the same exons in our probands, we estimate that 73% of disease causing splice disrupting DNMs occur within the CSS, while 27% are in non-canonical, near-splice positions.
Figure 3 – Clinical classifications of non-canonical near splice de novo mutations

Relationship between clinical classifications of 38 splice region de novo mutations (DNMs) in undiagnosed DDD probands and positive predictive values (PPVs) calculated using observed and expected numbers of DNMs in 7,833 probands.
Eight DNMs were selected for functional validation via a minigene vector system, including six likely diagnostic PolyPy variants, a PolyPy variant of uncertain clinical significance, and a likely diagnostic don+5 variant, where both the phenotype of the patient and that associated with the gene (MBD5) are nonspecific, along with two negative controls (untransmitted variants identified in unaffected parents within the same PolyPys as test variants). For six of the variants selected for validation, differences in splicing between the reference and mutant constructs were observed (Figure S3a and S3b). The five PolyPy variants generating alterations in splice products all generated a cryptic splice site upstream of the canonical splice site, causing retention of part of the intron, in four instances leading to a frameshift effect, and in one leading to the inclusion of two additional amino acids in the protein sequence. The don+5 variant caused the utilisation of a second “GT” site within the reference sequence as a splice donor site, again causing retention of intronic sequence within the transcript, leading to a frameshift effect. For the CHD7 variant, two splice products were observed, corresponding to the expected (wild type) splicing and the retention of 5bp intronic sequence. The MBD5 variant gave multiple splice isoforms, with retention of 12bp intronic sequence being the most prevalent, but normally spliced, 19bp intronic retention, and complete intron retention also observed. Figure S3 shows the predominant isoform observed for these variants. One of the likely diagnostic PolyPy mutations, the PolyPy mutation of uncertain significance, and both negative controls showed no difference in splicing between the reference and mutant constructs (Figure S3c and S3d).

Given the concordant signals of purifying selection, enrichment of DNMs and number of likely diagnostic variants in the don+5 site and PolyPy (PyPu) region, we looked at the distribution of observed DNMs in genes with respect to their probability of being LoF intolerant (using the pLI metric14, Figure 4). For synonymous variants, we observed no significant enrichment of DNMs in high pLI genes. For don+5 mutations, there is a clear excess of DNMs in genes most likely to be intolerant to LoF mutations in the DDD cohort, further supporting the likely pathogenicity of mutations in these positions. For the PolyPy PyPu mutations, although there is a nominally significant enrichment of DNMs in general, this does not show a significant skew towards high pLI genes in our cohort.
**Figure 4 – Enrichment of de novo mutations by probability of loss of function intolerance**

Enrichment (observed/expected) of de novo mutations (DNMs) by gene probability of loss of function intolerance (pLI), split in to sextiles for donor+5, pyrimidine to purine PolyPy, and synonymous sites. pLI scores encompassed by each sextile: 1 = 5.36E-91 - 0.000000605, 2 = 0.000000609 - 0.000558185, 3 = 0.000559475 - 0.027905143, 4 = 0.027908298 - 0.377456159, 5 = 0.377491926 - 0.919495985, 6 = 0.91955878 - 1.

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**Assessing splicing pathogenicity prediction tools**

The population genetic metrics of purifying selection and mutation enrichment metric for pathogenicity that we have derived provide an orthogonal approach to assessing the accuracy of splicing pathogenicity prediction tools. We assessed four splicing pathogenicity prediction tools: two recently published genome-wide ensemble learning methods: AdaBoost and RandomForest, Spidex (based on deep learning trained on RNA sequencing data), and the longer standing, widely used MaxEntScan\(^{27-29}\).
We divided the scores from each prediction tool, plus CADD, into 20 bins of equal mutation rate, to facilitate cross-method comparability. We calculated the MAPS for each bin of each of the scoring metrics for the splicing variants observed in the 13,750 DDD unaffected parents, and saw a strong positive correlation between pathogenicity metric and MAPS for all tools (Figure S5). AdaBoost had the highest absolute MAPS value for the top scoring bin, suggesting that it is best able to identify variants under the strongest purifying selection. The proportion of variants in the unaffected parents falling in genes with pLI > 0.9 broadly recapitulates this pattern, with fewer variants in high pLI genes in the highest scoring brackets for all metrics (Figure S4). We then looked at the distribution of scores for each tool for the 83 splicing DNMs observed in DDD probands in autosomal dominant DD-associated genes which were covered by all five scoring systems to compare performance of the metrics on mutations more likely to have a deleterious impact on splicing (Figure 6). Again, all metrics performed well, with the majority of DNMs being classified in the most deleterious score brackets. Here AdaBoost gave the highest area under the curve (AUC) value, suggesting it weighted these likely damaging variants as more deleterious than the other metrics comparatively.

Interestingly, when CSS positions were removed from the analysis, AdaBoost remained the tool with the highest AUC. The largest reduction in the AUC metric was seen for Spidex and CADD, indicating these tools may be least informative for positions outside of the CSS. Upon removal of the CSS positions from the analyses of MAPS and deficit of parental variants in high pLI genes, similar results were revealed, with the highest AdaBoost scores retaining strong signals of purifying selection but a marked reduction in signal from the highest Spidex scores (Figure S5 and Figure S6).

Taken together, these data show a strong relationship between the considered splicing pathogenicity scoring systems and the general landscape of purifying selection on splicing control, but demonstrate that the utility of these systems in identifying likely diagnostic variants is limited outside of the CSS. However, it is worth noting that the scores reflect the probability of a variant affecting splicing, rather than overt phenotype. The high scoring variants may be affecting the splicing of transcripts, albeit not sufficiently to cause disease.
Figure 5 – Selective constraint and pathogenicity scores

Mutability adjusted proportion of singletons (MAPS) calculated for pathogenicity score brackets (least to most severe) in 13,750 unaffected parents from the DDD project, with Spearman correlation coefficient.
Figure 6 – Pathogenicity scores for observed near splice site de novo mutations

Cumulative percentage of de novo mutations (DNMs) in known dominant DD genes with decreasing pathogenicity score bracket, shown with canonical splice site positions included (left) and excluded (right). * AUC = area under curve
Discussion

Our analyses, taken together, suggest the pathogenic contribution of non-canonical splice sites has been under-appreciated. We estimate that around 27% of splice disrupting pathogenic mutations within the DDD cohort are in non-canonical positions. In sites with pathogenic or likely pathogenic clinical significance in ClinVar\(^{31}\) overlapping with our splicing positions of interest (with non-synonymous consequences removed), we found 83.5% of variants fell within canonical positions, with just 16.5% in non-canonical positions. When adjusted for number of submissions as a proxy for allele count, this figure was 17.5%, perhaps indicative that recurrence strengthens evidence of pathogenicity. Both of these values are significantly below our estimate of 27% (\(p = 1.22 \times 10^{-15}\) and \(p = 2.2 \times 10^{-16}\) respectively), suggesting under-ascertainment of non-canonical splicing variants by around 35-40% in clinical databases.

Estimates of the relative contribution of canonical and non-canonical splice site mutations are sparse in the literature. When comparing canonical and non-canonical mutations within HGMD, Krawczak et al.\(^{33}\) stated canonical mutations accounted for 64% of mutations at donor sites and 77.4% of mutations at acceptor sites, giving an estimated non-canonical contribution of ~30% overall, while data taken from Caminsky et al.\(^{34}\) put this estimate at around 43%. These values are much closer to our 27% estimate than to the ClinVar proportion of ~17%, despite our approach focussing on DNMs and dominant disorders, whereas the other two studies did not discriminate on mode of inheritance and included recessive disorders, which can also be caused by non-canonical splicing mutations\(^ {35; 36}\) and exonic variants.

Our analysis of non-canonical splice position mutations did not include exonic missense variants\(^ {4; 5; 33}\), nor did it explicitly include branchpoints\(^ {37-40}\), splicing enhancers and suppressors\(^ {41; 42}\) or deep intronic mutations\(^ {43; 44}\). Detecting splice disrupting variants at these sites is even more challenging, as despite recent efforts\(^ {45-49}\), comprehensive catalogues of all branchpoints and ESE/ISE/ESS/ISS are currently unavailable, algorithms that predict the impact of mutations at such sites are not highly accurate, and some of these sites are not covered by exome sequencing. As such, our estimate of the contribution of non-canonical splicing position mutations is likely to be a lower bound. If the higher estimated non-canonical contributions from Krawczak and Caminsky are more accurate, our estimate of 35-40% under-ascertainment in clinical databases may be conservative, and the true extent of missed diagnoses may be even higher.

The nature of developmental disorders makes obtaining RNA samples from relevant tissues of patients (i.e. neural tissue) acutely problematic, so we investigated the effects on splicing of several
of the potentially diagnostic DNMs using a minigene vector system. We were able to demonstrate changes to splicing for five out of six likely diagnostic PolyPy variants as well as the likely diagnostic don+5 variant. We did not observe an effect on splicing for one likely diagnostic PolyPy variant, and one PolyPy variant of uncertain clinical significance. Although the accuracy of minigene assays when compared with patient RNA is generally high\textsuperscript{50-52}, known limitations of the system (e.g. lack of full endogenous genetic context\textsuperscript{53; 54}, and sensitivity to cell type utilised\textsuperscript{55}) mean we cannot definitively state that the effects seen in the minigene assay would be the same in the full genetic, developmental and cellular context within the patient.

We envisage that greater appreciation of the importance of near splice site mutations will increase diagnostic yields, as well as providing increased power for the detection of new genetic associations, both within the field of rare disease and beyond. We highlight two challenges to improving detection of pathogenic non-canonical splice site mutations.

First, many commonly used tools for annotating the likely functional impact of variants do not discriminate between different non-canonical splice site positions with very different likelihoods of being pathogenic. Moreover, commonly used annotation tools differ in the ways in which variants are annotated, with splicing variants displaying the highest level of disagreement between tools\textsuperscript{56}. This highlights the need for a more consistent and evidence based annotation of splicing variants. Of the positions shown in our analyses to be most damaging, don+5 sites are annotated by VEP\textsuperscript{19} and SnpEff\textsuperscript{57} as “splice_region_variant”, while most positions of the PolyPy are annotated as intronic, so are potentially easily overlooked. With Annovar\textsuperscript{58}’s default settings, only the CSSs are flagged as splicing variants, although with both Annovar and SnpEff, the user can optionally extend the region to be annotated as splice variants. We note that Ensembl have recently implemented a VEP plugin which allows greater granularity in splice region annotation (see web resources), including annotating the don+5 and other near-donor positions, as well as the PolyPy region. This type of increased granularity of splicing annotation should facilitate consideration of these variants in future studies.

Second, current tools that predict the pathogenicity of non-canonical splice site mutations have limited accuracy, and it is not clear how to translate the scores that they output into a likelihood of pathogenicity. The quantitative framework that we introduced here of estimating PPVs for different classes of mutations by comparing the number of observed mutations to the number expected under a well-calibrated null model of germline mutation has much more direct relevance to clinical interpretation. We propose that the scores generated by such splicing prediction tools could be
calibrated by performing analogous analyses of mutation enrichment to estimate PPVs for different bins of scores. As the size of trio-based cohorts increases, the accuracy of calibration will improve.

In summary, our results demonstrate a significant contribution of non-canonical splicing mutations to the genetic landscape of DDs, a finding which is highly likely to be recapitulated across other monogenic disorders and contexts. We demonstrate disparities in the control of splice site recognition between the two positions of the canonical splice-donor site, and the importance of other, non-canonical positions (particularly the don+5 site and pyrimidine-removing mutations in the PolyPy region). These inferences are supported by both population genetic investigations of purifying selection, as well as a disease based approach, considering the burden of DNMs in ~8,000 children with severe DDs. Mutations at some non-canonical splicing positions convey a risk of disease similar to that of protein truncating and missense mutations, but are commonly under-represented in existing databases of disease-causing variants.
Description of supplemental data

Supplemental data contains six figures and one table.

Conflicts of interest

M.E.H. is a co-founder of, consultant to, and holds shares in, Congenica Ltd, a genetics diagnostic company.

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Web resources

denovoFilter, https://github.com/jeremymcrae/denovoFilter

Gencode v19, https://www.gencodegenes.org/releases/19.html

MAPS, https://github.com/pjshort/dddMAPS

Ensembl’s VEP, https://www.ensembl.org/info/docs/tools/vep/index.html

DDG2P, http://www.ebi.ac.uk/gene2phenotype

ExAC, http://exac.broadinstitute.org/

HPO, http://compbio.charite.de/hpoweb/showterm?id=HP:0000118

SangerSeqR, http://bioconductor.org/packages/release/bioc/html/sangerseqR.html

ExPASy, https://web.expasy.org/translate/

CADD, http://cadd.gs.washington.edu/

ClinVar, https://www.ncbi.nlm.nih.gov/clinvar/

SpliceRegion.pm, https://github.com/Ensembl/VEP_plugins/blob/release/88/SpliceRegion.pm

OMIM, https://www.omim.org/
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**Table 1 – Diagnostic de novo mutations in non-canonical dinucleotide near splice positions**

Variant and proband information for 18 de novo likely diagnostic splice region variants identified in previously undiagnosed DDD probands in known dominant DD-associated genes (hg19 coordinates)

| chrom:pos_ref/alt | symbol | VEP annotation | Splice annotation | Associated disorder | HPO terms | HPO terms (translation) | Clinical classification |
|-------------------|--------|----------------|-------------------|--------------------|-----------|------------------------|------------------------|
| 7:42063221_G/C   | GLI3   | intron_v variant | acc-14            | Greig Cephalopolysyndactyly Syndrome | HP:0001841, HP:0010709, HP:0011304 | 2-4 finger syndactyly, Broad thumb, Preaxial foot polydactyly | Likely pathogenic, full contribution |
| 16:3819367_C/T   | CREBBP | intron_v variant | acc-13            | Rubinstein-Taybi Syndrome Type 1 | HP:0000028, HP:0000179, HP:0000248, HP:0000252, HP:0000347, HP:0000486, HP:0001263, HP:0001510, HP:0001831, HP:0002019, HP:0002205, HP:0004691, HP:0011304 | 2-3 toe syndactyly, Brachycephaly, Broad thumbs, Constipation, Cryptorchidism, Global developmental delay, Growth delay, Microcephaly, Micrognathia, Recurrent respiratory infections, Short toes, Strabismus, Thick lower lip vermilion | Likely pathogenic, full contribution |
| 22:24143120_T/G  | SMARCB1| intron_v variant | acc-11            | Rhabdoid Predisposition Syndrome 1 / Coffin-Siris Syndrome 3 | HP:0000294, HP:0000680, HP:0000696, HP:0000750, HP:0001263, HP:0001763, HP:0001999, HP:0002205, HP:0002213, HP:0007021, HP:0007096, HP:0010830, HP:0010877, HP:0100543, HP:0003045, HP:0002926, HP:000708, HP:0000545, HP:0002164, HP:0001212 | Unilateral strabismus, Delayed eruption of permanent teeth, Delayed eruption of primary teeth, Delayed speech and language development, Global developmental delay, Cognitive impairment, Impaired tactile sensation, Pain insensitivity, Pes planus, Recurrent respiratory infections, Fine hair, Low anterior hairline, Abnormal facial shape, Hypoplasia of the optic tract, Abnormality of the patella, | Likely pathogenic, full contribution |
| Chromosome Position | Gene | Variant Type | Change | Condition | Likely Pathogenicity |
|---------------------|------|--------------|--------|-----------|---------------------|
| 18:52895603_T/C     | TCF4 | intron_variant | acc-11 | Abnormality of thyroid physiology, Behavioural abnormality, Myopia , Nail dysplasia, Prominent fingertip pads | Abnormal facial shape, Amblyopia, Global developmental delay, Microcephaly, Myopia, Unilateral renal agenesis | Likely pathogenic, full contribution |
| 5:88025173_A/C      | MEF2C| splice_region_variant | acc-9  | Abnormality of the cerebral white matter, Prolonged neonatal jaundice, Recurrent hand flapping, Seizures, Severe global developmental delay, Thick lower lip vermilion | Likely pathogenic, full contribution |
| 9:130988306_G/A     | DNM1 | splice_region_variant | acc-8  | Delayed speech and language development, Postnatal microcephaly, Short stature | Likely pathogenic, full contribution |
| 8:61763045_G/A      | CHD7 | splice_region_variant | acc-7  | Cleft soft palate, Coloboma, Global developmental delay, Oral cleft, Proportionate short stature, Tetralogy of Fallot with pulmonary atresia and major aortopulmonary collateral arteries, obsolete Malformation of the heart and great vessels | Definitely pathogenic, full contribution |
| 17:38801875_T/C     | SMARCE1| splice_region_variant | acc-4  | Abnormality of the cerebral white matter, Prolonged neonatal jaundice, Recurrent hand flapping, Seizures, Severe global developmental delay, Thick lower lip vermilion | Likely pathogenic, full contribution |
| Chromosome | Gene Symbol | Mutation Type | Annotation | Description | Pathogenicity |
|------------|-------------|---------------|------------|-------------|--------------|
| 1:27097607_C/A | ARID1A | splice_region_variant | acc-3 | Coffin-Siris Syndrome 2 | Abnormality of the pinna, Cutis laxa, Deeply set eye, Generalized neonatal hypotonia, Hearing abnormality, Intrauterine growth retardation, Low-set ears, Micrognathia, Partial agenesis of the corpus callosum, Thick lower lip vermilion | Likely pathogenic, full contribution |
| 9:140728798_C/G | EHMT1 | splice_region_variant | acc-3 | 9q Subtelomeric Deletion Syndrome / Kleefstra Syndrome 1 | Supernumerary nipple, Gastroesophageal reflux, Pyloric stenosis, Moderate conductive hearing impairment, Truncal obesity, Brachycephaly, Enlarged tonsils, Hypoplastic toenails, Abnormal social behaviour, Cognitive impairment, Delayed speech and language development, Happy demeanor, Intellectual disability, severe, Premature birth following premature rupture of fetal membranes | Definitely pathogenic, full contribution |
| 2:223160248_T/C | PAX3 | splice_region_variant | don-1 | Waardenburg Syndrome, Type 1 / Craniofacial-Deafness-Hand Syndrome | Arthralgia, Blepharophimosis, Few cafe-au-lait spots, Freckles in sun-exposed areas, High palate, Hypertelorism, Localized hirsutism, Long eyelashes, Narrow nose, Upslanted palpebral fissure, High nasal bridge, Low-frequency sensorineural hearing impairment, Lacrimal duct obstruction, Narrow ear canal | Likely pathogenic, partial contribution |
| Chromosome Start/End | Gene | Splice Region Variant | Donor/ acceptor | Description | OMIM Numbers | Likely Pathogenic Status |
|----------------------|------|-----------------------|------------------|-------------|--------------|--------------------------|
| 2:166229861_A/G      | SCN2A| splice_region_variant | don+4            | Nonspecific Severe Id / Benign Familial Neonatal Infantile Seizures / Infantile Epileptic Encephalopathy | HP:0000717, HP:0001344, HP:0002342, HP:0001250 | Likely pathogenic, full contribution |
| 9:130422391_A/G      | STXBP1| splice_region_variant | don+4            | Angelman/Pitt Hopkins Syndrome-Like Disorder / Epileptic Encephalopathy Early Infantile Type 4 | HP:0001048, HP:0001252, HP:0002599, HP:0011344 | Likely pathogenic, full contribution |
| 22:41556731_G/A      | EP300| splice_region_variant | don+5            | Rubinstein-Taybi Syndrome Type 2 | HP:0000023, HP:0000213, HP:0000220, HP:0000322, HP:0000369, HP:0000414, HP:0000486, HP:0000490, HP:0000527, HP:0001263, HP:0001537, HP:0001771, HP:0005484, HP:0007993, HP:0008551, HP:0008850, HP:0100023, HP:0000717 | Likely pathogenic, full contribution |
| Chromosome | Gene | Splice Region | Donor | Description | OMIM Numbers | Conditions | Pathogenicity |
|------------|------|---------------|-------|-------------|---------------|------------|--------------|
| 2:149221493 | MBD5 | splice_region | don+5 | Ehmt1-Like Intellectual Disability | HP:0000252, HP:0000664, HP:0001601, HP:0002020 | Gastroesophageal reflux, Laryngomalacia, Microcephaly, Synophrys | Likely pathogenic, full contribution |
| 9:130427615 | STXB1 | splice_region | don+5 | Angelman/Pitt Hopkins Syndrome-Like Disorder / Epileptic Encephalopathy Early Infantile Type 4 | HP:0000733, HP:0002066, HP:0002378, HP:0002943, HP:0003763, HP:0007359, HP:0010864 | Bruxism, Focal seizures, Gait ataxia, Hand tremor, Intellectual disability, severe, Stereotypy, Thoracic scoliosis | Likely pathogenic, full contribution |
| 17:42956919 | EFTUD2 | splice_region | don+5 | Mandibulofacial Dysostosis With Microcephaly | HP:0000253, HP:0000286, HP:0000384, HP:0000396, HP:0000412, HP:0011343 | Epicanthus, Moderate global developmental delay, Overfolded helix, Preauricular skin tag, Progressive microcephaly, Protruding ear | Definitely pathogenic, full contribution |
| 20:61452890 | COL9A3 | splice_region | don+8 | Multiple Epiphyseal Dysplasia Type 3 | HP:0000729, HP:0000750, HP:0010529, HP:0000735, HP:0001382, HP:0008947, HP:0000736, HP:0000733 | Autistic behavior, Delayed speech and language development, Echolalia, Impaired social interactions, Joint hypermobility, Muscular hypotonia, Short attention span, Stereotypy | Likely pathogenic, partial contribution |
Table 2 – *De novo* mutations in non-canonical near splice positions not thought to be diagnostic

Genomic coordinates and annotations of 20 *de novo* splice region variants identified in undiagnosed DDD probands in known dominant DD-associated genes, deemed unlikely to be diagnostic based on lack of phenotypic match between proband and associated syndrome (hg19 coordinates)

| Variant          | Gene    | VEP annotation | Splice Annotation | Clinical classification |
|------------------|---------|----------------|-------------------|-------------------------|
| 18:42618432_G/T  | SETBP1  | intron_variant | acc-18            | Likely benign           |
| 3:38988415_AC/A  | SCN11A  | intron_variant | acc-17            | Likely benign           |
| 17:38240072_A/G  | THRA    | intron_variant | acc-16            | Likely benign           |
| 19:13387958_G/A  | CACNA1A | intron_variant | acc-16            | Likely benign           |
| 3:189456422_A/G  | TP63    | intron_variant | acc-9             | Likely benign           |
| 1:7309543_GT/GTT | CAMTA1  | splice_region_variant | acc-8        | Likely benign           |
| 16:30745810_C/G  | SRCAP   | splice_region_variant | acc-7        | Likely benign           |
| 16:29816431_G/A  | KIF22   | splice_region_variant | acc-5        | Likely benign           |
| 22:41543944_C/T  | EP300   | synonymous_variant | don-6        | Likely benign           |
| 10:94381235_G/T  | KIF11   | splice_region_variant | don+5        | Likely benign           |
| 16:2129206_G/A   | TSC2    | intron_variant | don+9             | Likely benign           |
| 2:158594942_G/T  | ACVR1   | intron_variant | don+10            | Likely benign           |
| 19:50912018_C/T  | POLD1   | synonymous_variant | acc-24        | Uncertain               |
| 3:41266439_T/G   | CTNNB1  | splice_region_variant | acc-6        | Uncertain               |
| 17:44159911_GC/G | KANSL1  | splice_region_variant | acc-3        | Uncertain               |
| 3:71021701_C/T   | FOXP1   | splice_region_variant | don+5        | Uncertain               |
| 6:157431700_G/A  | ARID1B  | splice_region_variant | don+5        | Uncertain               |
| X:41196724_T/G   | DDX3X   | splice_region_variant | don+6        | Uncertain               |
| 3:111366523_A/C  | CD96    | intron_variant | don+10            | Uncertain               |
| 8:117869033_G/C  | RAD21   | intron_variant | acc-23            | Uncertain               |